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CASA Synthesis & Single Dish Reduction
Reference Manual & Cookbook

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Jeff Kern – CASA Project Manager

http://casa.nrao.edu

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3.7 Multi-panel display of visibility versus channel (top), antenna array configuration (bottom left) and the resulting uv coverage (bottom right). The commands to make these three panels respectively are: 1) `plotxy('ngc5921.ms', xaxis='channel', datacolumn='data', field='0', subplot=211, plotcolor='r', plotsymbol='go')` 2) `plotxy('ngc5921.ms', xaxis='x', field='0', subplot=223, plotcolor='g', plotsymbol='r')`, 3) `plotxy('ngc5921.ms', xaxis='u', yaxis='v', field='0', subplot=224, plotcolor='b', plotsymbol='g')`. 

3.8 Plot of amplitude versus uv distance, before (left) and after (right) flagging two marked regions. The call was: `plotxy(vis='ngc5921.ms', xaxis='uvdist', field='1445*', plotcolor='blue', plotsymbol='o')`.

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4.11 Use of plotxy to display corrected data (red and blue points) and uv model fit data (green circles).

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5.2 Screen-shots of the interactive clean window during deconvolution of the VLA 6m Jupiter dataset. We start from the calibrated data, but before any self-calibration. In the initial stage (left), the window pops up and you can see it dominated by a bright source in the center. Next (right), we zoom in and draw a box around this emission. We have also at this stage dismissed the tape deck and Position Tracking parts of the display (§ 7.2) as they are not used here. We have also changed the iterations to 30 for this boxed clean. We will now hit the Next Action Continue Cleaning button (the green clockwise arrow) to start cleaning.
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7.4 The ‘Mouse Tool’ Bar allows you to assign how mouse buttons behave in the image display area. Initially, zooming, color adjustment, and rectangular regions are assigned to the left, middle and right mouse buttons. Click on a tool with a mouse button to assign that tool to that mouse button.

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7.8 The load tab of the Data Manager panel. This appears if you open the viewer without any infile specified, if you use select Open from the Data drop down menu, or click the Open (Folder) icon. You can access the save image or save region tabs from this view or by selecting Save as... from the Data drop down menu. The load tab shows all files in the current directory that can be loaded into the viewer — images, MS, CASA region files, and Display Panel State files.

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7.23 The toolbar for the Spectral Profile tool allows the user to save the spectrum, print or save the tool as an image, edit preferences (general, tool, legend), spectral smoothing, pan or zoom around the spectrum, select a range of interest, jump to a channel, or add a label.
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7.28 Line Overlays in the Spectral Profile Tool. The Line Overlay tab, shown at the bottom, allows users to query the CASA copy of the Spaltalogue spectral line database. Enter the redshift of your source (right panel), select and Astronomical Filter from the drop down menu, and use shift+click+drag to select a frequency range (or do so manually). The "Search" button will bring up the dialog seen at the left top part of the image, which can in turn be used to graph the candidate lines in the main Spectral Profile window (here CO v=0).

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7.32 The interface to the two dimensional fitting tool (Tools:Fit... or the blue circles icon). The interface allows you to specify and automatically generate (Find Sources) initial estimates, to specify the range of pixel values to be included in the fit, and to specify the output (log file, residual image, and visualization). Click Fit to start the fit.

7.33 Interactive creation of position-velocity cuts in the viewer. Use the P/V tool from the Mouse Toolbar to define a cut, then use the pV tool from the Region Manager Panel to adjust the cut (including the width). Click Generate P/V to build the position velocity cut and open it in a new Viewer Display Panel (from which it can be saved to disk).
7.34 The **Load Data - Viewer** panel as it appears if you select an MS. The only option available is to load this as a **Raster Image**. In this example, clicking on the **Raster Image** button would bring up the displays shown in Figure 7.2.

7.35 The MS for NGC4826 BIMA observations has been loaded into the viewer. We see the first of the **spw** in the Display Panel, and have opened up MS and Visibility Selections in the **Data Display Options** panel. The display panel raster is not full of visibilities because **spw 0** is continuum and was only observed for the first few scans. This is a case where the different spectral windows have different numbers of channels also.

7.36 The MS for NGC4826 from Figure 7.35, now with the **Display Axes** open in the **Data Display Options** panel. By default, **channels** are on the **Animation Axis** and thus in the tapedeck, while **spectral window** and **polarization** are on the **Display Axes** sliders.

7.37 The MS for NGC4826, continuing from Figure 7.36. We have now put **spectral window** on the **Animation Axis** and used the tapedeck to step to **spw 2**, where we see the data from the rest of the scans. Now **channels** is on a **Display Axes** slider, which has been dragged to show **Channel 33**.

7.38 Printing the display to a hardcopy of a file. From the **Viewer Print Manager**, located in top right here and accessed by the print icon or from the Data drop down menu, you can use the **Save** button to save an image or **Print** directly to a printer. To achieve the best results, it is often helpful to adjust the settings in the **Data Display Options** and **Viewer Canvas Manager**, shown at right.

7.39 Data selection in **msview**.

8.1 An example of figure using **plotter2**.

8.2 Multi-panel display of the scantable. Sub-panels are displayed per scan. There are two spectra in each scan indicating two polarization (RR and LL).

8.3 Two panel plot of the calibrated spectra. The GBT data have a separate scan for the SOURCE and REFERENCE positions so scans 20, 21, 22 and 23 result in these two spectra.

8.4 Calibrated spectrum with a line at zero (using histograms).

8.5 FLS3a HI emission. The display illustrates the visualization of the data cube (left) and the profile display of the cube at the cursor location (right); the Tools menu of the Viewer Display Panel has a Spectral Profile button which brings up this display. By default, it grabs the left-mouse button. Pressing down the button and moving in the display will show the profile variations.

8.6 Wiring diagram for the SD task **sdreduce**. The stages of processing within the task are shown, along with the parameters that control them.
8.7 The **Flag plotter**. The **bottom set of buttons** are the standard **matplotlib** toolbar. See the caption of Figure 3.5 for detailed descriptions. The **upper set of buttons in the lower left** are: 1) **region**. Press this to begin marking regions (rather than zooming or panning). 2) **panel**. Press this to begin marking panels to select the whole spectrum. 3, 4, 5, 6) **clear, flag, unflag, statistics**. Click on these to clear, flag, unflag, or calculate statistics of the data within the marked regions and spectra. 7) **notation**. Press this to begin editing notes on the plotter. 8, 9) +, −. Click to move to the next or previous page in a series of iterated plots. The page counter on their left shows the current page number. Finally, the **Quit** is on the bottom right.

8.8 The toolbars on **ASAP plotter**. The **bottom set of buttons** are the standard **matplotlib** toolbar. See the caption of Figure 3.5 for detailed descriptions. The **upper set of buttons** are: 1) **notation**. Press this to begin editing notes on the plotter. 2) **statistics**. Press this to begin printing statistics to the logger. 3, 4) +, −. Click to move to the next or previous page in a series of iterated plots. The page counter on their left shows the current page number. Finally, the **Quit** is on the bottom right.

8.9 The **Notation widget**.

8.10 Total power data display using **sdtpimaging**, with **calmode=`baseline`**. The top panel shows uncalibrated data versus row numbers. The middle panel shows baseline fitting of each scan (only shown here the last scan). The bottom panel shows the calibrated (baseline subtracted) data.
Chapter 1

Introduction

This document describes how to calibrate and image interferometric and single-dish radio astronomical data using the CASA (Common Astronomy Software Application) package. CASA is a suite of astronomical data reduction tools and tasks that can be run via the IPython interface to Python. CASA is being developed in order to fulfill the data post-processing requirements of the ALMA and EVLA projects, but also provides basic and advanced capabilities useful for the analysis of data from other radio, millimeter, and submillimeter telescopes.

You have in your hands the latest release of CASA. This package is under active development, and thus there are a number of caveats and limitations for the use of this package. See the release notes (§ 1.1) below for more information, and pay heed to the numerous ALERTs placed throughout this reference. You can expect regular updates and patches, as well as increasing functionality.

This user reference and cookbook is a task-based walkthrough of interferometric data reduction and analysis. In CASA, tasks represent the more streamlined operations that a typical user would carry out. The idea for having tasks is that they are simple to use, provide a more familiar interface, and are easy to learn for most astronomers who are familiar with radio interferometric data reduction (and hopefully for novice users as well). In CASA, the tools provide the full capability of the package, and are the atomic functions that form the basis of data reduction. These tools augment the tasks, or fill in gaps left by tasks that are under development but not yet available. See the CASA Toolkit Manual for more details on the tools (available from casa.nrao.edu). Note that in most cases, the tasks are Python interface scripts to the tools, but with specific, limited access to them and a standardized interface for parameter setting. The tasks and tools can be used together to carry out more advanced data reduction operations.

For the moment, the audience is assumed to have some basic grasp of the fundamentals of synthesis imaging, so details of how a radio interferometer or telescope works and why the data needs to undergo calibration in order to make synthesis images are left to other documentation — a good

This reference is broken down by the main phases of data analysis:

- data import, export, and selection (Chapter 2),
- examination and flagging of data (Chapter 3),
- interferometric calibration (Chapter 4),
- interferometric imaging (Chapter 5),
- image analysis (Chapter 6), and
- data and image visualization (Chapter 7).
- single dish data analysis (Chapter 8), and
- simulation (Chapter 9).

These are included for users that will be doing EVLA and ALMA telescope commissioning and software development.

The general appendices provide more details on what’s happening under the hood of CASA, as well as supplementary material on tasks, scripts, and relating CASA to other packages. These appendices include:

- obtaining and installing CASA (Appendix A),
- more details about Python and CASA (Appendix B),
- a list of used models, conventions, and reference frames (Appendix C),
- a description of the CASA region format (Appendix D),
- a discussion of the Hamaker-Bregman-Sault Measurement Equation (Appendix E),
- annotated scripts for typical data reduction cases (Appendix F), and
- CASA dictionaries to AIPS, MIRIAD, and CLIC (Appendix H).
- Writing your own CASA Task (Appendix I).

The CASA User Documentation includes:

- CASA User Reference & Cookbook — this document, a task-based data analysis walkthrough and instructions;
- CASA in-line help — accessed using help in the casapy interface;
• The **CASA Toolkit Reference Manual** — details on a specific task or tool does and how to use it.

• The **CASA Task Reference Manual** — the information from the inline help and task documentation, available online in HTML.

The CASA home page can be found at:

[http://casa.nrao.edu](http://casa.nrao.edu)

From there you can find documentation and assistance for the use of the package, including the User Documentation. You will also find information on how to obtain the latest release and receive user support.

There is also a CASAGuides Wiki

[http://casaguides.nrao.edu](http://casaguides.nrao.edu)

that contains helpful information on CASA startup, AIPS-to-CASA cheat sheet, example scripts of processing your data in CASA, along with hints and tricks to best use this package.

### 1.0.1 Reference for Publications

If you use CASA for any of your data reduction or analysis, you may use the following reference:


### 1.1 About This Release

CASA 4.3.0 is now available and the main feature improvements are listed below.

We occasionally issue patches and ‘stable’ versions of CASA. To get notified, please subscribe to the ‘casa-users’ mailing list. ‘Stable’ as well as CASA releases are available at [http://casa.nrao.edu](http://casa.nrao.edu) Releases will be announced via the ‘casa-announce’ mailing list. To subscribe, please visit [http://casa.nrao.edu](http://casa.nrao.edu). For feedback, and help please go to the NRAO helpdesk [http://help.nrao.edu](http://help.nrao.edu) for ALMA questions please use the ALMA helpdesk [http://help.almascience.org](http://help.almascience.org).
Note that in its current incarnation CASA is designed to support Karl G. Jansky VLA, ALMA, and older VLA data, as well as single dish data from ALMA as well as Nobeyama. Data from other telescopes, be it single dish or interferometers can be imported from uvfits, FITS-IDI, or sdfits formats into measurements sets (ms) or scantables in CASA. Given the variety of non-standard fits formats, we cannot guarantee that CASA will fully support data from all telescopes. However, efforts are made to support data formats from other facilities.

### 1.2 Obtaining CASA

CASA is available for the following operating systems:

- Linux
  - RedHat 5.11 and 6.6 (64-bit)
- Mac OS
  - Mac OS 10.8 (Mountain Lion; 64-bit)

The above OSs are the ones that we use to test the CASA package. Other flavors of Linux may work, too. We do not support Mac OS 10.9 or 10.10 at this stage.

The latest and previous releases can be downloaded from our CASA home page: [http://casa.nrao.edu](http://casa.nrao.edu), following the 'Obtaining CASA' link (direct link: [http://casa.nrao.edu/casa_obtaining.shtml](http://casa.nrao.edu/casa_obtaining.shtml)).

#### 1.2.1 What’s New in Release 4.3

Major improvements over the previous version of CASA include:

- **Data examination/editing/import/export**
  - `plotms` multi-panel plots for data and calibration files
  - `plotms` can now plot multiple files, including overplotting in the same panel
  - `plotms` can now write hardcopies of multi-page plots
  - `flagdata`: clipping of values based on WEIGHT and WEIGHT_SPECTRUM
  - `importasdm` with the 'lazy' option will now create an MS which is numerically identical to an MS imported with lazy=False. This means the both the row order and the values of all columns will be the same. It can also be used with all the allowed values of the option `wvr-corrected-data`.
  - formal introduction of UNDEFINED velocity frame
  - WEIGHT_SPECTRUM plotting in `plotms`
• Calibration
  – ability to fill/handle data using ephemeris to follow source as single target field
  – data selection in fluxscale
  – syspower is now associated with scan/field to allow more flexible selection
  – setting of polarization fraction and angle in setjy
  – higher spectral order parameters for component lists
  – more robust spectral index calculation in fluxscale
  – new mode to create a caltable for ionospheric corrections
  – wvrgcal: Several improvements concerning the handling of flagged data and NaN solutions. New parameters "wvrspw" and "spw" to control which WVR data is used and for which spectral windows solutions are written into the caltable.

• Data and Image Manipulation
  – Graphical plotting of data for the feather task to combine low and high resolution images. The new casafeather standalone GUI can plot the data as slices or radially averaged and allows scatter plots of the two images in uv-space to determine the relative scaling.
  – imsmooth can smooth to a common beam if the beam size varies as a function of frequency imsmooth can also use arbitrary image as smoothing kernel, in addition to Gaussian and boxcar kernels
  – Field selection for MODEL images in ft
  – ability to re-shape spws in mstransform
  – per spw rest frequencies in cvel
  – imregrid automatically transposes axes to template images
  – automated sorting of images by their frequency in ia.imageconcat()

• Data analysis
  – rotation measure calculation
  – more flexible specifications for position velocity cuts
  – support for complex-valued images in many image analysis tasks and tools
  – new task imrebin for binning of image pixels
  – new task specsmooth for 1-dimensinal smoothing of images
  – improved error estimates in imfit

• Viewer
  – logarithmic display of data pixel values for range selection
  – interactive display of the position on the position-velocity diagram and the position on the cut
loading of partial cubes and saving of images

• Single Dish
  – consistent flag handling in single dish tasks (see cookbook for details)
  – a new task sdbaseline2 is available for baseline subtraction, creation and application of baseline table.
  – sdimage and sdplot: the telescope position is used in conversions of frequency and direction frame.
  – data selection by raster scan row in sdflag and sdplot
  – data selection by raster map iteration in sdplot
  – for raster-scanned data, rasterutil module is now available to select individual raster rows and raster scans.

• Other
  – the telescope weights are now calculated according to bandwidth and integration time
  – spectral weights in applycal and mstransform
  – weights can be restored to their initial values
  – the async parameter was removed from all tasks
  – bugfix for the internal storage manager that in some cases caused flags to extend to good data

• Experimental tasks
  – cvel2 to provide the functionality of cvel based on the mstransform task
  – split2 to provide the functionality of split based on the mstransform task
  – hanningsmooth2 to provide the functionality of hanningsmooth based on the mstransform task
  – tclean refactored clean task
  – CASA has a new MPI parallelization framework and is currently testing use cases.

For known issues with this release please visit the CASA webpages:
http://casa.nrao.edu/release_ki.shtml

1.3 CASA Basics — Information for First-Time Users

This section assumes that CASA has been installed on your LINUX or OSX system. See Appendix for instructions on how to obtain and install CASA.
1.3.1 Before Starting CASA

First, you will most likely be starting CASA running from a working directory that has your data in it, or at least where you want your output to go. It is easiest to start from there rather than changing directories inside `casapy`. **Alert:** There is at least one task (`plotxy`) that fails if the path to your working directory contains spaces in its name, e.g. `/users/smyers/MyTest/` is fine, but `/users/smyers/My Test/` is not! Please use our new task `plotms` whenever possible and we may work on a better handling of spaces in path names.

If you have done a default installation under Linux using rpms, or on the Mac with the CASA application, then there should be a `sh` script called `casapy` in the `/usr/bin` area which is in your path. This shell will set up its environment and run the version of `casapy` that it points to. If this is how you set up the system, then you need to nothing further and can run `casapy`.

For internal NRAO users we keep different version of CASA, the latest “casapy” release, the “casapy-stable” “Stable” version that is more developed than the Release but without the full documentation (and no GUI testing). We also offer the “Test” version, “casapy-test”, which is produced on a roughly weekly basis with all the latest code but it underwent much less rigorous testing. Instructions how to run the different versions at NRAO can be found on our [http://casa.nrao.edu](http://casa.nrao.edu) webpages under the “CASA at NRAO” link for the different NRAO sites.

1.3.1.1 Environment Variables

Before starting up `casapy`, you should set or reset any `environment variables` needed, as CASA will adopt these on startup. For example, the `PAGER` environment variable determines how help is displayed in the CASA terminal window (see §1.3.8.3). The choices are `less`, `more`, and `cat`.

In `bash`, pick one of

```
PAGER=less
PAGER=more
PAGER=cat
```

followed by

```
export PAGER
```

In `csh` or `tcsh`, pick one of

```
setenv PAGER less
setenv PAGER more
setenv PAGER cat
```

The actions of these are as if you were using the equivalent Unix shell command to view the help material. See §1.3.8.3 for more information on these choices. We recommend using the `cat` option for most users, as this works smoothly both interactively and in scripts.
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Some CASA processes will work on a large number of temporary files. The OS, however, may have a built-in limit on the number of such files. We recommend to increase the limit to $> 1024$. A command like

```
ulimit -n 2048
```

should give CASA enough accessible files to run successfully.

1.3.1.2 Where is CASA?

Note that the path to the CASA installation, which contains the scripts and data repository, will also depend upon the installation. With a default installation under Linux this will probably be in

```
/usr/lib64/casapy/
```

If the unpacked tarball is placed somewhere else, one may add the `PATH` variable to include, e.g.

```
export PATH=$PATH:/<path>/casapy=<version>/bin
```

for bash and

```
set path = ($path /<path>/casapy=<version>/bin)
```

in csh shell.

In a Mac OSX default install it will likely be an application in the Applications folder.

You can find the location after initialized by looking at the `CASAPATH` environment variable. You can find it within `casapy` by

```
import os
print os.environ.get('CASAPATH').split()[0]
```

1.3.2 Starting CASA

After having run the appropriate `casainit` script, CASA is started by typing `casapy`

```
casapy
```
on the UNIX command line, e.g.

```
casapy
```
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After startup information, you should get an IPython
CASA <1>:
command prompt in the xterm window where you started CASA. CASA will take approximately 10
seconds to initialize at startup in a new working directory; subsequent startups are faster. CASA is active when you get a
CASA <1>
prompt in the command line interface. You will also see a logger GUI appear on your Desktop
(usually near the upper left).

You also have the option of starting CASA with various logger options (see §1.5.2.1). For example, if you are running remotely in a terminal window without an X11 connection, or if you just do not want to see the logger GUI, and want the logger messages to come to your terminal, do

    casapy --nologger --log2term

See §1.5.2 for information on the logger in general.

1.3.3 Ending CASA

You can exit CASA by typing quit. This will bring up the query

Do you really want to exit ([y]/n)?

to give you a chance in case you did not mean to exit. You can also quit using %exit or CTRL-D.

If you don’t want to see the question "Do you really want to exit [y]/n?", then just type
Exit or exit and CASA will stop right then and there.

1.3.4 What happens if something goes wrong?

ALERT: Please check the CASA Home Page for Release Notes and FAQ information including
a list of known problems. If you think you have encountered an unknown problem, please consult
the CASA HelpDesk (contact information on the CASA Home Page). See also the caveats to this
Release (§1.1) for pointers to our policy on User Support.

First, always check that your inputs are correct; use the

    help <taskname>

(§1.3.8.2) or

    help par.<parameter name>

(§1.3.8.4) to review the inputs/output.
1.3.5 Aborting CASA execution

If something has gone wrong and you want to stop what is executing, then typing \texttt{CTRL-C} (Control and C keys simultaneously) will usually cleanly abort the application. This will work if you are running a task synchronously. If this does not work on your system then try \texttt{CTRL-Z} to put the task or shell in the background, and then follow up with a \texttt{kill -9 <PID>} where you have found the relevant \texttt{casapy} process ID (PID) using \texttt{ps} (see §1.3.6 below).

If the problem causes CASA to crash, see the next sub-section.

See §1.4.2 for more information on running tasks.

\textbf{Alert:} \texttt{CTRL-C} while a tasks runs can corrupt your input data file, e.g. when a scratch column is filled while aborting. If in doubt, wait until the task has finished, delete the new files produced, and start again.

1.3.6 What happens if CASA crashes?

Usually, restarting \texttt{casapy} is sufficient to get you going again after a crash takes you out of the Python interface. Note that there may be spawned subprocesses still running, such as the \texttt{casaviewer} or the \texttt{logger}. These can be dismissed manually in the usual manner. After a crash, there may also be hidden processes. You can find these by listing processes, e.g. in Linux:

\begin{verbatim}
ps -elf | grep casa
\end{verbatim}

or on MacOSX (or other BSD Unix):

\begin{verbatim}
ps -aux | grep casa
\end{verbatim}

You can then kill these, for example using the Unix \texttt{kill} or \texttt{killall} commands. This may be necessary if you are running remotely using \texttt{ssh}, as you cannot logout until all your background processes are terminated. For example,

\begin{verbatim}
killall ipcontroller
\end{verbatim}

or

\begin{verbatim}
killall Python
\end{verbatim}

will terminate the most common post-crash zombies.
1.3.7 Python Basics for CASA

Within CASA, you use Python to interact with the system. This does not mean an extensive Python course is necessary - basic interaction with the system (assigning parameters, running tasks) is straightforward. At the same time, the full potential of Python is at the more experienced user’s disposal. Some further details about Python, IPython, and the interaction between Python and CASA can be found in Appendix B.

The following are some examples of helpful hints and tricks on making Python work for you in CASA.

1.3.7.1 Variables

Python variables are set using the `<parameter> = <value>` syntax. Python assigns the type dynamically as you set the value, and thus you can easily give it a nonsensical value, e.g.

\[
\begin{align*}
\text{vis} &= \text{\textquote{ngc5921.ms}} \\
\text{vis} &= 1
\end{align*}
\]

The CASA parameter system will check types when you run a task or tool, or more helpfully when you set inputs using `inp` (see below). CASA will check and protect the assignments of the global parameters in its namespace.

Note that Python variable names are case-sensitive:

\[
\begin{align*}
\text{CASA <109>: Foo} &= \text{\textquote{bar}} \\
\text{CASA <110>: foo} &= \text{\textquote{Bar}} \\
\text{CASA <111>: foo} &\quad \Rightarrow \\
\text{Out[111]} &= \text{\textquote{Bar}} \\
\text{CASA <112>: Foo} &\quad \Rightarrow \\
\text{Out[112]} &= \text{\textquote{bar}}
\end{align*}
\]

so be careful.

Also note that misspelling a variable assignment will not be noticed (as long as it is a valid Python variable name) by the interface. For example, if you wish to set `correlation='RR'` but instead type `correlation='RR'` you will find `correlation` unset and a new `correlation` variable set. Command completion (see §1.3.8.1) should help you avoid this.

1.3.7.2 Lists and Ranges

Sometimes, you need to give a task a list of indices. If these are consecutive, you can use the Python `range` function to generate this list:

\[
\begin{align*}
\text{CASA <1>: iflist}\&=\text{range}(4,8) \\
\text{CASA <2>: print iflist}
\end{align*}
\]
[4, 5, 6, 7]
CASA <3>: iflist=range(4)
CASA <4>: print iflist
[0, 1, 2, 3]

See Appendix B.4 for more information.

1.3.7.3 Indexes

As in C, Python indices are 0-based. For example, the first element in a list antlist would be antlist[0]:

CASA <113>: antlist=range(5)
CASA <114>: antlist
Out[114]: [0, 1, 2, 3, 4]
CASA <115>: antlist[0]
Out[115]: 0
Out[116]: 4

CASA also uses 0-based indexing internally for elements in the Measurement Set (MS – the basic construct that contains visibility and/or single dish data; see Chapter 2). Thus, we will often talk about Field or Antenna “ID”s which will be start at 0. For example, the first field in an MS would have FIELD_ID==0 in the MSselect syntax, and can be addressed as be indexed as field='0' in most tasks, as well as by name field='0137+331' (assuming that’s the name of the first field). You will see these indices in the MS summary from the task listobs.

1.3.7.4 Indentation

Python pays attention to the indentation of lines, as it uses indentation to determine the level of nesting in loops. Be careful when cutting and pasting: if you get the wrong indentation, then unpredictable things can happen (usually it just gives an error).

See Appendix B.3 for more information.

1.3.7.5 System shell access

If you want to access system commands from a script, use the os.system command (Appendix B.7.1).

In interactive mode, any input line beginning with a ’!’ character is passed verbatim (minus the ’!’ itself, of course) to the underlying operating system. Also, several common commands (ls, pwd, less) may be executed with or without the ’!’; although the cp command must use ’!’ and cd must be executed without the ’!’). For example:

CASA <5>: !rm -r mydata.ms
Note that if you want to access a Unix environment variable, you will need to prefix with a double $$ instead of a single $ — for example, to print the value of the $$PAGER variable, you would use

   CASA <6>: !echo $$PAGER

See Appendix B.7 for more information.

1.3.7.6 Executing Python scripts

You can execute Python scripts (ASCII text files containing Python or casapy commands) using the `execfile` command. For example, to execute the script contained in the file `myscript.py` (in the current directory), you would type

   CASA <7>: execfile('myscript.py')

or

   CASA <8>: execfile 'myscript.py'

which will invoke the IPython auto-parenthesis feature.

NOTE: in some cases, you can use the IPython `run` command instead, e.g.

   CASA <9>: run myscript.py

In this case, you do not need the quotes around the filename. This is most useful for re-initializing the task parameters, e.g.

   CASA <10>: run clean.last

(see § 1.4.5.8).

See Appendix B.12 for more information.

1.3.8 Getting Help in CASA

1.3.8.1 TAB key

At any time, hitting the `<TAB>` key will complete any available commands or variable names and show you a list of the possible completions if there’s no unambiguous result. It will also complete filenames in the current directory if no CASA or Python names match.

For example, it can be used to list the available functionality using minimum match; once you have typed enough characters to make the command unique, `<TAB>` will complete it.
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1.3.8.2 help <taskname>

Basic information on an application, including the parameters used and their defaults, can be obtained by typing `pdoc task`, `help task`, `help 'task'` or `task?`. The `pdoc` task currently gives the cleanest documentation format with the smallest amount of object-oriented (programmer) output. This inline help provides a one line description of the task and then lists all parameters, a brief description of the parameter, the parameter default, an example setting the parameter and any options if there are limited allowed values for the parameter.

For example:

Import VLA archive file(s) to a measurement set

Imports an arbitrary number of VLA archive-format data sets into a casa measurement set. If more than one band is present, they will be put in the same measurement set but in a separate spectral window. The task will handle old style and new style VLA (after July 2007) archive data and apply the tsys to the data and to the weights.

Keyword arguments:

archivefiles -- Name of input VLA archive file(s)
    default: none. Must be supplied
    example: archivefiles = 'AP314_A959519.xp1'
    example: archivefiles=['AP314_A950519.xp1','AP314_A950519.xp2']

vis -- Name of output visibility file
    default: none. Must be supplied.
    example: vis='NGC7538.ms'
    Will not over-write existing ms of same name.
    A backup flag-file version 'Original' will be made in vis.flagversions. See help flagmanager

bandname -- VLA Frequency band
    default: => '' = all bands
    example: bandname='K'
    Options: '4'=48-96 MHz,'P'=298-345 MHz,'L'=1.15-1.75 GHz,
    'C'=4.2-5.1 GHz,'X'=6.8-9.6 GHz,'U'=13.5-16.3 GHz,
    'K'=20.8-25.8 GHz,'Q'=38-51 GHz

frequencytol -- Tolerance in frequency shift in making spectral windows
    default: => 150000 (Hz). For Doppler shifted data, <10000 Hz may produce too many unnecessary spectral windows.
    example: frequencytol = 150000.0 (units = Hz)
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project -- Project name to import from archive files:
  default: '' => all projects in file
  example: project='AL519'
  project = 'al519' or AL519 will work. Do not include
  leading zeros; project = 'AL0519' will not work.
starttime -- Time after which data will be considered for importing
  default: '' => all: Date must be included.
  syntax: starttime = '2003/1/31/05:05:23'
stoptime -- Time before which data will be considered for importing
  default: '' => all: Date must be included.
  syntax: stoptime = '2003/1/31/08:05:23'
applytsys -- Apply data scaling and weight scaling by nominal
  sensitivity (~Tsys)
  default: True. Strongly recommended
autocorr -- import autocorrelations to ms
  default: => False (no autocorrelations)
antnamescheme -- 'old' or 'new' antenna names.
  default => 'new' gives antenna names
  'VA04' or 'EA13 for VLA telescope 04 and 13 (EVLA)
  'old' gives names '04' or '13'
keepblanks -- Should sources with blank names be filled into the data base
  default => false. Do not fill
  These scans are tipping scans (as of June 1, 2009) and should not
  be filled in the visibility data set.
evlabands -- Use the EVLA's center frequency and bandwidths for frequencies
  specified via wavelength or band.
  default => True.

You can also get the short help for a CASA tool method by typing 'help tool.method'.

CASA <46>: help ia.subimage
Help on built-in function subimage:

subimage(...)  
Create a (sub)image from a region of the image  
--- --- --- --- --- --- Parameters --- --- --- --- --- ---  
outfile: Output image file name. Default is unset.  
region: Region of interest. Default is whole image.  
mask: OTF mask, Boolean LEL expression or mask region. Default is none.  
dropdeg: Drop degenerate axes false  
overwrite: Overwrite (unprompted) pre-existing output file? false  
list: List informative messages to the logger true  
--- --- --- --- --- --- --- --- --- --- --- --- --- --- ---

outfile
region
mask
dropdeg = false
overwrite = false
list = true

----------------------------------------
For a full list of keywords associated with the various tools, see the CASA User Reference Manual.

1.3.8.3 help and PAGER

Your PAGER environment variable (§1.3.1) determines how help is displayed in the terminal window where you start CASA. If you set your bash environment variable PAGER=less (setenv PAGER less in csh) then typing help <taskname> will show you the help but the text will vanish and return you to the command line when you are done viewing it. Setting PAGER=more (setenv PAGER more) will scroll the help onto your command window and then return you to your prompt (but leaving it on display). Setting PAGER=cat (setenv PAGER cat) will give you the more equivalent without some extra formatting baggage and is the recommended choice.

If you have set PAGER=more or PAGER=less, the help display will be fine, but the display of 'taskname?' will often have confusing formatting content at the beginning (lots of ESC surrounding the text). This can be remedied by exiting casapy and doing an ’unset PAGER’ (unsetenv PAGER in [t]csh) at the Unix command line.

You can see the current value of the PAGER environment variable with CASA by typing:

```bash
echo $$PAGER
```

(note the double $$). This will show what command paging is pointed to.

1.3.8.4 help par.<parameter>

Typing help par.<parameter> provides a brief description of a given parameter <parameter>.

```
CASA <46>: help par.robust
Help on function robust in module parameter_dictionary:

robust()
  Brigg's robustness parameter.

  Options: -2.0 (close to uniform) to 2.0 (close to natural)
```

1.3.8.5 Python help

Typing help at the casapy prompt with no arguments will bring up the native Python help facility, and give you the help> prompt for further information; hitting <RETURN> at the help prompt returns you to the CASA prompt.

```
CASA <2>: help
--------> help()
```
Welcome to Python 2.5! This is the online help utility.

If this is your first time using Python, you should definitely check out the tutorial on the Internet at http://www.python.org/doc/tut/.

Enter the name of any module, keyword, or topic to get help on writing Python programs and using Python modules. To quit this help utility and return to the interpreter, just type "quit".

To get a list of available modules, keywords, or topics, type "modules", "keywords", or "topics". Each module also comes with a one-line summary of what it does; to list the modules whose summaries contain a given word such as "spam", type "modules spam".

help> keywords

Here is a list of the Python keywords. Enter any keyword to get more help.

```
and   else    import    raise
assert except in    return
break exec is    try
class finally lambda while
continue for    not    yield
def   from    or
del   global   pass
elif  if      print
```

help>

# hit <RETURN> to return to CASA prompt

You are now leaving help and returning to the Python interpreter. If you want to ask for help on a particular object directly from the interpreter, you can type "help(object)". Executing "help(‘string’)" has the same effect as typing a particular string at the help> prompt.

Further help in working within the Python shell is given in Appendix B.

1.4 Tasks and Tools in CASA

Originally, CASA consisted of a collection of tools, combined in the so-called toolkit. Since the majority of prospective users is far more familiar with the concept of tasks, an effort is underway to replace most - if not all - toolkit functionality by tasks.

While running CASA, you will have access to and be interacting with tasks, either indirectly by providing parameters to a task, or directly by running a task. Each task has a well defined
purpose, and a number of associated parameters, the values of which are to be supplied by the user. Technically speaking, tasks are built on top of tools - when you are running a task, you are running tools in the toolkit, though this should be transparent.

As more tasks are being written, and the functionality of each task is enhanced, there will be less and less reason to run tools in the toolkit. We are working toward a system in which direct access to the underlying toolkit is unnecessary for all standard data processing.

1.4.1 What Tasks are Available?

As mentioned in the introduction, tasks in CASA are python interfaces to the more basic toolkit. Tasks are executed to perform a single job, such as loading, plotting, flagging, calibrating, and imaging the data.

Basic information on tasks, including the parameters used and their defaults, can be obtained by typing `help <taskname>` or `<taskname>?` at the CASA prompt, where `<taskname>` is the name of a given task. As described above in §1.3.8.2, `help <taskname>` provides a description of the task and then lists all parameters, a brief description of the parameter, the parameter default, an example setting the parameter and any options if there are limited allowed values for the parameter.

To see what tasks are available in CASA, use `tasklist`, e.g.

```bash
--------> tasklist()
Available tasks, organized by category (experimental tasks in parenthesis ()
   deprecated tasks in curly brackets {}).
```

<table>
<thead>
<tr>
<th>Import/export</th>
<th>Information</th>
<th>Editing</th>
<th>Manipulation</th>
</tr>
</thead>
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<tr>
<td>exportsasdm</td>
<td>imhead</td>
<td>fixplanets</td>
<td>concat</td>
</tr>
<tr>
<td>exportfits</td>
<td>imreframe</td>
<td>fixvis</td>
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<tr>
<td>exportuvfits</td>
<td>imstat</td>
<td>flagcmd</td>
<td>cvel</td>
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<tr>
<td>importasdm</td>
<td>inval</td>
<td>flagdata</td>
<td>fixvis</td>
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<td>importfits</td>
<td>listcal</td>
<td>flagmanager</td>
<td>hanningsmooth</td>
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<td>importfitsidi</td>
<td>listfits</td>
<td>mview</td>
<td>imhead</td>
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<td>importmiriad</td>
<td>listhistory</td>
<td>plotms</td>
<td>msmoments</td>
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<td>importuvfits</td>
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<td>partition</td>
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<td>importvla</td>
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<td>listvis</td>
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<td>(importgart)</td>
<td>plotms</td>
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<td>testconcat</td>
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<td>virtualconcat</td>
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<td>visstat</td>
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<td>(asdmsummary)</td>
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<td>(cvel2)</td>
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<td>(listsdm)</td>
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<td>(hanningsmooth2)</td>
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<td>(makemask)</td>
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<td>(split2)</td>
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<td>(statwt)</td>
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<td>(uvcontsub3)</td>
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</tbody>
</table>
### Calibration

<table>
<thead>
<tr>
<th>Command</th>
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<tbody>
<tr>
<td>accum</td>
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<tr>
<td>applycal</td>
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<tr>
<td>bandpass</td>
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<td>blcal</td>
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<td>calstat</td>
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<td>clearcal</td>
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<td>delmod</td>
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<td>fixplanets</td>
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<td>fluxscale</td>
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<td>ft</td>
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<td>gaincal</td>
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<td>gencal</td>
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<td>initweights</td>
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<td>listcal</td>
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<td>plotants</td>
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<td>plotbandpass</td>
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<td>plotcal</td>
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<td>polcal</td>
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<td>predictcomp</td>
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<td>setjy</td>
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<td>smoothcal</td>
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<td>uvmodelfit</td>
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<td>uvsub</td>
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<td>wvrgcal</td>
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</tbody>
</table>

### Modeling

<table>
<thead>
<tr>
<th>Command</th>
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<tbody>
<tr>
<td>predictcomp</td>
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<tr>
<td>setjy</td>
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<td>uvcontsub</td>
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<td>uvmodelfit</td>
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<td>uvsub</td>
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</table>

### Imaging

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<tr>
<th>Command</th>
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<tr>
<td>clean</td>
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<td>deconvolve</td>
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<td>feather</td>
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<td>ft</td>
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<td>imcontsub</td>
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<td>(boxit)</td>
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<td>(csvclean)</td>
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<td>(tclean)</td>
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<td>(widebandpbcor)</td>
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<td>{mosaic}</td>
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<td>{widefield}</td>
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</table>

### Analysis

<table>
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<tr>
<th>Command</th>
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<tbody>
<tr>
<td>imcollapse</td>
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<td>imcontsub</td>
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<td>imfit</td>
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<td>imhead</td>
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<td>impbcor</td>
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<td>imrebin</td>
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<td>imregrid</td>
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<td>imsmooth</td>
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<td>imstat</td>
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<td>imsubimage</td>
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<td>imtrans</td>
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<td>inval</td>
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<td>listvis</td>
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<td>rmfit</td>
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<td>slsearch</td>
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<td>specsmooth</td>
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<td>splattotable</td>
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<tr>
<td>(specfit)</td>
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<td>(spxfit)</td>
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</table>

### Visualization

<table>
<thead>
<tr>
<th>Command</th>
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<tbody>
<tr>
<td>clearplot</td>
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<tr>
<td>imview</td>
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<td>msvview</td>
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<td>plotants</td>
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<td>plotbandpass</td>
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<td>plotcal</td>
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<td>plotsms</td>
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<td>plotuv</td>
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<td>viewer</td>
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<td>(plotweather)</td>
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### Simulation

<table>
<thead>
<tr>
<th>Command</th>
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<tbody>
<tr>
<td>simanalyze</td>
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<tr>
<td>simobserve</td>
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<tr>
<td>(simalma)</td>
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</tbody>
</table>

### Single dish

<table>
<thead>
<tr>
<th>Command</th>
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</thead>
<tbody>
<tr>
<td>asap_init</td>
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<tr>
<td>sdaverage</td>
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<tr>
<td>sdbaseline</td>
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<tr>
<td>sdbaseline2</td>
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<tr>
<td>sdbaselineold</td>
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<td>sdcal</td>
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<td>sdcal2</td>
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<td>sdcal2old</td>
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<td>sdcalold</td>
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<td>sdcoadd</td>
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<td>sdfit</td>
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<td>sdfitold</td>
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<td>sdflag</td>
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<td>sdflagold</td>
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<td>sdflagold</td>
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<td>sdgrid</td>
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<td>sdgridold</td>
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<td>sdimageing</td>
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<td>sdimageingold</td>
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<tr>
<td>sdimaging</td>
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<td>sdimagingold</td>
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<td>sdimprocess</td>
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### Utility

<table>
<thead>
<tr>
<th>Command</th>
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<tbody>
<tr>
<td>browsetable</td>
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<tr>
<td>caltabconvert</td>
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<tr>
<td>clearplot</td>
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<tr>
<td>clearstat</td>
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<tr>
<td>concat</td>
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<tr>
<td>conjugatevis</td>
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<tr>
<td>find</td>
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<tr>
<td>help</td>
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<tr>
<td>par.parameter</td>
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<tr>
<td>help</td>
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<tr>
<td>taskname</td>
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<tr>
<td>help</td>
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<tr>
<td>imview</td>
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<tr>
<td>msvview</td>
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<tr>
<td>plotsms</td>
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<tr>
<td>rmtables</td>
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<tr>
<td>startup</td>
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<tr>
<td>taskhelp</td>
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<tr>
<td>tasklist</td>
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<tr>
<td>testconcat</td>
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<tr>
<td>toolhelp</td>
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<tr>
<td>virtualconcat</td>
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</tbody>
</table>
The tasks with name in parentheses are experimental, those in curly brackets are deprecated and will be removed in future releases. The functionality of deprecated tasks is usually available in some other task (e.g., instead of mosaic one should use clean). In the above case, the user has not defined any task him/herself.

Typing taskhelp provides a one line description of all available tasks.

CASA <3>: taskhelp
--------> taskhelp()
Available tasks:

- accum : Accumulate incremental calibration solutions into a calibration table
- applycal : Apply calibrations solutions(s) to data
- asdmsummary : Summarized description of an ASDM dataset.
- autoclean : CLEAN an image with automatically-chosen clean regions.
- bandpass : Calculates a bandpass calibration solution
- bical : Calculate a baseline-based calibration solution (gain or bandpass)
- boxit : Box regions in image above given threshold value.
- browsetable : Browse a table (MS, calibration table, image)
- calstat : Displays statistical information on a calibration table
- caltabconvert : Convert old-style caltables into new-style caltables.

- clean : Invert and deconvolve images with selected algorithm
- clearcal : Re-initializes the calibration for a visibility data set
- clearplot : Clear the matplotlib plotter and all layers
- clearstat : Clear all autolock locks
- concat : Concatenate several visibility data sets.
- conjugatevis : Change the sign of the phases in all visibility columns.
- csvclean : This task does an invert of the visibilities and deconvolve in the image plane.
- cvel : regrid an MS to a new spectral window / channel structure or frame
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- cvel2: Regrid an MS or MMS to a new spectral window, channel structure or frame
- deconvolve: Image based deconvolver
- delmod: Deletes model representations in the MS
- exportasdm: Convert a CASA visibility file (MS) into an ALMA or EVLA Science Data Model
- exportfits: Convert a CASA image to a FITS file
- exportuvfits: Convert a CASA visibility data set to a UVFITS file
- feather: Combine two images using their Fourier transforms
- find: Find string in tasks, task names, parameter names
- fixplanets: Changes FIELD and SOURCE table entries based on user-provided direction or POINTING table
- fixvis: Recalculates (u, v, w) and/or changes Phase Center
- flagcmd: Flagging task based on batches of flag-commands
- flagdata: All-purpose flagging task based on data-selections and flagging modes/algorithms
- flagmanager: Enable list, save, restore, delete and rename flag version files
- fluxscale: Bootstrap the flux density scale from standard calibrators
- ft: Insert a source model a visibility set
- gaincal: Determine temporal gains from calibrator observations
- gencal: Specify Calibration Values of Various Types
- hanningsmooth: Hanning smooth frequency channel data to remove Gibbs ringing
- hanningsmooth2: Hanning smooth frequency channel data to remove Gibbs ringing
- imcollapse: Collapse image along one axis, aggregating pixel values along that axis
- imcontsub: Estimates and subtracts continuum emission from an image cube
- imfit: Fit one or more elliptical Gaussian components on an image region(s)
- imhead: List, get and put image header parameters
- immath: Perform math operations on images
- immoments: Compute moments from an image
- impbcor: Construct a primary beam corrected image from an image and a primary beam pattern
- importasdm: Convert an ALMA Science Data Model observation into a CASA visibility file (MS) or single-dish data format (Scantable)
- importevla: Convert an Science Data Model observation into a CASA Measurement Set
- importfits: Convert an image FITS file into a CASA image
- importfitsidi: Convert a FITS-IDI file to a CASA visibility data set
- importgmr: Convert a UVFITS file to a CASA visibility data set
- importmiriad: Convert a Miriad visibility file into a CASA MeasurementSet
- importuvfits: Convert a UVFITS file to a CASA visibility data set
- importvla: Import VLA archive file(s) to a measurement set
- impv: Construct a position-velocity image by choosing two points in the direction plane
- imrebin: Rebin an image by the specified integer factors
- imreframe: Change the frame in which the image reports its spectral values
- imregrid: Regrid an image onto a template image
- imsmooth: Smooth an image or portion of an image
- imstat: Displays statistical information from an image or image region
- imsubimage: Create a (sub)image from a region of the image
- imtrans: Reorder image axes
- imval: Get the data value(s) and/or mask value in an image
- imview: View an image
- initweights: Initializes weight information in the MS
- listcal: List antenna gain solutions
- listfits: List the HDU and typical data rows of a fits file
- listhistory: List the processing history of a dataset
- listobs: List the summary of a data set in the logger or in a file
- listpartition: List the summary of a multi-MS data set in the logger or in a file
- listsdm: Lists observation information present in an SDM directory.
listvis : List measurement set visibilities.
makebeam : Makes and manipulates image masks
mosaic : Create a multi-field deconvolved image with selected algorithm
mosmoments : Compute moments from an MS
mstransform : Split the MS, combine/separate/regrid spws and do channel and time averaging
msview : View a visibility data set
partition : Task to produce multi-MSs using parallelism
pclean : Invert and deconvolve images with parallel engines
plotants : Plot the antenna distribution in the local reference frame:
plotbandpass : Makes detailed plots of Tsys and bandpass solutions.
plotcal : An all-purpose plotter for calibration results
plotuv : Plots the baseline distribution
plotweather : Plot elements of the weather table; estimate opacity.
polcal : Determine instrumental polarization calibrations
predictcomp : Make a component list for a known calibrator
rmfit : Calculate rotation measure.
rmtables : 
sdaverage : ASAP SD task: averaging and smoothing of spectra
sdbaseline : Fit/subtract a spectral baseline
sdbaseline2 : Fit/subtract a spectral baseline
sdbaselineold : ASAP SD task: fit/subtract a spectral baseline
sdcal : ASAP SD calibration task
sdcal2 : ASAP SD calibration task
sdcal2old : ASAP SD task: generate and/or apply caltables
sdcalold : ASAP SD task: do data selection, calibration, and averaging
sdcoadd : Coadd multiple scantables into one
sdfit : Fit a spectral line
sdfitold : ASAP SD task: fit a spectral line
sdflag : ASAP SD spectral spectral/row flagging task
sdflag2old : ASAP SD spectral flagging task
sdflagmanager : ASAP SD task to manipulate flag version files
sdflagold : ASAP SD spectral flagging task
sdgrid : SD gridding task
sdgridold : SD task: gridding single dish data
sdimaging : SD task: imaging for total power and spectral data
sdimagingold : SD task: imaging for total power and spectral data
sdimprocess : Task for single-dish image processing
sdlist : list summary of single dish data
sdmath : ASAP SD task for simple arithmetic of spectra
sdmathold : ASAP SD task: do simple arithmetic (subtraction, addition, multiplication, and division)
sdplot : ASAP SD plotting task
sdplotold : ASAP SD plotting task
sdreduce : ASAP SD task: do sdcal, sdaverage, and sdbaseline in one task
sdreduceold : ASAP SD task: do sdcalold, sdsmoother, and sdbaselineold in one task
sdsave : Save the sd spectra in various format
sdsaveold : ASAP SD task: save the sd spectra in various format
sdscale : Scale the sd spectra
sdsmoother : ASAP SD task: do smoothing of spectra
sdstat : list statistics of spectral
sdstatold : ASAP SD task: list statistics of spectral region
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sdtpimaging : SD task: do a simple calibration (baseline subtraction) and imaging for total power data
setjy : Fills the model column with the visibilities of a calibrator
simalma : Simulation task for ALMA
simanalyzer : image and analyze measurement sets created with simobserve
simobserve : visibility simulation task
slinks : Search a spectral line table.
smoothcal : Smooth calibration solution(s) derived from one or more sources:
specfit : Fit 1-dimensional gaussians and/or polynomial models to an image or image region
specsMOOTH : Smooth an image region in one dimension
sphtotable : Convert a downloaded Splatalogue spectral line list to a casa table.
split : Create a visibility subset from an existing visibility set
split2 : Create a visibility subset from an existing visibility set
spxfit : Fit a 1-dimensional model(s) to an image(s) or region for determination of spectral index
ssoflux : Fills the model column with the visibilities of a calibrator
statwt : Reweight visibilities according to their scatter (Experimental)
tclean : Construct and deconvolve images
testconcat : Concatenate the subtables of several visibility data sets, not the MAIN bulk data.
uvcontsub : Continuum fitting and subtraction in the uv plane
uvcontsub3 : An experimental clone of uvcontsub
uvmodelfit : Fit a single component source model to the uv data
uvsub : Subtract/add model from/to the corrected visibility data.
viewer : View an image or visibility data set
virtualconcat : Concatenate several visibility data sets into a multi-MS
vishead : List, summary, get, and put metadata in a measurement set
visstat : Displays statistical information from a Measurement Set, or from a Multi-MS
widebandpbcor : Wideband PB-correction on the output of the MS-MFS algorithm
widefield : Wide-field imaging and deconvolution with selected algorithm
wvrgcal : Generate a gain table based on Water Vapour Radiometer data

Typing startup will provide the startup page displayed when entering CASA. The startup screen lists the various options to obtain help within CASA.

CASA <26>: startup
---------- startup()

For help use the following commands:
tasklist - Task list organized by category
taskhelp - One line summary of available tasks
help taskname - Full help for task
toolhelp - One line summary of available tools
help par.parametername - Full help for parameter name

1.4.2 Running Tasks and Tools

Tools are functions linked to the Python interface which must be called by name with arguments. Tasks have higher-level capabilities than tools. Tasks require input parameters which maybe be
specified when you call the task as a function, or be set as parameters in the interface. A task, like
a tool, is a function under Python and may be written in Python, C, or C++ (the CASA toolkit
is made up of C++ functions).

There are two distinct ways to run tasks. You can either set the global CASA parameters relevant
to the task and tell the task to “go”, or you can call the task as a function with one or more
arguments specified. These two invocation methods differ in whether the global parameter values
are used or not.

For example,

```
default('plotxy')
vis='ngc5921.ms'
xaxis='channel'
yaxis='amp'
datacolumn='data'
go
```

will execute `plotxy` with the set values for the parameters (see §1.4.5). Instead of using `go`
command (§1.4.5.3) to invoke the task, you can also call the task with no arguments, e.g.

```
default('plotxy')
vis='ngc5921.ms'
xaxis='channel'
yaxis='amp'
datacolumn='data'
plotxy()
```

which will also use the global parameter values.

Second, one may call tasks and tools by name with parameters set on the same line. Parameters
may be set either as explicit `<parameter>=<value>` arguments, or as a series of comma delimited
`<value>`s in the correct order for that task or tool. Note that missing parameters will use the
default values for that task. For example, the following are equivalent:

```
# Specify parameter names for each keyword input:
plotxy(vis='ngc5921.ms',xaxis='channel',yaxis='amp',datacolumn='data')
# when specifying the parameter name, order doesn't matter, e.g.:
plotxy(xaxis='channel',vis='ngc5921.ms',datacolumn='data',yaxis='amp')
# use parameter order for invoking tasks
plotxy('ngc5921.ms','channel','amp','data')
```

This non-use of globals when calling as a function is so that robust scripts can be written. One
need only cut-and-paste the calls and need not worry about the state of the global variables or
what has been run previously. It is also more like the standard behavior of function calls in Python
and other languages.

Tools can only be called in this second manner by name, with arguments (§1.4.6). Tools never use
the global parameters and the related mechanisms of `inp` and `go`.
1.4.2.1 Aborting Synchronous Tasks

If you are running CASA tasks synchronously, then you can usually use \texttt{CTRL-C} to abort execution of the task. If this does not work, try \texttt{CTRL-Z} followed by a \texttt{kill}. See §1.3.5 for more on these methods to abort CASA execution.

You may have to quit and restart CASA after an abort, as the internal state can get mixed up.

1.4.3 Getting Return Values

Some tasks and tools return a record (usually a Python dictionary) to the interface. For example, the \texttt{imstat} task (§6.9) returns a dictionary with the image statistics in it. To catch these return values into a Python variable, you MUST assign that variable to the task call, e.g.

\begin{verbatim}
xstat = imstat('ngc5921.clean.image')
\end{verbatim}

or

\begin{verbatim}
default('imstat')
imagename = 'ngc5921.clean.image'
xstat = imstat()
\end{verbatim}

Note that tools that return values work in the same way (§1.4.6).

You can print or use the return value in Python for controlling scripts. For example,

\begin{verbatim}
CASA <1>: xstat = imstat('ngc5921.clean.image')
CASA <2>: xstat
Out[2]:
{'blc': array([0, 0, 0, 0]),
 'blcf': '15:24:08.404, +04.31.59.181, I, 1.41281e+09Hz',
 'flux': array([ 4.15292207]),
 'max': array([ 0.05240594]),
 'maxpos': array([134, 134,  0, 38]),
 'maxposf': '15:21:53.976, +05.05.29.998, I, 1.41374e+09Hz',
 'mean': array([ 1.62978083e-05]),
 'medabsdevmed': array([ 0.00127287]),
 'median': array([ -1.10467618e-05]),
 'min': array([-0.0105249]),
 'minpos': array([160,  1,  0, 30]),
 'minposf': '15:21:27.899, +04.32.14.923, I, 1.41354e+09Hz',
 'npts': array([ 3014656.]),
 'quartile': array([ 0.00254587]),
 'rms': array([ 0.00201818]),
 'sigma': array([ 0.00201811]),
 'sum': array([ 49.1322855]),
 'sumsq': array([ 12.27880404]),
 'trc': array([255, 255,  0, 45]),
\end{verbatim}
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If you do not catch the return variable, it will be lost

```
imstat('ngc5921.clean.image')
```

or

```
default('imstat')
imagename = 'ngc5921.clean.image'
imstat()
```

and spewed to terminal. Note that go will trap and lose the return value, e.g.

```
default('imstat')
imagename = 'ngc5921.clean.image'
```

will not dump the return to the terminal either.

1.4.4 Running Tasks Asynchronously

By default, most tasks run synchronously in the foreground. Many tasks, particularly those that can take a long time to execute, have the `async` parameter. This allows the user to send the task to the background for execution.

1.4.5 Setting Parameters and Invoking Tasks

One can set parameters for tasks (but not for tools) by performing the assignment within the CASA shell and then inspecting them using the `inp` command:

```
CASA <30>: default(bandpass)
CASA <31>: vis = 'ngc5921.demo.ms'
CASA <32>: caltable = 'ngc5921.demo.bcal'
CASA <33>: field = '0'
CASA <34>: refant = '15'
CASA <35>: inp('bandpass')
# bandpass :: Calculates a bandpass calibration solution
vis = 'ngc5921.demo.ms' # Name of input visibility file
caltable = 'ngc5921.demo.bcal' # Name of output gain calibration table
field = '0' # Select field using field id(s) or field name(s)
spw = '' # Select spectral window/channels
```
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intent = '', # Select observing intent
selectdata = True # Other data selection parameters
timerange = '', # Select data based on time range
uvrange = '', # Select data within uvrange (default units meters)
antenna = '', # Select data based on antenna/baseline
scan = '', # Scan number range
observation = '', # Select by observation ID(s)
mselect = '', # Optional complex data selection (ignore for now)
solint = 'inf' # Solution interval in time[,freq]
combine = 'scan' # Data axes which to combine for solve (obs, scan, spw, and/or field)
refant = '15' # Reference antenna name(s)
minblperant = 4 # Minimum baselines _per antenna_ required for solve
minsnr = 3.0 # Reject solutions below this SNR (only applies for bandtype = B)
solnorm = False # Normalize average solution amplitudes to 1.0
bandtype = 'B' # Type of bandpass solution (B or BPOLY)
fillgaps = 0 # Fill flagged solution channels by interpolation
smo = [] # Point source Stokes parameters for source model.
append = False # Append solutions to the (existing) table
docallib = False # Callib or traditional cal apply parameters
gaintable = [] # Use callib or traditional cal apply parameters
gainfield = [] # Gain calibration table(s) to apply on the fly
interp = [] # Interpolation mode (in time) to use for each gaintable
spwmap = [] # Spectral windows combinations to form for gaintables(s)
parang = False # Apply parallactic angle correction

See §1.4.5.4 below for more details on the use of the inputs command.

All task parameters have global scope within CASA: the parameter values are common to all tasks and also at the CASA command line. This allows the convenience of not changing parameters that are shared between tasks but does require care when chaining together sequences of task invocations (to ensure proper values are provided).

If you want to reset the input keywords for a single task, use the default command (§1.4.5.2).

For example, to set the defaults for the bandpass task, type:

CASA <30>: default('bandpass')

as we did above.

To inspect a single parameter value just type it at the command line. Continuing the above example:

CASA <36>: combine
Out[14]: 'scan'

CASA parameters are just Python variables.

Parameters for a given task can be saved by using the saveinputs command (see §1.4.5.5) and restored using the execfile '<filename>' command. Note that if the task is successfully executed, then a <taskname>.last file is created in the working directory containing the parameter values (see §1.4.5.8).
We now describe the individual CASA task parameter interface commands and features in more detail.

1.4.5.1 The scope of parameters in CASA

All task parameters have global scope within CASA: the parameter values are common to all tasks and also at the CASA command line. This allows the convenience of not changing parameters that are shared between tasks but does require care when chaining together sequences of task invocations (to ensure proper values are provided). Tasks DO NOT change the values of the global parameters, nor does the invocation of tasks using the functional call with arguments change the globals.

This does mean that unless you do an explicit default of the task (§1.4.5.2), previously set values may be unexpectedly used if you do not inspect the inp carefully. For example, good practice is:

```python
default('imhead')
imagename = 'ngc5921.demo.cleaning.image'
mode = 'list'
imhead()
```

If you supply the task call with arguments, then these will be used for the values of those parameters (see above). However, if some but not all arguments are supplied, then those parameters not given as arguments will default and NOT use the current global values. Thus,

```python
imhead('ngc5921.demo.cleaning.image',mode='list')
```

will reproduce the above.

1.4.5.2 The default Command

Each task has a special set of default parameters defined for its parameters. You can use the default command to reset the parameters for a specified task (or the current task as defined by the taskname variable) to their default.

**Important Note:** The default command resets the values of the task parameters to a set of “defaults” as specified in the task code. Some defaults are blank strings ‘’ or empty lists [], others are specific numerical values, strings, or lists. It is important to understand that just setting a string parameter to an empty string ‘’ is not setting it to its default! Some parameters do not have a blank as an allowed value. See the help for a particular task to find out its default. If ‘’ is the default or an allowed value, it will say so explicitly.

For example, suppose we have been running CASA on a particular dataset, e.g.
and now we wish to switch to a different one. We can reset the parameter values using default:

```
CASA <41>: default
----------> default()
```

```
CASA <42>: inp
----------> inp()
```

It is good practice to use default before running a task if you are unsure what state the CASA global variables are in.

**ALERT:** You currently can only reset ALL of the parameters for a given task to their defaults. In an upcoming update we will allow the default command to take a second argument with a specific parameter to default its value.

### 1.4.5.3 The go Command

You can execute a task using the go command, either explicitly

```
CASA <44>: go listobs
----------> go(listobs)
Executing: listobs()
...
or implicitly if `taskname` is defined (e.g. by previous use of `default` or `inp`)

CASA <45>: `taskname` = 'clean'
CASA <46>: `go`
--------------- `go()`
Executing: `clean()`
...

You can also execute a task simply by typing the `taskname`.

CASA <46>: `clean`
--------------- `clean()`
Executing: `clean()`
...

The `go` command can also be used to launch a different task without changing the current `taskname`, without disrupting the `inp` process on the current task you are working on. For example

```
default 'gaincal' # set current task to gaincal and default
vis = 'n5921.ms' # set the working ms
... # set some more parameters
go listobs # launch listobs w/o changing current task
inp # see the inputs for gaincal (not listobs!)
```

**ALERT:** Doing `go listobs(vis='foo.ms')` will currently change the `taskname`, and will change `vis`, which might not be what is desired.

### 1.4.5.4 The `inp` Command

You can set the values for the parameters for tasks (but currently not for tools) by performing the assignment within the CASA shell and then inspecting them using the `inp` command. This command can be invoked in any of three ways: via function call `inp('<taskname>')` or `inp(<taskname>)`, without parentheses `inp 'taskname'` or `inp <taskname>`, or using the current `taskname` variable setting with `inp`. For example,

CASA <1>: `inp('clean')`
...
CASA <2>: `inp 'clean`
---------- `inp('clean')`
...
CASA <3>: `inp(clean)
...
CASA <4>: `inp clean`
---------- `inp(clean)
...
CASA <5>: `taskname` = 'clean'
CASA <6>: `inp`
---------- `inp()`
all do the same thing.

When you invoke the task inputs via `inp`, you see a list of the parameters, their current values, and a short description of what that parameters does. For example, starting from the default values,

```
CASA <18>: inp('clean')
# clean :: Deconvolve an image with selected algorithm
vis = '' # name of input visibility file
imagename = '' # Pre-name of output images
field = '' # Field Name
spw = '' # Spectral windows:channels: '' is all
selectdata = False # Other data selection parameters
mode = 'mfs' # Type of selection (mfs, channel, velocity, frequency)
niter = 500 # Maximum number of iterations
gain = 0.1 # Loop gain for cleaning
threshold = '0.0mJy' # Flux level to stop cleaning. Must include units
psfmode = 'clark' # method of PSF calculation to use during minor cycles
imagermode = '' # Use csclean or mosaic. If '', use psfmode
multiscale = [] # multi-scale deconvolution scales (pixels)
interactive = False # use interactive clean (with GUI viewer)
mask = [] # cleanbox(es), mask image(s), and/or region(s)
imsize = [256, 256] # x and y image size in pixels
cell = ['1.0arcsec', '1.0arcsec'] # x and y cell size. default unit arcsec
phasedcenter = '' # Image phase center: position or field index
restfreq = '' # rest frequency to assign to image (see help)
stokes = 'I' # Stokes params to image (eg I,IV, QU, IQUV)
weighting = 'natural' # Weighting of uv (natural, uniform, briggs, ...)
uvtaper = False # Apply additional uv tapering of visibilities.
modelimage = '' # Name of model image(s) to initialize cleaning
restoringbeam = [''] # Output Gaussian restoring beam for CLEAN image
pbcor = False # Output primary beam-corrected image
minpb = 0.1 # Minimum PB level to use
```

Figure 1.1 shows how this will look to you on your terminal. Note that some parameters are in boldface with a gray background. This means that some values for this parameter will cause it to expand, revealing new sub-parameters to be set.

CASA uses color and font to indicate different properties of parameters and their values:

```
<table>
<thead>
<tr>
<th>Text Font</th>
<th>Text Color</th>
<th>Highlight</th>
<th>Indentation</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameters:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>plain</td>
<td>black</td>
<td>none</td>
<td>none</td>
<td>standard parameter</td>
</tr>
<tr>
<td>bold</td>
<td>black</td>
<td>grey</td>
<td>none</td>
<td>expandable parameter</td>
</tr>
<tr>
<td>plain</td>
<td>green</td>
<td>none</td>
<td>yes</td>
<td>sub-parameter</td>
</tr>
<tr>
<td>Values:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>plain</td>
<td>black</td>
<td>none</td>
<td>none</td>
<td>default value</td>
</tr>
<tr>
<td>plain</td>
<td>blue</td>
<td>none</td>
<td>none</td>
<td>non-default value</td>
</tr>
<tr>
<td>plain</td>
<td>red</td>
<td>none</td>
<td>none</td>
<td>invalid value</td>
</tr>
</tbody>
</table>
```
Figure 1.1: Screen shot of the default CASA inputs for task clean.

Figure 1.2 shows what happens when you set some of the clean parameters to non-default values. Some have opened up sub-parameters, which can now be seen and set. Figure 1.3 shows what happens when you set a parameter, in this case vis and mode, to an invalid value. Its value now appears in red. Reasons for invalidation include incorrect type, an invalid menu choice, or a filename that does not exist. For example, since vis expects a filename, it will be invalidated (red) if it is set to a non-string value, or a string that is not the name of a file that can be found. The mode='happy' is invalid because it’s not a supported choice (‘mfs’, ‘channel’, ‘velocity’, or ‘frequency’).

1.4.5.5 The saveinputs Command

The saveinputs command will save the current values of a given task parameters to a Python (plain ascii) file. It can take up to two arguments, e.g.

```python
def saveinputs(taskname, outfile)
```

The first is the usual taskname parameter. The second is the name for the output Python file. If there is no second argument, for example,

```python
saveinputs('clean')
```

a file with name <taskname>.saved (in this case ‘clean.saved’ will be created or overwritten if extant. If invoked with no arguments, e.g.

```python
saveinputs
```
it will use the current values of the `taskname` variable (as set using `inp <taskname>` or `default <taskname>`). You can also use the `taskname` global parameter explicitly,

```
saveinputs(taskname, taskname+'_1.save')
```

For example, starting from default values

```
CASA <1>: default('listobs')
CASA <2>: vis='ngc5921.demo.ms'
CASA <3>: saveinputs
CASA <4>: !more 'listobs.saved'
```

```
taskname = "listobs"
vis = "ngc5921.demo.ms"
selectdata = True
spw = 
field = 
antenna = 
array = 
```
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Figure 1.3: The clean inputs where one parameter has been set to an invalid value. This is drawn in red to draw attention to the problem. This hapless user probably confused the 'hogbom' clean algorithm with Harry Potter.

observation = ""
verbose = True
listfile = ""

#listobs(vis="ngc5921.demo.ms",selectdata=True,spw="",field="",
antenna="",uvrange="",timerange="",correlation="",scan="",intent="",
feed="",array="",observation="",verbose=True,listfile="")

To read these back in, use the Python execfile command. For example,

CASA <5>: execfile('listobs.saved')

and we are back.

An example save to a custom named file:

CASA <6>: saveinputs('listobs','ngc5921_listobs.par')

You can also use the CASA tget command (see § 1.4.5.6 below) instead of the Python execfile to restore your inputs.
1.4.5.6 The tget Command

The `tget` command will recover saved values of the inputs of tasks. This is a convenient alternative to using the Python `execfile` command (see above).

Typing `tget` without a taskname will recover the saved values of the inputs for the current task as given in the current value of the `taskname` parameter.

Adding a task name, e.g. `tget <taskname>` will recover values for the specified task. This is done by searching for 1) a `<taskname>.last` file (see §1.4.5.8 below), then for 2) a `<taskname>.saved` file (see §1.4.5.5 above), and then executing the Python in these files.

For example,

```plaintext
default('gaincal') # set current task to gaincal and default
tget # read saved inputs from gaincal.last (or gaincal.saved)
inp # see these inputs!
tget bandpass # now get from bandpass.last (or bandpass.saved)
inp # task is now bandpass, with recovered inputs
```

1.4.5.7 The tput Command

The `tput` command will save the current parameter values of a task to its `<taskname>.last` file. This is a shorthand to `saveinputs` and is a counterpart to `tget`.

Typing `tput` without a taskname will save the values of the inputs for the current task as given in the current value of the `taskname` parameter.

Adding a task name, e.g. `tput <taskname>` will save the values for the specified task.

For example,

```plaintext
default('gaincal') # set current task to gaincal and default
tget # read saved inputs from gaincal.last (or gaincal.saved)
inp # see these inputs!
vis = 'new.ms' # change the vis parameter
tput # save back to the gaincal.last file for later use
```

1.4.5.8 The .last file

Whenever you successfully execute a CASA task, a Python script file called `<taskname>.last` will be written (or over-written) into the current working directory. For example, if you ran the `listobs` task as detailed above, then

```plaintext
CASA <14>: vis = 'ngc5921.ms'
CASA <15>: verbose = True
CASA <16>: listobs()
```
CASA <17>: `!more 'listobs.last'`

IPython system call: more listobs.last

taskname = "listobs"
vis = "ngc5921.ms"
verbose = True
listfile = ""
#listobs(vis="ngc5921.ms",verbose=False,listfile="")

You can restore the parameter values from the save file using

CASA <18>: execfile('listobs.last')

or

CASA <19>: run listobs.last

Note that the .last file is generally not created until the task actually finished (successfully), so it is often best to manually create a save file beforehand using the `saveinputs` command if you are running a critical task that you strongly desire to have the inputs saved for.

### 1.4.6 Tools in CASA

The CASA **toolkit** is the foundation of the functionality in the package, and consists of a suite of functions that are callable from Python. The tools are used by the tasks, and can be used by advanced users to perform operations that are not available through the tasks.

It is beyond the scope of this reference to describe the toolkit in detail. Occasionally, examples will be given that utilize the tools (e.g. §[6.25](#)). In short, tools are always called as functions, with any parameters that are not to be defaulted given as arguments. For example:

```python
ia.open('ngc5921.chan21.clean.cleanbox.mask')
ia.calcmask('ngc5921.chan21.clean.cleanbox.mask">0.5', 'mymask')
ia.summary()
ia.close()
```

uses the **image** tool (ia) to turn a clean mask image into an image mask. Tools never use the CASA global parameters.

To find what tools are available, use the `toolhelp` command:

CASA <4>: toolhelp
--------- toolhelp()

**Available tools:**

`af`: Agent flagger utilities
at : Juan Pardo ATM library
cn : Calibration analysis utilities
cb : Calibration utilities
cl : Component list utilities
cp : Cal solution plotting utilities
cs : Coordinate system utilities
cu : Class utilities
dc : Deconvolver utilities
fi : Fitting utilities
fn : Functional utilities
ia : Image analysis utilities
im : Imaging utilities
me : Measures utilities
ms : MeasurementSet (MS) utilities
msmd : MS metadata accessors
mt : MS transformer utilities
qa : Quanta utilities
pm : PlotMS utilities
po : Imagepol utilities
rg : Region manipulation utilities
sl : Spectral line import and search
sm : Simulation utilities
tb : Table utilities (selection, extraction, etc)
tr : Table plotting utilities
vp : Voltage pattern/primary beam utilities
---
pl : pylab functions (e.g., pl.title, etc)
sd : Single dish utilities
---

You can find much more information about the toolkit in the CASA User Reference Manual:

http://casa.nrao.edu/docs/casaref/CasaRef.html

1.5 Getting the most out of CASA

There are some other general things you should know about using CASA in order to make things go smoothly during your data reduction.

1.5.1 Your command line history

Your command line history is automatically maintained and stored as ipython.log in your local directory. This file can be edited and re-executed as appropriate using the execfile '<filename>' feature.

You can also use the “up-arrow” and “down-arrow” keys for command line recall in the casapy interface. If you start typing text, and then use “up-arrow”, you will navigate back through commands matching what you typed.
1.5.2 Logging your session

The output from CASA commands is sent to the file `casapy-YYYYMMDD-HHMMSS.log` in your local directory, where `YYYYMMDD-HHMMSS` are the UT date and time when CASA was started up. New starts of CASA create new log files.

![The CASA Logger GUI window under Linux. Note that under MacOSX a stripped down logger will instead appear as a Console.](image)

The output contained in `casapy-YYYYMMDD-HHMMSS.log` is also displayed in a separate window using the `casalogger`. Generally, the logger window will be brought up when `casapy` is started. If you do not want the logger GUI to appear, then start `casapy` using the `--nologger` option,

```
casapy --nologger
```

which will run CASA in the terminal window. See § 1.5.2.1 for more startup options.

**ALERT:** Due to problems with Qt under MacOSX, we had to replace the GUI `qtcasalogger` with a special stripped down one that uses the Mac Console. This still has the important capabilities such as showing the messages and cut/paste. The following description is for the Linux version and thus should mostly be disregarded on OSX. On the Mac, you treat this as just another console window and use the usual mouse and hot-key actions to do what is needed.

The CASA logger window for Linux is shown in Figure 1.4. The main feature is the display area for the log text, which is divided into columns. The columns are:

- **Time** — the time that the message was generated. Note that this will be in local computer time (usually UT) for `casapy` generated messages, and may be different for user generated messages;
- **Priority** — the *Priority Level* (see below) of the message;
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Figure 1.5: Using the Search facility in the casalogger. Here we have specified the string ‘apply’ and it has highlighted all instances in green.

- **Origin** — where within CASA the message came from. This is in the format Task:Tool:Method (one or more of the fields may be missing depending upon the message);

- **Message** — the actual text.

The casalogger GUI has a range of features, which include:

- **Search** — search messages by entering text in the Search window and clicking the search icon. The search currently just matches the exact text you type anywhere in the message. See Figure 1.5 for an example.

- **Filter** — a filter to sort by message priority, time, task/tool of origin, and message contents. Enter text in the Filter window and click the filter icon to the right of the window. Use the pull-down at the left of the Filter window to choose what to filter. The matching is for the exact text currently (no regular expressions). See Figure 1.6 for an example.

- **View** — show and hide columns (Time, Priority, Origin, Message) by checking boxes under the View menu pull-down. You can also change the font here.

- **Insert Message** — insert additional comments as “notes” in the log. Enter the text into the “Insert Message” box at the bottom of the logger, and click on the Add (+) button, or choose to enter a longer message. The entered message will appear with a priority of “NOTE” with the Origin as your username. See Figure 1.7 for an example. **ALERT:** This message currently will not be inserted into the correct (or user controllable) order into the log.

- **Copy** — left-click on a row, or click-drag a range of rows, or click at the start and shift click at the end to select. Use the Copy button or Edit menu Copy to put the selected rows into the clipboard. You can then (usually) paste this where you wish. **ALERT:** this does not work
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Figure 1.6: Using the casalogger Filter facility. The log output can be sorted by Priority, Time, Origin, and Message. In this example we are filtering by Origin using 'clean', and it now shows all the log output from the clean task.

routinely in the current version. You are best off going to the casapy-YYYYMMDD-HHMMSS.log file if you want to grab text.

- **Open** — There is an Open function in the File menu, and an Open button, that will allow you to load old casalogger files.

Other operations are also possible from the menu or buttons. Mouse “flyover” will reveal the operation of buttons, for example.

It is possible to change the file that the logging is directed to. Per default it is 'casapy-YYYYMMDD-HHMMSS.log'. But starting CASA with the option `--logfile`

casapy --logfile otherfile.log

will redirect the output of the logger to the file 'otherfile.log' (see also Sect. 1.5.2.1). The log file can also be changed during a CASA session. Type

CASA <15>: casalog.setlogfile('otherfile.log')

and you will redirect the output to the 'otherfile.log' file. However, the logger GUI will still be monitoring the previous 'casapy-YYYYMMDD-HHMMSS.log' file. To change it to the new file, go on File - Open and select the new log file, in our case 'otherfile.log'.

1.5.2.1 Startup options for the logger

One can specify logger options at the startup of casapy on the command line:
Figure 1.7: CASA Logger - Insert facility: The log output can be augmented by adding notes or comments during the reduction. The file should then be saved to disk to retain these changes.

```
casapy <logger option>
```

These options are:

- `--log2term`   == logging message go to terminal
- `--nologfile`  == no casapy.log logfile is produced
- `--logfile <filename>` == use specified name for logfile instead of casapy.log
- `--nologger`   == do not bring up GUI logger (see above)
- `--nolog` (is deprecated use --nologger)

For example, to not bring up a GUI but send the message to your terminal, do

```
casapy --nologger --log2term
```

while

```
casapy --logfile mynewlogfile.log
```

will start casapy with logger messages going to the file `mynewlogfile.log`.

### 1.5.2.2 Setting priority levels in the logger

Logger messages are assigned a `Priority Level` when generated within CASA. The current levels of Priority are:

1. **SEVERE** — errors;
2. **WARN** — warnings;
3. **INFO** — basic information every user should be aware of or has requested;
4. **INFO1** — information possibly helpful to the user;
5. **INFO2** — details the power user might want to see;
6. **INFO3** — even more details;
7. **INFO4** — lowest level of non-debugging information;
8. **DEBUGGING** — most “important” debugging messages;
9. **DEBUG1** — more details;
10. **DEBUG2** — lowest level of debugging messages.

The “debugging” levels are intended for the developers use.

There is a threshold for which these messages are written to the `casapy-YYYYMMDD-HHMMSS.log` file and are thus visible in the logger. By default, only messages at level **INFO** and above are logged. The user can change the threshold using the `casalog.filter` method. This takes a single string argument of the **level** for the threshold. The **level** sets the lowest priority that will be generated, and all messages of this level or higher will go into the `casapy-YYYYMMDD-HHMMSS.log` file.

Some examples:

```
  casalog.filter('INFO')  # the default
  casalog.filter('INFO2') # should satisfy even advanced users
  casalog.filter('INFO4') # all INFOx messages
  casalog.filter('DEBUG2') # all messages including debugging
```

**WARNING:** Setting the threshold to **DEBUG2** will put lots of messages in the log!

### 1.5.3 Where are my data in CASA?

Interferometric data are filled into a so-called Measurement Set (or MS). In its logical structure, the MS looks like a generalized description of data from any interferometric or single dish telescope. Physically, the MS consists of several tables in a directory on disk.

Tables in CASA are actually directories containing files that are the sub-tables. For example, when you create a MS called `AM675.ms`, then the name of the directory where all the tables are stored will be called `AM675.ms/`. See Chapter 2 for more information on Measurement Set and Data Handling in CASA.
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The data that you originally get from a telescope can be put in any directory that is convenient to you. Once you "fill" the data into a measurement set that can be accessed by CASA, it is generally best to keep that MS in the same directory where you started CASA so you can get access to it easily (rather than constantly having to specify a full path name).

When you generate calibration solutions or images (again these are in table format), these will also be written to disk. It is a good idea to keep them in the directory in which you started CASA.

1.5.3.1 How do I get rid of my data in CASA?

Note that when you delete a measurement set, calibration table, or image, which are in fact directories, you must delete this and all underlying directories and files. If you are not running casapy, this is most simply done by using the file delete method of the operating system you started CASA from. For example, when running CASA on a Linux system, in order to delete the measurement set named AM675.ms type:

   CASA <5>: !rm -r AM675.ms

from within CASA. The ! tells CASA that a system command follows (see §1.3.7.5), and the -r makes sure that all subdirectories are deleted recursively.

It is convenient to prefix all MS, calibration tables, and output files produced in a run with a common string. For example, one might prefix all files from VLA project AM675 with AM675, e.g. AM675.ms, AM675.cal, AM675.clean. Then,

   CASA <6>: !rm -r AM675*

will clean up all of these.

In scripts, the ! escape to the OS will not work. Instead, use the os.system() function (Appendix B.7.1) to do the same thing:

   os.system('rm -r AM675*')

If you are within casapy, then the CASA system is keeping a cache of tables that you have been using and using the OS to delete them will confuse things. For example, running a script that contains rm commands multiple times will often not run or crash the second time as the cache gets confused. The clean way of removing CASA tables (MS, caltables, images) inside casapy is to use the rmtables task:

   rmtables('AM675.ms')

and this can also be wildcarded

   rmtables('AM675*')
(though you may get warnings if it tries to delete files or directories that fit the name wildcard that are not CASA tables).

**ALERT:** Some CASA processes lock the file and forget to give it up when they are done (plotxy is usually the culprit). You will get WARNING messages from rmtables and your script will probably crash second time around as the file isn’t removed. The safest thing is still to exit casapy and start a new session for multiple runs.

### 1.5.4 What’s in my data?

The actual data is in a large **MAIN** table that is organized in such a way that you can access different parts of the data easily. This table contains a number of “rows”, which are effectively a single timestamp for a single spectral window (like an IF from the VLA) and a single baseline (for an interferometer).

There are a number of “columns” in the MS, the most important of which for our purposes is the **DATA** column — this contains the original visibility data from when the MS was created or filled. There are other helpful “scratch” columns which hold useful versions of the data or weights for further processing: the **CORRECTED_DATA** column, which is used to hold calibrated data and an optional **MODEL_DATA** column, which may hold the Fourier inversion of a particular model image. The creation and use of the scratch columns is generally done behind the scenes, but you should be aware that they are there (and when they are used). We will occasionally refer to the rows and columns in the MS.

More on the contents of the MS can be found in §2.1.

### 1.5.5 Data Selection in CASA

We have tried to make the CASA task interface as uniform as possible. If a given parameter appears in multiple tasks, it should, as far as is possible, mean the same thing and be used in the same way in each. There are groups of parameters that appear in a number of tasks to do the same thing, such as for data selection.

The parameters **field**, **spw**, and **selectdata** (which if True expands to a number of sub-parameters) are commonly used in tasks to select data on which to work. These common data selection parameters are described in §2.3.

### 1.6 From Loading Data to Images

The subsections below provide a brief overview of the steps you will need to load data into CASA and obtain a final, calibrated image. Each subject is covered in more detail in Chapters 2 through 6.
An end-to-end workflow diagram for CASA data reduction for interferometry data is shown in Figure 1.8. This might help you chart your course through the package. In the following subsections, we will chart a rough course through this process, with the later chapters filling in the individual boxes.

Note that single-dish data reduction (for example with the ALMA single-dish system) follows a similar course. This is detailed in Chapter 8.

**1.6.1 Loading Data into CASA**

The key data and image import tasks are:
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• importuvfits — import visibility data in UVFITS format (§2.2.5);
• importvla — import data from VLA that is in export format (§2.2.3);
• importasdm — import ALMA data in ASDM format (§2.2.1);
• importevla — import JVLA/EVLA data in SDM format (§2.2.2);
• importfits — import a FITS image into a CASA image format table (§6.24).

These are used to bring in your interferometer data, to be stored as a CASA Measurement set (MS), and any previously made images or models (to be stored as CASA image tables).

The data import tasks will create a MS with a path and name specified by the vis parameter. See §1.5.3 for more information on MS in CASA. The measurement set is the internal data format used by CASA, and conversion from any other native format is necessary for most of the data reduction tasks.

Once data is imported, there are other operations you can use to manipulate the datasets:

• concat — concatenate multiple MSs into a given or a new MS (§2.2.12)

Data import, export, concatenation, and selection detailed in Chapter 2.

1.6.1.1 VLA: Filling data from VLA archive format

VLA data in “archive” format are read into CASA from disk using the importvla task (see §2.2.3). This filler supports the new naming conventions of EVLA antennas when incorporated into the old VLA system.

Note that future data from the EVLA in ASDM format will use a different filler. This will be made available in a later release.

1.6.1.2 Filling data from UVFITS format

For UVFITS format, use the importuvfits task. A subset of popular flavors of UVFITS (in particular UVFITS as written by AIPS) is supported by the CASA filler. See §2.2.5 for details.

1.6.1.3 Loading FITS images

For FITS format images, such as those to be used as calibration models, use the importfits task. Most, though not all, types of FITS images written by astronomical software packages can be read in.

See §6.24 for more information.
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1.6.1.4 Concatenation of multiple MS

Once you have loaded data into measurement sets on disk, you can use the tasks `concat` or `virtualconcat` to combine them.

See §2.2.12 for details.

1.6.2 Data Examination, Editing, and Flagging

The main data examination and flagging tasks are:

- `listobs` — summarize the contents of a MS (§2.2.7);
- `flagmanager` — save and manage versions of the flagging entries in the measurement set (§3.2);
- `plotms` — interactive X-Y plotting and flagging of visibility data (§3.3.1);
- `plotxy` — interactive X-Y plotting and flagging of visibility data (§3.3.2). **note:** `plotxy` is slower than `plotms` and will eventually be phased out, `plotxy` is still useful to create scripted hardcopy output, this functionality will likely be available in `plotms` in the next release);
- `flagdata` — flagging (and unflagging) of specified data (§3.4);
- `viewer` — the CASA viewer can display (as a raster image) MS data, with some editing capabilities (§7);

These tasks allow you to list, plot, and/or flag data in a CASA MS.

There will eventually be tasks for “automatic” flagging to data based upon statistical criteria. Stay tuned.

Examination and editing of synthesis data is described in Chapter 3.

Visualization and editing of an MS using the `casaviewer` is described in Chapter 7.

1.6.2.1 Interactive X-Y Plotting and Flagging

The principal tool for making X-Y plots of visibility data is `plotms` (see §3.3.1). Amplitudes and phases (among other things) can be plotted against several x-axis options.

Interactive flagging (i.e., “see it – flag it”) is possible on the `plotms` X-Y displays of the data (§3.3.1.5). Since flags are inserted into the measurement set, it is useful to backup (or make a copy) of the current flags before further flagging is done, using `flagmanager` (§3.2). Copies of the flag table can also be restored to the MS in this way.
1.6.2.2 Flag the Data Non-interactively

The `flagdata` task ([§3.4](#)) will flag the visibility data set based on the specified data selections. The `listobs` task ([§2.2.7](#)) may be run (e.g. with `verbose=True`) to provide some of the information needed to specify the flagging scope. `flagdata` also contains autoflagging routines.

1.6.2.3 Viewing and Flagging the MS

The CASA viewer can be used to display the data in the MS as a (grayscale or color) raster image. The MS can also be edited. Use of the viewer on an MS is detailed in [§7.5](#).

1.6.3 Calibration

The major calibration tasks are:

- `setjy` — Computes the model visibilities for a specified source given a flux density or model image, knows about standard calibrator sources ([§4.3.5](#));
- `initweights` — if necessary, supports (re-)initialization of the data weights, including an option for enabling spectral weight accounting ([§4.3.1](#));
- `gencal` — Creates a calibration table for known delay and antenna position offsets ([§4.3.6](#));
- `bandpass` — Solves for frequency-dependent (bandpass) complex gains ([§4.4.2](#));
- `gaincal` — Solves for time-dependent (frequency-independent) complex gains ([§4.4.3](#));
- `fluxscale` — Bootstraps the flux density scale from standard calibrators ([§4.4.4](#));
- `polcal` — polarization calibration ([§4.4.5](#));
- `applycal` — Applies calculated calibration solutions ([§4.6.1](#));
- `clearcal` — Re-initializes calibrated visibility data in a given measurement set ([§4.6.3](#));
- `listcal` — Lists calibration solutions ([§4.5.2](#));
- `plotcal` — Plots (and optionally flags) calibration solutions ([§4.5.1](#));
- `uvcontsub` — carry out uv-plane continuum subtraction for spectral-line data ([§4.7.6](#));
- `split` — write out a new (calibrated) MS for specified sources ([§4.7.1](#));
- `cvel` — Regrid a spectral MS onto a new frequency channel system ([§4.7.7](#)).
During the course of calibration, the user will specify a set of calibrations to pre-apply before solving for a particular type of effect, for example gain or bandpass or polarization. The solutions are stored in a calibration table (subdirectory) which is specified by the user, not by the task: care must be taken in naming the table for future use. The user then has the option, as the calibration process proceeds, to accumulate the current state of calibration in a new cumulative table. Finally, the calibration can be applied to the dataset.

Synthesis data calibration is described in detail in Chapter 4.

1.6.3.1 Prior Calibration

The setjy task calculates absolute fluxes for measurement set base on known calibrator sources. This can then be used in later calibration tasks. Currently, setjy knows the flux density as a function of frequency for several standard EVLA flux calibrators and solar system objects, and the value of the flux density can be manually inserted for any other source. If the source is not well-modeled as a point source, then a model image of that source structure can be used (with the total flux density scaled by the values given or calculated above for the flux density). Models are provided for the standard VLA calibrators.

Antenna gain-elevation curves (e.g. for the EVLA antennas) and atmospheric optical depth corrections (applied as an elevation-dependent function) may be pre-applied before solving for the bandpass and gains. CASA v4.1 was the last version where these specialized calibration were supported by explicit parameters in the calibration tasks (gaincurve and opacity). As of v4.2, these parameters have been removed, and gain curves and opacity are supported via gencal, which will generate standard calibration tables describing these effects, much as other a priori effects (Tsys, switched power, etc.) are supported.

See §4.3 for more details.

1.6.3.2 Bandpass Calibration

The bandpass task calculates a bandpass calibration solution: that is, it solves for gain variations in frequency as well as in time. Since the bandpass (relative gain as a function of frequency) generally varies much more slowly than the changes in overall (mean) gain solved for by gaincal, one generally uses a long time scale when solving for the bandpass. The default ‘B’ solution mode solves for the gains in frequency slots consisting of channels or averages of channels.

A polynomial fit for the solution (solution type 'BPOLY’) may be carried out instead of the default frequency-slot based ‘B’ solutions. This single solution will span (combine) multiple spectral windows.

Bandpass calibration is discussed in detail in §4.4.2.

If the gains of the system are changing over the time that the bandpass calibrator is observed, then you may need to do an initial gain calibration (see next step).
1.6.3.3 Gain Calibration

The **gaincal** task determines solutions for the time-based complex antenna gains, for each spectral window, from the specified calibration sources. A solution interval may be specified. The default 'G' solution mode solves for antenna-based gains in each polarization in specified time solution intervals. The 'T' solution mode is the same as 'G' except that it solves for a single solution shared by both polarizations.

A spline fit for the solution (solution type 'GSPLINE') may be carried out instead of the default time-slot based 'G' solutions.

See §4.4.3 for more on gain calibration.

1.6.3.4 Polarization Calibration

The **polcal** task will solve for any unknown polarization leakage and cross-hand phase terms ('D' and 'X' solutions). The 'D' leakage solutions will work on sources with no polarization and sources with known (and supplied, e.g., using **smodel**) polarization. For sources with unknown polarization tracked through a range in parallactic angle on the sky, using poltype 'D+QU', which will first estimate the calibrator polarization for you.

The solution for the unknown cross-hand polarization phase difference 'X' term requires a polarized source with known linear polarization (Q,U).

Frequency-dependent (i.e., per channel) versions of all of these modes are also supported (poltypes 'Df', 'Df+QU', and 'Xf').

See §4.4.5 for more on polarization calibration.

1.6.3.5 Examining Calibration Solutions

The **plotcal** task (§4.5.1) will plot the solutions in a calibration table. The xaxis choices include time (for gaincal solutions) and channel (e.g. for bandpass calibration). The **plotcal** interface and plotting surface is similar to that in **plotxy**. Eventually, **plotcal** will allow you to flag and unflag calibration solutions in the same way that data can be edited in **plotxy**.

The **listcal** task (§4.5.2) will print out the calibration solutions in a specified table.

1.6.3.6 Bootstrapping Flux Calibration

The **fluxscale** task bootstraps the flux density scale from “primary” standard calibrators to the “secondary” calibration sources. Note that the flux density scale must have been previously established on the “primary” calibrator(s), typically using **setjy**, and of course a calibration table containing valid solutions for all calibrators must be available.

See §4.4.4 for more.
1.6.3.7 Correcting the Data

The final step in the calibration process, `applycal` may be used to apply several calibration tables (e.g., from `gaincal` or `bandpass`, along with prior calibration tables). The corrections are applied to the DATA column of the visibility, writing the CORRECTED_DATA column which can then be plotted (e.g. in `plotxy`), split out as the DATA column of a new MS, or imaged (e.g. using `clean`). Any existing corrected data are overwritten.

See §4.6.1 for details.

1.6.3.8 Splitting the Data

After a suitable calibration is achieved, it may be desirable to create one or more new measurement sets containing the data for selected sources. This can be done using the `split` task (§4.7.1).

Further imaging and calibration (e.g. self-calibration) can be carried out on these split Measurement Sets.

1.6.3.9 UV Continuum subtraction

For spectral line data, continuum subtraction can be performed in the image domain (imcontsub) or in the uv domain. For the latter, there are two tasks available: uvcontsub subtracts polynomial of desired order from each baseline, defined by line-free channels.

1.6.3.10 Transforming the Data to a new frame

If you want to transform your dataset to a different frequency and velocity frame than the one it was observed in, then you can use the `cvel` task (§4.7.7). Alternatively, you can do the regridding during the imaging process in `clean` without running `cvel` before.

1.6.4 Synthesis Imaging

The key synthesis imaging tasks are:

- `clean` — Calculates a deconvolved image based on the visibility data, using one of several clean algorithms (§5.3);
- `feather` — Combines a single dish and synthesis image in the Fourier plane (§5.5).

Most of these tasks are used to take calibrated interferometer data, with the possible addition of a single-dish image, and reconstruct a model image of the sky. **Alert:** The `clean` task is now even more powerful and incorporates the functionality of previous specialized tasks such as `mosaic` and `widefield`.

See Chapter 5 for more on synthesis imaging.
1.6.4.1 Cleaning a single-field image or a mosaic

The CLEAN algorithm is the most popular and widely-studied method for reconstructing a model image based on interferometer data. It iteratively removes at each step a fraction of the flux in the brightest pixel in a defined region of the current “dirty” image, and places this in the model image. The clean task implements the CLEAN algorithm for single-field data. The user can choose from a number of options for the particular flavor of CLEAN to use.

Often, the first step in imaging is to make a simple gridded Fourier inversion of the calibrated data to make a “dirty” image. This can then be examined to look for the presence of noticeable emission above the noise, and to assess the quality of the calibration by searching for artifacts in the image. This is done using clean with niter=0.

The clean task can jointly deconvolve mosaics as well as single fields, and also has options to do wide-field and wide-band multi-frequency synthesis imaging.

See §5.3 for an in-depth discussion of the clean task.

1.6.4.2 Feathering in a Single-Dish image

If you have a single-dish image of the large-scale emission in the field, this can be “feathered” into the image obtained from the interferometer data. This is carried out using the feather tasks as the weighted sum in the uv-plane of the gridded transforms of these two images. While not as accurate as a true joint reconstruction of an image from the synthesis and single-dish data together, it is sufficient for most purposes.

See §5.5 for details on the use of the feather task.

1.6.5 Self Calibration

Once a calibrated dataset is obtained, and a first deconvolved model image is computed, a “self-calibration” loop can be performed. Effectively, the model (not restored) image is passed back to another calibration process (on the target data). This refines the calibration of the target source, which up to this point has had (usually) only external calibration applied. This process follows the regular calibration procedure outlined above.

Any number of self-calibration loops can be performed. As long as the images are improving, it is usually prudent to continue the self-calibration iterations.

This process is described in §5.9.

1.6.6 Data and Image Analysis

The key data and image analysis tasks are:

- imhead — summarize and manipulate the “header” information in a CASA image (§6.2);
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• imcontsub — perform continuum subtraction on a spectral-line image cube (§ 6.4);
• immath — perform mathematical operations on or between images (§ 6.6);
• immoments — compute the moments of an image cube (§ 6.7);
• imstat — calculate statistics on an image or part of an image (§ 6.9);
• imval — extract values of one or more pixels, as a spectrum for cubes, from an image (§ 6.10);
• imfit — simple 2D Gaussian fitting of single components to a region of an image (§ 6.5);
• imregrid — regrid an image onto the coordinate system of another image (§ 6.13);
• viewer — there are useful region statistics and image cube plotting capabilities in the viewer (§ 7).

1.6.6.1 What’s in an image?

The imhead task will print out a summary of image “header” keywords and values. This task can also be used to retrieve and change the header values.

See § 6.2 for more.

1.6.6.2 Image statistics

The imstat task will print image statistics. There are options to restrict this to a box region, and to specified channels and Stokes of the cube. This task will return the statistics in a Python dictionary return variable.

See § 6.9 for more.

1.6.6.3 Image values

The imval task will values from an image. There are options to restrict this to a box region, and to return specified channels and Stokes of the cube as a spectrum. This task will return these values in a Python dictionary return variable which can then be operated on in the casapy environment.

See § 6.9 for more.

1.6.6.4 Moments of an image cube

The immoments task will compute a “moments” image of an input image cube. A number of options are available, from the traditional true moments (zero, first, second) and variations thereof, to other images such as median, minimum, or maximum along the moment axis.

See § 6.7 for details.
1.6.6.5 Image math

The `immath` task will allow you to form a new image by mathematical combinations of other images (or parts of images). This is a powerful, but tricky, task to use.

See § 6.6 for more.

1.6.6.6 Regridding an Image

It is occasionally necessary to regrid an image onto a new coordinate system. The `imregrid` task can be used to regrid an input image onto the coordinate system of an existing template image, creating a new output image.

See § 6.13 for a description of this task.

1.6.6.7 Displaying Images

To display an image use the `viewer` task. The viewer will display images in raster, contour, or vector form. Blinking and movies are available for spectral-line image cubes. To start the viewer, type:

```
viewer
```

Executing the `viewer` task will bring up two windows: a viewer screen showing the data or image, and a file catalog list. Click on an image or ms from the file catalog list, choose the proper display, and the image should pop up on the screen. Clicking on the wrench tool (second from left on upper left) will obtain the data display options. Most functions are self-documenting.

The viewer can be run outside of `casapy` by typing `casaviewer`.

See § 7 for more on viewing images.

1.6.7 Getting data and images out of CASA

The key data and image export tasks are:

- `exportuvfits` — export a CASA MS in UVFITS format (§ 2.2.5);
- `exportfits` — export a CASA image table as FITS (§ 6.24).

These tasks can be used to export a CASA MS or image to UVFITS or FITS respectively. See the individual sections referred to above for more on each.
Chapter 2

Visibility Data Import, Export, and Selection

To use CASA to process your data, you first will need to get it into a form that is understood by the package. These are “measurement sets” for synthesis (and single dish) data, and “image tables” for images.

There are a number of tasks used to fill telescope-specific data, to import/export standard formats, to list data contents, and to concatenate multiple datasets. These are:

- `asdmsummary` — list the contents of a archive file in ASDM format ($\S$ 2.2.1)
- `importasdm` — import of ALMA data in ASDM format ($\S$ 2.2.1)
- `importevla` — import of Jansky VLA data and flags in ASDM format ($\S$ 2.2.2)
- `importuvfits` — import visibility data in UVFITS format ($\S$ 2.2.5.1)
- `importfitsidi` — import visibility data in the FITS-IDI format ($\S$ 2.2.5.2)
- `importvla` — import data from VLA that is in export format ($\S$ 2.2.3)
- `importmiriad` — import data from MIRIAD visibilities ($\S$ 2.2.4)
- `exportuvfits` — export a CASA MS in UVFITS format ($\S$ 2.2.5.3)
- `listobs` — summarize the contents of a MS ($\S$ 2.2.7)
- `listpartition` — List the summary of a multi-MS data set in the logger or in a file (($\S$ 2.2.8)
- `listvis` — list the data in a MS ($\S$ 2.2.9)
- `vishead` — list and change the metadata contents of a MS ($\S$ 2.2.10)
- `visstat` — statistics on data in a MS ($\S$ 2.2.11)
CHAPTER 2. VISIBILITY DATA IMPORT, EXPORT, AND SELECTION

- **concat** — concatenate two or more MS into a new MS (§ 2.2.12)
- **virtualconcat** — concatenate two or more MS or multi-MS into a new multi-MS (§ 2.2.12 and chapter 10)

In CASA, there is a standard syntax for selection of data that is employed by multiple tasks. This is described in § 2.3.

There are also tasks for the import and export of image data using FITS:

- **listfits** — show the header content of any FITS file (§ 2.1)
- **importfits** — import a FITS image into a CASA image format table (§ 6.24)
- **exportfits** — export a CASA image table as FITS (§ 6.24)

## 2.1 CASA Measurement Sets

Data is handled in CASA via the table system. In particular, visibility data are stored in a CASA table known as a Measurement Set (MS). Details of the physical and logical MS structure are given below, but for our purposes here an MS is just a construct that contains the data. An MS can also store single dish data (essentially a set of auto-correlations of a 1-element interferometer), though there are also data formats more suitable for single-dish spectra (see § 8).

A full description of the Measurement Set can be found at [http://casa.nrao.edu/Memos/229.html](http://casa.nrao.edu/Memos/229.html).

Note that images are handled through special image tables, although standard FITS I/O is also supported. Images and image data are described in a separate chapter.

The headers of any FITS files can be displayed in the logger with the **listfits** task:

```python
# listfits :: List the HDU and typical data rows of a fits file:
fitsfile = '' # Name of input fits file
```

Unless your data was previously processed by CASA or software based upon its predecessor aips++, you will need to import it into CASA as an MS. Supported formats include some “standard” flavors of UVFITS, the VLA “Export” archive format, and most recently, the ALMA Science Data Model (ASDM) format. These are described below in § 2.2.

Once in Measurement Set form, your data can be accessed through various tools and tasks with a common interface. The most important of these is the **data selection interface** (§ 2.3) which allows you to specify the subset of the data on which the tasks and tools will operate.

### 2.1.1 Under the Hood: Structure of the Measurement Set
CHAPTER 2. VISIBILITY DATA IMPORT, EXPORT, AND SELECTION

It is not necessary that a casual CASA user know the specific details on how the data in the MS is stored and the contents of all the sub-tables. However, we will occasionally refer to specific “columns” of the MS when describing the actions of various tasks, and thus we provide the following synopsis to familiarize the user with the necessary nomenclature. You may skip ahead to subsequent sections if you like!

All CASA data files, including Measurement Sets, are written into the current working directory by default, with each CASA table represented as a separate sub-directory. MS names therefore need only comply with UNIX file or directory naming conventions, and can be referred to from within CASA directly, or via full path names.

An MS consists of a MAIN table containing the visibility data, and associated sub-tables containing auxiliary or secondary information. The tables are logical constructs, with contents located in the physical table.* files on disk. The MAIN table consists of the table.* files in the main directory of the ms-file itself, and the other tables are in the respective subdirectories. The various MS tables and sub-tables can be seen by listing the contents of the MS directory itself (e.g. using Unix ls), or via the browsetable task (§3.6).

See Fig 2.1 for an example of the contents of a MS directory. Or, from the casapy prompt,

CASA <1>: ls ngc5921.ms
IPython system call: ls -F ngc5921.ms
ANTENNA POLARIZATION table.f1 table.f3_TSM1 table.f8
DATA_DESCRIPTION PROCESSOR table.f10 table.f4 table.f8_TSM1
FEED SORTED_TABLE table.f10_TSM1 table.f5 table.f9
FIELD SOURCE table.f11 table.f5_TSM1 table.f9_TSM1
FLAG_CMD SPECTRAL_WINDOW table.f11_TSM1 table.f6 table.info
HISTORY STATE table.f2 table.f6_TSM0 table.lock
OBSERVATION table.dat table.f2_TSM1 table.f7
POINTING table.f0 table.f3 table.f7_TSM1

Note that the MAIN table information is contained in the table.* files in this directory. Each of the sub-table sub-directories contain their own table.dat and other files, e.g.

CASA <2>: ls ngc5921.ms/SOURCE
IPython system call: ls -F ngc5921.ms/SOURCE
table.dat table.f0 table.f0i table.info table.lock

Each “row” in a table contains entries for a number of specified “columns”. For example, in the MAIN table of the MS, the original visibility data is contained in the DATA column — each “cell” contains a matrix of observed complex visibilities for that row at a single time stamp, for a single baseline in a single spectral window. The shape of the data matrix is given by the number of channels and the number of correlations (voltage-products) formed by the correlator for an array.

Table 2.1 lists the non-data columns of the MAIN table that are most important during a typical data reduction session. Table 2.2 lists the key data columns of the MAIN table of an interferometer MS. The MS produced by fillers for specific instruments may insert special columns, such as
Figure 2.1: The contents of a Measurement Set. These tables compose a Measurement Set named `ngc5921.demo.ms` on disk. This display is obtained by using the **File:Open** menu in `browsetable` and left double-clicking on the `ngc5921.demo.ms` directory.

**ALMA_PHASE_CORR**, **ALMA_NO_PHAS_CORR** and **ALMA_PHAS_CORR_FLAG_ROW** for ALMA data filled using the `importasdm` filler (§2.2.1). These columns are visible in `browsetable` and are accessible from the toolkit in the `ms` tool (e.g. the `ms.getdata` method) and from the `tb` “table” tool (e.g. using `tb.getcol`).

Note that when you examine table entries for IDs such as `FIELD_ID` or `DATA_DESC_ID`, you will see 0-based numbers.

The MS can contain a number of “scratch” columns, which are used to hold useful versions of other columns such as the data or weights for further processing. The most common scratch columns are:

- **CORRECTED_DATA** — used to hold calibrated data for imaging or display;

- **MODEL_DATA** — holds the Fourier inversion of a particular model image for calibration or imaging. This column is optional.
CHAPTER 2. VISIBILITY DATA IMPORT, EXPORT, AND SELECTION

Table 2.1: Common columns in the MAIN table of the MS.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANTENNA1</td>
<td>First antenna in baseline</td>
</tr>
<tr>
<td>ANTENNA2</td>
<td>Second antenna in baseline</td>
</tr>
<tr>
<td>FIELD_ID</td>
<td>Field (source no.) identification</td>
</tr>
<tr>
<td>DATA_DESC_ID</td>
<td>Spectral window number, polarization identifier pair (IF no.)</td>
</tr>
<tr>
<td>ARRAY_ID</td>
<td>Subarray number</td>
</tr>
<tr>
<td>OBSERVATION_ID</td>
<td>Observation identification</td>
</tr>
<tr>
<td>POLARIZATION_ID</td>
<td>Polarization identification</td>
</tr>
<tr>
<td>SCAN_NUMBER</td>
<td>Scan number</td>
</tr>
<tr>
<td>TIME</td>
<td>Integration midpoint time</td>
</tr>
<tr>
<td>UVW</td>
<td>UVW coordinates</td>
</tr>
</tbody>
</table>

The creation and use of the scratch columns is generally done behind the scenes, but you should be aware that they are there (and when they are used).

Table 2.2: Commonly accessed MAIN Table data-related columns. Note that the columns ALMA_PHASE_CORR, ALMA_NO_PHAS_CORR and ALMA_PHAS_CORR_FLAG_ROW are specific to ALMA data filled using the importasdm filler.

<table>
<thead>
<tr>
<th>Column</th>
<th>Format</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>DATA</td>
<td>Complex(N_c, N_f)</td>
<td>complex visibility data matrix (= ALMA_PHASE_CORR by default)</td>
</tr>
<tr>
<td>FLAG</td>
<td>Bool(N_c, N_f)</td>
<td>cumulative data flags</td>
</tr>
<tr>
<td>WEIGHT</td>
<td>Float(N_c)</td>
<td>weight for a row</td>
</tr>
<tr>
<td>WEIGHT_SPECTRUM</td>
<td>Float(N_c, N_f)</td>
<td>individual weights for a data matrix</td>
</tr>
<tr>
<td>ALMA_PHASE_CORR</td>
<td>Complex(N_c, N_f)</td>
<td>on-line phase corrected data (Not in VLA data)</td>
</tr>
<tr>
<td>ALMA_NO_PHAS_CORR</td>
<td>Bool(N_c, N_f)</td>
<td>data that has not been phase corrected (Not in VLA data)</td>
</tr>
<tr>
<td>ALMA_PHAS_CORR_FLAG_ROW</td>
<td>Bool(N_c, N_f)</td>
<td>flag to use phase-corrected data or not (not in VLA data)</td>
</tr>
<tr>
<td>MODEL_DATA</td>
<td>Complex(N_c, N_f)</td>
<td>Scratch: created by calibrator or imager tools</td>
</tr>
<tr>
<td>CORRECTED_DATA</td>
<td>Complex(N_c, N_f)</td>
<td>Scratch: created by calibrator or imager tools</td>
</tr>
</tbody>
</table>

Data flags can be set in the MS, too. Whenever a flag is set, the data will be ignored in all processing steps but not physically deleted from the MS. The flags are channel-based and stored in the MS FLAG subtable. Backups can be stored in the ‘MS.flagversions’ file that can be accessed
via the flagmanager (§3.2).

The most recent specification for the MS is Aips++ MeasurementSet definition version 2.0
(http://casa.nrao.edu/Memos/229.html).

2.2 Data Import and Export

There are a number of tasks available to bring data in various forms into CASA as a Measurement Set:

- ALMA Science Data Model format data can be imported into CASA (importasdm)
- JVLA Science Data Model format data, including online flags, can be imported into CASA (importevla)
- VLA Archive format data can be imported into CASA (importvla)
- UVFITS format can be imported into and exported from CASA (importuvfits, importfitsidi, and exportuvfits)

2.2.1 ALMA: Filling of Science Data Model (ASDM) data

The ALMA and JVLA projects have agreed upon a common archival science data model (ASDM) format, and have jointly developed the software to fill this data into CASA. In the ASDM format, the bulk of the data is contained in large binary data format (BDF) tables, with the meta-data and ancillary information in XML tables. This is structured as a directory, like the MS, and was designed to be similar to the MS to facilitate conversion.

The content of an ASDM can be listed with the task asdmsummary:

```
# asdmsummary :: Summarized description of an ASDM dataset.
asdm = '' # Name of input ASDM directory
```

with an output that contains the list and positions of the antennas, followed by the parameters of each scan like observation time, source name, frequency and polarization setup:

```
Input ASDM dataset : TDEM0008.sb3373760.eb3580330.55661.22790537037
=====================================================================
ASDM dataset : TDEM0008.sb3373760.eb3580330.55661.22790537037
=====================================================================
```

Under the Hood:
The importasdm task is just an interface to the stand-alone asdm2MS application. To find out the command-line arguments to this application, do asdm2MS --help.
Exec Block : ExecBlock_0
Telescope : JVLA
Configuration name : B
Observer name : Dr. Juergen Ott


27 antennas have been used in this exec block.

<table>
<thead>
<tr>
<th>Id</th>
<th>Name</th>
<th>Make Station</th>
<th>Diameter</th>
<th>X</th>
<th>Y</th>
<th>Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antenna_0</td>
<td>ea01</td>
<td>UNDEFINED</td>
<td>W36</td>
<td>-1606841.96</td>
<td>-5042279.689</td>
<td>3551913.017</td>
</tr>
<tr>
<td>Antenna_1</td>
<td>ea02</td>
<td>UNDEFINED</td>
<td>E20</td>
<td>-1599340.8</td>
<td>-5043150.965</td>
<td>3554065.219</td>
</tr>
<tr>
<td>Antenna_2</td>
<td>ea03</td>
<td>UNDEFINED</td>
<td>E36</td>
<td>-1596127.728</td>
<td>-5045193.751</td>
<td>3552662.421</td>
</tr>
<tr>
<td>Antenna_3</td>
<td>ea04</td>
<td>UNDEFINED</td>
<td>W28</td>
<td>-1604865.649</td>
<td>-5042190.04</td>
<td>3552962.365</td>
</tr>
<tr>
<td>Antenna_4</td>
<td>ea05</td>
<td>UNDEFINED</td>
<td>W08</td>
<td>-1601614.091</td>
<td>-5042001.653</td>
<td>3554662.509</td>
</tr>
<tr>
<td>Antenna_5</td>
<td>ea06</td>
<td>UNDEFINED</td>
<td>N24</td>
<td>-1600930.06</td>
<td>-5040316.397</td>
<td>3557330.397</td>
</tr>
<tr>
<td>Antenna_6</td>
<td>ea07</td>
<td>UNDEFINED</td>
<td>E32</td>
<td>-1597053.116</td>
<td>-5044604.687</td>
<td>3553068.987</td>
</tr>
<tr>
<td>Antenna_7</td>
<td>ea08</td>
<td>UNDEFINED</td>
<td>N28</td>
<td>-1600863.684</td>
<td>-5039885.318</td>
<td>3557965.319</td>
</tr>
<tr>
<td>Antenna_8</td>
<td>ea09</td>
<td>UNDEFINED</td>
<td>E24</td>
<td>-1598663.09</td>
<td>-5043581.392</td>
<td>3553767.029</td>
</tr>
<tr>
<td>Antenna_9</td>
<td>ea10</td>
<td>UNDEFINED</td>
<td>N32</td>
<td>-1600781.039</td>
<td>-5039347.456</td>
<td>3558761.542</td>
</tr>
<tr>
<td>Antenna_10</td>
<td>ea11</td>
<td>UNDEFINED</td>
<td>E04</td>
<td>-1601068.79</td>
<td>-5042051.91</td>
<td>3554824.835</td>
</tr>
<tr>
<td>Antenna_11</td>
<td>ea12</td>
<td>UNDEFINED</td>
<td>E08</td>
<td>-1600801.926</td>
<td>-5042219.366</td>
<td>3554706.448</td>
</tr>
<tr>
<td>Antenna_12</td>
<td>ea14</td>
<td>UNDEFINED</td>
<td>W12</td>
<td>-1602044.903</td>
<td>-5042025.824</td>
<td>3554427.832</td>
</tr>
<tr>
<td>Antenna_13</td>
<td>ea15</td>
<td>UNDEFINED</td>
<td>W24</td>
<td>-1604008.742</td>
<td>-5042135.828</td>
<td>3553403.707</td>
</tr>
<tr>
<td>Antenna_14</td>
<td>ea16</td>
<td>UNDEFINED</td>
<td>N12</td>
<td>-1601110.052</td>
<td>-5041488.079</td>
<td>3555697.439</td>
</tr>
<tr>
<td>Antenna_15</td>
<td>ea17</td>
<td>UNDEFINED</td>
<td>W32</td>
<td>-1605808.656</td>
<td>-5042230.082</td>
<td>3552459.202</td>
</tr>
<tr>
<td>Antenna_16</td>
<td>ea18</td>
<td>UNDEFINED</td>
<td>N16</td>
<td>-1601061.961</td>
<td>-5041175.88</td>
<td>3556608.022</td>
</tr>
<tr>
<td>Antenna_17</td>
<td>ea19</td>
<td>UNDEFINED</td>
<td>W04</td>
<td>-1601315.893</td>
<td>-5041985.32</td>
<td>3554808.305</td>
</tr>
<tr>
<td>Antenna_18</td>
<td>ea20</td>
<td>UNDEFINED</td>
<td>N36</td>
<td>-1600690.606</td>
<td>-5038758.734</td>
<td>3559632.061</td>
</tr>
<tr>
<td>Antenna_19</td>
<td>ea21</td>
<td>UNDEFINED</td>
<td>E12</td>
<td>-1600416.51</td>
<td>-5042462.45</td>
<td>3554536.041</td>
</tr>
<tr>
<td>Antenna_20</td>
<td>ea22</td>
<td>UNDEFINED</td>
<td>N04</td>
<td>-1601173.979</td>
<td>-5041902.658</td>
<td>3554987.518</td>
</tr>
<tr>
<td>Antenna_21</td>
<td>ea23</td>
<td>UNDEFINED</td>
<td>E16</td>
<td>-1599926.104</td>
<td>-5042772.967</td>
<td>3554319.789</td>
</tr>
<tr>
<td>Antenna_22</td>
<td>ea24</td>
<td>UNDEFINED</td>
<td>W16</td>
<td>-1602592.854</td>
<td>-5042054.997</td>
<td>3554140.7</td>
</tr>
<tr>
<td>Antenna_23</td>
<td>ea25</td>
<td>UNDEFINED</td>
<td>N20</td>
<td>-1601004.709</td>
<td>-5040802.809</td>
<td>3556610.133</td>
</tr>
<tr>
<td>Antenna_24</td>
<td>ea26</td>
<td>UNDEFINED</td>
<td>W20</td>
<td>-1603249.685</td>
<td>-5042091.404</td>
<td>3553797.803</td>
</tr>
<tr>
<td>Antenna_25</td>
<td>ea27</td>
<td>UNDEFINED</td>
<td>E28</td>
<td>-1597899.903</td>
<td>-5044068.676</td>
<td>3553432.445</td>
</tr>
<tr>
<td>Antenna_26</td>
<td>ea28</td>
<td>UNDEFINED</td>
<td>N08</td>
<td>-1601147.94</td>
<td>-5041733.837</td>
<td>3556235.956</td>
</tr>
</tbody>
</table>

Number of scans in this exec Block : 234

scan #1 from 2011-04-10T05:28:13.200000000 to 2011-04-10T05:33:35.500000256
Intents : OBSERVE_TARGET
Sources : 1331+305=3C286
Subscan #1 from 2011-04-10T05:28:13.200000000 to 2011-04-10T05:33:35.500000256
Intent : UNSPECIFIED
Number of integrations : 322

Binary data in uid:///evla/bdf/1302413292901
Number of integrations : 322
Time sampling : INTEGRATION
Correlation Mode : CROSS_AND_AUTO
Spectral resolution type : FULL_RESOLUTION
Atmospheric phase correction : AP_UNCORRECTED
SpectralWindow_0 : numChan = 256, frame = TOPO,
firstChan = 8484000000, chandWidth = 125000 x Polarization_0 : corr = RR,LL

scan #2 from 2011-04-10T05:33:35.500000256 to 2011-04-10T05:35.200000000
CHAPTER 2. VISIBILITY DATA IMPORT, EXPORT, AND SELECTION

The importasdm task will fill SDM1.2 and SDM1.3 format data into a CASA visibility data set (MS). ALMA data was recorded in SDM1.2 format from October 2009 until May 2011. Since May 2011, ALMA is using the SDM 1.3 format. In particular all science data from cycle 0 will be in SDM1.3. The JVLA also started using SDM1.2 in October 2009 and continues to use this format as of October 2011. importasdm can read all of the above formats. The parameter useversion can be used to enable the options process_syspower, process_caldevice, and process_pointing.

The default inputs of importasdm are:

```bash
# importasdm :: Convert an ALMA Science Data Model observation into a
# CASA visibility file (MS) or single-dish data format (Scantable)

asdm    = ''  # Name of input asdm directory (on disk)
vis     = ''  # Root name of the ms to be created. Note the .ms is NOT
              # added
singledish = False  # Set true to output single-dish data format
corr_mode = 'all'  # specifies the correlation mode to be considered on input.
                  # A quoted string containing a sequence of ao, co, ac,or
                  # all separated by whitespaces is expected
srt     = 'all'  # specifies the spectral resolution type to be considered
                # on input. A quoted string containing a sequence of fr,
                # ca, bw, or all separated by whitespaces is expected
time_sampling = 'all'  # specifies the time sampling (INTEGRATION and/or
                       # SUBINTEGRATION) to be considered on input. A quoted
                       # string containing a sequence of i, si, or all separated
                       # by whitespaces is expected
ocorr_mode = 'ca'  # output data for correlation mode AUTO_ONLY (ao) or
                  # CROSS_ONLY (co) or CROSS_AND_AUTO (ca)
compression = False  # Flag for turning on data compression
lazy       = False  # Make the MS DATA column read the ASDM Binary data
                 # directly (faster import, smaller MS)
```
asis = '', # Creates verbatim copies of the ASDMtables in the output measurement set. Value given must be a string of table names separated by spaces; A * wildcard is allowed.

wvr_corrected_data = 'no', # Specifies which values are considered in the SDM binary data to fill the DATA column in the MAIN table of the MS. Expected values for this option are: no, for uncorrected data (default), yes, for the corrected data, and both, for for corrected and uncorrected data. Note if both is selected two measurement sets are created, one with uncorrected data and the other with corrected data.

scans = '', # processes only the specified scans. This value is a semicolon separated list of scan specifications. A scan specification consists in an exec block index followed by the : character; followed by a comma separated list of scan indexes or scan index ranges. A scan index is relative to the exec block it belongs to. Scan indexes are 1-based while exec blocks are 0-based. "0:1" or "2:2~6" or "0:1,1:2~6,8;2:3:24~30" "1,2" are valid values for the option. "3:" alone will be interpreted as, all the scans of the exec block#3. An scan index or a scan index range not preceded by an exec block index will be interpreted as, all the scans with such indexes in all the exec blocks. By default all the scans are considered.

ignore_time = False, # All the rows of the tables Feed, History, Pointing, Source, SysCal, CalDevice, SysPower, and Weather are processed independently of the time range of the selected exec block / scan.

process_syspower = True, # The SysPower table is processed if and only if this parameter is set to true.

process_caldevice = True, # The CalDevice table is processed if and only if this parameter is set to true.

process_pointing = True, # The Pointing table is processed if and only if this parameter is set to true. If set to False, the POINTING table is empty in the resulting MS

process_flags = True, # Create online flags in the FLAG_CMD sub-table.

tbuff = 0.0, # Time padding buffer (seconds)

applyflags = False, # Apply the flags to the MS.

savecmds = False, # Save flag commands to an ASCII file

outfile = '', # Name of ASCII file to save flag commands

flagbackup = True, # Back up flag column before applying flags.

verbose = False, # Output lots of information while the filler is working

overwrite = False, # Over write an existing MS

showversion = False, # Report the version of asdm2MS being used

useversion = 'v3', # Version of asdm2MS to be used ('v3' (default, should work for all data))

bdfflags = False, # Set the MS FLAG column according to the ASDM_binary flags

with_pointing_correction = False, # add (ASDM::Pointing::encoder -
# ASDM::Pointing::pointingDirection) to the value to be
# written in MS::Pointing::direction

If `scans` is set, then `importasdm` processes only the scans specified in the option's value. This value is a semicolon separated list of scan specifications. A scan specification consists in an exec block index followed by the character ':' followed by a comma separated list of scan indexes or scan index ranges. A scan index is relative to the exec block it belongs to. Scan indexes are 1-based while exec blocks are 0-based. The expressions

```
"0:1"
"2:2~6"
"0:1,1:2~6,8;2:,3:24~30"
"1,2"
"3:"
```

are all valid values for the selection. The "3:" selector will be interpreted as 'all the scans of the exec block 3'. An scan index or a scan index range not preceded by an exec block index will be interpreted as 'all the scans with such indexes in all the exec blocks'. By default all the scans are considered.

When `process_flags=True` the task will create online flags based on the `Flag.xml`, `Antenna.xml` and `SpectralWindow.xml` files and copy them to the `FLAG_CMD` sub-table of the MS. The flags will NOT be applied unless the parameter `applyflags` is set to `True`. Optionally, the flags can also be saved to an external ASCII file if `savecmds` is set to `True`.

When `bdfflags=True` the task will apply online flags contained in the ASDM BDF data by calling the executable `bdflags2MS` which the user can also do from the OS prompt. This is recommended for ALMA data.

If `singledish=True`, output data format is scantable (single-dish data format, see [8]) instead of MS. In that case, you must specify name or id of the antenna that you want to obtain data. This can be done by using antenna parameter that is defined as a subparameter of singledish. For single-dish mode, only auto-correlation data are filled, i.e. `ocorr_mode` is forcibly set to 'ao'.

### 2.2.1.1 Import of ASDM data with option lazy=True

With release 4.3, the parameter 'lazy' (default = `False`) is fully tested and operational. If the default value `False` is chosen, `importasdm` will (as in previous versions) fill the visibilities into a newly created `DATA` column of the MS converting them from their binary format in the ASDM to the CASA Table format.

If, however, `lazy` is set to `True`, the task will create the `DATA` column with an ALMA data-specific storage manager, the `(asdmstman)`, which enables CASA to directly read the binary data from the ASDM with on-the-fly conversion. No redundant copy of the raw data is created.

This procedure has the advantage that it saves more than 60% disk space and at least in some cases makes the access to the `DATA` column ≥ 10% faster because the data I/O volume is decreased.
For the same reason, it also accelerates the import itself by ca. a factor 2. The acceleration is particularly large in the applycal task and here particularly on standard SATA disks.

E.g., if your ASDM has a size of 36 GB, the import with default parameters will turn this into an MS of 73 GB size (total disk space consumption = 36 GB + 73 GB = 109 GB). With lazy=True, the imported MS has a size of only 2 GB (total disk space consumption = 36 GB + 2 GB = 38 GB). I.e. your total disk space savings are ca. 65%. Even when you compare to the case where you delete the ASDM after normal import, the solution with lazy import and keeping the ASDM will save you ca. 48% disk space (in the example above 38 GB compared to 73 GB).

The only caveats are the following:

1. You must not delete your ASDM. You can, however, move it but you have to update the reference stored in the MS. Symbolic links will work. See below on how to use the tool method ms.asdmref() to manipulate the ASDM reference.

2. The lazily imported DATA column is read-only. But in any normal data reduction, the DATA column (as opposed to CORRECTED_DATA) is treated as read-only anyway.

The lazily imported MS is numerically identical with the traditionally imported MS and so are all results derived from the MSs. The setting lazy=True might be made the default setting in future CASA releases.

An important additional tool to manipulate lazily imported MSs is the new method ms.asdmref() in the ms tool. If the MS is imported from an ASDM with option lazy=True, the DATA column of the MS is virtual and directly reads the visibilities from the ASDM. A reference to the original ASDM is stored with the MS. If the ASDM needs to be moved to a different path, the reference to it in the MS needs to be updated. This can be achieved with ms.asdmref().

The method takes one argument: abspath. When called with abspath equal to an empty string (default), the method just reports the currently set ASDM path or an empty string if the ASDM path was not set, i.e. the MS was not lazily imported.

If you want to move the referenced ASDM to a different path, you can set the new absolute path by providing it as the value of abspath to the method.

```python
ms.open('uid___A12345_X678_X910.ms',False)
ms.asdmref('/home/alma/myanalysis/uid___A12345_X678_X910')
ms.close()
```

will set the new location of the referenced ASDM to /home/alma/myanalysis/uid___A12345_X678_X910.

Note that the lazily imported MS can be moved without any restrictions independently from the referenced ASDM as long as the absolute path to the ASDM remains accessible, even across file systems.

### 2.2.2 Jansky VLA: Filling of Science Data Model (ASDM) data
CHAPTER 2. VISIBILITY DATA IMPORT, EXPORT, AND SELECTION

The `importevla` task will fill SDM data from the Jansky VLA (or ALMA) into a MS, along with online flagging data contained in the Flag.xml SDM table. Otherwise, it behaves as `importasdm` but with a streamlined parameter set.

The default inputs are:

```plaintext
# importevla :: Convert an Science Data Model observation into a CASA Measurement Set
asdm      = 'asdm'  # Name of input asdm directory (on disk)
vis       = 'vis'   # Root name of the ms to be created. Note the .ms # is NOT added
ocorr_mode = 'co'   # Fill correlation mode AUTO_ONLY (ao), # CROSS_ONLY (co) or CROSS_AND_AUTO (ca)
compression = False  # Flag for turning on data compression
asis      = 'asis'  # Create verbatim copies of these SDM tables in # the MS.
scans     = ''      # List of scans to fill (default is all scans).
verbose   = False   # Output lots of information while the filler is # working
overwrite = False  # Over write an existing MS
online    = True    # Create online flags
tbuff     = 0.0     # Time padding buffer (in seconds)
flagzero  = True    # Create flag commands for zero points
flagpol   = True    # Create flag commands for cross-hand # correlations
shadow    = True    # Create flag commands for shadowed data
tolerance = 0.0     # Amount of shadow allowed (in meters)
addantenna = ''    # File name or dictionary with additional antenna # names, positions and diameters
applyflags = False  # Apply flag commands to MS
savecmds  = False  # Save flag commands to an ASCII file
flagbackup = True  # Back up flag column before applying flags
```

**Under the Hood:**
The `importevla` task is a modified version of the `importasdm` task, that includes import of online flags from the Flag.xml table into the FLAG_CMD MS table, and a streamlined set of parameters.

**ALERT:** If you want to use your JVLA online flags then you must use `importevla` rather than `importasdm`. The `flagcmd` task will process these flags. Also, if you have run `importevla` in CASA 3.3 or earlier, the flag syntax will be processed by the task `oldflagcmd`.

The default action of `importevla` is to construct the FLAG_CMD MS table based on the settings of `online`, `flagzero`, and `shadow` (and sub-parameters). If `applyflags=True` then these flags will be applied after filling. We recommend you use the `flagcmd` task after filling to examine these flags and then apply.

See `importasdm` (§ 2.2.1) for a description of the common parameters. Some differences:

Note that `importevla` automatically loads in VLA switched power information (unlike in previous versions).
The online parameter controls creation of online flags from the Flag.xml SDM table. The tbuff parameter adds a time “buffer” padding for these flags in both directions to deal with timing mismatches. **ALERT:** For JVLA data taken before April 2011, you should set tbuff to a value (in seconds) equal to \(1.5 \times\) the integration time.

The flagzero parameter controls creation of clipping commands to flag visibilities with amplitudes that are exact zeros. If flagpol=True then it will flag the cross-hands (e.g. RL and LR) as well, which might result in low but correct values of these correlations being thrown out (but can catch erroneous zeros also). **ALERT:** This facility is provided as the JVLA correlator, particularly in 2010, occasionally produces visibilities with zero or very small values that need to get flagged out.

The shadow parameter turns on creation of flag commands to remove antenna time ranges where they are shadowed by other antennas in the array. By default it will flag based on the antenna diameter, but if you want more lenient or conservative flagging then set the tolerance sub-parameter, where the shadowed antennas are flagged for all baselines that are shorter than \(r_1 + r_2 - \text{tolerance}\) (the radii are those for the antennas as listed in the ANTENNA sub-table). addantenna can be a file that defines the positions of antennas that are on the ground but do not appear in the MS. They can still shadow antennas in the array.

savecmds will save all flagging commands in the flagdata and flagcmd syntax (\S3.4 and 3.5) to a file to be applied later or for bookkeeping.

A flag backup can be performed using the flagbackup parameter. It saves all current flags to the ‘*.flagversions’ file of the MS, before all new flags are applied.

### 2.2.3 VLA: Filling data from archive format (importvla)

VLA data in archive format (i.e., as downloaded from the VLA data archive) are read into CASA from disk using the importvla task. The inputs are:

```
# importvla :: import VLA archive file(s) to a measurement set:

archivefiles = '' # Name of input VLA archive file(s)
vis = '' # Name of output visibility file
bandname = '' # VLA frequency band name:''=>obtain all bands in archive files
frequencytol = 150000.0 # Frequency shift to define a unique spectral window (Hz)
project = '' # Project name: '' => all projects in file
starttime = '' # start time to search for data
stoptime = '' # end time to search for data
applytsys = True # apply nominal sensitivity scaling to data & weights
autocorr = False # import autocorrelations to ms, if set to True
antnamescheme = 'new' # 'old' or 'new'; 'VA04' or '4' for ant 4
keepblanks = False # Fill scans with empty source names (e.g. tipping scans)?
evlabands = False # Use updated eVLA frequencies and bandwidths
```

The main parameters are archivefiles to specify the input VLA Archive format file names, and vis to specify the output MS name.
ALERT: The scaling of VLA data both before and after the June 2007 Modcomp-turnoff is fully supported, based on the value of applytsys.

The NRAO Archive is located at:

- [https://archive.nrao.edu](https://archive.nrao.edu)

Note that archivefiles takes a string or list of strings, as there are often multiple files for a project in the archive.

For example:

```plaintext
archivefiles = ['AP314_A950519.xp1','AP314_A950519.xp2']
vis = 'NGC7538.ms'
```

The importvla task allows selection on the frequency band. Suppose that you have 1.3 cm line observations in K-band and you have copied the archive data files AP314_A95019.xp* to your working directory and started casapy. Then,

```plaintext
default('importvla')
archivefiles = ['AP314_A950519.xp1','AP314_A950519.xp2','AP314_A950519.xp3']
vis = 'ngc7538.ms'
bandname = 'K'
frequencytol = 10e6
importvla()
```

If the data is located in a different directory on disk, then use the full path name to specify each archive file, e.g.:

```plaintext
archivefiles=['/home/rohir2/jmcnulli/ALMATST1/Data/N7538/AP314_A950519.xp1',
             '/home/rohir2/jmcnulli/ALMATST1/Data/N7538/AP314_A950519.xp2',
             '/home/rohir2/jmcnulli/ALMATST1/Data/N7538/AP314_A950519.xp3']
```

**Important Note:** importvla will import the on-line flags (from the VLA system) along with the data. Shadowed antennas will also be flagged. The flags will be put in the MAIN table and thus available to subsequent tasks and tools. If you wish to revert to unflagged data, use flagmanager (§3.2) to save the flags (if you wish), and then use flagdata (§3.4) with mode='manualflag' and unflag=True to toggle off the flags.

The other parameters are:

### 2.2.3.1 Parameter applytsys

The applytsys parameter controls whether the nominal sensitivity scaling (based on the measured TSYS, with the weights scaled accordingly using the integration time) is applied to the visibility amplitudes or not. If True, then it will be scaled so as to be the same as AIPS FILLM (i.e. approximately in deciJanskys). Note that post-Modcomp data is in raw correlation coefficient and
will be scaled using the TSYS values, while Modcomp-era data had this applied online. In all cases importvla will do the correct thing to data and weights based on an internal flag in the VLA Archive file, either scaling it or unscaling based on your choice for applytsys.

If applytsys=True and you see strange behavior in data amplitudes, it may be due to erroneous TSYS values from the online system. You might want to then fill with applytsys=False and look at the correlation coefficients to see if the behavior is as expected.

2.2.3.2 Parameter bandname

The bandname indicates the VLA Frequency band(s) to load, using the traditional bandname codes. These are:

- '4' = 48-96 MHz
- 'P' = 298-345 MHz
- 'L' = 1.15-1.75 GHz
- 'C' = 4.2-5.1 GHz
- 'X' = 6.8-9.6 GHz
- 'U' = 13.5-16.3 GHz
- 'K' = 20.8-25.8 GHz
- 'Q' = 38-51 GHz
- '' = all bands (default)

Note that as the transition from the VLA to JVLA progresses, the actual frequency ranges covered by the bands will expand, and additional bands will be added (namely 'S' from 1-2 GHz and 'A' from 26.4-40 GHz).

2.2.3.3 Parameter frequencytol

The frequencytol parameter specifies the frequency separation tolerated when assigning data to spectral windows. The default is frequencytol=150000 (Hz). For Doppler tracked data, where the sky frequency changes with time, a frequencytol < 10000 Hz may produce too many unnecessary spectral windows.
2.2.3.4 Parameter project

You can specify a specific project name to import from archive files. The default ‘’ will import data from all projects in file(s) archivefiles.

For example for VLA Project AL519:

```
project = 'AL519' # this will work
project = 'al519' # this will also work
```

while project='AL0519' will NOT work (even though that is what queries to the VLA Archive will print it as - sorry!).

2.2.3.5 Parameters starttime and stoptime

You can specify start and stop times for the data, e.g.:

```
starttime = '1970/1/31/00:00:00'
stoptime = '2199/1/31/23:59:59'
```

Note that the blank defaults will load all data fitting other criteria.

2.2.3.6 Parameter autocorr

Note that autocorrelations are filled into the data set if autocorr=True. Generally for the VLA, autocorrelation data is not useful, and furthermore the imaging routine will try to image the autocorrelation data (it assumes it is single dish data) which will swamp any real signal. Thus, if you do fill the autocorrelations, you will have to flag them before imaging.

2.2.3.7 Parameter antnamescheme

The antnamescheme parameter controls whether importvla will try to use a naming scheme where JVLA antennas are prefixed with EA (e.g. 'EA16') and old VLA antennas have names prefixed with VA (e.g. 'VA11'). Our method to detect whether an antenna is JVLA is not yet perfected, and thus unless you require this feature, simply use antnamescheme='old'.

2.2.3.8 Parameter evlabands

The evlabands=True option is provided to allow users to access JVLA frequencies outside the standard VLA tunings (e.g. the extended C-band above 6 GHz). ALERT: use of this option for standard VLA data will cause unexpected associations, such as X-band data below 8 GHz being extracted to C-band (as the JVLA C-band is 4–8 GHz). Use with care.
2.2.4 Import MIRIAD visibilities (importmiriad)

The task importmiriad allows one to import visibilities in the MIRIAD data format to be converted to a MS. The task has mainly be tested on data from the ATCA and CARMA telescopes and the inputs are:

```
# importmiriad :: Convert a Miriad visibility file into a CASA MeasurementSet
mirfile = '' # Name of input Miriad visibility file
vis = '' # Name of output MeasurementSet
tsys = False # Use the Tsys to set the visibility weights
spw = 'all' # Select spectral windows
vel = '' # Select velocity reference (TOPO,LSRK,LSRD)
linecal = False # (CARMA) Apply line calibration
wide = 'all' # (CARMA) Select wide window averages
debug = 0 # Display increasingly verbose debug messages
```

For details, please contact ATCA or CARMA staff.

2.2.5 UVFITS Import and Export

The UVFITS format is not exactly a standard, but is a popular archive and transport format nonetheless. CASA supports UVFITS files written by the AIPS FITTP task, and others.

UVFITS is supported for both import and export.

2.2.5.1 Import using importuvfits

To import UVFITS format data into CASA, use the importuvfits task:

```
CASA <1>: inp(importuvfits)
fitsfile = '' # Name of input UVFITS file
vis = '' # Name of output visibility file (MS)
antnamescheme = 'old' # For VLA only; 'new' or 'old'; 'VA04' or '04' for VLA ant 4
```

This is straightforward, since all it does is read in a UVFITS file and convert it as best it can into a MS.

For example:

```
importuvfits(fitsfile='NGC5921.fits',vis='ngc5921.ms')
```

**ALERT:** CARMA data can be loaded into CASA. However,
2.2.5.2 Import using importfitsidi

Some uvfits data is written in the FITS-IDI standard. Those files can be imported into CASA with the importfitsidi task:

```python
# importfitsidi :: Convert a FITS-IDI file to a CASA visibility data set
fitsidifile = [''] # Name(s) of input FITS-IDI file(s)
vis = '' # Name of output visibility file (MS)

Example:

importuvfits(fitsidifile='NGC1300.fits', vis='NGC1300.ms')
```

2.2.5.3 Export using exportuvfits

The exportuvfits task will take a MS and write it out in UVFITS format. The defaults are:

```python
# exportuvfits :: Convert a CASA visibility data set (MS) to a UVFITS file
vis = '' # Name of input visibility file
fitsfile = '' # Name of output UVFITS file

datacolumn = 'corrected' # which data to write (data, corrected, model)
field = '' # Field name list
spw = '' # Spectral window and channel selection
antenna = '' # antenna list to select
time = '' # time range selection
nchan = -1 # Number of channels to select
start = 0 # Start channel
width = 1 # Channel averaging width (value>1 indicates averaging)
writesyscal = False # Write GC and TY tables
multisource = True # Write in multi-source format
combinespw = True # Combine spectral windows (True for AIPS)
writestation = True # Write station name instead of antenna name
```

For example:
exportuvfits(vis='ngc5921.split.ms',
fitsfile='NGC5921.split.fits',
multisource=False)

The MS selection parameters field, spw, antenna, and timerange follow the standard selection syntax described in §2.3.

**ALERT:** The nchan, start, and width parameters will be superseded by channel selection in spw. Currently, there is a time parameter rather than timerange.

The datacolumn parameter chooses which data-containing column of the MS (see §2.1.1) is to be written out to the UV FITS file. Choices are: 'data', 'corrected', and 'model'.

There are a number of special parameters that control what is written out. These are mostly here for compatibility with AIPS.

The writesyscal parameter toggles whether GC and TY extension tables are written. These are important for VLBA data, and for JVLA data. **ALERT:** Not yet available.

The multisource parameter determines whether the UV FITS file is a multi-source file or a single-source file, if you have a single-source MS or choose only a single source. Note: the difference between a single-source and multi-source UVFITS file here is whether it has a source (SU) table and the source ID in the random parameters. Some programs (i.e. difmap) only accept single-source files. If you select more than one source in fields, then the multisource parameter will be overridden to be True regardless.

The combinespw parameter allows, if some conditions are met, exporting all of spectral windows (SpW) as a set of ”IF”s in a single ”FREQID” setup instead of giving each SpW its own FREQID in the FITS file. In this context an IF (Intermediate Frequency) is a specialization of an SpW, where each IF in a UV FITS file must have the same number of channels and polarizations, each channel must have the same width, and each IF must be present (even if flagged) throughout the entire observation. If these conditions are not met the data must be exported using multiple FREQIDs, the UV FITS equivalent of a general SpW. This matters since many (sub)programs will work with multiple IFs, but not multiple FREQIDs. For example, a UV FITS file with multiple FREQIDs can be read by AIPS, but you may find that you have to separate the FREQIDs with SPLIT before you can do very much with them. Therefore combinespw=True should be True if possible. Typically MSes where each band was observed simultaneously can be exported with combinespw=True. MSes where the tuning changed with time, e.g. 10 minutes at 4.8 GHz followed by 15 minutes at 8.4 GHz, should be exported to multiple UV FITS files using spw to select one tuning (set of simultaneous SpWs) per file.

The multisource parameter determines whether the UV FITS file is a multi-source file or a single-source file, if you have a single-source MS or choose only a single source. Note: the difference between a single-source and multi-source UVFITS file here is whether it has a source (SU) table and the source ID in the random parameters. If you select more than one source in fields, then the multisource parameter will be overridden to be True regardless.

The combinespw parameter allows combination of all spectral windows at one time. If True, then all spectral windows must have the same shape. For AIPS to read an exported file, then set combinespw=True.
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The writestation parameter toggles the writing of the station name instead of antenna name.

2.2.6 Handling Measurement Set metadata and data

There are tasks provided for basic listing and manipulation of Measurement Set data and metadata. These include:

- **listobs** — summarize the contents of a MS ([2.2.7](#))
- **vishead** — list and change the metadata contents of a MS ([2.2.10](#))
- **visstat** — statistics on data in a MS ([2.2.11](#))
- **concat** — concatenate two or more MS into a new MS ([2.2.12](#))

2.2.7 Summarizing your MS (listobs)

Once you import your data into a CASA Measurement Set, you can get a summary of the MS contents with the listobs task.

The inputs are:

```bash
# listobs :: List the summary of a data set in the logger or in a file
vis = 'day2_TDEM0003_10s_norx' # Name of input visibility file (MS)
selectdata = True # Data selection parameters
field = '' # Field names or field index
# numbers: '' ==> all, field='0~2,3C286'
spw = '' # spectral-window/frequency/channel
# numbers: '' ==> all, antenna='3,VA04'
antenna = '' # antenna/baselines:
# time range: '' ==> all, timerange='09:14:0~09:54:0'
timerange = '' # time range: '' ==> all, timerange='09:14:0~09:54:0'
correlation = '' # Select data based on correlation
correlation = '' # Select data based on correlation
scan = '' # Select data based on observation intent: '' ==> all
intent = '' # Select data based on observation intent: '' ==> all
feed = '' # multi-feed numbers: Not yet implemented
array = '' # (sub)array numbers: '' ==> all
uvrange = '' # uv range: '' ==> all; uvrange
# = '0~100klambda', default units=meters
observation = '' # Select data based on observation ID: '' ==> all
verbose = True # Name of disk file to write output: '' ==> to terminal
listfile = '' # List unflagged row counts?
listunfl = False # If true, it can have significant negative performance impact.
```

The summary (of the selected data) will be written to the logger, to the `casapy-YYYYMMDD-HHMMSS.log` file, and optionally to a file specified in the listfile parameter. For example,
listobs('n5921.ms')

results in the logger messages:

listobs(vis="day2_TDEM0003_10s_norx", selectdata=True, spw="", field="", antenna="", uvrange="", timerange="", correlation="", scan="", intent="", feed="", array="", observation="", verbose=True, listfile="", listunfl=False)

================================================================================
MeasurementSet Name: /Users/jott/casa/casatest/casa4.0/irc/day2_TDEM0003_10s_norx MS Version 2
================================================================================
Observer: Mark J. Mark Claussen      Project: T.B.D.
Observation: EVLA
Data records: 290218      Total integration time = 10016 seconds
Observed from 26-Apr-2010/03:21:56.0 to 26-Apr-2010/06:08:52.0 (UTC)

ObservationID = 0      ArrayID = 0
Date          TimeRange (UTC)        Scan       FldId         FieldName       nRows       SpwIds        Average Interval(s)         Sc
26-Apr-2010/03:21:51.0 - 03:23:21.0 5 2 J0954+1743 2720       [0, 1]          [10, 10]   
03:23:39.0 - 03:28:25.0 6 3 IRC+10216 9918       [0, 1]          [10, 10]   
03:28:38.0 - 03:29:54.0 7 2 J0954+1743 2700       [0, 1]          [10, 10]   
03:30:08.0 - 03:34:53.5 8 3 IRC+10216 9918       [0, 1]          [10, 10]   
...
(nRows = Total number of rows per scan)

Fields: 4

<table>
<thead>
<tr>
<th>ID</th>
<th>Code</th>
<th>Name</th>
<th>RA</th>
<th>Decl</th>
<th>Epoch</th>
<th>SrcId</th>
<th>nRows</th>
</tr>
</thead>
<tbody>
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<td>2</td>
<td>D</td>
<td>J0954+1743</td>
<td>09:54:56.823626</td>
<td>+17.43.31.22243</td>
<td>J2000</td>
<td>2</td>
<td>65326</td>
</tr>
<tr>
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<td>09:47:57.382000</td>
<td>+13.16.40.65999</td>
<td>J2000</td>
<td>3</td>
<td>208242</td>
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<td>F</td>
<td>J1229+0203</td>
<td>12:29:06.699729</td>
<td>+02.03.08.59820</td>
<td>J2000</td>
<td>5</td>
<td>10836</td>
</tr>
<tr>
<td>7</td>
<td>E</td>
<td>J1331+3030</td>
<td>13:31:08.287984</td>
<td>+30.30.32.95886</td>
<td>J2000</td>
<td>7</td>
<td>5814</td>
</tr>
</tbody>
</table>

Spectral Windows: (2 unique spectral windows and 1 unique polarization setups)

<table>
<thead>
<tr>
<th>SpwID</th>
<th>Name</th>
<th>#Chans</th>
<th>Frame</th>
<th>ChanWid(kHz)</th>
<th>TotBW(kHz)</th>
<th>Corrs</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Subband:0</td>
<td>64</td>
<td>TPDU</td>
<td>36387.229</td>
<td>125.000</td>
<td>8000.0 RR RL LR LL</td>
</tr>
<tr>
<td>1</td>
<td>Subband:0</td>
<td>64</td>
<td>TPDU</td>
<td>36304.542</td>
<td>125.000</td>
<td>8000.0 RR RL LR LL</td>
</tr>
</tbody>
</table>

Sources: 10

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>SpwId</th>
<th>RestFreq(MHz)</th>
<th>SysVel(km/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>J1008+0730</td>
<td>0</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>0</td>
<td>J1008+0730</td>
<td>1</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>2</td>
<td>J0954+1743</td>
<td>0</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>2</td>
<td>J0954+1743</td>
<td>1</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>3</td>
<td>IRC+10216</td>
<td>0</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>3</td>
<td>IRC+10216</td>
<td>1</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>5</td>
<td>J1229+0203</td>
<td>0</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>5</td>
<td>J1229+0203</td>
<td>1</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>7</td>
<td>J1331+3030</td>
<td>0</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
<tr>
<td>7</td>
<td>J1331+3030</td>
<td>1</td>
<td>0.03639232</td>
<td>-0.026</td>
</tr>
</tbody>
</table>

Antennas: 19:

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>Station</th>
<th>Diam. (m)</th>
<th>Long.</th>
<th>Lat.</th>
<th>Offset from array center (m)</th>
<th>ITRF Geodesy</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>East</td>
<td>North</td>
</tr>
<tr>
<td>0</td>
<td>ea01</td>
<td>W09</td>
<td>25.0</td>
<td>-107.37.25.2</td>
<td>+33.53.51.0</td>
<td>-521.9407</td>
<td>-332.7782</td>
</tr>
<tr>
<td>1</td>
<td>ea02</td>
<td>E02</td>
<td>25.0</td>
<td>-107.37.04.4</td>
<td>+33.54.01.1</td>
<td>9.8247</td>
<td>-20.4292</td>
</tr>
<tr>
<td>2</td>
<td>ea03</td>
<td>E09</td>
<td>25.0</td>
<td>-107.36.45.1</td>
<td>+33.53.53.6</td>
<td>506.0591</td>
<td>-251.8666</td>
</tr>
<tr>
<td>3</td>
<td>ea04</td>
<td>W01</td>
<td>25.0</td>
<td>-107.37.05.9</td>
<td>+33.54.00.5</td>
<td>-27.3562</td>
<td>-41.3030</td>
</tr>
<tr>
<td>4</td>
<td>ea05</td>
<td>W08</td>
<td>25.0</td>
<td>-107.37.21.6</td>
<td>+33.53.53.0</td>
<td>-432.1158</td>
<td>-272.1493</td>
</tr>
<tr>
<td>5</td>
<td>ea07</td>
<td>N06</td>
<td>25.0</td>
<td>-107.37.06.9</td>
<td>+33.54.10.3</td>
<td>-54.0667</td>
<td>263.8720</td>
</tr>
</tbody>
</table>
using the (default) `verbose=True` option. The most useful extra information that `verbose=True` gives is the list of the scans in the dataset.

### 2.2.8 MMS summary (listpartition)

Similar to `listobs`, `listpartition` shows the summary of a multi-measurement set (MMS).

### 2.2.9 Listing MS data (listvis)

The `listvis` task will print to the terminal (or file) listing of the data in your MS. The inputs are:

```python
# listvis :: List measurement set visibilities.

vis = '' # Name of input visibility file
options = 'ap' # List options: ap only
datacolumn = 'data' # Column to list: data, float_data, corrected, model, residual
field = '' # Field names or index to be listed: ''==>all
spw = '*') # Spectral window: channels: '\*'=>all, spw='1:5~57'
selectdata = False # Other data selection parameters
observation = '' # Select by observation ID(s)
average = '' # Averaging mode: ==>none (Not yet implemented)
showflags = False # Show flagged data (Not yet implemented)
pagerows = 50 # Rows per page
listfile = '' # Output file
```

For example:

```
Units of columns are: Date/Time(YYMMDD/HH:MM:SS UT), UVDist(wavelength), Phase(deg), UVW(m)

WEIGHT: 7
FIELD: 2
SPW: 0
```

Date/Time: 2010/04/26

<table>
<thead>
<tr>
<th>RR:</th>
<th>RL:</th>
<th>LR:</th>
<th>LL:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intrf UVDist</td>
<td>Chn</td>
<td>Amp</td>
<td>Phs Wt F Amp</td>
</tr>
<tr>
<td>2010/04/26/</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
2.2.10 Listing and manipulating MS metadata (vishead)

The vishead task is provided to access keyword information in the Measurement Set. The default inputs are:

```python
# vishead :: List, get, and put metadata in a measurement set
vis = ''     # Name of input visibility file
mode = 'list' # options: list, summary, get, put
listitems = [] # items to list ([] for all)
```

The mode = 'summary' option just gives the same output as listobs.

For mode = 'list' the options are: 'telescope', 'observer', 'project', 'field', 'freq_group_name', 'spw_name', 'schedule', 'schedule_type', 'release_date'.

```python
CASA <29>: vishead('ngc5921.demo.ms',mode='list',listitems=[])  
Out[29]:
{'cal_grp': (array([[-1, -1, -1], dtype=int32), {}),
 'field': (array([['1331+3050002_0', '1445+09900002_0', 'N5921_2'],
                    dtype='|S16'), {}),
 'fld_code': (array([['C', 'A', ''],
                     dtype='|S2'), {}),
 'freq_group_name': (array([['none'],
                            dtype='|S5'), {}),
 'log': ({'r1': False}, {}),
 'observer': (array([['TEST'],
                     dtype='|S5'), {}),
 'project': (array([[''],
                    dtype='|S1'), {}),
 'ptcs': ({'r1': array([[0.53248521]],
                       'r2': array([[2.74392758]],
                       [[0.17412604]]),
```

Type Q to quit, A to toggle long/short list, or RETURN to continue [continue]:

**ALERT:** We are working on improving the format of the listvis output.
'r3': array([[-2.26020138]],
            [[ 0.08843002]]),
            {'MEASINFO': {'Ref': 'J2000', 'type': 'direction'},
             'QuantumUnits': array([['rad', 'rad']],
              dtype='|S4'))},
'release_date': (array([ 4.30444800e+09]),
            {'MEASINFO': {'Ref': 'TAI', 'type': 'epoch'},
             'QuantumUnits': array([['s']],
              dtype='|S2'))},
'schedule': ({'r1': False}, {}),
'schedule_type': (array([['']],
              dtype='|S1'), {}),
'source_name': (array(['1331+30500002_0', '1445+09900002_0', 'N5921_2'],
            dtype='|S16'),
            {}),
'spw_name': (array(['none'],
            dtype='|S5'), {}),
'telescope': (array(['VLA'],
            dtype='|S4'), {})}

You can use mode='get' to retrieve the values of specific keywords, and likewise mode='put' to change them. The inputs are:

mode = 'get'  # options: list, summary, get, put
     hdkey = ''    # keyword to get/put
     hdindex = ''  # keyword index to get/put, counting from zero. ==>all

and

# vishead :: List, summary, get, and put metadata in a measurement set
mode = 'put'  # options: list, summary, get, put
     hdkey = ''    # keyword to get/put
     hdindex = ''  # keyword index to get/put, counting from zero. ==>all
     hdvalue = ''  # value of hdkey

For example, a common operation is to change the Telescope name (e.g. if it is unrecognized), e.g.

CASA <36>: vishead('ngc5921.demo.ms',mode='get',hdkey='telescope')
Out[36]:
(array(['VLA'],
          dtype='|S4'), {})

CASA <37>: vishead('ngc5921.demo.ms',mode='put',hdkey='telescope',hdvalue='JVLA')

CASA <38>: vishead('ngc5921.demo.ms',mode='get',hdkey='telescope')
Out[38]:
(array(['JVLA'],
          dtype='|S5'), {})
2.2.11 MS statistics (visstat)

ALERT: This is still a prototype task.

The visstat task is provided to obtain simple statistics for a Measurement Set, useful in regression tests.

The inputs are:

```plaintext
# visstat :: Displays statistical information from a measurement set
vis = '' # Name of input visibility file
axis = 'amp' # Which values to use
datacolumn = 'data' # Which data column to use (data, corrected, model)

useflags = True # Take flagging into account?
spw = '' # spectral-window/frequency/channel
field = '' # Field names or field index numbers: ''==>all, field='0~2,3C286'
selectdata = True # More data selection parameters (antenna, timerange etc)

antenna = '' # antenna/baselines: ''==>all, antenna = '3,VA04'
timerange = '' # time range: ''==>all, timerange='09:14:0~09:54:0'
correlation = '' # Select data based on correlation
scan = '' # scan numbers: ''==>all
array = '' # (sub)array numbers: ''==>all
uvrange = '' # uv range: ''==>all; uvrange = '0~100klambda', default units=meters
```

Running this task returns a record (Python dictionary) with the statistics, which can be captured in a Python variable. For example,

CASA <42>: mystat = visstat('ngc5921.demo.ms',axis='amp',datacolumn='corrected',field='0')

CASA <43>: mystat
Out[43]:
{'CORRECTED': {'max': 51.938671112060547,
'mean': 14.796444141750133,
'medabsdevmed': 0.28020858764648438,
'median': 14.764373779296875,
'min': 0.81362706422805786,
'npts': 514916.0,
'quartile': 0.56053066253662109,
'rms': 14.829294204711914,
'stddev': 0.98650836609147285,
'sum': 7618925.8316934109,
'sumsq': 113234125.12642419,
'var': 0.97319875636846753}}

CASA <44>: print mystat['CORRECTED']['stddev']
0.986508366091

The options for axis are:
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axis='amplitude'  # or ('amp')
axis='phase'
axis='imag'
axis='scan_number'
axis='flag'

The phase of a complex number is in radians with range \((-\pi, \pi)\).

2.2.12 Concatenating multiple datasets (concat)

Once you have your data in the form of CASA Measurement Sets, you can go ahead and process your data using the editing, calibration, and imaging tasks. In some cases, you will most efficiently operate on single MS for a particular session (such as calibration). Other tasks will (eventually) take multiple Measurement Sets as input. For others, it is easiest to combine your multiple data files into one.

If you need to combine multiple datasets, you can use the `concat` task. The default inputs are:

```plaintext
# concat :: Concatenate several visibility data sets.
vis = ''  # Name of input visibility files to be concatenated
concatvis = ''  # Name of output visibility file
freqtol = ''  # Frequency shift tolerance for considering data as the same spwid
dirtol = ''  # Direction shift tolerance for considering data as the same field
respectname = False  # If true, fields with a different name are not merged even if their direction agrees
timesort = False  # If true, sort by TIME in ascending order
copypointing = True  # Copy all rows of the POINTING table.
visweightscale = []  # List of the weight scaling factors to be applied to the individual MSs
```

The `vis` parameter will take a list of one or more MS. Usually, this will contain all the MS to combine. `concat` will presort the visibilities in time.

With `visweightscale`, a list of weights can be manually specified for the respective input data sets. They will be applied at the time of the combination. To determine the appropriate weights for this procedure, one can inspect the weights (`Wt axis parameter`) of the input datasets in `plotms`.

The `concatvis` parameter contains the name of the output MS. If this points to an existing file on disk, then the MS in `vis` will appended to it, otherwise a new MS file is created to contain the concatenated data. Be careful here!

The `timesort` parameter can be used to make sure the output MS is in time order (e.g. if your input MS have concurrent times). This can possibly speed up some subsequent calibration operations.

Furthermore, the parameter `copypointing` can be used to control whether the POINTING table will be carried along in the concatenation process or if the output MS should not contain a POINTING table. This table is quite large for some data (e.g. ALMA) and is mainly needed for mosaic
imaging. If you are certain that you will not need it, you can save time and disk space by setting `copypointing` to `False`. Also, as stated in the ephemeris handling section 4.7.11.3 `concat` will correctly merge fields which use the same ephemeris.

The parameters `freqtol` and `dirtol` control how close together in frequency and angle on the sky spectral windows or field locations need to be before calling them the same.

**ALERT:** Note that if multiple frequencies or pointings are combined using `freqtol` or `dirtol`, then the data are not changed (i.e. not rephased to the single phase center). Use of these parameters is intended to be tolerant of small offsets (e.g. planets tracked which move slightly in J2000 over the course of observations, or combining epochs observed with slightly different positions).

For example:

```python
default('concat')
vis = ['n4826_16apr.split.ms','n4826_22apr.split.ms']
concatvis = 'n4826_tboth.ms'
freqtol = '50MHz'
visweightscale=['1',2']
concat()
```

combines the two days in `n4826_16apr.split.ms` and `n4826_22apr.split.ms` into a new output MS called `n4826_tboth.ms`, and the second MS is weighted twice the first one.

**ALERT:** Note that if you are concatenating MSs which use antennas which were moved between observations, the MS definition does only foresee a unique antenna ID, but not a unique name(!). The moved antenna will appear twice in the antenna list under the same name but on different stations and with two different IDs. The pair (`NAME@STATION`) will be the unique identifier.

If you would like to only concatenate the subtables of several MSs, not the bulk visibility data, you can use the task `testconcat` instead of `concat` to save time and disk space. `testconcat` has the same parameters as `concat`. It produces an output MS with the concatenated subtables and an empty Main table.

Furthermore, the task `virtualconcat` permits to concatenate MSs into a multi-MS (MMS, see chapter 10) which is much faster as the data is moved into the MMS rather than copied and only some reindexing is done. The bulk data is not rewritten. If you want to keep a copy of the original MSs, set the parameter `keepcopy` of `virtualconcat` to True. The creation of that copy will of course consume some of the time you saved by doing a virtual concatenation. Otherwise `virtualconcat` offers the same functionality as `concat`.

### 2.3 Data Selection

Once in MS form, subsets of the data can be operated on using the tasks and tools. In CASA, there are three common data selection parameters used in the various tasks: `field`, `spw`, and `selectdata`. In addition, the `selectdata` parameter, if set to `True`, will open up a number of other sub-parameters for selection. The selection operation is unified across all the tasks. The
available *selectdata* parameters may not be the same in all tasks. But if present, the same parameters mean the same thing and behave in the same manner when used in any task.

For example:

```python
field = '' # field names or index of calibrators ''==>all
spw = '' # spectral window:channels: ''==>all
selectdata = False # Other data selection parameters
```

versus

```python
field = '' # field names or index of calibrators ''==>all
spw = '' # spectral window:channels: ''==>all
selectdata = True # Other data selection parameters
timerange = '' # time range: ''==>all
uvrange = '' # uv range''=all
antenna = '' # antenna/baselines: ''==>all
scan = '' # scan numbers: Not yet implemented
msselect = '' # Optional data selection (Specialized. but see help)
```

The following are the general syntax rules and descriptions of the individual selection parameters of particular interest for the tasks:

### 2.3.1 General selection syntax

Most of the selections are effected through the use of selection strings. This sub-section describes the general rules used in constructing and parsing these strings. Note that some selections are done through the use of numbers or lists. There are also parameter-specific rules that are described under each parameter.

All lists of basic selection specification-units are comma separated lists and can be of any length. White-spaces before and after the commas (e.g. '3C286, 3C48, 3C84') are ignored, while white-space within sub-strings is treated as part of the sub-string (e.g. '3C286, VIRGO A, 3C84').

All integers can be of any length (in terms of characters) composed of the characters 0–9. Floating point numbers can be in the standard format (DIGIT.DIGIT, DIGIT. or .DIGIT) or in the mantissa-exponent format (e.g. 1.4e9). Places where only integers make sense (e.g. IDs), if a floating point number is given, only the integer part is used (it is truncated).

Range of numbers (integers or real numbers) can be given in the format 'N0~N1'. For integer ranges, it is expanded into a list of integers starting from N0 (inclusive) to N1 (inclusive). For real numbers, it is used to select all values present for the appropriate parameter in the Measurement Set between N0 and N1 (including the boundaries). Note that the '~' character is used rather than the more obvious '-' in order to accommodate hyphens in strings and minus signs in numbers.

Wherever appropriate, units can be specified. The units are used to convert the values given to the units used in the Measurement Set. For ranges, the unit is specified only once (at the end) and applies to both the range boundaries.
2.3.1.1 String Matching

String matching can be done in three ways. Any component of a comma separated list that cannot be parsed as a number, a number range, or a physical quantity is treated as a regular expression or a literal string. If the string does not contain the characters '*', '{', '}' or '?', it is treated as a literal string and used for exact matching. If any of the above mentioned characters are part of the string, they are used as a regular expression. As a result, for most cases, the user does not need to supply any special delimiters for literal strings and/or regular expressions. For example:

```
field = '3'    # match field ID 3 and not select field named "3C286".

field = '3*'   # used as a pattern and matched against field names. If
               # names like "3C84", "3C286", "3020+2207" are found,
               # all will match. Field ID 3 will not be selected
               # (unless of course one of the above mentioned field
               # names also correspond to field ID 3!).

field = '30*'  # will match only with "3020+2207" in above set.
```

However, if it is required that the string be matched exclusively as a regular expression, it can be supplied within a pair of '/' as delimiters (e.g. '/.+/BAND.+/' ). A string enclosed within double quotes (""") is used exclusively for pattern matching (patterns are a simplified form of regular expressions - used in most UNIX commands for string matching). Patterns are internally converted to equivalent regular expressions before matching. See the Unix command "info regex", or visit [http://www.regular-expressions.info](http://www.regular-expressions.info) for details of regular expressions and patterns.

Strings can include any character except the following:

```
  , ; " / NEWLINE
```

(since these are part of the selection syntax). Strings that do not contain any of the characters used to construct regular expressions or patterns are used for exact matches. Although it is highly discouraged to have name in the MS containing the above mentioned reserved characters, if one does choose to include the reserved characters as parts of names etc., those names can only be matched against quoted strings (since regular expression and patterns are a super-set of literal strings – i.e., a literal string is also a valid regular expression).

This leaves '"', '*', '{', '}' or '?' as the list of printable character that cannot be part of a name (i.e., a name containing this character can never be matched in a MSSelection expression). These will be treated as pattern-matching even inside double double quotes ("""). There is currently no escape mechanism (e.g. via a backslash).

Some examples of strings, regular expressions, and patterns:

- The string 'LBAND' will be used as a literal string for exact match. It will match only the exact string LBAND.
- The wildcarded string '*BAND*' will be used as a string pattern for matching. This will match any string which has the sub-string BAND in it.

Some examples of strings, regular expressions, and patterns:
• The string '"*BAND*"' will also be used as a string pattern, matching any string which has the sub-string BAND in it.

• The string '/.+BAND.+/' will be used as a regular expression. This will also match any string which as the sub-string BAND in it. (the .+ regex operator has the same meaning as the * wildcard operator of patterns).

2.3.2 The field Parameter

The field parameter is a string that specifies which field names or ids will be processed in the task or tool. The field selection expression consists of comma separated list of field specifications inside the string.

Field specifications can be literal field names, regular expressions or patterns (see §2.3.1.1). Those fields for which the entry in the NAME column of the FIELD MS sub-table match the literal field name/regular expression/pattern are selected. If a field name/regular expression/pattern fails to match any field name, the given name/regular expression/pattern are matched against the field code. If still no field is selected, an exception is thrown.

Field specifications can also be given by their integer IDs. IDs can be a single or a range of IDs. Field ID selection can also be done as a boolean expression. For a field specification of the form '>ID', all field IDs greater than ID are selected. Similarly for '<ID' all field IDs less than the ID are selected.

For example, if the MS has the following observations:

<table>
<thead>
<tr>
<th>FIELDID</th>
<th>SPWID</th>
<th>NChan</th>
<th>Pol</th>
<th>NRows</th>
<th>Source Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>127</td>
<td>RR</td>
<td>10260</td>
<td>0530+135</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>127</td>
<td>RR</td>
<td>779139</td>
<td>05582+16320</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>127</td>
<td>RR</td>
<td>296190</td>
<td>05309+13319</td>
</tr>
<tr>
<td>3</td>
<td>0</td>
<td>127</td>
<td>RR</td>
<td>58266</td>
<td>0319+415</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>127</td>
<td>RR</td>
<td>32994</td>
<td>1331+305</td>
</tr>
<tr>
<td>5</td>
<td>1</td>
<td>1</td>
<td>RR,RL,LL,RR</td>
<td>23166</td>
<td>KTIP</td>
</tr>
</tbody>
</table>

one might select

field = '0~2,KTIP' # FIELDID 0,1,2 and field name KTIP
field = '0530+135' # field 0530+135
field = '05*'     # fields 0530+135,05582+16320,05309+13319

2.3.3 The spw Parameter

The spw parameter is a string that indicates the specific spectral windows and the channels within them to be used in subsequent processing. Spectral window selection ('SPwSEL') can be given as
a spectral window integer ID, a list of integer IDs, a spectral window name specified as a literal string (for exact match) or a regular expression or pattern.

The specification can be via frequency ranges or by indexes. A range of frequencies are used to select all spectral windows which contain channels within the given range. Frequencies can be specified with an optional unit — the default unit being Hz. Other common choices for radio and mm/sub-mm data are kHz, MHz, and GHz. You will get the entire spectral windows, not just the channels in the specified range. You will need to do channel selection (see below) to do that.

The `spw` can also be selected via comparison for integer IDs. For example, `'>ID'` will select all spectral windows with ID greater than the specified value, while `'<ID'` will select those with ID lesser than the specified value.

Spectral window selection using strings follows the standard rules:

```
spw = '1'       # SPWID 1
spw = '1,3,5'   # SPWID 1,3,5
spw = '0~3'     # SPWID 0,1,2,3
spw = '0~3,5'   # SPWID 0,1,2,3 and 5
spw = '<3,5'    # SPWID 0,1,2,3 and 5
spw = '*'       # All spectral windows
spw = '1412~1415MHz'   # Spectral windows containing 1412-1415MHz
```

In some cases, the spectral windows may allow specification by name. For example,

```
spw = '3mmUSB, 3mmLSB'   # choose by names (if available)
```

might be meaningful for the dataset in question.

Note that the order in which multiple `spws` are given may be important for other parameters. For example, the `mode = 'channel'` in `clean` uses the first `spw` as the origin for the channelization of the resulting image cube.

### 2.3.3.1 Channel selection in the `spw` parameter

Channel selection can be included in the `spw` string in the form `'SPWSEL:CHANSEL'` where `CHANSEL` is the channel selector. In the end, the spectral selection within a given spectral window comes down to the selection of specific channels. We provide a number of shorthand selection options for this. These `CHANSEL` options include:

- **Channel ranges**: `'START~STOP'`
- **Frequency ranges**: `'FSTART~FSTOP'`
- **Velocity ranges**: `'VSTART~VSTOP'` (not yet available)
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- **Bandwidth percentages:** 'PSTART~PSTOP' or 'PWIDTH' (not yet available)
- **Channel striding/stepping:** 'START~STOP~STEP' or 'FSTART~FSTOP~FSTEP'

The most common selection is via channel ranges 'START~STOP' or frequency ranges 'FSTART~FSTOP':

\[
\text{spw} = '0:13\sim53' \quad \text{# spw 0, channels 13-53, inclusive}
\]
\[
\text{spw} = '0:1413\sim1414MHz' \quad \text{# spw 0, 1413-1414MHz section only}
\]

All ranges are inclusive, with the channel given by, or containing the frequency or velocity given by, START and STOP plus all channels between included in the selection. You can also select the spectral window via frequency ranges 'FSTART~FSTOP', as described above:

\[
\text{spw} = '1413\sim1414MHz:1413\sim1414MHz' \quad \text{# channels falling within 1413\sim1414MHz}
\]
\[
\text{spw} = '*:1413\sim1414MHz' \quad \text{# does the same thing}
\]

You can also specify multiple spectral window or channel ranges, e.g.

\[
\text{spw} = '2:16, 3:32\sim34' \quad \text{# spw 2, channel 16 plus spw 3 channels 32-34}
\]
\[
\text{spw} = '2:1\sim3;57\sim63' \quad \text{# spw 2, channels 1-3 and 57-63}
\]
\[
\text{spw} = '1\sim3:10\sim20' \quad \text{# spw 1-3, channels 10-20}
\]
\[
\text{spw} = '*:4\sim56' \quad \text{# all spw, channels 4-56}
\]

Note the use of the wildcard in the last example.

A step can be also be included using '^STEP' as a postfix:

\[
\text{spw} = '0:10\sim100^2' \quad \text{# chans 10,12,14,...,100 of spw 0}
\]
\[
\text{spw} = '*:4' \quad \text{# chans 0,4,8,... of all spw}
\]
\[
\text{spw} = '100\sim150GHz^10GHz' \quad \text{# closest chans to 100,110,...,150GHz}
\]

A step in frequency or velocity will pick the channel in which that frequency or velocity falls, or the nearest channel.

### 2.3.4 The selectdata Parameters

The selectdata parameter, if set to True (default), will expand the inputs to include a number of sub-parameters, given below and in the individual task descriptions (if different). If selectdata = False, then the sub-parameters are treated as blank for selection by the task.

The common selectdata expanded sub-parameters are:
2.3.4.1 The antenna Parameter

The **antenna** selection string is a semi-colon (‘;’) separated list of baseline specifications. A baseline specification is of the form:

- ’ANT1’ — Select all baselines including the antenna(s) specified by the selector ANT1.
- ’ANT1&’ — Select only baselines between the antennas specified by the selector ANT1.
- ’ANT1&ANT2’ — Select only the cross-correlation baselines between the antennas specified by selector ANT1 and antennas specified by selector ANT2. Thus ’ANT1&’ is an abbreviation for ’ANT1&ANT1’.
- ’ANT1&&ANT2’ — Select only auto-correlation and cross-correlation baselines between antennas specified by the selectors ANT1 and ANT2. Note that this is what the default `antenna=''` gives you.
- ’ANT1&&&’ — Select only autocorrelations specified by the selector ANT1.

The selectors **ANT1** and **ANT2** are comma-separated lists of antenna integer-IDs or literal antenna names, patterns, or regular expressions. The **ANT** strings are parsed and converted to a list of antenna integer-IDs or IDs of antennas whose name match the given names/pattern/regular expression. Baselines corresponding to all combinations of the elements in lists on either side of ampersand are selected.

Integer IDs can be specified as single values or a range of integers. When items of the list are parsed as literal strings or regular expressions or patterns (see §2.3.1 for more details on strings). All antenna names that match the given string (exact match)/regular expression/pattern are selected.

**Alert:** Just for antenna selection, a user supplied integer (or integer list) is converted to a string and matched against the antenna name. If that fails, the normal logic of using an integer as an integer and matching it with antenna index is done. Note that currently there is no method for specifying a pure index (e.g. a number that will not first be checked against the name).

The comma is used only as a separator for the list of antenna specifications. The list of baselines specifications is a semi-colon separated list, e.g.

```
antenna = '1~3 & 4~6 ; 10&11'
```

will select baselines between antennas 1,2,3 and 4,5,6 (‘1&4’, ‘1&5’, …, ‘3&6’) plus baseline ‘10&11’.

The wildcard operator (‘*’) will be the most often used pattern. To make it easy to use, the wildcard (and only this operator) can be used without enclosing it in quotes. For example, the selection

```
antenna = 'VA*'
```
will match all antenna names which have ‘VA’ as the first 2 characters in the name (irrespective of what follows after these characters).

There is also a negation operator “!” that can be used to de-select antennas or baselines.

Some examples:

- `antenna='''` # shows blank autocorr pages
- `antenna='*&&*'` # does not show the autocorrs
- `antenna='*&&&'` # show both auto and cross-cor (default)
- `antenna='*&&&'` # shows only autocorrs
- `antenna='5&&&'` # shows non-auto baselines with AN 5
- `antenna='5&&&&'` # AN 5 and 6 autocor
- `antenna='5&&&;6&&*'` # AN 5 autocor plus cross-cors to AN 6
- `antenna='!5'` # baselines not involving AN 5

**Antenna numbers as names:** Needless to say, naming antennas such that the names can also be parsed as a valid token of the syntax is a bad idea. Nevertheless, antenna names that contain any of the reserved characters and/or can be parsed as integers or integer ranges can still be used by enclosing the antenna names in double quotes ("ANT"). E.g. the string

`antenna = '10~15,21,VA22'`

will expand into an antenna ID list 10,11,12,13,14,15,21,22 (assuming the index of the antenna named 'VA22' is 22). If, however, the antenna with ID index 50 is named '21', then the string

`antenna = '10~15,21,VA22'`

will expand into an antenna ID list of 10,11,12,13,14,15,50,22. *Keep in mind that numbers are FIRST matched against names, and only against indices if that matching fails.* There is currently no way to force a selection to use the index, and if there an antenna with that name it will select that.

Read elsewhere (e.g. info regex under Unix) for details of regular expression and patterns.

**Antenna stations** Instead of antenna names, the antenna station names are also accepted by the selection syntax., e.g. 'N15' for the JVLA.

**ANT@STATION sections syntax** Sometimes, data from multiple array configurations are stored in a single MS. But some antennas may have been moved during reconfiguration and the 'ANT@STATION' syntax can distinguish between them. 'ANT' is the antenna name or index and 'STATION' is the antenna station name, e.g., 'EA12@W03' selects antenna EA012 but only at times when it is positioned on station W03. Wildcards are accepted, e.g. 'EA12@*' selects all visibilities from antenna EA12, and '*@W03' would select all antennas that are located on station 'W03' during any observations included in the MS.
2.3.4.2 The scan Parameter

The scan parameter selects the scan ID numbers of the data. There is currently no naming convention for scans. The scan ID is filled into the MS depending on how the data was obtained, so use this with care.

Examples:

- `scan = '3'`  # scan number 3.
- `scan = '1~8'`  # scan numbers 1 through 8, inclusive
- `scan = '1,2,4,6'`  # scans 1,2,4,6
- `scan = '<9'`  # scans <9 (1-8)

NOTE: ALMA and VLA/JVLA number scans starting with 1 and not 0. You can see what the numbering is in your MS using the listobs task with `verbose=True` (see §2.2.7).

2.3.4.3 The timerange Parameter

The time strings in the following (T0, T1 and dT) can be specified as YYYY/MM/DD/HH:MM:SS.FF. The time fields (i.e., YYYY, MM, DD, HH, MM, SS and FF), starting from left to right, may be omitted and they will be replaced by context sensitive defaults as explained below.

Some examples:

1. `timerange='T0~T1'`: Select all time stamps from T0 to T1. For example:

   ```python
timerange = '2007/10/09/00:40:00 ~ 2007/10/09/03:30:00'
```

   Note that fields missing in T0 are replaced by the fields in the time stamp of the first valid row in the MS. For example,

   ```python
timerange = '09/00:40:00 ~ 09/03:30:00'
```

   where the YY/MM/ part of the selection has been defaulted to the start of the MS.

   Fields missing in T1, such as the date part of the string, are replaced by the corresponding fields of T0 (after its defaults are set). For example:

   ```python
timerange = '2007/10/09/22:40:00 ~ 03:30:00'
```

   does the same thing as above.

2. `timerange='T0'`: Select all time stamps that are within an integration time of T0. For example,

   ```python
timerange = '2007/10/09/23:41:00'
```

   Integration time is determined from the first valid row (more rigorously, an average integration time should be computed). Default settings for the missing fields of T0 are as in (1).
3. timerange='T0+dT': Select all time stamps starting from T0 and ending with time stamp T0+dT. For example,

\[
\text{timerange} = '23:41:00+01:00:00'
\]

picks an hour-long chunk of time.
Defaults of T0 are set as usual. Defaults for dT are set from the time corresponding to MJD=0. Thus, dT is a specification of length of time from the assumed nominal ”start of time”.

4. timerange='>T0': Select all times greater than T0. For example,

\[
\begin{align*}
\text{timerange} &= '>2007/10/09/23:41:00' \\
\text{timerange} &= '>23:41:00' \quad \# \text{Same thing without day specification}
\end{align*}
\]

Default settings for T0 are as above.

5. timerange='<T1': Select all times less than T1. For example,

\[
\text{timerange} = '<2007/10/09/23:41:00'
\]

Default settings for T1 are as above.

An ultra-conservative selection might be:

\[
\text{timerange} = '1960/01/01/00:00:00~2020/12/31/23:59:59'
\]

which would choose all possible data!

2.3.4.4 The uvrange Parameter

Rows in the MS can also be selected based on the uv-distance or physical baseline length that the visibilities in each row correspond to. This uvrange can be specified in various formats.

The basic building block of uv-distance specification is a valid number with optional units in the format N[UNIT] (the unit in square brackets is optional). We refer to this basic building block as UVDIST. The default unit is meter. Units of length (such as 'm' and 'km') select physical baseline distances (independent of wavelength). The other allowed units are in wavelengths (such as 'lambda', 'klambda' and 'Mlambda' and are true uv-plane radii

\[
r_{uv} = \sqrt{u^2 + v^2}.
\]

If only a single UVDIST is specified, all rows, the uv-distance of which exactly matches the given UVDIST, are selected.

UVDIST can be specified as a range in the format 'N0~N1[UNIT]' (where N0 and N1 are valid numbers). All rows corresponding to uv-distance between N0 and N1 (inclusive) when converted the specified units are selected.
UVDIST can also be selected via comparison operators. When specified in the format ‘>UVDIST’, all visibilities with uv-distances greater than the given UVDIST are selected. Likewise, when specified in the format ‘<UVDIST’, all rows with uv-distances less than the given UVDIST are selected.

Any number of above mentioned uv-distance specifications can be given as a comma-separated list.

Examples:

```plaintext
uvrange = '100~200km' # an annulus in physical baseline length
uvrange = '24~35Mlambda, 40~45Mlambda' # two annuli in units of mega-wavelengths
uvrange = '< 45klambda' # less than 45 kilolambda
uvrange = '> 0lambda' # greater than zero length (no auto-corrs)
uvrange = '100km' # baselines of length 100km
uvrange = '100klambda' # uv-radius 100 kilolambda
```

### 2.3.4.5 The correlation Parameter

The correlation parameter will select between different correlation products. They can be either the correlation ID or values such as 'XX', 'YY', 'XY', 'YX', 'RR', 'LL', 'RL', 'LR'.

### 2.3.4.6 The intent Parameter

intent is the scan intent that was specified when the observations were set up. They typically describe what was intended with a specific scan, i.e. a flux or phase calibration, a bandpass, a pointing, an observation of your target, or something else or a combination. The format for the scan intents of your observations are listed in the logger when you run `listobs`. Minimum matching with wildcards will work, like '*BANDPASS*'. This is especially useful when multiple intents are attached to scans.

### 2.3.4.7 The observation Parameter

The observation parameter can select between different observation IDs. They will be assigned to parts of a combined data set during a run of `concat`. Each input MS will receive its own observation id in the process.

### 2.3.4.8 The msselect Parameter

More complicated selections within the MS structure are possible using the Table Query Language (TaQL). This is accessed through the msselect parameter.

Note that the TaQL syntax does not follow the rules given in §2.3.1 for our other selection strings. TaQL is explained in more detail in Aips++ NOTE 199 — Table Query Language (http://aips2.nrao.edu/docs/notes/199/199.html). This will eventually become a CASA document. The specific columns of the MS are given in the most recent MS specification document: Aips++ NOTE 229 — MeasurementSet definition version 2.0 (http://aips2.nrao.edu/docs/notes/229/229.html).
Most selection can be carried out using the other selection parameters. However, these are merely shortcuts to the underlying TaQL selection. For example, field and spectral window selection can be done using \texttt{msselect} rather than through \texttt{field} or \texttt{spw}:

\begin{verbatim}
msselect='FIELD_ID == 0' # Field id 0 only
msselect='FIELD_ID <= 1' # Field id 0 and 1
msselect='FIELD_ID IN [1,2]' # Field id 1 and 2
msselect='FIELD_ID==0 && DATA_DESC_ID==3' # Field id 0 in spw id 3 only
\end{verbatim}

\textbf{ALERT:} The \texttt{msselect} style parameters will be phased out of the tasks. TaQL selection will still be available in the Toolkit.
Chapter 3

Data Examination and Editing

3.1 Plotting and Flagging Visibility Data in CASA

The tasks available for plotting and flagging of data are:

- **flagmanager** — manage versions of data flags (§ 3.2)
- **plotms** — create X-Y plots of data in MS and calibration tables, flag data (§ 3.3.1)
- **plotxy** — older X-Y plotter with some functionalities not yet implemented in plotms (§ 3.3.2)
- **flagdata** — Data Flagging (§ 3.4)
- **flagcmd** — manipulate and apply flags using FLAG_CMD table (§ 3.5)
- **browsetable** — browse data in any CASA table (including a MS) (§ 3.6)
- **plotants** — create simple plots of antenna positions (§ 3.3.3)
- **plotuv** — plotting of uv-coverages (§ 3.3.4)

The following sections describe the use of these tasks.

Information on other related operations can be found in:

- **listobs** — list summary of a MS (§ 2.2.7)
- **listvis** — list data in a MS (§ 2.2.9)
- **selectdata** — general data selection syntax (§ 2.3)
- **viewer** — use the casaviewer to display the MS as a raster image, and flag it (§ 7)
3.2 Managing flag versions with flagmanager

The flagmanager task will allow you to manage different versions of flags in your data. These are stored inside a CASA flagversions table, under the name of the MS `<msname>.flagversions`. For example, for the MS `jupiter6cm.usecase.ms`, there will need to be `jupiter6cm.usecase.ms.flagversions` on disk. This is created on import (by importvla or importuvfits) or when flagging is first done on an MS without a .flagversions (e.g. with plotxy).

By default, when the .flagversions is created, this directory will contain a flags.Original in it containing a copy of the original flags in the MAIN table of the MS so that you have a backup. It will also contain a file called FLAG_VERSION_LIST that has the information on the various flag versions there. The flagversions are cumulative, i.e. a specific version number contains all the flags from the lower version numbers, too.

The inputs for flagmanager are:

```
vis = '' # Name of input visibility file (MS)
mode = 'list' # Flag management operation (list,save,restore,delete)
```

The mode='list' option will list the available flagversions from the `<msname>.flagversions` file. For example:

```
CASA <102>: default('flagmanager')
CASA <103>: vis = 'jupiter6cm.usecase.ms'
CASA <104>: mode = 'list'
CASA <105>: flagmanager()
```

MS : /home/imager-b/smyers/Oct07/jupiter6cm.usecase.ms

main : working copy in main table
Original : Original flags at import into CASA
flagautocorr : flagged autocorr
xyflags : Plotxy flags

The mode parameter expands the options. For example, if you wish to save the current flagging state of vis=<msname>,

```
mode = 'save' # Flag management operation (list,save,restore,delete)
versionname = '' # Name of flag version (no spaces)
comment = '' # Short description of flag version
merge = 'replace' # Merge option (replace, and, or)
```

with the output version name specified by versionname. For example, the above xyflags version was written using:

```
default('flagmanager')
vis = 'jupiter6cm.usecase.ms'
mode = 'save'
versionname = 'xyflags'
comment = 'Plotxy flags'
flagmanager()
```
and you can see that there is now a sub-table in the flagversions directory

CASA <106>: ls jupiter6cm.usecase.ms.flagversions/
IPython system call: ls -F jupiter6cm.usecase.ms.flagversions/
flags.flagautocorr flags.Original flags.xyflags FLAG_VERSION_LIST

It is recommended that you use this facility regularly to save versions during flagging.

You can restore a previously saved set of flags using the mode='restore' option:

```python
mode = 'restore'  # Flag management operation (list, save, restore, delete)
versionname = ''  # Name of flag version (no spaces)
merge = 'replace'  # Merge option (replace, and, or)
```

The merge sub-parameter will control how the flags are restored. For merge='replace', the flags in versionname will replace those in the MAIN table of the MS. For merge='and', only data that is flagged in BOTH the current MAIN table and in versionname will be flagged. For merge='or', data flagged in EITHER the MAIN or in versionname will be flagged.

The mode='delete' option can be used to remove versionname from the flagversions:

```python
mode = 'delete'  # Flag management operation (list, save, restore, delete)
versionname = ''  # Name of flag version (no spaces)
```

3.3 X-Y Plotting and Editing of the Data

There are three main X-Y plotting tasks in CASA:

- **plotms** — create X-Y plots of data in MS, flag data (§3.3.1)
- **plotxy** — older X-Y plotter with some functionalities not yet implemented in plotms (§3.3.2)
- **plotants** — create simple plots of antenna positions (§3.3.3)

3.3.1 MS Plotting and Editing using plotms

The principal way to get X-Y plots of visibility data and calibration tables is the plotms task. This task also provides editing capability. Plotms is a GUI-style plotter, based on Qt. It can either be started as a task within CASA (plotms) or from outside CASA (type casaplotms on the command line).

The current inputs to the plotms task are:
CASA <13>: inp plotms
-------------> inp(plotms)

# plotms :: A plotter/interactive flagger for visibility data.

vis = ''  # input MS (or CalTable) (blank for none)
gridrows = 1  # Number of subplot rows (default 1).
gridcols = 1  # Number of subplot columns (default 1).
rowindex = 0  # Row location of the plot (0-based, default 0)
colindex = 0  # Column location of the plot (0-based, default 0)
plotindex = 0  # Index to address a subplot (0-based, default 0)
xaxis = ''  # plot x-axis (blank for default/current)
yaxis = ''  # plot y-axis (blank for default/current)
selectdata = True  # data selection parameters
field = ''  # field names or field index numbers (blank for all)
spw = 'all'  # spectral windows:channels (blank for all)
timerange = ''  # time range (blank for all)
vurange = ''  # uv range (blank for all)
antenna = ''  # antenna/baselines (blank for all)
scan = ''  # scan numbers (blank for all)
correlation = ''  # correlations (blank for all)
array = ''  # (sub)array numbers (blank for all)
observation = ''  # Select by observation ID(s)
msselect = ''  # MS selection (blank for all)

averagedata = True  # data averaging parameters
avgchannel = ''  # average over channel? (blank = False, otherwise value in channels)
avgtime = ''  # average over time? (blank = False, other value in seconds)
avgscan = False  # only valid if time averaging is turned on. average over scans?
avgfield = False  # only valid if time averaging is turned on. average over fields?
avgbaseline = False  # average over all baselines? (mutually exclusive with avgantenna)
avgantenna = False  # average by per-antenna? (mutually exclusive with avgbaseline)
avgspw = False  # average over all spectral windows?
scalar = False  # Do scalar averaging?
transform = False  # transform data in various ways?
extendflag = False  # have flagging extend to other data points?
iteraxis = ''  # the axis over which to iterate
customsymbol = False  # set a custom symbol(s) for unflagged points
coloraxis = ''  # selects which data to use for colorizing
customflaggedsymbol = False  # set a custom plot symbol for flagged points
plotrange = []  # plot axes ranges: [xmin,xmax,ymin,ymax]
title = ''  # Title written along top of plot
xlabel = ''  # Text for horizontal axis. Blank for default.
ylabel = ''  # Text for vertical axis. Blank for default.
showmajorgrid = False  # Show major grid lines (horiz and vert.)
showminorgrid = False  # Show minor grid lines (horiz and vert.)
showlegend = False  # Show a legend on the plot.
plotfile = ''      # Name of plot file to save automatically.
showgui = True    # Show GUI

All of these parameters can also be set or modified from inside the `plotms` window. Note that, if the `vis` parameter is set to the name of a measurement or calibration table set here, when you start up `plotms`, the *entire* measurement set will be plotted, which can be time consuming. It is probably best to leave all parameters blank for now, setting them as needed inside the `plotms` GUI.

![Image of PlotMS GUI]

Figure 3.1: A freshly-started `plotms` GUI window. Note that the **Plots > Data** tab is selected, which is discussed in §3.3.1, 3.3.6 and 3.3.8.
3.3.1.1 Loading and Selecting Data

When `plotms` is first started, a window will appear as in Figure 3.1. It will, by default, display the Plots tab (as chosen from the tabs at the top of the `plotms` window—e.g., Plots, Flagging, Tools...) and the Plots > Data tab (as chosen from the tabs on the far left side of the `plotms` window—e.g., Data, Axes, Pages, Transform...). First, a measurement set should be loaded by clicking on Browse near the top of the Plots tab, and selecting a .ms directory (just select the directory itself; do not descend into the .ms directory). A plot can now be made of the measurement set by clicking on Plot—but beware, this would plot the entire measurement set, and could take quite some time! It is probably better to select a subset of the measurement set using the Selection windows in the Plots > Data tab before clicking Plot.

The options for data selection are:

- field
- spw
- timerange
- uvrange
- antenna
- scan
- corr
- array
- observation
- msselect

These are described in § 2.3. Note that, unlike when setting data selection parameters from the CASA command line, no quotation marks are needed around strings.

Once you have selected the desired subset of data, if you click Plot, `plotms` will by default plot amplitude versus time. See the next section for information about other possible axes.

For a given data selection, `plotms` will only load the data once. This speeds up plotting considerably when changing plot parameters such as different axes, colors etc. Sometimes, however, the data changes on disk, e.g., when other data processing tasks were applied. To force `plotms` to reload the data, checkmark the little force reload box left to the Plot button or press the SHIFT key while clicking the Plot button.
3.3.1.2 A Brief Note Regarding plotms Memory Usage

In order to provide a wide range of flexible interactive plotting options while minimizing the I/O burden, plotms caches the data values for the plot (along with a subset of relevant meta-info) in as efficient a manner as possible. For plots of large numbers of points, the total memory requirement can be quite large. plotms attempts to predict the memory it will require (typically 5 or 6 bytes per plotted point when only one axis is a data axis, depending upon the data shapes involved), and will complain if it believes there is insufficient memory to support the requested plot. For most practical interactive purposes (plots that load and draw in less than a few or a few 10s of minutes), there is usually not a problem on typical modern workstations (attempts to plot large datasets on small laptops might be more likely to encounter problems here).

The absolute upper limit on the number of simultaneously plotted points is currently set by the ability to index the points in the cache. For modern 64 bit machines, this is about 4.29 billion points (requiring around 25GB of memory). (Such plots are not especially useful interactively, since the I/O and draw become prohibitive.)

In general, it is usually most efficient to plot data in modest chunks of not more than a few hundred million points or less, either using selection or averaging. Note that all iterations are (currently) cached simultaneously for iterated plots, so iteration is not a way to manage memory use. A few hundred million points tends to be the practical limit of interactive plotms use w.r.t. information content and utility in the resulting plots, especially when you consider the number of available pixels on your screen.

Since datasets are growing very large, options for plotting arbitrarily large numbers of points—probably in a non-interactive mode—are under consideration for a future release.

3.3.1.3 Plot Axes

The X and Y axes of a plot are selected by clicking on the **Plots > Axes** tab on the left side of the plotms window, and choosing an entry from the drop-down menus below **X Axis** and **Y Axis** (see Figure 3.2). Possible axes are:

- **Scan** — The scan number, as listed by listobs (§ 2.2.7) or the data summary in plotms (§ 3.3.1.8).
- **Field** — The field number, as listed by listobs (§ 2.2.7) or the plotms data summary (§ 3.3.1.8).
- **Time** — The time at which the visibility was observed, given in terms of calendar year (yyyy/mm/dd/hh:mm:ss.ss).
- **Interval** — The integration time in seconds.
- **Spw** — The spectral window number. The characteristics of each spectral window are listed in listobs (§ 2.2.7) or the plotms data summary (§ 3.3.1.8).
- **Channel** — The spectral channel number.
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Figure 3.2: The \textbf{Plots > Axes} tab in the \texttt{plotms} GUI window, used to make a plot of \textbf{Amp} versus Channel.

- **Frequency** — Frequency in units of GHz. The frame for the frequency (e.g., topocentric, barycentric, LSRK) can be set in the \textbf{Plots \textgreater Transform} tab (§3.3.1.9).

- **Velocity** — Velocity in units of km s\(^{-1}\), as defined by the \textbf{Frame}, \textbf{Velocity Defn}, and \textbf{Rest Freq} parameters in the \textbf{Plots \textgreater Transform} tab (§3.3.1.9).

- **Corr** — Correlations which have been assigned integer IDs: 5 = RR; 6 = RL; 7 = LR; and 8 = LL.

- **Antenna1** — The first antenna in a baseline pair; for example, for baseline 2-4, Antenna1 = 2. Antennae are numbered according to the antenna IDs listed in \texttt{listobs} (§2.2.7) or the \texttt{plotms} data summary (§3.3.1.8).

- **Antenna2** — The second antenna in a baseline pair; for baseline 2-4, Antenna2 = 4. Antennae are numbered according to the antenna IDs listed in \texttt{listobs} (§2.2.7) or the \texttt{plotms} data summary (§3.3.1.8).
• **Antenna** — Antenna ID for plotting antenna-based quantities. Antennae are numbered according to the antenna IDs listed in `listobs` (§2.2.7) or the `plotms` data summary (§3.3.1.8).

• **Baseline** — The baseline number.

• **Row** — The MS data row number.

• **Real** and **Imag** — The real and imaginary parts of the visibility in units which are proportional to Jansky (for data which are fully calibrated, the units should be Jy).

• **UVDist** — Projected baseline separations in units of meters. Note that **UVDist** is not a function of frequency.

• **UVwave** — Projected baseline separations in units of the observing wavelength (lambda, not kilolambda). **UVDist\_L** is a function of frequency, and therefore, there will be a different data point for each frequency channel.

• **U, V, and W** — $u, v,$ and $w$ in units of meters.

• **Uwave, Vwave, and Wwave** — $u, v,$ and $w$ in units of wavelengths lambda.

• **Amp** — Data amplitudes in units which are proportional to Jansky (for data which are fully calibrated, the units should be in Jy).

• **Phase** — Data phases in units of degrees.

• **Wt** and **Wt*Amp** — the weight of the visibility and the product of the weight and the amplitude

• **WtSp** — WEIGHT\_SPECTRUM column, i.e. a weight per channel

• **Flag** and **FlagRow** — Data which are flagged have Flag $= 1$, whereas unflagged data are set to Flag $= 0$, **FlagRow** is the MS row number. Note that, to display flagged data, you will have to click on the Plots > Display tab and choose a Flagged Points Symbol (§3.3.1.7).

• **Azimuth** and **Ant-Azimuth** — Azimuth in units of degrees. **Azimuth** plots a fiducial value for the entire array, while **Ant-Azimuth** plots the azimuth for each individual antenna (their azimuths will differ by small amounts, because each antenna is located at a slightly different longitude, latitude, and elevation).

• **Elevation** and **Ant-Elevation** — Elevation in units of degrees. **Elevation** is a representative value for the entire array, while **Ant-Elevation** is the elevation for each individual antenna (their elevations will differ by small amounts, because each antenna is located at a slightly different longitude, latitude, and elevation).

• **HourAngle** — Hour angle in units of hours. This is a fiducial value for the entire array.

• **ParAngle** and **Ant-ParAng** — Parallactic angle in units of degrees. **ParAngle** is the fiducial parallactic angle for all antennae in the array, while **Ant-ParAng** plots the parallactic angle for each individual antenna (their parallactic angles will differ by small amounts, because each antenna is located at a slightly different longitude, latitude, and elevation).
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- **Row** — Data row number. A row number corresponds to a unique time, baseline, and spectral window in the measurement set.

- **FlagRow** — In some tasks, if a whole data row is flagged, then FlagRow will be set to 1 for that row. Unflagged rows have FlagRow = 0. However, note that some tasks (like plotms) may flag a row, but not set FlagRow = 1. It is probably better to plot Flag than FlagRow for most applications.

- **GainAmp, GainPhase, GainReal, GainImag** — are the amplitude, phase, real and imaginary part of the calibration tables for regular complex gain tables.

- **Delay** — The delay of a delay calibration table

- **Opac** — Opacity values of a Opacity calibration table

- **SwPower** — Switched Power values for VLA switched power calibration tables

- **Tsys** — Tsys for ALMA Tsys calibration tables

If the data axis selected from the drop-down menu is already stored in the cache (therefore implying that plotting will proceed relatively quickly), an “X” will appear in the checkbox next to In Cache?. If the data shall be reloaded from disk, the “force reload” checkmark should be set at the bottom of this display.

For relevant data axes like **Amp** and **Phase**, the user will be presented with the option to plot raw data or calibrated data. This can be selected via a drop-down menu called Data Column, located directly under the drop-down menu for X or Y Axis selection (see the Y axis in Figure 3.2). To plot raw data, select “data”; to plot calibrated data, select “corrected”. Note that this choice will only have an impact on a plot if a calibration table has been applied to the measurement set (see applycal, Sect.[4.6.1]).

If a data model has been applied to the measurement set (e.g., with setjy, Sect.[4.3.5]) it can be plotted by selecting “model” from the Data Column menu. Residuals can be plotted via “corrected-model”, “data-model”, “data/model”, and “corrected/model”.

3.3.1.4 Tools

Various tools—selectable as icon buttons at the bottom of the plotms window—can be used to zoom, edit, and locate data. The icon buttons can be seen at the bottom of Figures 3.1 and 3.2 and are, from left to right:

- **Zoom** — The “magnifying glass” button (1st on left) lets you draw a box around a region of the plot (left-click on one corner of the box, and drag the mouse to the opposite corner of the desired box), and then zooms in on this box.

- **Pan** — The “four-arrow” button (2nd from left) lets you pan around a zoomed plot.
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• **Annotate** — The 3rd button from the left is chosen from a drop-down menu to either **Annotate Text** (“T with a green diamond” button) or **Annotate Rectangle** (“pencil” button). In the **Annotate Text** environment, click on a location in the plot where text is desired; a window will pop up, allowing you to type text in it. When you click the **OK** button, this text will appear on the plot. **Annotate Rectangle** simply lets you draw a box on the plot by left-clicking and dragging the mouse. By clicking on the **Annotator** tab near the top of the **plotms** window, different fonts, colors, line styles, etc. can be selected for annotations.

• **Home** — The “house” button (5th from left) returns to the original zoom level.

• **Stack Back** and **Stack Forward** — The left and right arrow buttons (4th and 6th from left) step through the zoom settings you’ve visited.

• **Mark Regions** — The “box with a green diamond” button (7th from left) lets you mark a region for flagging, unflagging, or locating. Left-click on one corner of the desired region, and then drag the mouse to set the opposite corner of the region. You can mark multiple boxes before performing an operation on them.

• **Clear Regions** — Clicking on the “box with a red circle” button (8th from left) will clear all regions which have been marked using **Mark Regions**.

• **Flag** — Click on the “flag” button (9th from left) to flag all points in the marked regions.

• **Unflag** — Click on the “crossed-out flag” button (10th from left) to unflag any flagged points that would be in the marked regions (even if invisible).

• **Locate** — The “magnifying glass on a sheet of paper” button (11th from left) will print out information to the command line about points in the marked regions.

• **Hold Drawing** — If the “hold drawing” button (rightmost, or 12th from left) is depressed, and if new plot axes are selected from the **Plots > Axes** tab, these new data will be cached but not plotted. When the button is clicked on again and un-depressed, it will automatically plot the data that was last requested. This can be particularly useful when changing the size of the **plotms** window.

Under the **Options** tab near the top of the **plotms** window one can select the layout of the page. For multiple plots per page, one can select the grid layout, i.e. the number of rows and columns that determine the number of sub-plots. The **when changing plot axes, clear any existing regions or annotations** checkbox determines when regions and annotation are deleted from the plot. The **Tool Button Style** drop-down menu determines if icons and/or text represent the buttons at the bottom of the **plotms** window.

It is possible to hide these icons by going to the **View > Toolbars** menu at the top of the **plotms** window and un-depressing the **Tools** option (except for **Hide Drawing**, which is hidden by clicking on **View > Toolbars > Display**). In addition, the above tools can also be accessed by clicking on the **Tools** tab near the top of the **plotms** window (just below the **View** menu).
The Tools tab also enables one additional tool, the Tracker. To use Tracker, click on the Hover and/or Display checkbox, and place your mouse over the plot. Tracker will output the X and Y position of your mouse, either as text superimposed on the plot near your mouse (if Hover is selected) or in the blank window in the Tools tab (if Display is selected). Pressing the SPACE bar will copy the lines into the larger white box below to the right. This can be repeated many times and a log of positions and values will be created. The content in the box can then be easily copied and pasted into any other application that is used for data analysis. The Clear button wipes out the content of the box for a fresh start into new scientific adventures.

3.3.1.5 Interactive Flagging in plotms

Interactive flagging, on the principle of “see it — flag it”, is possible on the X-Y display of the data plotted by plotms. The user can use the cursor to mark one or more regions, and then flag, unflag, or list the data that falls in these zones of the display.

Using the row of icons buttons at the bottom of the plotms window (§3.3.1.4), click on the Mark Regions button (which will appear to depress), then mark a region by left-clicking and dragging the mouse (each click and drag will mark an additional region). You can get rid of all your regions by clicking on the Clear Regions. Once regions are marked, you can then click on one of the other buttons to take action:

1. Flag — flag the points in the region(s),
2. Unflag — unflag flagged points in the region(s),
3. Locate — spew out a list of the points in the region(s) to the command line (Warning: this could be a long list!).

Figure 3.3 shows an example of marking regions and then clicking the Flag button. Whenever you click on a button, that action occurs without requiring an explicit disk-write. If you quit plotms and re-enter, you will see your previous edits.

A table with the name <msname>.flagversions (where vis=<msname>) will be created in the same directory if it does not exist already. It is recommended that you save important flagging stages using the flagmanager task (§3.2).

Flags can also be extended with options in the Flagging tab, found near the top of the plotms window. Flag extension enables the user to plot a subset of the data and extend the flagging to a wider set. In this release, the only functional extensions are over channel and correlation.

By checking the boxes next to Extend Flags and Channel, flagging will be extended to other channels in the same spw as the displayed point. For example, if spw='0:0' and channel 0 is displayed, then flagging will extend to all channels in spw 0.

By checking the boxes next to Extend Flags and Correlation, flags will be extended beyond the correlations displayed. Currently the only option is to extend to All correlations, implying that all
correlations will be flagged, e.g. with RR displayed, the correlations RR, RL, LR, and LL will all be flagged.

**WARNING:** use of flag extensions may lead to deletion of much more data than desired. Be careful!

### 3.3.1.6 Averaging Data

The Plots > Data tab enables averaging of the data in order to increase signal-to-noise of the plotted points or to increase plotting speed. The options for **Averaging** are:

- channel
- time
- all baselines or per antenna
- all spectral windows
- scalar

The box next to a given **Averaging** mode needs to be checked for that averaging to take effect.

For example, to average \(n\) channels together, the user would click on the box next to **Channels** so that an “X” appears in it, and then type the number \(n\) in the empty box. When the user next clicks on **Plot**, every \(n\) channels will then be averaged together and the total number of channels plotted will be decreased by a factor of \(n\).

Time averaging is a little trickier, as it is controlled by three fields. If the checkbox next to **Time** under **Averaging** is clicked on, a blank box with units of seconds will become active, along with
two additional checkboxes: \textbf{Scan} and \textbf{Field}. If averaging is desired over a relatively short interval (say, 30 seconds, shorter than the scan length), a number can simply be entered into the blank box and, when the data are replotted, the data will be time averaged. Clicking on the \textbf{Scan} or \textbf{Field} checkbox in this case will have no impact on the time averaging.

These checkboxes become relevant if averaging over a relatively long time—say the entire observation, which consists of multiple scans—is desired. Regardless of how large a number is typed into the \textbf{Time} averaging blank box, only data within individual scans will be averaged together. In order to average data across scan boundaries, the \textbf{Scan} checkbox must be clicked on and the data replotted. Finally, clicking on the \textbf{Field} checkbox enables the averaging of multiple fields together in time.

Clicking on the \textbf{All Baselines} checkbox will average all baselines in the array together. Alternatively, the \textbf{Per Antenna} box may be checked, which will average all baselines for a given antenna together. In this case, all baselines are represented twice; baseline 3-24 will contribute to the averages for both antenna 3 and antenna 24. This can produce some rather strange-looking plots if the user also selects on antenna—say, if the user requests to plot only antenna 0 and then averages \textbf{Per Antenna}. In this case, an average of all baselines including antenna 0 will be plotted, but each individual baseline including antenna 0 will also be plotted (because the presence of baselines 0-1, 0-2, 0-3, etc. trigger \textbf{Per Antenna} averaging to try and compute averages for antennae 1, 2, 3, etc. Therefore, baseline 0-1 will contribute to the average for antenna 0, but it will also singlehandedly be the average for antenna 1.)

Spectral windows can be averaged together by checking the box next to \textbf{All Spectral Windows}. This will result in, for a given channel \( n \), all channels \( n \) from the individual spectral windows being averaged together.

Finally, the default mode is vector averaging, where the complex average is formed by averaging the real and imaginary parts of the relevant visibilities. If \textbf{Scalar} is chosen, then the amplitude of the average is formed by a scalar average of the individual visibility amplitudes.

When averaging, \texttt{plotms} will prefer unflagged data. I.e., if an averaging bin contains any unflagged data at all, only the average of the unflagged will be shown. For averaging bins that contain \textit{only} unflagged data, the average of that unflagged data will be shown. When flagging on a plot of averaged data, the flags will be applied to the unaveraged data in the MS.

### 3.3.1.7 Plot Symbols

Plot symbols are selected in the \textbf{Plots} > \textbf{Display} tab. Most fundamentally, the user can choose to plot unflagged data and/or flagged data. By default, unflagged data is plotted (the circle next to \textbf{Default} is checked under \textbf{Unflagged Points Symbol}), and flagged data is not plotted (the circle next to \textbf{None} is checked under \textbf{Flagged Points Symbol}). We note here that plotting flagged data on an averaged plot is undertaken at the user’s own risk, as the distinction between flagged points and unflagged points becomes blurred if data are averaged over a dimension that is partially flagged. Take, for example, a plot of amplitude versus. time where all channels are averaged together, but some channels have been flagged due to RFI spikes. In creating the average, \texttt{plotms}
will skip over the flagged channels and only use the unflagged ones. The averaged points will be considered unflagged, and the flagged data will not appear on the plot at all.

A selection of **None** produces no data points, **Default** results in data points which are small circles (blue for unflagged data and red for flagged data), and **Custom** allows the user to define a plot symbol. If **Custom** plot symbols are chosen, the user can determine the symbol size by typing a number in the blank box next to **px** or by clicking on the adjacent up or down arrows. Symbol shape can be chosen from the drop-down menu to be either “circle”, “square”, “diamond”, or “pixel” (note than “pixel” only has one possible size). “autoscaling” attempts to adjust the size of the points from dots to circles of different sizes, depending on how many points are plotted. Symbol color can be chosen by typing a hex color code in the blank box next to **Fill**: (e.g., “ff00ff”), or by clicking on the ... button and selecting a color from the pop-up GUI. The adjacent drop-down menu provides options for how heavily the plot symbol is shaded with this color, from heaviest to lightest: “fill”, “mesh1”, “mesh2”, “mesh3”, and “no fill”. Finally, the plot symbol can be outlined in black (if **Outline: Default** is checked) or not (if **Outline: None** is checked). Note that if “no fill” and **Outline: None** are selected, the plot symbols will be invisible.

Finally, unflagged data points can be given informative symbol colors using the **Colorize** parameter. By checking the box next to **Colorize** and selecting a data dimension from the drop-down menu, the data will be plotted with colors that vary along that dimension. For example, if “corr” is chosen from the **Colorize** menu, “RR”, “LL”, “RL”, and “LR” data will each be plotted with a different color. Note that, currently, colorize and plotting flagged data appear to be incompatible; a plot can only include one of these special features at a time.

### 3.3.1.8 Summarizing Data

Information about the measurement set can be obtained from within **plotms** by clicking on the **Summary** button, found at the top menu bar. If “All” is chosen from the pull-down menu next to **Summary**, listobs-style output about scans, correlator configurations, and antennae will be written to the command line from which **plotms** was started. For more detail, click on the **Verbose** checkbox. For a specific subset of the data, choose a selection from the pull-down menu like “Antenna” or “Field”.

### 3.3.1.9 Defining Frequency and Velocity

If the user plans to plot **Frequency**, the reference frame must be defined. By default, the plotted frequency is simply that observed at the telescope. However, transformations can be made by choosing a **Frame** from the drop-down menu in the **Plots > Transform** tab. Frequency reference frames can be chosen to be:

- **LSRK** — local standard of rest (kinematic)
- **LSRD** — local standard of rest (dynamic)
- **BARY** — barycentric
• GEO — geocentric
• TOPO — topocentric
• GALACTO — galactocentric
• LGROUP — Local Group
• CMB — cosmic microwave background dipole

Velocity is affected by the user’s choice of Frame, but it is also impacted by the choice of velocity definition and spectral line rest frequency. The velocity definition is chosen from the Velocity Defn drop-down menu in the Plots > Trans tab, offering selections of Radio, True, or Optical.

For more information on frequency frames and spectral coordinate systems, see the paper by Greisen et al. (A&A, 446, 747, 2006)\textsuperscript{1}.

Finally, the spectral line’s rest frequency in units of MHz should be typed into the blank box next to Rest Freq in the Plots > Trans tab. You can use the me.spectralline tool method to turn transition names into frequencies

CASA <16>: me.spectralline('HI')
Out[17]:
{'m0': {'unit': 'Hz', 'value': 1420405751.786},
 'refer': 'REST',
 'type': 'frequency'}

For a list of known lines in the CASA measures system, use the toolkit command me.linelist().

For example:

CASA <21>: me.linelist()
Out[21]: 'HI H186A H185A H184A H183A H182A H181A H180A H179A H178A H177A H176A H175A H174A H173A H172A H171A H170A H169A H168A H167A H166A H165A H164A H163A H162A H161A H160A... He182A He181A He180A He179A He178A He177A He176A He175A He174A He173A He172A He171A He170A He169A He168A He167A He166A He165A He164A He163A He162A He161A He160A He159A He158A He157A... C186A C185A C184A C183A C182A C181A C180A C179A C178A C177A C176A C175A C174A C173A C172A C171A C170A C169A C168A C167A C166A C165A C164A C163A C162A C161A C160A C159A C158A C157A... NH3_11 NH3_22 NH3_33 NH3_44 NH3_55 NH3_66 NH3_77 NH3_88 NH3_99 NH3_1010 NH3_1111 NH3_1212 OH1612 OH1665 OH1667 OH1720 OH4660 OH4750 OH4765 OH5523 OH6016 OH6030 OH6035 OH6049 OH13433 OH13434 OH13441 OH13442 OH23817 OH23826 CH3OH6.7 CH3OH44 H2022 H2CD4.8 C0_1_0 C0_2_1 C0_3_2 C0_4_3 C0_5_4 C0_6_5 C0_7_6 C0_8_7 13CO_1_0 13CO_2_1 13CO_3_2 13CO_4_3 13CO_5_4 13CO_6_5 13CO_7_6 13CO_8_7 13CO_9_8 C18O_1_0 C18O_2_1 C18O_3_2 C18O_4_3 C18O_5_4 C18O_6_5 C18O_7_6 C18O_8_7 C18O_9_8 C19O_1_0 C19O_2_1 C19O_3_2 C19O_4_3 C19O_5_4 C19O_6_5 C19O_7_6 C19O_8_7 C19O_9_8 CS_1_0 CS_2_1 CS_3_2 CS_4_3 CS_5_4 CS_6_5 CS_7_6 CS_8_7 CS_9_8 CS_10_9 CS_11_10 CS_12_11 CS_13_12 CS_14_13 CS_15_14 CS_16_15 CS_17_16 CS_18_17 CS_19_18 CS_12_19 S10_1_0 S10_2_1 S10_3_2 S10_4_3 S10_5_4 S10_6_5 S10_7_6 S10_8_7 S10_9_8 S10_10_9 S10_11_10 S10_12_11 S10_13_12 S10_14_13 S10_15_14 S10_16_15 S10_17_16 S10_18_17 S10_19_18 S10_20_19 S10_21_20 S10_22_21 S10_23_22'\textsuperscript{1}Also at http://www.aoc.nrao.edu/\textasciitilde greisen/scs.ps
3.3.1.10 Shifting the Phase Center

The plot’s phase center can be shifted in the Plots > Trans tab. Enter the X and Y shifts in units of arcseconds in the blank boxes under Phase center shift.

3.3.1.11 Plot Ranges

The X and Y ranges of the plot can be set in the Plots > Axes tab. By default, the circle next to Automatic will be checked, and the ranges will be auto-scaled. To define the range, click on the circle below Automatic and enter a minimum and maximum value in the blank boxes (as for the X Axis in Figure 3.2). Note that if identical values are placed in the blank boxes (xmin=xmax and/or ymin=ymax), then the values will be ignored and a best guess will be made to auto-range that axis.

3.3.1.12 Plot Labels

The plot and axes labels which are displayed in the plot window are set in the Plots > Canvas tab. To change the plot title, under Canvas Title, click on the circle next to the blank box and enter the desired text. To change the X- and Y-axis labels, similarly click on the circles next to the blank boxes under Show X Axis and Show Y Axis and type the desired text in the blank box. To display these new labels, simply click the Plot button.

The user can determine the locations of axis labels in the Plots > Axes tab. The X-axis label switches from the bottom to the top of the plot depending on what is selected for Attach to. Similarly for the Y-Axis, the user can choose to attach axis labels and tick marks to the Top or Bottom (note that the axis labels have been attached to the Bottom and Right in Figure 3.2). Finally, axis labels can be removed all together by unchecking the boxes next to Show X Axis and Show Y Axis on the Plots > Canvas tab.

3.3.1.13 Grid Lines

A grid of lines can be superimposed on the plot using Grid Lines in the Plots > Canvas tab. “Major” grid lines are drawn at the locations of major tick marks, while “minor” grid lines are drawn at minor tick marks.

Grid line colors, thicknesses, and styles are selected independently for the “major” and “minor” grid lines. Desired line thickness should be typed into the blank boxes just to the right of the Major and Minor labels. Colors are set by clicking on the ... buttons. The blank boxes to the left of the ... buttons will then contain the hex codes for the selected colors (e.g., “808080”). Line styles can also be selected from the drop-down menus to the right of ... buttons.
3.3.1.14 Legend

A plot symbol legend can be added to the plot by clicking on the checkbox next to **Legend** in the **Plots > Canvas** tab. However, given the current functionalities of **plotms**, a symbol legend is of very limited use. This option is useful when overplotting data.

3.3.1.15 The Options Tab

A few miscellaneous options are available in the **Options** tab, the last tab at the top of the **plotms** window. The **Tool Button Style** drop-down menu determines if icons and/or text represent the buttons in the toolbar near the bottom of the **plotms** window.

The **Log Events** drop-down menu determines how verbose **plotms** is in documenting its actions on the command line.

There is a checkbox that determines the persistence of regions and annotations on new plots, labelled **When changing plot axes, clear any existing regions and annotations**.

A useful option is the **fixed size for cached image** checkbox. It determines how large the dots in the panel are with respect to the screen resolution. The values influence how the data is redrawn on the panel. When the **Screen resolution** is selected, the **plotms** window can be resized without redrawing on the canvas – a considerable speedup for large data sets. The penalty is that the dots of the data points are the size of a pixel on the screen, which may be very small for high resolution monitors.

Finally, the **File chooser history limit** determines the number of remembered directories in the file loading pop-up of the **Browse** selection of the **Data** tab.

3.3.1.16 Iteration

In many cases, it is desirable to iterate through the data that were selected in the **Data** tab. A typical example is to display a single baseline in a time vs. amplitude plot and then proceed to the next baselines step by step. This can be done via the **Page** tab on the left hand side of **plotms**. A drop-down menu allows you to select the parameter to be iterated on, such as baseline or spw (press **plot** after changing your selection). The plot titles in the main panel in **plotms** show which data slice is currently displayed. To proceed to the next plot use the **green buttons** below the main panel. The different button symbols let you to proceed panel by panel or to jump to the first or last panel directly. The number of plots per page can be selected under **Options-> Grid**, the last of the top row of tabs (corresponding to the **gridrows** and **gridcols** in the command line interface).

There are two scaling options for the axes: **Global** and **Self**. **Global** will use a common axis range based on data loaded with the selection criteria specified in the **Data** tab. **Self** readjusts the axes scaling to the data for each individual panel of the iteration.

Below, one can invoke multiple panels per display by selecting the number of rows and columns to be displayed on the canvas.
3.3.1.17 Overplotting

Different values of the same dataset can be shown at the same time. E.g. to add a second y-axis, press the “Add Y Axis Data” button under the “Axes” tab. Then select the parameters for the newly created axis by selecting from the now available “Y Axis Data” drop-down menu. If the two y-axes have the same units, they can be displayed both on the same axis. If they are different, e.g. Amplitude and Elevation (both versus time; see Fig. 3.4), one axis should be attached to the left and the other to the right hand side of the plot. Using more than a single y-axis data is also reflected in the “Display” tab where a drop-down menu appears in order to select multiple y-axis options.

![Figure 3.4: Overplotting in plotms](image)

Figure 3.4: Overplotting in plotms: Two different y-axes have been chosen for this plot, amplitude and elevation.

In the plotms input interface, you can overplot by invoking plotms more than once with clearplot=F. Each run of plotms corresponds to a plot to go on top of previous ones.

3.3.1.18 Plotting Multiple Data Sets

plotms can also plot more than a single dataset in separate panels. To do so, press “Add Plot”
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next to the “Plot” button. This will bring up a new data window where the plot parameters are defined. Right-click options are used to “Minimize”, “Maximize”, or “Close” these panels which helps to keep a better overview on the individual datasets. If Options-> Grid is selected to have more than a single panel, the different datasets will be shown side by side.

When plotms is run from the command line, the location of the plots can be defined as follows. gridcols and gridrows define the number of plots on the screen. To define the location where a subplot is to appear on this grid, use colindex and rowindex. If one uses a plotindex, this will be used as a label to address the plot. Each call of plotms with the same plotindex will overplot on the subplot where plotindex was defined the first time. Here is an example on multiple plotms calls:

#Plot in the second column, first row of a 2x2 grid and define this plot as plotindex=0
plotms(vis='vis1.ms', gridrows=2, gridcols=2, colindex=1, rowindex=0)

#Overplot in the same panel using a different axis and symbol for the second plot.
plotms(vis='vis2.ms', clearplots=False, plotindex=1, rowindex=0, colindex=1, gridrows=2, gridcols=2, yaxislocation='right', symbolshape='circle')

#Define a second plot and give it a label plotindex=2, in the lower right corner of the grid.
plotms(vis='vis1.ms', clearplots=False, plotindex=2, rowindex=1, colindex=1, gridrows=2, gridcols=2)

#Move the plot with the overplot one panel to the left. This requires clearing
#the plots and rerunning the script specifications with the new plot locations.
plotms(vis='vis1.ms', gridrows=2, gridcols=2, colindex=0, rowindex=0, symbolshape='diamond')
plotms(vis='vis2.ms', clearplots=False, plotindex=1, rowindex=0, colindex=0, gridrows=2, gridcols=2, yaxislocation='right', symbolshape='circle')
plotms(vis=self.ms, clearplots=False, plotindex=2, rowindex=1, colindex=1, gridrows=2, gridcols=2)

3.3.1.19 Saving your plot

You can save a copy of a plot to file in the Plots > Export tab. Click the Browse button for a GUI-based selection of the directory and file name to which the plot will be saved. The file format can also be determined in this GUI by the suffix given to the filename: .png (PNG), .jpg (JPG), .ps (PS), .pdf (PDF), and txt (TEXT). Alternatively, the file format can be selected from the Format drop-down menu located just below the Browse button. In this case, plotms will add a suffix to the file name depending on the format chosen.

ALERT: The plot files produced by the PS and PDF options can be large and time-consuming to export. The JPG is the smallest.

The exported plot resolution can be manipulated using the High Resolution, DPI, and Size options.

Click on Export to create the file, you may select to either plot only the current page or all pages (filenames will be automatically incremented).
the TEXT format will not save an image but the data points themselves. This allows one to dump the current plot into a file that is used in other programs for further processing. The reported data is the same as when using the locate button in plotms and the format looks like:

```
# x y chan scan field ant1 ant2 ant1name ant2name time freq spw corr offset currchunk irel
# Real Imag None None None None None None None MJD(seconds) GHz None None None None None
0.282938 0.0387583 31 5 2 1 12 ea02@E02 ea21@E01 477868956.000 36.30479452 1 RR 26 0 26
0.263241 -0.00806298 31 7 2 1 12 ea02@E02 ea21@E01 477869356.000 36.30479452 1 RR 29 1 28
0.258207 0.0301206 31 9 2 1 12 ea02@E02 ea21@E01 477869745.000 36.30479452 1 RR 30 2 28
0.311155 -0.0180511 31 11 2 1 12 ea02@E02 ea21@E01 477870133.250 36.30479452 1 RR 31 3 28
0.284589 -0.0628808 31 13 2 1 12 ea02@E02 ea21@E01 477870522.500 36.30479452 1 RR 32 4 28
```

where x and y are the two plotted axes and the other columns contain additional information such as the baselines or frequencies. The three last columns offset, corrchunk, and irel are internal data management items for plotms and you most likely will never use them.

### 3.3.1.20 Exiting plotms

To exit the plotms GUI, select Quit from the File menu at the top of the plotms window. You can also dismiss the window by killing it with the “X” on the frame.

Alternatively, you can just leave it alone, and plotms will keep running in the background. If the data file changes in the background, you can force reloading the data via the ‘force reload’ checkbox next to the ‘Plot’ button. Alternatively, press SHIFT while clicking on ‘Plot’ for the same purpose.

### 3.3.2 Plotting and Editing using plotxy

**Inside the Toolkit:** Access to matplotlib is also provided through the pl tool. See below for a description of the pl tool functions.

**ALERT:** The plotxy code is fragile and slow, and is being replaced by the plotms (§ 3.3.1). We retain plotxy in this release as not all functionality is available yet in plotms.

Plotxy is a tool for visualizing and editing visibility data. Unlike plotms, it is useful in scripting, as it can non-interactively produce a hardcopy plot (see § 3.3.2.13). It also has multi-plot (§ 3.3.2.8), iteration (§ 3.3.2.3), and overplotting (§ 3.3.2.4) functionality—unlike plotms in the current release. Plotxy uses the matplotlib plotting library to display its plots. You can find information on matplotlib at [http://matplotlib.sourceforge.net/](http://matplotlib.sourceforge.net/).

To bring up this plotter use the plotxy task. The inputs are:

```
# plotxy :: X-Y plotter/interactive flagger for visibility data
vis = '' # Name of input visibility
xaxis = 'time' # X-axis: def = 'time': see help for options
yaxis = 'amp' # Y-axis: def = 'amp': see help for options
```
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Figure 3.5: The plotxy plotter, showing the Jupiter data versus uv-distance. You can see bad data in this plot. The **bottom set of buttons** on the lower left are: 1,2,3) **Home, Back, and Forward**. Click to navigate between previously defined views (akin to web navigation). 4) **Pan**. Click and drag to pan to a new position. 5) **Zoom**. Click to define a rectangular region for zooming. 6) **Subplot Configuration**. Click to configure the parameters of the subplot and spaces for the figures. 7) **Save**. Click to launch a file save dialog box. The **upper set of buttons in the lower left** are: 1) **Mark Region**. Press this to begin marking regions (rather than zooming or panning). 2,3,4) **Flag, Unflag, Locate**. Click on these to flag, unflag, or list the data within the marked regions. 5) **Next**. Click to move to the next in a series of iterated plots. Finally, the **cursor readout** is on the bottom right.

```python
datacolumn = 'data'  # data (raw), corrected, model, residual (corrected - model)
selectdata = False  # Other data selection parameters
spw = ''  # spectral window:channels: ''==>all, spw='1:5-57'
field = ''  # field names or index of calibrators: ''==>all
averagemode = ''  # Select averaging type: 'vector', 'scalar'
```
restfreq = '' # a frequency quanta or transition name. see help for options
extendflag = False # Have flagging extend to other data points?
subplot = 111 # Panel number on display screen (yxn)
plotsymbol = '.' # Options include . : , o ^ v > < s + x D d 2 3 4 h H | _
plotcolor = 'darkcyn' # Plot color
plotrange = [-1, -1, -1, -1] # The range of data to be plotted (see help)
multicolor = 'corr' # Plot in different colors: Options: none, both, chan, corr
selectplot = False # Select additional plotting options (e.g, fontsize, title, etc)
overplot = False # Overplot on current plot (if possible)
showflags = False # Show flagged data?
interactive = True # Show plot on gui?
figfile = '' # ''= no plot hardcopy, otherwise supply name

**ALERT:** The `plotxy` task expects all of the scratch columns to be present in the MS, even if it is not asked to plot the contents. If you get an error to the effect "Invalid Table operation: Table: cannot add a column" then use `clearcal()` to force these columns to be made in the MS. Note that this will clear anything in all scratch columns (in case some were actually there and being used).

Setting `selectdata=True` opens up the selection sub-parameters:

```
selectdata = True # Other data selection parameters
    antenna = '' # antenna/baselines: ''==>all, antenna = '3,VA04'
    timerange = '' # time range: ''==>all
    correlation = '' # correlations: default = ''
    scan = '' # scan numbers: Not yet implemented
    feed = '' # multi-feed numbers: Not yet implemented
    array = '' # array numbers: Not yet implemented
    uvrange = '' # uv range''==>all; uvrange = '0~100kl' (default unit=meters)
```

These are described in § 2.3.

Averaging is controlled with the set of parameters

```
averagemode = 'vector' # Select averaging type: vector, scalar
timebin = '0' # Length of time-interval in seconds to average
crossscans = False # Have time averaging cross scan boundaries?
crossbls = False # have averaging cross over baselines?
crossarrays = False # have averaging cross over arrays?
stackspw = False # stack multiple spw on top of each other?
width = '1' # Number of channels to average
```

See § 3.3.2.9 below for more on averaging.

You can extend the flagging beyond the data cell plotted:

```
extendflag = True # Have flagging extend to other data points?
extendcorr = '' # flagging correlation extension type
extendchan = '' # flagging channel extension type
extendspw = '' # flagging spectral window extension type
extendant = '' # flagging antenna extension type
extendtime = '' # flagging time extension type
```
See §3.3.2.11 below for more on flag extension.

The restfreq parameter can be set to a transition or frequency:

\[
\text{restfreq} = \text{`HI'} \quad \# \text{a frequency quanta or transition name. see help for options}
\]

frame = 'LSRK' # frequency frame for spectral axis. see help for options
doppler = 'RADIO' # doppler mode. see help for options

See §3.3.2.12 below for more on setting rest frequencies and frames.

Setting selectplot=True will open up a set of plotting control sub-parameters. These are described in §3.3.2.2 below.

The interactive and figfile parameters allow non-interactive production of hardcopy plots. See §3.3.2.13 for more details on saving plots to disk.

The iteration, overplot, plotrange, plotsymbol, showflags and subplot parameters deserve extra explanation, and are described below.

For example:

\[
\text{plotxy(vis='jupiter6cm.ms',} \\
\text{xaxis='uvdist',} \\
\text{yaxis='amp',} \\
\text{field='JUPITER',} \\
\text{selectdata=True,} \\
\text{correlation='RR,LL',} \\
\text{selectplot=True,} \\
\text{title = 'Jupiter 6cm uncalibrated'})
\]

The plotter resulting from these settings is shown in figure 3.5.

\textbf{ALERT:} The plotxy task still has a number of issues. The averaging has been greatly speeded up in this release, but there are cases where the plots will be made incorrectly. In particular, there are problems plotting multiple spw at the same time. There are sometimes also cases where data that you have flagged in plotxy from averaged data is done so incorrectly. This task is under active development for the next cycle to fix these remaining problems, so users should be aware of this.

\textbf{ALERT:} Another know problem with (plotxy) is that it fails if the path to your working directory contains spaces in its name, e.g. /users/smyers/MyTest/ is fine, but /users/smyers/My Test/ is not!

3.3.2.1 GUI Plot Control

You can use the various buttons on the plotxy GUI to control its operation – in particular, to determine flagging and unflagging behaviors.

There is a standard row of buttons at the bottom. These include (left to right):

- \textbf{Home} — The “house” button (1st on left) returns to the original zoom level.
• **Step** — The left and right arrow buttons (2nd and 3rd from left) step through the zoom settings you’ve visited.

• **Pan** — The “four-arrow button” (4th from left) lets you pan in zoomed plot.

• **Zoom** — The most useful is the “magnifying glass” (5th from the left) which lets you draw a box and zoom in on the plot.

• **Panels** — The “window-thingy” button (second from right) brings up a menu to adjust the panel placement in the plot.

• **Save** — The “disk” button (last on right) saves a .png copy of the plot to a generically named file on disk.

In a row above these, there are a set of other buttons (left to right):

• **Mark Region** — If depressed lets you draw rectangles to mark points in the panels. This is done by left-clicking and dragging the mouse. You can Mark multiple boxes before doing something. Clicking the button again will un-depress it and forget the regions. ESC will remove the last region marked.

• **Flag** — Click this to Flag the points in a marked region.

• **Unflag** — Click this to Unflag any flagged point that would be in that region (even if invisible).

• **Locate** — Print out some information to the logger on points in the marked regions.

• **Next** — Step to the next plot in an iteration.

• **Quit** — Exit plotcal, clear the window and detach from the MS.

These buttons are shared with the plotcal tool.

### 3.3.2.2 The selectplot Parameters

These parameters work in concert with the native matplotlib functionality to enable flexible representations of data displays.

Setting `selectplot=True` will open up a set of plotting control sub-parameters:

```makefile
selectplot = True  # Select additional plotting options (e.g., fontsize, title, etc)
markersize = 5.0  # Size of plotted marks
linewidth = 1.0  # Width of plotted lines
skipnrows = 1  # Plot every nth point
newplot = False  # Replace the last plot or not when overplotting
clearpanel = 'Auto'  # Specify if old plots are cleared or not
title = ''  # Plot title (above plot)
xlabels = ''  # Label for x-axis
```
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ylabels = '' # Label for y-axis
fontsize = 10.0 # Font size for labels
windowsize = 5.0 # Window size: not yet implemented

The markersize parameter will change the size of the plot symbols. Increasing it will help legibility when doing screen shots. Decreasing it can help in congested plots. The linewidth parameter will do similar things to the lines.

The skipnrows parameter, if set to an integer \( n \) greater than 1, will allow only every \( n \)th point to be plotted. It does this, as the name suggests, by skipping over whole rows of the MS, so beware (channels are all within the same row for a given spw). Be careful flagging on data where you have skipped points! Note that you can also reduce the number of points plotted via averaging (§3.3.2.9) or channel striding in the spw specification (§2.3.3).

The newplot toggle lets you choose whether or not the last layer plotted is replaced when overplot=True, or whether a new layer is added.

The clearpanel parameter turns on/off the clearing of plot panels that lie under the current panel layer being plotted. The options are: 'none' (clear nothing), 'auto' (automatically clear the plotting area), 'current' (clear the current plot area only), and 'all' (clear the whole plot panel).

The title, xlabels, and ylabels parameters can be used to change the plot title and axes labels.

The fontsize parameter is useful in order to enlarge the label fonts so as to be visible when making plots for screen capture, or just to improve legibility. Shrinking can help if you have lots of panels on the plot also.

The windowsize parameter is supposed to allow adjustments on the window size. ALERT: This currently does nothing, unless you set it below 1.0, in which case it will produce an error.

3.3.2.3 The iteration parameter

Under Page one can select plot iterations parameters. There are currently four iteration options available: 'field', 'antenna', and 'baseline'. If one of these options is chosen, the data will be split into separate plot displays for each value of the iteration axis (e.g., for the VLA, the 'antenna' option will get you 27 displays, one for each antenna). An example use of iteration:

```python
# choose channel averaging, every 5 channels
plotxy('n5921.ms','channel',subplot=221,iteration='antenna',width='5')
```

The results of this are shown in Figure 3.6. Note that this example combines the use of width, iteration and subplot.

NOTE: If you use iteration='antenna' or 'baseline', be aware if you have set antenna selection. You can also control whether you see auto-correlations or not using the appropriate syntax, e.g. antenna='*&&*', or antenna='*&&&' (§2.3.4.1).
3.3.2.4 The overplot parameter

The overplot parameter toggles whether the current plot will be overlaid on the previous plot or subpanel (via the subplot setting, § section:edit.plot.plotxy.subplot) or will overwrite it. The default is False and the new plot will replace the old.

The overplot parameter interacts with the newplot sub-parameter (see § 3.3.2.2).

See § 3.3.2.7 for an example using overplot.
3.3.2.5 The plotrange parameter

The plotrange parameter can be used to specify the size of the plot. The format is \([\text{xmin}, \text{xmax}, \text{ymin}, \text{ymax}]\). The units are those on the plot. For example,

\[
\text{plotrange} = \[-20,100,15,30\]
\]

Note that if \(\text{xmin}=\text{xmax}\) and/or \(\text{ymin}=\text{ymax}\), then the values will be ignored and a best guess will be made to auto-range that axis.

Unfortunately, the units for the time axis must be in Julian seconds. This is somewhat inconvenient as the usual time parameter is given in Julian days. To calculate the Julian seconds the \text{me.epoch} tool can be used. An example: For 02:00 UT on 2012/05/22, the value of MJD seconds can be calculated via

\[
86400*(\text{me.epoch('utc','2012/05/22')}['m0']['value']+2/24.)
\]

which results in \(4844368800.0\).

3.3.2.6 The plotsymbol parameter

The plotsymbol parameter defines both the line or symbol for the data being drawn as well as the color; from the matplotlib online documentation (e.g., type \text{pl.plot?} for help):

The following line styles are supported:
- : solid line
-- : dashed line
-. : dash-dot line
.: : dotted line
, : points
: : pixels
o : circle symbols
^ : triangle up symbols
v : triangle down symbols
< : triangle left symbols
> : triangle right symbols
s : square symbols
+ : plus symbols
x : cross symbols
D : diamond symbols
d : thin diamond symbols
1 : tripod down symbols
2 : tripod up symbols
3 : tripod left symbols
4 : tripod right symbols
h : hexagon symbols
H : rotated hexagon symbols
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- \( p \) : pentagon symbols
- \( | \) : vertical line symbols
- \( \_ \) : horizontal line symbols
- \( \text{steps} \) : use gnuplot style 'steps' # kwarg only

The following color abbreviations are supported:
- \( b \) : blue
- \( g \) : green
- \( r \) : red
- \( c \) : cyan
- \( m \) : magenta
- \( y \) : yellow
- \( k \) : black
- \( w \) : white

In addition, you can specify colors in many weird and wonderful ways, including full names 'green', hex strings '#008000', RGB or RGBA tuples (0,1,0,1) or grayscale intensities as a string '0.8'. Line styles and colors are combined in a single format string, as in 'bo' for blue circles.

### 3.3.2.7 The `showflags` parameter

The `showflags` parameter determines whether only unflagged data (showflags=False) or flagged (showflags=True) data is plotted by this execution. The default is False and will show only unflagged “good” data.

Note that if you want to plot both unflagged and flagged data, in different colors, then you need to run `plotxy` twice using `overplot` (see §3.3.2.4) the second time, e.g.

```
> plotxy(vis="myfile", xaxis='uvdist', yaxis='amp')
> plotxy(vis="myfile", xaxis='uvdist', yaxis='amp', overplot=True, showflags=True)
```

### 3.3.2.8 The `subplot` parameter

The `subplot` parameter takes three numbers. The first is the number of y panels (stacking vertically), the second is the number of xpanels (stacking horizontally) and the third is the number of the panel you want to draw into. For example, `subplot=212` would draw into the lower of two panels stacked vertically in the figure.

An example use of subplot capability is shown in Fig 3.7. These were drawn with the commands (for the top, bottom left, and bottom right panels respectively):

```
plotxy('n5921.ms','channel', # plot channels for the n5921.ms data set
    field=0', # plot only first field
    datacolumn='corrected', # plot corrected data
    plotcolor='', # over-ride default plot color
    plotsymbol='go', # use green circles
    subplot=211) # plot to the top of two panels
```
plotxy(‘n5921.ms’,’x’,
    field=’0’,
    datacolumn=’corrected’,
    subplot=223,
    plotcolor=’’,
    plotsymbol=’r.’)  # plot antennas for n5921.ms data set
    # plot only first field
    # plot corrected data
    # plot to 3rd panel (lower left) in 2x2 grid
    # over-ride default plot color
    # red dots

plotxy(‘n5921.ms’,’u’,’v’,
    field=’0’,
    datacolumn=’corrected’,
    subplot=224,
    plotcolor=’’,
    plotsymbol=’b,’)  # plot uv-coverage for n5921.ms data set
    # plot only first field
    # plot corrected data
    # plot to the lower right in a 2x2 grid
    # over-ride default plot color
    # blue, somewhat larger dots
    # NOTE: You can change the gridding
    # and panel size by manipulating
    # the ny x nx grid.

See also §3.3.2.3 above, and Figure 3.6 for an example of channel averaging using iteration and subplot.

### 3.3.2.9 Averaging in plotxy

The averaging parameters and sub-parameters are:

```python
averagemode = ’vector’ # Select averaging type: vector, scalar
timebin = ’0’ # length of time in seconds to average, default=’0’, or: ’all’
crossscans = False # have time averaging cross over scans?
crossbls = False # have averaging cross over baselines?
crossarrays = False # have averaging cross over arrays?
stackspw = False # stack multiple spw on top of each other?
width = ’1’ # number of channels to average, default: ’1’, or: ’all’, ’allspw’
```

The choice of `averagemode` controls how the amplitudes are calculated in the average. The default mode is `’vector’`, where the complex average is formed by averaging the real and imaginary parts of the relevant visibilities. If `’scalar’` is chosen, then the amplitude of the average is formed by a scalar average of the individual visibility amplitudes.

Time averaging is effected by setting the `timebin` parameter to a value larger than the integration time. Currently, `timebin` takes a string containing the averaging time in seconds, e.g.

```python
timebin = ’60.0’
```

to plot one-minute averages.

Channel averaging is invoked by setting `width` to a value greater than 1. Currently, the averaging `width` is given as a number of channels.
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Figure 3.7: Multi-panel display of visibility versus channel (top), antenna array configuration (bottom left) and the resulting uv coverage (bottom right). The commands to make these three panels respectively are: 1) `plotxy('ngc5921.ms', xaxis='channel', datacolumn='data', field='0', subplot=211, plotcolor="", plotsymbol='go')` 2) `plotxy('ngc5921.ms', xaxis='x', field='0', subplot=223, plotsymbol='r.')`, 3) `plotxy('ngc5921.ms', xaxis='u', yaxis='v', field='0', subplot=224, plotsymbol='b', figfile='ngc5921_multiplot.png')`.

By default, the averaging will not cross scan boundaries (as set in the import process). However, if `crossscans=True`, then averaging will cross scans.

Note that data taken in different sub-arrays are never averaged together. Likewise, there is no way to plot data averaged over field.

### 3.3.2.10 Interactive Flagging in plotxy
Interactive flagging, on the principle of “see it — flag it”, is possible on the X-Y display of the data plotted by `plotxy`. The user can use the cursor to mark one or more regions, and then flag, unflag, or list the data that falls in these zones of the display.

There is a row of buttons below the plot in the window. You can punch the **Mark Region** button (which will appear to depress), then mark a region by left-clicking and dragging the mouse (each click and drag will mark an additional region). You can get rid of all your regions by clicking again on the **Mark Region** button (which will appear to un-depress), or you can use the **ESC** key to remove the last box you drew. Once regions are marked, you can then click on one of the other buttons to take action:

1. **Flag** — flag the points in the region(s),
2. **Unflag** — unflag flagged points in the region(s),
3. **Locate** — spew out a list of the points in the region(s) to the logger (Warning: this could be a long list!).

Whenever you click on a button, that action occurs without forcing a disk-write (unlike previous versions). If you quit `plotxy` and re-enter, you will see your previous edits.

![Figure 3.8: Plot of amplitude versus uv distance, before (left) and after (right) flagging two marked regions. The call was: `plotxy(vis='ngc5921.ms', xaxis='uvdist', field='1455*').`]

A table with the name `<msname>.flagversions` (where vis=<msname>) will be created in the same directory if it does not exist already.

It is recommended that you save important flagging stages using the **flagmanager** task (§ 3.2).
3.3.2.11 Flag extension in plotxy

Flag extension is controlled using extendflag=T and its sub-parameters:

```
extendflag = True # Have flagging extend to other data points?
extendcorr = '' # flagging correlation extension type
extendchan = '' # flagging channel extension type
extendspw = '' # flagging spectral window extension type
extendant = '' # flagging antenna extension type
extendtime = '' # flagging time extension type
```

The use of extendflag enables the user to plot a subset of the data and extend the flagging to a wider set.

**ALERT:** Using the extendflag options will greatly slow down the flagging in plotxy. You will see a long delay after hitting the Flag button, with lots of logger messages as it goes through each flag. Fixing this requires a refactoring of plotxy which is underway starting in Patch 4 development.

Setting extendchan='all' will extend the flagging to other channels in the same spw as the displayed point. For example, if spw='0:0' and channel 0 is displayed, then flagging will extend to all channels in spw 0.

The extendcorr sub-parameter will extend the flagging beyond the correlations displayed. If extendcorr='all', then all correlations will be flagged, e.g. with RR displayed RR,RL,LR,LL will be flagged. If extendcorr='half', then the extension will be to those correlations in common with that show, e.g. with RR displayed then RR,RL,LR will be flagged.

Setting extendspw='all' will extend the flagging to all other spw for the selection. Using the same example as above, with spw='0:0' displayed, then channel 0 in ALL spw will be flagged. Note that use of extendspw could result in unintended behavior if the spw have different numbers of channels, or if it is used in conjunction with extendchan.

**WARNING:** use of the following options, particularly in conjunction with other flag extensions, may lead to deletion of much more data than desired. Be careful!

Setting extendant='all' will extend the flagging to all baselines that have antennas in common with those displayed and marked. For example, if antenna='1&2' is shown, then ALL baselines to BOTH antennas 1 and 2 will be flagged. Currently, there is no option to extend the flag to ONLY baselines to the first (or second) antenna in a displayed pair.

Setting extendtime='all' will extend the flagging to all times matching the other selection or extension for the data in the marked region.

### 3.3.2.12 Setting rest frequencies in plotxy

The restfreq parameter can be set to a transition or frequency and expands to allow setting of frame information. For example,
restfreq = 'HI'  # a frequency quanta or transition name. see help for options
frame = 'LSRK'  # frequency frame for spectral axis. see help for options
doppler = 'RADIO'  # doppler mode. see help for options

Examples of transitions include:

```
restfreq='1420405751.786Hz'  # 21cm HI frequency
restfreq='HI'  # 21cm HI transition name
restfreq='115.2712GHz'  # CO 1-0 line frequency
```

For a list of known lines in the CASA measures system, use the toolkit command `me.linelist()`. For example:

```
CASA <14>: me.linelist()
H2CO HE110A HE138B HI OH1612 OH1665 OH1667 OH1720'
```

**ALERT:** The list of known lines in CASA is currently very restricted, and will be increased in upcoming releases (to include lines in ALMA bands for example).

You can use the `me.spectralline` tool method to turn transition names into frequencies

```
CASA <16>: me.spectralline('HI')
Out[17]:
{'m0': {'unit': 'Hz', 'value': 1420405751.786},
  'refer': 'REST',
  'type': 'frequency'}
```

(not necessary for this task, but possibly useful).

The `frame` sub-parameter sets the frequency frame. The allowed options can be listed using the `me.listcodes` method on the `me.frequency()` method, e.g.

```
CASA <17>: me.listcodes(me.frequency())
Out[17]:
{'extra': array([],
  dtype='|S1'),
  'normal': array(['REST', 'LSRK', 'LSRD', 'BARY', 'GEO', 'TOPO', 'GALACTO', 'LGROUP',
    'CMB'],
    dtype='|S8'))
```

The `doppler` sub-parameter likewise sets the Doppler system. The allowed codes can be listed using the `me.listcodes` method on the `me.doppler()` method,

```
CASA <18>: me.listcodes(me.doppler())
Out[18]:
{'extra': array([],
  dtype='|S1'),
  'normal': array(['RADIO', 'Z', 'RATIO', 'BETA', 'GAMMA', 'OPTICAL', 'TRUE',
    'RELATIVISTIC'],
    dtype='|S13'))
```
For most cases the 'RADIO' Doppler system is appropriate, but be aware of differences.
For more information on frequency frames and spectral coordinate systems, see the paper by Greisen et al. (A&A, 446, 747, 2006)\textsuperscript{2}

3.3.2.13 Printing from \texttt{plotxy}

There are two ways to get hardcopy plots in \texttt{plotxy}.

The first is to use the “disk save” icon from the interactive plot GUI to print the current plot. This will bring up a sub-menu GUI that will allow you to choose the filename and format. The allowed formats are \texttt{.png} (PNG), \texttt{.eps} (EPS), and \texttt{svg} (SVG). If you give the filename with a suffix (\texttt{.png}, \texttt{.eps}, or \texttt{svg}) it will make a plot of that type. Otherwise it will put a suffix on depending on the format chosen from the menu.

\textbf{ALERT:} The plot files produced by the EPS option can be large, and the SVG files can be very large. The PNG is the smallest.

The second is to specify a \texttt{figfile}. You probably want to disable the GUI using \texttt{interactive=False} in this case. The type of plot file that is made will depend upon the filename suffix. The allowed choices are \texttt{.png} (PNG), \texttt{.eps} (EPS), and \texttt{svg} (SVG).

This latter option is most useful from scripts. For example,

\begin{verbatim}
default('plotxy')
vis = 'ngc5921.ms'
field = '2'
spw = '',
xaxis = 'uvdist'
yaxis = 'amp'
interactive=False
figfile = 'ngc5921.uvplot.amp.png'
plotxy()
\end{verbatim}

will plot amplitude versus uv-distance in PNG format. No \texttt{plotxy} GUI will appear.

\textbf{ALERT:} if you use this option to print to \texttt{figfile} with an \texttt{iteration} set, you will only get the first plot.

3.3.2.14 Exiting \texttt{plotxy}

You can use the \texttt{Quit} button to clear the plot from the window and detach from the MS. You can also dismiss the window by killing it with the X on the frame, which will also detach the MS.

You can also just leave it alone. The plotter pretty much keeps running in the background even when it looks like it’s done! You can keep doing stuff in the plotter window, which is where the

\textsuperscript{2}Also at \url{http://www.aoc.nrao.edu/~egreisen/scs.ps}
overplot parameter comes in. Note that the \texttt{plotcal} task (§4.5.1) will use the same window, and can also overplot on the same panel.

If you leave \texttt{plotxy} running, beware of (for instance) deleting or writing over the MS without stopping. It may work from a memory version of the MS or crash.

### 3.3.2.15 Example session using \texttt{plotxy}

The following is an example of interactive plotting and flagging using \texttt{plotxy} on the Jupiter 6cm continuum VLA dataset. This is extracted from the script \texttt{jupiter6cm\_usecase.py} available in the script area.

This assumes that the MS \texttt{jupiter6cm\_usecase.ms} is on disk with \texttt{flagautocorr} already run.

```python
default('plotxy')
vis = 'jupiter6cm.usecase.ms'

# The fields we are interested in: 1331+305, JUPITER, 0137+331
selectdata = True

# First we do the primary calibrator
field = '1331+305'

# Plot only the RR and LL for now
correlation = 'RR LL'

# Plot amplitude vs. uvdist
xaxis = 'uvdist'
yaxis = 'amp'
multicolor = 'both'

# The easiest thing is to iterate over antennas
iteration = 'antenna'

plotxy()

# You'll see lots of low points as you step through RR LL RL LR
# A basic clip at 0.75 for RR LL and 0.055 for RL LR will work
# If you want to do this interactively, set
# iteration = ''

plotxy()

# You can also use flagdata to do this non-interactively
# (see below)

# Now look at the cross-polar products
correlation = 'RL LR'
```
# Now do calibrator 0137+331
field = '0137+331'
correlation = 'RR LL'
xaxis = 'uvdist'
spw = ''
iteration = ''
antenna = ''

plotxy()

# You’ll see a bunch of bad data along the bottom near zero amp
# Draw a box around some of it and use Locate
# Looks like much of it is Antenna 9 (ID=8) in spw=1

xaxis = 'time'
spw = '1'
correlation = ''

# Note that the strings like antenna='9' first try to match the
# NAME which we see in listobs was the number '9' for ID=8.
# So be careful here (why naming antennas as numbers is bad).
antenna = '9'

plotxy()

# YES! the last 4 scans are bad. Box ʼem and flag.

# Go back and clean up
xaxis = 'uvdist'
spw = ''
antenna = ''
correlation = 'RR LL'

plotxy()

# Box up the bad low points (basically a clip below 0.52) and flag

# Note that RL,LR are too weak to clip on.

# Finally, do JUPITER
field = 'JUPITER'
correlation = ''
iteration = ''
xaxis = 'time'

plotxy()}
# Here you will see that the final scan at 22:00:00 UT is bad
# Draw a box around it and flag it!

# Now look at what’s left
correlation = 'RR LL'
xaxis = 'uvdist'
spw = '1'
antenna = ''
iteration = 'antenna'

plotxy()

# As you step through, you will see that Antenna 9 (ID=8) is often
# bad in this spw. If you box and do Locate (or remember from
# 0137+331) it’s probably a bad time.

# The easiest way to kill it:

antenna = '9'
iteration = ''
xaxis = 'time'
correlation = ''

plotxy()

# Draw a box around all points in the last bad scans and flag ’em!

# Now clean up the rest
xaxis = 'uvdist'
correlation = 'RR LL'
antenna = ''
spw = ''

# You will be drawing many tiny boxes, so remember you can
# use the ESC key to get rid of the most recent box if you
# make a mistake.

plotxy()

# Note that the end result is we’ve flagged lots of points
# in RR and LL. We will rely upon imager to ignore the
# RL LR for points with RR LL flagged!

3.3.3 Plotting antenna positions using plotants

This task is a simple plotting interface (to the plotxy functionality) to produce plots of the antenna positions (taken from the ANTEANNA sub-table of the MS).
The inputs to plotants are:

```
# plotants :: Plot the antenna distribution in the local reference frame:
vis    = ''    # Name of input visibility file (MS)
figfile = ''    # Save the plotted figure to this file
```

### 3.3.4 Plotting uv-coverages plotuv

A simple way to plot uv-coverages is offered by the task plotuv:

```
# plotuv :: Plot the baseline distribution
vis    = ''    # Name of input visibility file (MS)
field  = ''    # Select field using ID(s) or name(s)
antenna = ''    # Select data based on antenna/baseline
spw    = ''    # Select spectral window/channels
observation = '' # Select by observation ID(s)
array  = ''    # Select (sub)array(s) by array ID number
maxnpts = 100000 # Maximum number of points per plot.
colors  = ['r', 'y', 'g', 'b'] # a list of matplotlib color codes
symb   = ','    # A matplotlib plot symbol code
ncycles = 1      # How many times to cycle through colors per plot.
figfile = ''    # Save the plotted figure(s) using this name
```

plotuv provides basic selection of data as well as plotting style options. The difference to plotms is that plotuv is also plotting the Hermitian conjugates of the visibilities which produces the familiar symmetric plots. This is a remedy to the restriction in plotms to allow flagging of data. This is achieved via a unambiguous link from a displayed data point to a visibility. Plotting Hermitian conjugates would break this rule in plotms and plotuv is used instead to plot Hermitian conjugates.

### 3.4 Data Flagging using flagdata

flagdata can flag measurement sets and calibration tables with an elaborate selection syntax. It also contains auto-flagging routines.

For a full description of flagdata please visit:

[http://www.aoc.nrao.edu/~rurvashi/FlaggerDocs/FlaggerDocs.html](http://www.aoc.nrao.edu/~rurvashi/FlaggerDocs/FlaggerDocs.html)

The inputs to flagdata are:

```
# flagdata :: All-purpose flagging task based on data-selections and flagging modes/algorithms.
vis    = ''    # Name of MS file or calibration table to flag
mode   = 'manual'    # Flagging mode
field  = ''    # Field names or field index
    # numbers: ''  ==> all, field='0^2,3C286'
```
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spw = '' # Spectral-window/frequency/channel: '' ==> all, spw='0:17~19'
atenna = '' # Antenna/baselines: '' ==> all, antenna = '3,VA04'
timerange = '' # Time range: '' ==> all, timerange='09:14:0~09:54:0'
correlation = '' # Correlation: '' ==> all, correlation='XX,YY'
scan = '' # Scan numbers: '' ==> all
intent = '' # Observation intent: '' ==> all, intent='CAL*POINT*'
array = '' # (Sub)array numbers: '' ==> all
uvrange = '' # UV range: '' ==> all;
               # uvrange = '0~100klambda', default units=meters
observation = '' # Observation ID: '' ==> all
feed = '' # Multi-feed numbers: Not yet implemented
autocorr = False # Flag auto-correlations

action = 'apply' # Action to perform in MS
          # and/or in inpfile (none/apply/calculate)
display = '' # Display data and/or
               # end-of-MS reports at runtime (data/report/both).
flagbackup = True # Back up the state of flags before the run

savepars = False # Save the current parameters
               # to the FLAG_CMD table or to a file

vis can take a measurement set or calibration table. Data selection for calibration tables is limited

by field, scan, time, antenna, spw, and observation. Since calibration tables do not have
a FLAG_CMD table, parameter settings, if requested, can only be saved in external files.

The mode parameter (§3.4.2) selects the flagging algorithm and the following are available:

list = list of flagging commands to apply to MS
manual = flagging based on specific selection parameters
clip = clip data according to values
quack = remove/keep specific time range at scan beginning/end
shadow = remove antenna-shadowed data
elevation = remove data below/above given elevations
tfcrop = automatic identification of outliers on the time-freq plane
rflag = automatic detection of outliers based on sliding-window RMS filters
extend = extend and/or grow flags beyond what the basic algorithms detect
summary = report the amount of flagged data
unflag = unflag the specified data

Flagging will only be applied to the data selection that is performed with the usual selection
parameters (§2.3). The dataset is iterated-through in chunks (small pieces of data) consisting
of one field, one spw, and a user-defined timerange (default is one scan). In addition to the
typical antenna, spw, timerange, etc. selections, we would like to point out some addition of the
correlation syntax for modes clip, tfcrop, and rflag. One can combine correlation products
with simple mathematical expressions

'ABS', 'ARG', 'RE', 'IM', 'NORM'
followed by the polarization products (using an underscore in between “_”)

'ALL', 'I', 'XX', 'YY', 'RR', 'LL', 'WVR'

'WVR' refers to the water vapour radiometer of ALMA data. Note that the operators ABS, ARG, RE, etc. are written only once as the first value. If more than one correlation is given, the operator will be applied to all of them. An example would be

correlation='RE_XX,XY'

which would select all real XX and XY polarization for flagging.

3.4.1 The action parameter

The keyword action controls whether the actual flagging commands will be applied or not and the options are the empty string ‘’, ‘apply’ and ‘calculate’. apply is likely the most popular one as it applies the flags to the MS:

```
action = 'apply'  # Action to perform in MS and/or in inpfile
display = ''  # Display data and/or end-of-MS reports at runtime
flagbackup = True  # Back up the state of flags before the run
```

flagbackup specifies if a backup of the current flags should be saved in the ‘*.flagversions’ file. display can be ‘’, ‘data’, ‘report’, ‘both’ where the empty string ‘’ will report no individual flagging statistics, whereas ‘data’ launches an interactive GUI to display data and flags for each chunk to browse through. The plots are time-frequency planes and both old and new flags are being overlaid for all correlations per baseline. In the GUI, one can step through all chunks for inspection and if the flagging is unsatisfactory, one can exit without applying the flags. If the flagging is acceptable, it is also possible to continue flagging without viewing all chunks (the number of chunks can be very large for typical JVLA and ALMA data sets. display='report' lists the flagging statistics at the end of the procedure on the screen and both starts the GUI and reports all statistics at the end.

action='calculate' calculates the flags but does not write them to the MS or calibration table. This is useful if one would like to inspect the computed flags in the GUI without a straight application:

```
action = 'calculate'  # Action to perform in MS and/or in inpfile
display = ''  # Display data and/or end-of-MS reports at runtime
```
The empty string action='' will do nothing and is useful when the commands themselves shall only be written to the FLAG_CMD sub-table or to an external file using the savepars parameter to specify the filename.

savepars will save the flagging commands to a file that can be later used for input in flagdata via mode='list'. It also shares the flagcmd syntax and can be used there. The file name is specified by outfile and, if empty, the FLAG_CMD table in the MS will be populated. A REASON can be given by the reason keyword which may be useful for bookkeeping as well as for unflagging data that are marked by specific REASON keywords.

3.4.2 Flagging Modes

3.4.2.1 Manual Flag/Unflag

mode = 'manual'  # Flagging mode (list/manual/clip/shadow/quack/...)
field = ''  # Field names or field index numbers: '' ==> all, field=0~2,3C286'
spw = ''  # Spectral-window/frequency/channel: '' ==> all, spw=0:17~19'
antenna = ''  # Antenna/baselines: '' ==> all, antenna = '3,VA04'
timerange = ''  # Time range: '' ==> all, timerange=09:14:0~09:54:0'
correlation = ''  # Correlation: '' ==> all, correlation='XX,YY'
scan = ''  # Scan numbers: '' ==> all
intent = ''  # Observation intent: '' ==> all, intent='CAL*POINT*,
array = ''  # (Sub)array numbers: '' ==> all
uvrange = ''  # UV range: '' ==> all; uvrange = 0~100klambda,
observation = ''  # Observation ID: '' ==> all
feed = ''  # Multi-feed numbers: Not yet implemented
autocorr = False  # Flag auto-correlations

The 'manual' mode is the most straight-forward of all modes. All visibilities that are selected by the various data selection parameters will be flagged or unflagged, depending on the action parameter. autocorr is a shorthand for antenna='*&&&' to flag all auto correlations in the data.

3.4.2.2 List

mode = 'list'  # Flagging mode (list/manual/clip/shadow/quack/...)
inpfile = ''  # Input ASCII file, list of files or Python list of strings with flag commands.
reason = 'any'  # Select by REASON types
A list of ag commands can be provided through a file or a list of files, specified by the `inpfile` parameter. Each input line may contain a flagging `mode` with data selection parameters as well as parameters that are specific to that `mode`. All parameters that are not set will be reset to their default values (default `mode` is `'manual'`). Each line of this file or list of strings will be taken as a command to the `flagdata` task. This `mode=list` is similar to the task `flagcmd` with the `inpmode=list` option.

An example for such a file would be:

```plaintext
mode='shadow'
mode='clip' clipminmax=[0,5] correlation='ABS_ALL'
mode='quack' quackmode='end' quackinterval=1.0
antenna='ea01' timerange='00:00:00~01:00:00'
antenna='ea11' timerange='00:00:00~03:00:00' spw='0~4'
```

Alternatively, this can be issued in the task directly like:

```plaintext
flagdata(vis='vis',mode='list',
    inpfile=["mode='shadow',
             "mode='clip' clipminmax=[0,5] correlation='ABS_ALL',
             "mode='quack' quackmode='end' quackinterval=1.0",
             "antenna='ea01' timerange='00:00:00~01:00:00',
             "antenna='ea11' timerange='00:00:00~03:00:00' spw='0~4'"])
```

or via a variable

```plaintext
cmds=["mode=shadow,
       "mode=clip clipminmax=[0,5] correlation=ABS_ALL",
       "mode=quack quackmode=end quackinterval=1.0",
       "antenna=ea01 timerange=00:00:00~01:00:00",
       "antenna=ea11 timerange=00:00:00~03:00:00 spw=0~4"]
```

```plaintext
flagdata(vis='vis',mode='list', inpfile=cmds)
```

The syntax needs to be written with quotes e.g. `mode=manual antenna=ea10`. There should be no space between `key=value`. Spaces are used to separate pairs of parameters, not commas.

### 3.4.2.3 Clip

```plaintext
mode = 'clip'  # Flagging mode (list/manual/clip/shadow/quack/
               # elevation/tfcrop/rflag/extend/unflag/summary)

... datacolumn = 'DATA'  # Data column on which to operate
                       # (data,corrected,model,residual)
clipminmax = []  # Range to use for clipping
clipoutside = True  # Clip outside the range, or within it
channelavg = False  # Average over channels (scalar average)
clipzeros = False  # Clip zero-value data
```
in addition to the regular selection parameters, `mode='clip'` also has an option to select between a
number of scratch columns in `datacolumn`. This includes the usual `DATA`, `CORRECTED`, etc., and also
clipping based on data weights `WEIGHT`, `WEIGHT_SPECTRUM` as well as other MS columns. `clipminmax`
selects the range of values to be clipped – usually this is combined with `clipoutside=True` to clip
everything but the values covered in `clipminmax`. The data can also be averaged over the selected
`spw` channel ranges by setting `channelavg=True`. `clip` will also flag ‘NaN’, ‘inf’, and ‘-inf’ values
by default and can flag exact zero values (these are sometimes produced by the JVLA correlator)
using the `clipzeros` parameter.

Note: For modes `clip`, `tfcrop` and `rflag`, channel-ranges can be excluded from flagging by
selecting ranges such as `spw='0:05;1063'`. This is a way to protect known spectral-lines from
being flagged by the autoflag algorithms.

3.4.2.4 Shadow

```python
mode = 'shadow' # Flagging mode (list/manual/clip/shadow/quack/
toleration = 0.0 # Amount of shadow allowed (in meters)
addantenna = '' # File name or dictionary with additional antenna names,
```

This option flags shadowed antennas, i.e. when one antenna blocks part of the aperture of a second
antenna that is behind the first one. Shadowing can be gradual and the criterion for a shadow flag
is when a baseline is shorter than `radius_1 + radius_2 – tolerance` (where the radii of the antennae
are taken from the MS antenna subtable); see Fig. 3.9. `addantenna` may be used to account for
shadowing when antennas are not listed in the MS but are physically present. Please read the
`flagdata` inline help for the syntax of this option.

![Figure 3.9: This figure shows the geometry used to compute shadowed antennas.](image)

3.4.2.5 Quack

```python
mode = 'quack' # Flagging mode (list/manual/clip/shadow/quack/
```
quack is used to remove data at scan boundaries. quackinterval specifies the time in seconds to be flagged, and quackmode can be 'beg' to flag the quackinterval at the beginning of each selected scan, 'endb' at the end of scan. 'tail' flags all but the beginning of scan and 'end' all but the end of scan. The quackincrement is either True or False, depending if one wishes to flag the quackinterval from the first unflagged data in the scan, or from the scan boundaries independent of data being already flagged or not.

3.4.2.6 Elevation

mode = 'elevation' # Flagging mode (list/manual/clip/shadow/quack/ # elevation/tfcrop/rflag/extend/unflag/summary)
...
lowerlimit = 0.0 # Lower limiting elevation (in degrees)
upperlimit = 90.0 # Upper limiting elevation (in degrees)

Flagging based on the elevation of the antennae. This may be useful to avoid data taken at very low elevations or close to transit and the lowerlimit and upperlimit parameters specify the range of good elevations.

3.4.2.7 Tfcrop

mode = 'tfcrop' # Flagging mode (list/manual/clip/shadow/quack/ # elevation/tfcrop/rflag/extend/unflag/summary # )
...
ntime = 'scan' # Time-range to use for each chunk (in seconds # or minutes)
combinescans = False # Accumulate data across scans.
datacolumn = 'DATA' # Data column on which to operate # (data,corrected,model,residual)
timecutoff = 4.0 # Flagging thresholds in units of deviation # from the fit
freccutoff = 3.0 # Flagging thresholds in units of deviation # from the fit
timefit = 'line' # Fitting function for the time direction # (poly/line)
TFCrop is an autoflag algorithm that detects outliers on the 2D time-frequency plane, and can operate on un-calibrated data (non bandpass-corrected). The original implementation of this algorithm is described in NCRA Technical Report 202 (Oct 2003).

The algorithm iterates through the data in chunks of time. For each chunk, the result of user-specified visibility-expressions are organized as 2D time-frequency planes, one for each baseline and correlation-expression result, and the following steps are performed.

1. Calculate a bandshape template: Average the data across time, to construct an average bandpass. Construct an estimate of a clean bandpass (without RFI) via a robust piece-wise polynomial fit to the average bandpass shape.

   Note: A robust fit is computed in up to 5 iterations. It begins with a straight line fit across the full range, and gradually increases to 'maxnpieces' number of pieces with third-order polynomials in each piece. At each iteration, the stddev between the data and the fit is computed, values beyond N-stddev are flagged, and the fit and stddev are re-calculated with the remaining points. This stddev calculation is adaptive, and converges to a value that reflects only the data and no RFI. At each iteration, the same relative threshold is applied to detect flags, and this results in a varying set of flagging thresholds, that allows deep flagging only when the fit represents the true data best. Iterations stop when the stddev changes by less than 10%, or when 5 iterations are completed.

   The resulting clean bandpass is a fit across the base of RFI spikes.

2. Divide out this clean bandpass function from all timesteps in the current chunk. Now, any data points that deviate from a mean of 1 can be considered RFI. This step helps to separate narrow-band RFI spikes from a smooth but varying bandpass, in situations where a simple range-based clipping will flag good sections of the bandpass.

3. Perform iterative flagging (robust flagging) of points deviating from a value of 1.

   Flagging is done in up to 5 iterations. In each iteration, for every timestep, calculate the stddev of the bandpass-flattened data, flag all points further than N times stddev from the fit, and recalculate the stddev. At each iteration, the same relative threshold is applied to detect flags. Optionally, use sliding-window based statistics to calculate additional flags.
4. Repeat steps 1 and 3, but in the other direction (i.e. average the data across frequency, calculate a piece-wise polynomial fit to the average time-series, and find flags based on deviations w.r.to this fit.)

The default parameters of the tfcrop implementation are optimized for strong narrow-band RFI (see, e.g. Fig. 3.10). With broad-band RFI, the piece-wise polynomial can sometimes model it as part of the band-shape, and therefore not detect it as RFI. In this case, reducing the maximum number of pieces in the polynomial can help. This algorithm usually has trouble with noisy RFI that is also extended in time of frequency, and additional statistics-based flagging is recommended (via the 'usewindowstats' parameter). It is often required to set up parameters separately for each spectral-window.

If frequency ranges of known astronomical spectral lines are known a-priori, they can be protected from automatic flagging by de-selecting those frequency-ranges via the 'spw' data-selection parameter.

The extendflag parameter will clean up small portions of data between flagged data points along time and/or frequency when more than 50% of all timeranges or 80% of all channels are already flagged. It will also extend the flags to the other polarizations. Alternatively, mode='extend' can be used (Fig. 3.12).

![Figure 3.10: This screenshot represents a run where 'tfcrop' was run on a spw='9' with mainly narrow-band RFI. RIGHT : An example of protecting a spectral line (in this case, demonstrated on an RFI spike) by setting the spw-selection to spw='0:0 45;53 63'. In both figures, the top row indicates the data before flagging, and the bottom row after flagging.](image)

3.4.2.8 Rflag
RFlag is an autoflag algorithm based on a sliding window statistical filter. The RFlag algorithm was originally developed by Eric Greisen in AIPS (31DEC11). AIPS documentation: Subsection E.5 of the AIPS cookbook (Appendix E: Special Considerations for JVLA data calibration and imaging in AIPS).

In RFlag, the data is iterated-through in chunks of time, statistics are accumulated across time-chunks, thresholds are calculated at the end, and applied during a second pass through the dataset.

The CASA implementation also optionally allows a single-pass operation where statistics and thresholds are computed and also used for flagging, within each time-chunk (defined by 'ntime' and 'combinescans').

For each chunk, calculate local statistics, and apply flags based on user supplied (or auto-calculated) thresholds.

1. Time analysis (for each channel)
   
   (a) Calculate local rms of real and imag visibilities, within a sliding time window
   
   (b) Calculate the median rms across time windows, deviations of local rms from this median, and the median deviation
   
   (c) Flag if local rms is larger than timedevscale x (medianRMS + medianDev)

2. Spectral analysis (for each time)
   
   (a) Calculate avg of real and imag visibilities and their rms across channels

http://www.aips.nrao.edu/cook.html#CEE
(b) Calculate the deviation of each channel from this avg, and the median-deviation
(c) Flag if deviation is larger than freqdevscale x medianDev

The extendflag parameter will clean up small portions of data between flagged data points along
time and/or frequency when more than 50% of all timeranges or 80% of all channels are already
flagged. It will also extend the flags to the other polarizations. Alternatively, mode='extend' can
be used.

Some examples (also see Fig.3.11):

1. Calculate thresholds automatically per scan, and use them to find flags. Specify scale-factor
   for time-analysis thresholds, use default for frequency.

   flagdata('my.ms', mode='rflag', spw='9', timedevscale=4.0, writeflags=True)

2. Supply noise-estimates to be used with default scale-factors.

   flagdata(vis='my.ms', mode='rflag', spw='9', timedev=0.1, freqdev=0.5, writeflags=True);

Two-passes. This replicates the usage pattern in AIPS.

- The first pass saves commands in an output text files, with auto-calculated thresholds. Thresholds are returned from rflag only when writeflags=False (calc-only mode). The user can edit this file before doing the second pass, but the python-dictionary structure must be preserved.
- The second pass applies these commands (writeflags=True).

   flagdata(vis='my.ms', mode='rflag', spw='9,10',
            timedev='tdevfile.txt', freqdev='fdevfile.txt', writeflags=False);
   flagdata(vis='my.ms', mode='rflag', spw='9,10',
            timedev='tdevfile.txt', freqdev='fdevfile.txt', writeflags=True);

### 3.4.2.9 Extend

```python
mode = 'extend' # Flagging mode (list/manual/clip/shadow/quack/el
evate/tfcrop/rflag/extend/unflag/summary)
field = '' # Field names or field index numbers: '' ==> all,
# field='0~2,3C286'
spw = '' # Spectral-window/frequency/channel: '' ==> all,
# spw='0:17~19'
antenna = '' # Antenna/baselines: '' ==> all, antenna
# = '3,VA04'
timerange = '' # Time range: '' ==>
# all,timerange='09:14:0~09:54:0'
correlation = '' # Correlation: '' ==> all, correlation='XX,YY'
```
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![Figure 3.11: Example of rflag on narrow-band RFI](image)

Although the modes `tfcrop` and `rflag` already have `extendflag` parameters, some autoflagging algorithms may still leave small islands of unflagged data behind, data that are surrounded by flagged visibilities in the time-frequency space. Although the algorithm may deem these visibilities as good ones, they are frequently affected by low-level RFI that spills from the adjacent, flagged points and one may wish to clean those up.
ntime specifies the time ranges over which to clean up, e.g. ‘1.5min’ or ‘scan’ which checks on all data within a scan. To span time ranges larger than scans, one can set combinescans to True.

extendpols=True would extend all flags to all polarization products when at least one of them is flagged.

growtime flags the entire time range for a flagged channel, when a certain fraction of flagged time intervals is exceeded.

growfreq is similar but extends the flags in frequency when a given fraction of channels is already flagged.

growaround checks for flagged data points in the time-frequency domain that neighbor a datum. The threshold is four data points. If more surrounding points are flagged, the central datum will be flagged, too.

flagneartime flags adjacent data points along the time axis, around a flagged datum

flagnearfreq flags neighboring channels.

For an example, see Fig. 3.12

Figure 3.12: This screenshot represents a run where 'tfcrop' was run only on 'ABS_RR' (top row) and followed by an extension along time and correlations (bottom row).

3.4.2.10 Unflag

mode = 'unflag'  # Flagging mode (list/manual/clip/shadow/quack/
#           elevation/tfcrop/rflag/extend/unflag/summary
# )
field = '' # Field names or field index numbers: ''==>all, field='0~2,3C286'
spw = '' # spectral-window/frequency/channel antenna = 'ea01' # antenna/baselines: ''==>all, antenna 
# = '3,VA04'timerange = '' # time range:
correlation = '' # Select data based on correlation scan = '' # scan numbers: ''==>all intent = '' # Select data based on observation intent:
feed = '' # multi-feed numbers: Not yet implemented array = '' # (sub)array numbers: ''==>all uvr = False # uv range: ''==>all; uvr = '0~100klambda', basecnt = False # Print summary counts per baseline

The selection data will be unflagged.

### 3.4.2.11 Summary

mode = 'summary' # Flagging mode (list/manual/clip/shadow/quack/ 
# elevation/tfcrop/rflag/extend/unflag/summary 
# )

... 

minrel = 0.0 # minimum number of flags (relative)
maxrel = 1.0 # maximum number of flags (relative)
minabs = 0 # minimum number of flags (absolute)
maxabs = -1 # maximum number of flags (absolute). Use a 
# negative value to indicate infinity.
spwchan = False # Print summary of channels per spw 
spwcorr = False # Print summary of correlation per spw 
basecnt = False # Print summary counts per baseline

This mode reports the number of rows and data points that are flagged. The selection of reported 
points can be restricted (see inline help for details).

mode='summary' can also report back a dictionary if the task is run as 

```
s = flagdata(..., mode='summary')
```

with a variable assigned, here 's'.

### 3.5 Command-based flagging using flagcmd

The task flagcmd will flag the visibility data set or calibration table based on a specified set of 
flagging commands using a flagging syntax (see § 3.5.3). These commands can be input from the
FLAG_CMD MS table, from a Flag.xml SDM table, from an ascii file, or from input python strings. Facilities for manipulation, listing, or plotting of these flags are also provided.

The inputs to flagcmd are:

```python
# flagcmd :: Flagging task based on batches of flag-commands
vis = ''  # Name of MS file or calibration table to flag
inpmode = 'table'  # Input mode for flag commands(table/list/xml)
inpfie = ''  # Source of flag commands
tablrows = []  # Rows of inpfile to read
reason = 'any'  # Select by REASON types
useapplied = False  # Select commands whose rows
                    # have APPLIED column set to True
action = 'apply'  # Action to perform in MS and/or in inpfile
flagbackup = True  # Automatically backup the
                    # FLAG column before execution
savepars = False  # Save flag commands to the MS or to a file
```

The default input mode is `inpmode='table'` which directs the task to input flag commands from the FLAG_CMD internal MS table. See §3.5.1 for more options.

The default operation mode is `action='apply'` directing the task to apply relevant flagging commands to the MS data main table. See §3.5.2 for more options.

See §3.5.3 for a description of the flagging command syntax.

It is possible to flag calibration tables using flagcmd, although we recommend using the flagdata task for this.

When using flagcmd to flag calibration tables, only the `apply` and `list` actions are supported. Because calibration tables do not have a FLAG_CMD sub-table, the default `inpmode='table'` can only be used if an MS is given in the `inpfile` parameter so that flags from the MS are applied to the calibration table directly. Otherwise, the flag commands must be given using `inpmode='list'`, either from a file or from a list of strings.

### 3.5.1 Input modes inpmode

The `inpmode` parameter selects options for the input mode for the flagging commands.

Available `inpmode` options are:

- `'table'` — input from MS table (§3.5.1.1)
- `'list'` — input from ASCII file or from a list of strings (§3.5.1.2)
- `'xml'` — input from XML table (§3.5.1.3)
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3.5.1.1 Input mode 'table'

The default input mode is inpmode='table' which directs the task to input flag commands from a FLAG_CMD MS table. This has the sub-parameters:

```python
inpmode = 'table' # Input mode for flag commands(table/list/xml)
inpfie = '' # Source of flag commands
tablerows = [] # Rows of inpfile to read
reason = 'any' # Select by REASON types
useapplied = False # Select commands whose rows have APPLIED column set to True
```

If inpfile = '' then it will look for the FLAG_CMD table in the MS given by vis. You can use this sub-parameter to direct the task to look directly at another table.

The tablerows sub-parameter is a simple Python list of the row numbers of the table to consider in processing flags. The default is all rows.

The useapplied sub-parameter toggles whether only flag commands marked as not having been applied are considered (the default), or to allow (re)processing using all commands.

The reason sub-parameter selects the REASON type to process. The default 'any' means all commands, note that reason='' would only select flags who have a blank REASON column entry.

One use case is to read the flag commands from the FLAG_CMD of an MS and apply them to a calibration table given in the parameter vis. Example:

```python
flagcmd(vis='cal-X54.B1', inpmode='table',
inpfie='uid___A002_X2a5c2f_X54.ms', action='apply')
```

3.5.1.2 Input flag mode 'list'

This mode allows one to insert a list of strings with flagging commands, the name of a le or a list of filenames that contains these commands equivalent to the mode='list' in flagdata (§3.4.2.2).

E.g. a file flags.txt that contains

scan='1~3' mode='manual'
mode='clip' clipminmax=[0,2] correlation='ABS_XX' clipoutside=False
spw='9' mode='tfcrop' correlation='ABS_YY' ntime=51.0
mode='extend' extendpols=True

can be called via

```python
flagcmd(vis, inpmode='list', inpfile='flags.txt')
```

Alternatively, the individual flagging commands can be directly provided in the call itself like
CHAPTER 3. DATA EXAMINATION AND EDITING

inpfile= ["scan='1~3' mode='manual'",
          "mode='clip' clipminmax=[0,2] correlation='ABS_XX' clipoutside=False",
          "spw='9' mode='tfcrop' correlation='ABS_YY' ntime=51.0",
          "mode='extend' extendpols=True"]

3.5.1.3 Input flag mode ‘xml’

The input mode inpmode='xml' directs the task to input flag commands from a XML SDM online flagging Flag.xml file. When set this opens the sub-parameters:

inpmode = 'xml' # Input mode for flag commands(table/list/xml)
            tbuff = 0.0 # Time buffer (sec) to pad flags
            ants ='' # Allowed flag antenna names to select by
            reason = 'any' # Select by REASON types

This mode will look for a file called Flag.xml inside the MS directory specified under vis. Note that if the data was filled from the SDM using importevla ([2.2.2]) then the relevant XML file will have been copied to the MS already.

The tbuff sub-parameter sets a padding buffer (in seconds) to the begin and end times of the online flags in the XML file. As in importevla, the online flag time buffer tbuff is specified in seconds, but in fact should be keyed to the intrinsic online integration time to allow for events (like slewing) that occur within an integration period. This is particularly true for JVLA data, where a tbuff value of 0.5× to 1.5× the integration time is needed. For example, if data were taken with 1-second integrations, then at least tbuff=0.5 should be used, likewise tbuff=5 for 10-second integrations. Note: For JVLA data you should use 1.5× (e.g. tbuff=15 for 10-second integrations) for data taken in early 2011 or before due to a timing error. We do not yet know what ALMA data will need for padding (if any).

The ants sub-parameter selects the antennas from which online flags will be selected (default is all antennas). For example, ants='ea01' is a valid choice for JVLA data.

The reason sub-parameter selects by the REASON field in the Flag.xml file. The default 'any' means all commands. Note that reason='' would only select flags who have a blank REASON field entry.

3.5.2 Operation types action

The action selects options for operating on the selected flags and possibly the data.

Available action options are:

- 'apply' — apply flag commands to data ([3.5.2.1])
- 'unapply' — unapply flags in data ([3.5.2.2])
- 'list' — list and/or save flag commands ([3.5.2.3])
• 'plot’ — plot flag commands (§ 3.5.2.4)
• 'clear’ — clear rows from FLAG_CMD table (§ 3.5.2.5)
• 'extract’ — extract internal flag dictionary (§ 3.5.2.6)

3.5.2.1 Apply flags — optype option 'apply’

The default operation mode is action='apply’ directing the task to apply relevant flagging commands to the vis data main table.

```python
action = 'apply'  # Action to perform in MS and/or in inpfile
flagbackup = True  # Automatically backup the FLAG column before execution
```

The flagbackup toggle sets whether a new copy of the MS main table FLAG column is written to the .flagversions backup directory for that MS before the requested flagging operation.

3.5.2.2 Unapply flags — action option 'unapply’

The unapply option allows unflagging of data based on the selected flag commands. This choice opens the sub-parameters:

```python
action = 'unapply'  # Action to perform in MS and/or in inpfile
flagbackup = True  # Automatically backup the FLAG column before execution
```

As in action='apply’, it is possible to make a backup to the *.flagversions file by using flagbackup=True.

In order to guarantee that only the data selected in the command is unapplied, the framework will first unapply the selected rows and then re-apply the overlapping data that got unapplied in the first pass. This is a true unapply action, but it will take longer to process because it will re-apply all the remaining commands that have APPLIED = True!

3.5.2.3 List flags — action='list’

The 'list’ option will give a listing of the flagging commands. This choice opens the sub-parameters:
action = 'list' # Action to perform in MS and/or in inpfile
savepars = True # Save flag commands to the MS or to a file
outfile = '' # Name of output file to save commands

This action lists the commands on the screen without applying them. One can save the flagging script to a file specified in the outfile parameter when savepars=True. If outfile is empty, it will save the commands to the MS given in vis.

The format of the listing output depends on the source of the flagging commands. A set of flagging commands specified through inpmode='list' will be listed directly. The flagging commands extracted through inpmode='table' will reflect the columns in the table:

'Row', 'Timerange', 'Reason', 'Type', 'Applied', 'Lev', 'Sev', 'Command'

while commands from inpmode='xml' will be shown with the SDM XML table fields:

'Key', 'FlagID', 'Antenna', 'Reason', 'Timerange'

### 3.5.2.4 Plot flags — action='plot'

The 'plot' option will produce a graphical plot of flags of time versus antenna. This choice opens the sub-parameters:

action = 'plot' # Action to perform in MS and/or in inpfile
plotfile = '' # Name of output file to save plot

This is only useful for online flags or general flag commands that are specified by antenna plus timerange using the standard REASON codes that are known SDM Flag.xml enumerations.

If the plotfile sub-parameter is non-blank, then a plotfile will be made with that name instead of appearing in a matplotlib plotter window on the users workstation.

**ALERT:** The plotted enumerations are currently only those known to be allowed JVLA online flags as of 15 April 2011, and include:

'FOCUS', 'SUBREFLECTOR', 'OFF SOURCE', 'NOT IN SUBARRAY'

with all others being plotted as 'Other'.
3.5.2.5 Clear flags — action='clear'  

The 'clear' action will delete selected rows from the FLAG_CMD MS table. This choice opens the sub-parameters:

\[
\begin{align*}
\text{action} & = \text{'clear'} \quad \# \text{ Action to perform in MS and/or in inpfile} \\
\text{clearall} & = \text{False} \quad \# \text{ Delete all rows from FLAG_CMD} \\
\text{rowlist} & = \text{[]} \quad \# \text{ FLAG_CMD rows to clear}
\end{align*}
\]

The rowlist sub-parameter is a simple Python list of the row numbers of the table to consider in processing flags. The default is a blank list which indicates the desire to clear all rows.

In either case, if clearall=False then nothing will happen by default as a safeguard. If clearall=True, then a blank list will direct the deletion of the selected rows from the table.

**ALERT:** Use this option with care. You can easily mess up the FLAG_CMD table.

3.5.2.6 Extract Flag Commands— action='extract'  

The 'extract' option will return the internal flagging dictionary to python:

\[
\begin{align*}
\text{action} & = \text{'extract'} \quad \# \text{ Action to perform in MS and/or in inpfile} \\
\end{align*}
\]

The value can be returned to a variable like:

\[
\text{myflagd} = \text{flagcmd(vis=msfile,useapplied=True,action='extract')}
\]

3.5.3 Flagging command syntax  

A flagging command syntax has been devised to populate the COMMAND column of the FLAG_CMD table and to direct the operation of the flagcmd task.

The syntax is similar to flagdata, so please check help flagdata for more info.

You can also use help flagcmd inside casapy for this syntax guide also.

Commands are a string (which may contain internal "strings") consisting of KEY=VALUE pairs separated by whitespace (see examples below).

**NOTE:** There should be no whitespace between KEY=VALUE or within each KEY or VALUE, since the simple parser first breaks command lines on whitespace, then on "=".

Each key should only appear once on a given command line/string

There is an implicit "mode" for each command, with the default being 'manual' if not given.

Comment lines can start with '#' and will be ignored.
1. Data selection parameters (used by all flagging modes, see also §2.3)

```
timerange='',
antenna='',
spw='',
correlation='',
field='',
scan='',
feed='',
array='',
uvrange='',
intent='',
observation='',
```

Note: a command consisting only of selection key-value pairs is a basic "manual" operation, i.e. flag the data meeting the selection.

2. Modes specific parameters with default values (for further details, refer to the task flagdata, §3.4.2).

(a) Mode manual

```
autocorr=False
```

(b) Mode clip

(c) Mode manual

```
datacolumn='DATA'
clipminmax=[]
clipoutside=True
channelavg=False
clipzeros=False
```

(d) Mode shadow

```
tolerance=0.0
addantenna=''
```

(e) Mode quack

```
quackinterval=0.0
quackmode='beg'
quackincrement=False
```

(f) Mode elevation

```
lowerlimit=0.0
upperlimit=90.0
```

(g) Mode tfcrop

```
time='scan'
combine-scans=False
datacolumn='DATA'
timecutoff=4.0
```
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freqcutoff=3.0
timefit='line'
freqfit='poly'
maxnpieces=7
flagdimension='freqtime'
usewindowstats='none'
halfwin=1

(h) Mode extend

ntime='scan'
combinescans=False
extendpols=True
growtime=50.0
growfreq=50.0
growaround=False
flagneartime=False
flagnearfreq=False

(i) Mode rflag

ntime='scan'
combinescans=False
datacolumn='DATA'
winsize=3
timedev='',
freqdev='',
timedevscale=5.0
freqdevscale=5.0
spectralmax=1000000.0
spectralmin=0.0

(j) Mode unflag

3. Basic elaboration options for online and interface use

id=',' # flag ID tag (not necessary)
reason=',' # reason string for flag
flagtime=',' # a timestamp for when this flag was generated (for user history use)

NOTE: there is no flagtime column in FLAG_CMD at this time, but we will propose to add this as an optional column

NOTE: These are currently ignored and not used

4. Extended elaboration options for online and interface use Note: these are FLAG_CMD columns, but their use is not clear but included here for compatibility and future expansion

level=N # flagging "level" for flags with same reason
severity=N # Severity code for the flag, on a scale of 0-10 in order of increasing severity; user specified
3.6 Browse the Data

The `browsetable` task is available for viewing data directly (and handles all CASA tables, including Measurement Sets, calibration tables, and images). This task brings up the CASA Qt `casabrowser`, which is a separate program. You can launch this from outside `casapy`.

The default inputs are:

```plaintext
# browsetable :: Browse a table (MS, calibration table, image)

tablename = '' # Name of input table
```

Currently, its single input is the `tablename`, so an example would be:

```plaintext
browsetable('ngc5921.ms')
```

For an MS such as this, it will come up with a browser of the `MAIN` table (see Fig 3.13). If you want to look at sub-tables, use the tab `table keywords` along the left side to bring up a panel with the sub-tables listed (Fig 3.14), then choose (left-click) a table and `View:Details` to bring it up (Fig 3.15). You can left-click on a cell in a table to view the contents.

Note that one useful feature is that you can Edit the table and its contents. Use the `Edit table` choice from the `Edit` menu, or click on the `Edit` button. Be careful with this, and make a backup copy of the table before editing!

Use the `Close Tables and Exit` option from the `Files` menu to quit the `casabrowser`.

There are a lot of features in the `casabrowser` that are not fully documented here. Feel free to explore the capabilities such as plotting and sorting!

**ALERT:** You are likely to find that the `casabrowser` needs to get a table lock before proceeding. Use the `clearstat` command to clear the lock status in this case.
Figure 3.13: **browsetable**: The browser displays the main table within a frame. You can scroll through the data (x=columns of the MAIN table, and y=the rows) or select a specific page or row as desired. By default, 1000 rows of the table are loaded at a time, but you can step through the MS in batches.
Figure 3.14: **browseTable**: You can use the tab for **Table Keywords** to look at other tables within an MS. You can then double-click on a table to view its contents.
Figure 3.15: **browsetable**: Viewing the SOURCE table of the MS.
Chapter 4

Synthesis Calibration

This chapter explains how to calibrate interferometer data within the CASA task system. Calibration is the process of determining the net complex correction factors that must be applied to each visibility in order to make them as close as possible to what an idealized interferometer would measure, such that when the data is imaged an accurate picture of the sky is obtained. This is not an arbitrary process, and there is a philosophy behind the CASA calibration methodology (see §4.2.1 for more on this). For the most part, calibration in CASA using the tasks is not too different than calibration in other packages such as AIPS or Miriad, so the user should not be alarmed by cosmetic differences such as task and parameter names!

4.1 Calibration Tasks

Alert: The calibration table format changed in CASA 3.4. CASA 4.2 is the last version that will support the caltabconvert function that provides conversions from the pre-3.4 caltable format to the modern format; it will be removed for CASA 4.3. In general, it is best to recalculate calibration using CASA 3.4 or later.

Alert: In CASA 4.2 the gaincurve and opacity parameters have been removed from all calibration tasks (as advertised in 4.1). These calibration types are supported via the gencal task.

Alert: As part of continuing development of a more flexible and improved interface for specifying calibration for apply, a new parameter has been introduced in applycal and the solving tasks: docallib. This parameter toggles between use of the traditional calibration apply parameters (gaintable, gainfield, interp, spwmap, and calwt), and a new callib parameter which currently provides access to the experimental cal library mechanism, wherein calibration instructions are stored in a file. The default is docallib=False in 4.3, and this reveals the traditional apply parameters which continue to work as always. Since callib is still under development and testing, and does not yet support all calibration types, general users should continue to use the traditional parameters via docallib=False. The rest of this chapter is written assuming docallib=False.
The standard set of calibration solving tasks (to produce calibration tables) are:

- **bandpass** — complex bandpass (B) calibration solving, including options for channel-binned or polynomial solutions (§ 4.4.2),
- **gaincal** — complex gain (G,T) and delay (K) calibration solving, including options for time-binned or spline solutions. (§ 4.4.3),
- **polcal** — polarization calibration including leakage and angle (§ 4.4.5),
- **blcal** — baseline-based complex gain or bandpass calibration (§ 4.4.6).

There are helper tasks to create, manipulate, and explore calibration tables:

- **accum** — Accumulate incremental calibration solutions into a cumulative cal table (§ 4.5.5) (ALERT: The accum task is generally no longer recommended for most calibration scenarios. Please write to the NRAO CASA helpdesk if you need support using accum.),
- **applycal** — Apply calculated calibration solutions (§ 4.6.1),
- **clearcal** — Re-initialize the calibration for a visibility dataset (§ 4.6.3),
- **fluxscale** — Bootstrap the flux density scale from standard calibration sources (§ 4.4.4),
- **listcal** — List calibration solutions (§ 4.5.2),
- **plotcal** — Plot calibration solutions (§ 4.5.1),
- **setjy** — Compute model visibilities with the correct flux density for a specified source (§ 4.3.5),
- **smoothcal** — Smooth calibration solutions derived from one or more sources (§ 4.5.4),
- **split** — Write out new MS containing calibrated data from a subset of the original MS (§ 4.7.1).

There are some development versions of calibration and utility tasks that are recently added to the suite:

- **calstat** — Statistics of calibration solutions (§ 4.5.3),
- **cvel** — Regrid a spectral MS onto a new frequency channel system (§ 4.7.7),
- **gencal** — Create a calibration tables from metadata such as antenna position offsets, gain-curves and opacities (§ 4.3.6),
- **wvrgecal** — Generate a gain table based on Water Vapour Radiometer data (for ALMA use - § 4.3.8),
- **hanningsmooth** — Apply a Hanning smoothing filter to spectral-line uv data (§ 4.7.3),
CHAPTER 4. SYNTHESIS CALIBRATION

- **mstransform** — *experimental* Task to combine *cvel*, *hanningsmooth*, *split* operations in a single step (§4.7.4).
- **uvcontsub** — Carry out uv-plane continuum fitting and subtraction (§4.7.6).
- **uvmodelfit** — Fit a component source model to the uv data (§4.7.8).
- **uvsub** — Subtract the transform of a model image from the uv data (§4.7.5).
- **statwt** — Recalculate the data weights based on their scatter (§4.7.9).
- **conjugatevis** — Change the signs of visibility phases (§4.7.10).

These are not yet full-featured, and may have only rudimentary controls and options.

The following sections outline the use of these tasks in standard calibration processes.

Information on other useful tasks and parameter setting can be found in:

- **listobs** — summary of a MS (§2.2.7).
- **listvis** — list data in a MS (§2.2.9).
- **plotms** — prototype next-generation X-Y plotting and editing (§3.3.1).
- **plotxy** — previous generation X-Y plotting and editing (§3.3.2).
- **plotweather** — plot the weather information of an MS and calculate atmospheric opacities (§4.3.4.1).
- **flagdata** — non-interactive data flagging (§3.4).
- **data selection** — general data selection syntax (§2.3).

4.2 The Calibration Process — Outline and Philosophy

A work-flow diagram for CASA calibration of interferometry data is shown in Figure 4.1. This should help you chart your course through the complex set of calibration steps. In the following sections, we will detail the steps themselves and explain how to run the necessary tasks and tools.

This can be broken down into a number of discrete phases:

- **Calibrator Model Visibility Specification** — set model visibilities for calibrators, either unit point source visibilities for calibrators with unknown flux density or structure (generally, sources used for calibrators are approximately point-like), or visibilities derived from *a priori* images and/or known or standard flux density values.
Figure 4.1: Flow chart of synthesis calibration operations. Not shown are use of table manipulation and plotting tasks `accum`, `plotcal`, and `smoothcal` (see Figure 4.2).
• **Prior Calibration** — set up previously known calibration quantities that need to be pre-applied, such as antenna gain-elevation curves, atmospheric models, delays, and antenna position offsets. Use the `setjy` task (§4.3.5) for calibrator flux densities and models, and use `gencal` (§4.3.6) for antenna position offsets, gain curves, antenna efficiencies, and opacities;

• **Bandpass Calibration** — solve for the relative gain of the system over the frequency channels in the dataset (if needed), having pre-applied the prior calibration. Use the `bandpass` task (§4.4.2);

• **Gain Calibration** — solve for the gain variations of the system as a function of time, having pre-applied the bandpass (if needed) and prior calibration. Use the `gaincal` task (§4.4.3);

• **Polarization Calibration** — solve for polarization leakage terms and linear polarization position angle (§4.4.5);

• **Establish Flux Density Scale** — if only some of the calibrators have known flux densities, then rescale gain solutions and derive flux densities of secondary calibrators. Use the `fluxscale` task (§4.4.4);

• **Manipulate, Accumulate, and Iterate** — if necessary, accumulate different calibration solutions (tables), smooth, and interpolate/extrapolate onto different sources, bands, and times. Use the `accum` (§4.5.5) and `smoothcal` (§4.5.4) tasks;

• **Examine Calibration** — at any point, you can (and should) use `plotcal` (§4.5.1) and/or `listcal` (§4.5.2) to look at the calibration tables that you have created;

• **Apply Calibration to the Data** — this can be forced explicitly by using the `applycal` task (§4.6.1), and can be undone using `clearcal` (§4.6.3);

• **Post-Calibration Activities** — this includes the determination and subtraction of continuum signal from line data, the splitting of data-sets into subsets (usually single-source), and other operations (such as model-fitting). Use the `uvcontsub` (§4.7.6), `split` (§4.7.1), and `uvmodelfit` (§4.7.8) tasks.

The flow chart and the above list are in a suggested order. However, the actual order in which you will carry out these operations is somewhat fluid, and will be determined by the specific data-reduction use cases you are following. For example, you may need to do an initial **Gain Calibration** on your bandpass calibrator before moving to the **Bandpass Calibration** stage. Or perhaps the polarization leakage calibration will be known from prior service observations, and can be applied as a constituent of Prior Calibration.

### 4.2.1 The Philosophy of Calibration in CASA

Calibration is not an arbitrary process, and there is a methodology that has been developed to carry out synthesis calibration and an algebra to describe the various corruptions that data might be subject to: the Hamaker-Bregman-Sault Measurement Equation (ME), described in Appendix E. The user need not worry about the details of this mathematics as the CASA software does that for
you. Anyway, it’s just matrix algebra, and your familiar scalar methods of calibration (such as in AIPS) are encompassed in this more general approach.

There are a number of “physical” components to calibration in CASA:

- **data** — in the form of the Measurement Set (§2.1). The MS includes a number of columns that can hold calibrated data, model information, and weights;

- **calibration tables** — these are in the form of standard CASA tables, and hold the calibration solutions (or parameterizations thereof);

- **task parameters** — sometimes the calibration information is in the form of CASA task parameters that tell the calibration tasks to turn on or off various features, contain important values (such as flux densities), or list what should be done to the data.

At its most basic level, Calibration in CASA is the process of taking “uncalibrated” data, setting up the operation of calibration tasks using parameters, solving for new calibration tables, and then applying the calibration tables to form “calibrated” data. Iteration can occur as necessary, with the insertion of other non-calibration steps (e.g. imaging to generate improved source models for “self-calibration”).

### 4.2.2 Keeping Track of Calibration Tables

The calibration tables are the currency that is exchanged between the calibration tasks. The “solver” tasks (gaincal, bandpass, blcal, polcal) take in the MS (which may have a calibration model attached) and previous calibration tables, and will output an “incremental” calibration table (it is incremental to the previous calibration, if any). This table can then be smoothed using smoothcal if desired.

You can optionally accumulate the incremental calibration onto previous calibration tables with accum, which will then output a cumulative calibration table. This task will also interpolate onto a different time scale. See §4.5.5 for more on accumulation and interpolation.

Figure 4.2 graphs the flow of these tables through the sequence

```
solve => smooth => accumulate
```

Note that this sequence applied to separate types of tables (e.g. ’B’, ’G’) although tables of other types can be previous calibration input to the solver.

The final set of cumulative calibration tables is what is applied to the data using applycal. You will have to keep track of which tables are the intermediate incremental tables, and which are cumulative, and which were previous to certain steps so that they can also be previous to later steps until accumulation. This can be a confusing business, and it will help if you adopt a consistent table naming scheme (see Figure 4.2 for an example naming scheme).
Figure 4.2: Chart of the table flow during calibration. The parameter names for input or output of the tasks are shown on the connectors. Note that from the output solver through the accumulator only a single calibration type (e.g. `B`, `G`) can be smoothed, interpolated or accumulated at a time. `accum` is optional (and not recommended as of v4.0). The final set of cumulative calibration tables of all types (accumulated or as a list of caltables) are then input to `applycal` as shown in Figure 4.1.

### 4.2.3 The Calibration of traditional VLA data in CASA

CASA supports the calibration of traditional VLA data that is imported from the Archive through the `importvla` task. See §2.2.3 for more information.

**ALERT:** Data taken both before and after the Modcomp turn-off in late June 2007 will be handled automatically by `importvla`. You do not need to set special parameters to do so, and it will obey the scaling specified by `applytsys`.

You can also import VLA data in UVFITS format with the `importuvfits` task (§2.2.5.1). However, in this case, you must be careful during calibration in that some prior or previous calibrations (see
below) may or may not have been done in AIPS and applied (or not) before export.

For example, the default settings of AIPS FILLM will apply VLA gaincurve and approximate (weather-based) atmospheric optical depth corrections when it generates the extension table CL1. If the data is exported immediately using FITTP, then this table is included in the UVFITS file. However, CASA is not able to read or use the AIPS SN or CL tables, so that prior calibration information is lost and must be applied during calibration here (i.e. using gaincurve=True and setting the opacity parameter).

On the other hand, if you apply calibration in AIPS by using the SPLIT or SPLAT tasks to apply the CL tables before exporting with FITTP, then this calibration will be in the data itself. In this case, you do not want to re-apply these calibrations when processing in CASA.

4.2.4 Loading Jansky VLA data in CASA

Jansky VLA data can be loaded into CASA either via importevla or by using the task importasdm. Both tasks will convert ASDM raw data files into measurement sets. importasdm will convert the data itself and the majority of the metadata. importevla will run importasdm followed by Jansky VLA-specific corrections, like the application of the on-line flags (e.g. times when the subreflector was not in place or the an antenna was not on source), an option to clip values that are exactly zero (as of 2010, such values still may appear in the VLA raw data), and flagging for shadowing.

4.3 Preparing for Calibration

There are a number of “a priori” calibration quantities that may need to be initialized or estimated before further calibration solving is carried out. These include

- **weight initialization** — if desired, initialization of spectral weights (by default, unchannelized weight accounting is used, and no special action is required)
- **system temperature correction** — turn correlation coefficient into correlated flux density (necessary for some telescopes),
- **gain curves** — antenna gain-elevation dependence,
- **atmospheric optical depth** — attenuation of the signal by the atmosphere, including correcting for its elevation dependence.
- **flux density models** — establish the flux density scale using “standard” calibrator sources, with models for resolved calibrators,
- **delays** — antenna-based delay offsets,
- **antenna position errors** — offsets in the positions of antennas assumed during correlation.
- **ionosphere** — dispersive delay and Faraday effects arising from signal transmission through the magnetized plasma of the ionosphere.
These are pre-determined effects and should be applied (if known) as priors when solving for other calibration terms, and included in the final application of all calibration. If unknown, then they will be solved for or subsumed in other calibration such as bandpass or gains.

We now deal with these in turn.

4.3.1 Weight initialization and WEIGHT_SPECTRUM

See Appendix F for a more complete description of weight accounting in CASA.

CASA 4.3 introduces initial experimental support for spectral weights. At this time, this is mainly relevant to ALMA processing for which spectral $T_{\text{sys}}$ corrections, which faithfully reflect spectral sensitivity, are available. In most other cases, sensitivity is, to a very good approximation, channel-independent after bandpass calibration (and often also before), except perhaps at the very edges of spectral windows (and for which analytic expressions of the sensitivity loss are generally unavailable). Averaging of data with channel-dependent flagging which varies on sufficiently short timescales will also generate channel-dependent net weights (see split2 or mstransform for more details).

By default, CASA’s weight accounting scheme maintains unchannelized weight information that is appropriately updated when calibration is applied. In the case of spectral calibrations ($T_{\text{sys}}$ and bandpass), an appropriate spectral average is used for the weight update. This spectral average is formally correct for weight update by bandpass. For $T_{\text{sys}}$, traditional treatments used a single measurement per spectral window; ALMA has implemented spectral $T_{\text{sys}}$ to better track sensitivity as a function of channel, and so should benefit from spectral weight accounting as described here, especially where atmospheric emission lines occur. If spectral weight accounting is desired, users must re-initialize the spectral weights using the initweights task:

\[
\text{initweight(vis='mydata.ms', dobt=True, dowtsp=True)}
\]

In this task, the dobt parameter controls whether channel bandwidth and integration time information are used to initialize the weight information (the default is dobt=T, which uses bandwidth and integration time; if dobt=F, the weight information will be initialized to unity). The dowtsp parameter controls whether (T) or not (F) the spectral weights (WEIGHT_SPECTRUM column) are initialized. The default is dowtsp=False, wherein only the non-spectral weights (WEIGHT column) will be initialized. If the spectral weights have been initialized, then downstream processing that supports spectral weights will use/update them. In v4.3, this includes applycal, clean, and split2/mstransform; use of spectral weights in calibration solving (e.g., gaincal and other solve tasks) is scheduled for the v4.4 release.

Note that importasdm and importevla currently initialize the non-spectral weights using channel bandwidth and integration time information (equivalent to initweights(vis='mydata.ms', dobt=T, dowtsp=F)). In general, it only makes sense to run initweights on a raw dataset which has not yet been calibrated, and it should only be necessary if the filled weights are inappropriate, or if spectral weight accounting is desired in subsequent processing. It is usually not necessary to re-initialize the weight information when redoing calibration from scratch (the raw weight information is preserved in the
SIGMA/SIGMA Spectral columns. (Re-)initializing the weight information for data that has already been calibrated (with calwt=T, presumably) is formally incorrect and is not recommended.

When combining datasets from different epochs, it is generally preferable to have used the same version of CASA (most recent is best), and with the same weight information conventions and calwt settings. Doing so will minimize the likelihood of arbitrary weight imbalances that might lead to net loss of sensitivity, and maximize the likelihood that real differences in per-epoch sensitivity (e.g., due to different weather conditions and instrumental setups) will be properly accounted for. Modern instruments support more variety in bandwidth and integration time settings, and so use of these parameters in weight initialization is preferred (c.f. use of unit weight initialization, which has often been the traditional practice).

**Alert:** Full and proper weight accounting for the EVLA formally depends on the veracity of the switched power calibration scheme (§4.3.7). As of late 2014, use of the EVLA switched power is not yet recommended for general use, and otherwise uniform weights are carried through the calibration process. As such, spectral weight accounting is not yet meaningful. Facilities for post-calibration estimation of spectral weights are planned for a future release.

### 4.3.2 System Temperature and Switched-Power Corrections

Some telescopes, including the old VLA, ALMA, and the VLBA, record the visibilities in the form of raw correlation coefficient with weights proportional to the number of bits correlated. The correlation coefficient is the fraction of the total signal that is correlated, and thus multiplication by the system temperature ($T_{sys}$) and the antenna gain (in Jy/K) will produce visibilities with units of correlated flux density. ALMA records $T_{sys}$ (K) information in the MS which can be extracted as a caltable using gencal with calmode='tsys', and applied to data to yield units of K. Calibration to flux density in Jy is achieved via reference to sources of known power.

**Alert:** Note that the old VLA system did this initial calibration on-line. The modern VLA does not record normalized visibilities. Instead, the correlations are delivered in raw engineering units that are proportional to power. The actual total power received is continuously monitored during the observation, with a calibration signal of known temperature (K) switched in at a rate of 10 Hz. This is the so-called “switched-power” calibration system on the VLA. This enables a continuous record of the $T_{sys}$ (K), as well as net electronic gain variation of each antenna’s receiving system. The correlator requantizer gain is also monitored. These data are recorded in MS subtables and appropriate calibration factors can be derived from them by gencal with caltype='swpow', and stored in a caltable for application. This calibration is not a “$T_{sys}$” calibration of the traditional sort; the switched-power gain is used to correct the visibility amplitude, and the $T_{sys}$ is used to set the weights. This system is still being commissioned (as of early 2014). Observations using 8-bit sampling are usually reasonably calibrated; 3-bit-sampled switched-power data are subject to compression effects that are not yet completely understood, and the switched power calibration is not recommended (instead, correction only by the requantizer gain is recommended, using caltype='rq').

See §4.3.6 for more information on use of gencal.
4.3.3 Antenna Gain-Elevation Curve Calibration

Large antennas (such as the 25-meter antennas used in the VLA and VLBA) have a forward gain and efficiency that changes with elevation. Gain curve calibration involves compensating for the effects of elevation on the amplitude of the received signals at each antenna. Antennas are not absolutely rigid, and so their effective collecting area and net surface accuracy vary with elevation as gravity deforms the surface. This calibration is especially important at higher frequencies where the deformations represent a greater fraction of the observing wavelength. By design, this effect is usually minimized (i.e., gain maximized) for elevations between 45 and 60 degrees, with the gain decreasing at higher and lower elevations. Gain curves are most often described as 2nd- or 3rd-order polynomials in zenith angle.

Gain curve calibration has been implemented in CASA for the modern VLA and old VLA (only), with gain curve polynomial coefficients available directly from the CASA data repository. To make gain curve and antenna efficiency corrections for VLA data, use $\text{gencal}$ with $\text{caltable}='\text{gceff}'$. See §4.3.6 for more information on use of $\text{gencal}$.

**ALERT:** If you are not using VLA data, do not use gaincurve corrections. A general mechanism for incorporating gaincurve information for other arrays will be made available in future releases. The gain-curve information available for the VLA is time-dependent (on timescales of months to years, at least for the higher frequencies), and CASA will automatically select the date-appropriate gain curve information. Note, however, that the time-dependence was poorly sampled prior to 2001, and so gain curve corrections prior to this time should be considered with caution.

4.3.4 Atmospheric Optical Depth Correction

The troposphere is not completely transparent. At high radio frequencies (>15 GHz), water vapor and molecular oxygen begin to have a substantial effect on radio observations. According to the physics of radiative transmission, the effect is threefold. First, radio waves from astronomical sources are absorbed (and therefore attenuated) before reaching the antenna. Second, since a good absorber is also a good emitter, significant noise-like power will be added to the overall system noise. Finally, the optical path length through the troposphere introduces a time-dependent phase error. In all cases, the effects become worse at lower elevations due to the increased air mass through which the antenna is looking. In CASA, the opacity correction described here compensates only for the first of these effects, tropospheric attenuation, using a plane-parallel approximation for the troposphere to estimate the elevation dependence.

To make opacity corrections in CASA, an estimate of the zenith opacity is required (see observatory-specific chapters for how to measure zenith opacity). This is then supplied to the $\text{caltype}='\text{opac}'$ parameter in $\text{gencal}$ which creates a calibration table with all the information. E.g. for data with two spectral windows, the inputs are like:

```
gencal(vis='dataset.ms',
       caltable='opacity.cal',
       caltype='opac',
       spw='0,1',
       parameter=[0.0399,0.037])
```
CHAPTER 4. SYNTHESIS CALIBRATION

If you do not have an externally supplied value for opacity, for example from a VLA tip procedure, then you should either use an average value for the telescope, or leave it at zero and let your gain calibration compensate as best it can (e.g. that your calibrator is at the same elevation as your target at approximately the same time. As noted above, there are no facilities yet to estimate this from the data (e.g. by plotting \(T_{sys}\) vs. elevation).

Below, we give instructions for determining opacity for Jansky VLA data from weather statistics and VLA observations where tip-curve data is available. It is beyond the scope of this cookbook to provide information for other telescopes.

4.3.4.1 Determining opacity corrections for modern VLA data

For the VLA site, weather statistics and/or seasonal models that average over many years of weather statistics prove to be reasonable good ways to estimate the opacity at the time of the observations. The task plotweather calculates the opacity as a mix of both actual weather data and seasonal model. It has the following inputs:

```python
# plotweather :: Plot elements of the weather table; estimate opacity.
vis = '' # MS name
seasonal_weight = 0.5 # weight of the seasonal model
doPlot = True # set this to True to create a plot
```

The task plots the weather statistics if doPlot=T, like shown in Figure 4.3. The bottom panel displays the calculated opacities for the run as well as a seasonal model. The parameter seasonal_weight can be adjusted to calculate the opacities as a function of the weather data alone seasonal_weight=0, only the seasonal model seasonal_weight=1, or a mix of the two (values between 0 and 1). Calculated opacities are shown in the logger output, one for each spectral window. plotweather can also assign a python variable to a list of calculated opacities (one entry for each spw) when run as:

```python
myTau = plotweather(vis='myvladata.ms')
```

In this example, myTau will be returned with a list of per-spw opacities, e.g. `myTau=[0.02,0.03]` and can later be used as input for gencal in caltype='opac' in the parameter setting, e.g.,

```python
# opac for spws 0,1 in myTau
gencal(vis='myvladata.ms',caltype='opac',spw='0,1',parameter=myTau)
```

Note that it is important to explicitly specify the spws that are covered by the opacity values stored in myTau. For most modern VLA data there will be more than two spws, probably.

See § 4.3.6 for more information on use of gencal.
Figure 4.3: The weather information for a MS as plotted by the task `plotweather`.

Weather Summary for AS1039.sb1382796_2_000.55368.51883247685.ms

- Solar el
- Wind (m/s)
- Temp.
- Dew point
- PWP (mm)

2010/06/21/12:28:25 to 13:26:54
4.3.4.2 Determining opacity corrections for VLA data

For VLA data, zenith opacity can be measured at the frequency and during the time observations are made using a VLA tipping scan in the observe file. Historical tipping data are available at:

http://www.vla.nrao.edu/astro/calib/tipper

Choose a year, and click Go to get a list of all tipping scans that have been made for that year.

If a tipping scan was made for your observation, then select the appropriate file. Go to the bottom of the page and click on the button that says Press here to continue. The results of the tipping scan will be displayed. Go to the section called 'Overall Fit Summary' to find the fit quality and the fitted zenith opacity in percent. If the zenith opacity is reported as 6%, then the actual zenith optical depth value is $\text{opacity}=0.060$ for gaincal and other calibration tasks.

If there were no tipping scans made for your observation, then look for others made in the same band around the same time and weather conditions. If nothing is available here, then at K and Q bands you might consider using an average value (e.g. 6% in reasonable weather). See the VLA memo


for more on the atmospheric optical depth correction at the VLA, including plots of the seasonal variations.

4.3.5 Setting the Flux Density Scale using (setjy)

When solving for visibility-plane calibration, CASA calibration applications compare the observed DATA column with the Fourier transform of calibrator model when it is provided (if no model is specified, a point source at the phase center is assumed).

The setjy task is used to set the proper flux density and attaches a model image (if specified) of the calibrator to the MS. For sources that are recognized flux calibrators (listed in Tables 4.1 and 4.2, see also §C.1), setjy can calculate the flux densities as a function of frequency (and time, for Solar System objects). Otherwise, the flux densities should be manually specified (standard='manual').

For the VLA, the default source models are customarily point sources defined by the 'Baars', 'Perley 90', 'Perley-Taylor 99', 'Perley-Butler 2010', time-variable 'Perley-Butler 2013', or 'Scaife-Heald 2012' flux density scales (§C.1.1 'Perley-Butler 2010' is the current standard by default), or point sources of unit flux density if the flux density is unknown. In fact, the model can be any image in Jy/pixel units (models typically generated by the clean task).

Optionally, the MODEL column can be filled with the Fourier transform of (option usescratch=T is setjy, ft, and clean). But for most measurement sets, the performance and data storage requirements are less demanding without the MODEL_DATA column.

The inputs for setjy are:
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# setjy :: Fills the model column with the visibilities of a calibrator
vis = '' # Name of input visibility file
field = '' # Field name(s)
spw = 'all' # Spectral window identifier (list)
selectdata = True # Other data selection parameters
timerange = '' # Time range to operate on (for usescratch=T)
scan = '' # Scan number range (for usescaratch=T)
intent = '' # Observation intent
observation = '' # Observation ID range (for usescratch=T)
scalebychan = True # scale the flux density on a per channel basis or else on a per spw basis
standard = 'Perley-Butler 2010' # Flux density standard
model = '' # File location for field model
listmodels = False # List the available modimages for VLA calibrators or Tb models for Solar System objects
usescratch = False # Will create if necessary and use the MODEL_DATA

Table 4.1: Recognized Flux Density Calibrators. Note that the VLA uses J2000 calibrator names. CASA accepts all strings that contain the names below. E.g. 'PKS 1934-638' will be recognized

<table>
<thead>
<tr>
<th>3C Name</th>
<th>B1950 Name</th>
<th>J2000 Name</th>
<th>Alt. J2000 Name</th>
<th>Standards</th>
</tr>
</thead>
<tbody>
<tr>
<td>3C48</td>
<td>0134+329</td>
<td>0137+331</td>
<td>J0137+3309</td>
<td>1,3,4,5,6,7</td>
</tr>
<tr>
<td>3C123</td>
<td>0433+295</td>
<td>0437+296</td>
<td>J0437+2940</td>
<td>2</td>
</tr>
<tr>
<td>3C138</td>
<td>0518+165</td>
<td>0521+166</td>
<td>J0521+1638</td>
<td>1,3,4,5,6</td>
</tr>
<tr>
<td>3C147</td>
<td>0538+498</td>
<td>0542+498</td>
<td>J0542+4951</td>
<td>1,3,4,5,6,7</td>
</tr>
<tr>
<td>3C196</td>
<td>0809+483</td>
<td>0813+482</td>
<td>J0813+4813</td>
<td>1,2,7</td>
</tr>
<tr>
<td>3C286</td>
<td>1328+307</td>
<td>1331+305</td>
<td>J1331+3030</td>
<td>1,2,3,4,5,6,7</td>
</tr>
<tr>
<td>3C295</td>
<td>1409+524</td>
<td>1411+522</td>
<td>J1411+5212</td>
<td>1,2,3,4,5,6,7</td>
</tr>
<tr>
<td>–</td>
<td>1934-638</td>
<td>–</td>
<td>J1939-6342</td>
<td>1,3,4,5,6</td>
</tr>
<tr>
<td>3C380</td>
<td>1828+487</td>
<td>1829+487</td>
<td>J1829+4845</td>
<td>7</td>
</tr>
</tbody>
</table>


By default the setjy task will cycle through all fields spectral windows and channels, (one solution per spw with scalebychan = False), setting the flux density either to 1 Jy (unpolarized), or if the source is recognized as one of the calibrators in the above table, to the flux density (assumed unpolarized) appropriate to the observing frequency. For example, to run setjy on a measurement set called data.ms:

```python
setjy(vis='data.ms')  # This will set all fields and spectral windows
```
Table 4.2: 'Butler-JPL-Horizons 2012' recognized Solar System Objects for Flux Calibration

<table>
<thead>
<tr>
<th>Planets</th>
<th>Moons</th>
<th>Asteroids</th>
</tr>
</thead>
<tbody>
<tr>
<td>Venus(^1), Mars(^2), Jupiter(^3), Uranus(^4), Neptune(^5)</td>
<td><em>Jupiter</em>: Io, Europa, Ganymede, Callisto</td>
<td>Ceres, Pallas(^8), Vesta(^8), Juno(^8)</td>
</tr>
</tbody>
</table>

\(^1\) Venus: model for \(\sim\) 300 MHz to 350 GHz, no atmospheric lines (CO, H\(_2\)O, HDO, etc.)
\(^2\) Mars: tabulated as a function of time and frequency (30 - 1000 GHz) based on Rudy et al. (1988), no atmospheric lines (CO, H\(_2\)O, HDO, etc.)
\(^3\) Jupiter: model for 30-1020 GHz, does not include synchrotron emission
\(^4\) Uranus: model for 60-1800 GHz, contains no rings or synchrotron.
\(^5\) Neptune: model for 2-2000 GHz, the broad CO absorption line is included, but contains no rings or synchrotron.
\(^7\) Titan: model for 53.3-1024.1 GHz, include many spectral lines
\(^8\) not recommended (The temperature is not yet adjusted for varying distance from the Sun. The model data can be scaled after running setjy, but it is an involved process.)

Details are described in ALMA Memo 594 available on [https://science.nrao.edu/facilities/alma/aboutALMA/Technology/ALMA_Memo_Series/alma594/abs594](https://science.nrao.edu/facilities/alma/aboutALMA/Technology/ALMA_Memo_Series/alma594/abs594).

Models of available calibrator sources can be listed by setting `listmodels=True`. `setjy` will then come up with all images that are in the paths where calibrator models for known telescopes are stored. It will also show all images in the working directory - any image there could potentially be a calibrator model. If the calibrator model is found by `listmodels` it can be used in the `modimage` parameter without a path.

The `fluxdensity` parameter can be used to specify the flux of the calibrator in all Stokes parameters. It it thus a list of values \([I,Q,U,V]\), e.g. \(['12Jy', '13mJy', '0Jy', '0Jy']\). In addition, a spectral index can be specified via `spix`, a reference frequency `reffreq` (using the definition: \(S = fluxdensity \times \frac{freq}{reffreq}^{spix}\)), as well as a polarization index (`polindex`), angle (`polangle`) and a rotation measure (`rotmeas`).

Most calibrator sources are based on radio emission from quasars and jets. The spectral indices of these sources are such that at (sub)mm wavelengths the majority of these sources become too weak and variable to be reliable flux estimators. Alternatives are thermal objects such as planets, moons, and asteroids. Those sources, however, are all Solar System objects, which implies that they move and may be (strongly) resolved. The recommended `standard='Butler-JPL-Horizons 2012'` (for more information on the implemented models, see ALMA Memo 594 soon available on [https://science.nrao.edu/facilities/alma/aboutALMA/Technology/ALMA_Memo_Series/alma594/abs594](https://science.nrao.edu/facilities/alma/aboutALMA/Technology/ALMA_Memo_Series/alma594/abs594)) option of `setjy` includes flux density calibration using Solar System objects. For 'Butler-JPL-Horizons 2012' CASA currently supports the objects listed in Table 4.2 to be applied to ALMA data. These names are recognized when they are used in the 'field' parameter in `setjy`. In that case, `setjy` will obtain the geocentric distance and angular diameter at the time of the observation from a (JPL–Horizons) ephemeris and
calculate model visibilities. Currently the objects are modeled as uniform temperature disks, but
effects like primary beam attenuation and limb darkening will be accounted for soon. Note that
this model may oversimplify the real structure, in particular asteroids.


```python
setjy(vis='c0104I', field='MARS', spw='0 2', standard='Butler-JPL-Horizons 2012')
```

Tip: Running casalog.filter('INFO1') before running setjy with a Solar System object may send
the logger a reference to the temperature measurement. Use casalog.filter('INFO') to restore
the normal logging level.

The source model will be attached to the MS and applied to all calibration steps when usescratch=False.
usescratch=True fills the MODEL_DATA column with the Fourier transform of the model. As of
CASA 3.4. we found that under some circumstances, creation of the MODEL column may prevent
memory issues and if tasks fail, we recommend to set usescratch=True. Note that currently setjy
will not transform a full-Stokes model image such that all polarizations are applied correctly. You
need to use ft for this.

To limit this operation to certain fields and spectral windows, use the field and/or spw parameters,
which take the usual data selection strings (§2.3). For example, to set the flux density of the first
field (all spectral windows)

```python
setjy(vis='data.ms',field='0')
```
or to set the flux density of the second field in spectral window 17

```python
setjy(vis='data.ms',field='1',spw='17')
```
The full-polarization flux density (I,Q,U,V) may also be explicitly provided:

```python
setjy(vis='data.ms',
     field='1',spw='16', # Run setjy on field id 1, spw id 17
     fluxdensity=[3.5,0.2,0.13,0.0]) # and set I,Q,U,V explicitly
```

**ALERT:** The apparent brightness of objects in the Solar System will vary with time because of
the Earth’s varying distance to them, if nothing else. If the field *index* of a flux calibrator spans
several days, you should run setjy more than once, limiting each run to a suitable timerange by
using the timerange, scan, and/or observation selection parameters. Note that it is the field index
that matters, not the name. Typically concat assigns moving objects a new field index for each
observation, so usually it is not necessary to select a time range in setjy. However, it is worth
checking with listobs, especially for planets.

### 4.3.5.1 Using Calibration Models for Resolved Sources

For observations of solar system objects using the *Butler-JPL-Horizons 2010* and *Butler-JPL-
Horizons 2012* models (§4.3.5) setjy will know and apply the flux distribution across the extended
structure of the calibrators.
For other sources, namely VLA calibrator sources, a flux density calibrator can be resolved at the observing frequency and the point source model generated by \texttt{setjy} will not be appropriate. If available, a model image of the resolved source at the observing frequency may be used to generate the appropriate visibilities using the \texttt{modimage} parameter (or in older versions explicitly with the \texttt{ft} task). To use this, provide \texttt{modimage} with the path to the model image. Remember, if you just give the file name, it will assume that it is in the current working directory. Note also that \texttt{setjy} using a model image will only operate on that single source, thus you would run it multiple times (with different field settings) for different sources.

Otherwise, you may need to use the \texttt{uvrange} selection (§4.4.1.2) in the calibration solving tasks to exclude the baselines where the resolution effect is significant. There is not hard and fast rule for this, though you should consider this if your calibrator is shows a drop of more than 10% on the longest baselines (use \texttt{plotxy}, §3.3.2, to look at this). You may need to do \texttt{antenna} selection also, if it is heavily resolved and there are few good baselines to the outer antennas. Note that \texttt{uvrange} may also be needed to exclude the short baselines on some calibrators that have extended flux not accounted for in the model. \textbf{Note:} the calibrator guides for the specific telescopes usually indicate appropriate min and max for \texttt{uvrange}. For example, see the \textit{VLA Calibration Manual} at:

\url{http://www.vla.nrao.edu/astro/calib/manual/}

for details on the use of standard calibrators for the E/VLA.

Model images for some flux density calibrators are provided with CASA:

- Red Hat Linux RPMs 32bit (RHE4, Fedora 6): located in 
  /usr/lib/casapy/data/nrao/VLA/CalModels
- Red Hat Linux RPMs 64bit (RHE4, Fedora 6): located in 
  /usr/lib64/casapy/data/nrao/VLA/CalModels
- MAC OSX .dmg: located in 
  /Applications/CASA.app/Contents/Resources/casa-data/nrao/VLA/CalModels
- NRAO-AOC casapy-test: 
  /home/casa/data/nrao/VLA/CalModels

\textit{e.g.}, these are found in the \texttt{data/nrao/VLA/CalModels} sub-directory of the CASA installation. For example, just point to the repository copy, \textit{e.g.}

\texttt{modimage = '/usr/lib/casapy/data/nrao/VLA/CalModels/3C48_C.im'}

\textit{or if you like, you can copy the ones you wish to use to your working directory.}

The models available are:

\begin{verbatim}
3C138_L.im  3C147_L.im  3C286_L.im  3C48_L.im
3C138_C.im  3C147_C.im  3C286_C.im  3C48_C.im
\end{verbatim}
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3C138_X.im  3C147_X.im  3C286_X.im  3C48_X.im
3C138_U.im  3C147_U.im  3C286_U.im  3C48_U.im
3C138_K.im  3C147_K.im  3C286_K.im  3C48_K.im
3C138_Q.im  3C147_Q.im  3C286_Q.im  3C48_Q.im

(more calibrator models for the VLA are available at
https://science.nrao.edu/facilities/vla/data-processing/models). These are all un-reconvolved
images of AIPS CC lists. It is important that the model image not be one convolved with a finite
beam; it must have units of Jy/pixel (not Jy/beam).

Note that setjy will rescale the flux in the models for known sources (e.g. those in Table 4.1) to
match those it would have calculated. It will thus extrapolated the flux out of the frequency band
of the model image to whatever spectral windows in the MS are specified (but will use the structure
of the source in the model image).

**ALERT:** The reference position in the modimage is currently used by setjy when it does the
Fourier transform, thus differences from the positions for the calibrator in the MS will show up
as phase gradients in the uv-plane. If your model image position is significantly different but you
don’t want this to affect your calibration, then you can doctor either the image header using imhead
(§ 6.2) or in the MS (using the ms tool) as appropriate. In an upcoming release we will put in a
toggle to use or ignore the position of the modimage. Note that this will not affect the flux scaling
(only put in erroneous model phases); in any event small position differences, such as those arising
by changing epoch from B1950 to J2000 using regridimage (§ 6.13), will be inconsequential to the
calibration.

This illustrates the use of uvrange for a slightly resolved calibrator:

```python
# Import the data
importvla(archivefiles='AS776_A031015.xp2', vis='ngc7538_XBAND.ms',
    freqtol=10000000.0, bandname='X')

# Flag the ACs
flagautocorr('ngc7538_XBAND.ms')

# METHOD 1: Use point source model for 3C48, plus uvrange in solve

# Use point source model for 3C48
setjy(vis='ngc7538_XBAND.ms',field='0');

# Limit 3C48 (fieldid=0) solutions to uvrange = 0-40 klambda
gaincal(vis='ngc7538_XBAND.ms', caltable='cal.G', field='0',
    solint=60.0, refant='10', selectdata=True, uvrange='0~40klambda',
    append=False)

# Append phase-calibrator’s solutions (no uvrange) to the same table
gaincal(vis='ngc7538_XBAND.ms', caltable='cal.G', field='2',
    solint=60.0, refant='10', selectdata=True, uvrange='',
    append=True)

# Fluxscale
```
while the following illustrates the use of a model:

```python
# METHOD 2: use a resolved model copied from the data repository
# for 3C48, and no uvrange
# (NB: detailed freq-dep flux scaling TBD)

# Copy the model image 3C48_X.im to the working directory first!

setjy(vis='ngc7538_XBAND.ms', field='0', modimage='3C48_X.im')

# Solutions on both calibrators with no uvrange
gaincal(vis='ngc7538_XBAND.ms', caltable='cal.G2', field='0,2',
        solint=60.0, refant='10',
        append=False)

# Fluxscale
fluxscale(vis='ngc7538_XBAND.ms', caltable='cal.G2', reference=['0137+331'],
          transfer=['2230+697'], fluxtable='cal.G2flx', append=False)
```

# Both methods give 2230 flux densities ~0.7 Jy, in good agreement with
# AIPS

4.3.6 Correction for delay and antenna position offsets using gencal

The gencal task provides a means of specifying antenna-based calibration values manually. The values are put in designated tables and can be applied to the data on-the-fly in solving tasks and using applycal.

The `gencal` task has the inputs:

```python
# gencal :: Specify Calibration Values of Various Types
vis = '' # Name of input visibility file
caltab = '' # The new/existing calibration table
caltyp = 'tecim' # The calibration type: 'amp', 'ph', 'sbd', 'mbd', 'antpos', 'antposvla', 'tsys', 'evlagain', 'opac', 'gc', 'gceff', 'eff', 'te', 'cim'
infile = '' # Input ancilliary file
spw = 'all' # Calibration spw(s) selection
antenna = '1' # Calibration antenna(s) selection
pol = '' # Calibration polarization(s) selection
parameter = [] # The calibration values
```

Current antenna-based gencal options (caltyp) are:

- `amp` — amplitude correction
• 'ph' — phase correction
• 'sbd' — single-band delay (phase-frequency slope for each spw)
• 'mbd' — multi-band delay (phase-frequency slope over all spw)
• 'antpos' — ITRF antenna position corrections for the Jansky VLA (automatic parameter lookup is supported)
• 'antposvla' — old VLA-centric antenna position corrections
• 'tsys' — Tsys from the SYSCAL table (ALMA)
• 'evlagain' — VLA switched-power gains (experimental; equal to 'swpow')
• 'swpow' — VLA switched power (equal to 'evlagain')
• 'rq' — VLA requantizer gains
• 'swp/rq' — VLA switched-power gains divided by requantizer gain
• 'opac' — Tropospheric opacity
• 'gc' — Gain curve (zenith-angle-dependent gain) (VLA only) (auto-lookup only)
• 'gceff' — Antenna efficiency (sqrt(K/Jy)) (VLA only) (auto-lookup only)
• 'eff' — Gain curve and efficiency (VLA only) (auto-lookup only)
• 'tecim' — Total electron content to derive dispersive delays

The calibration parameter specifications cannot be time-variable in the present implementation (though some of them will introduce implicit time-dependence upon evaluation in the apply). Calibration values can be assigned to each spw, antenna and pol selection, where applicable. The list of calibration values specified in parameter must conform to the range of spectral windows, antennas, and polarizations specified in spw, antenna and pol, with the values specified in order of the specified polarizations (fastest), antennas, and spectral windows (slowest). If any of spw, antenna, or pol are left unspecified (empty strings), the values specified in parameter will be assumed applicable to all values of the unspecified data axes. The output caltable will otherwise assume nominal calibration values for unspecified spectral windows, antennas, and polarizations. Note that antenna position corrections formally do not have spectral-window or polarization dependence; such specifications should not be used with 'antpos'.

The same caltable can be specified for multiple runs of gencal, in which case the specified parameters will be incorporated cumulatively. E.g., amplitude parameters (caltype='amp') multiply and phase-like parameters ('ph', 'sbd', 'mbd', 'antpos') add. Parameters for 'amp' and 'ph' corrections can be incorporated into the same caltable (in separate runs), but each of the other types require their own unique caltable. A mechanism for specifying manual corrections via a text file will be provided in the future.
Two kinds of delay corrections are supported. For \texttt{caltype=\textquotesingle sbd\textquotesingle}, the specified delays (in nanoseconds) will be applied locally to each spectral window, referring the derived phase corrections to each spectral window’s reference frequency (where the phase correction will be zero). The phases in each spectral window will nominally be flattened, but any phase offsets between spectral windows will remain. (These can be corrected using \texttt{caltype=\textquotesingle phase\textquotesingle}, or via ordinary spectral-window-dependent phase calibration.) For \texttt{caltype=\textquotesingle mbd\textquotesingle}, the evaluated phase corrections are referred to zero frequency. This causes a correction that is coherent over many spectral windows. If the data are already coherent over many spectral windows and share a common multi-band delay (e.g., VLA data, per baseband), \texttt{caltype=\textquotesingle mbd\textquotesingle} corrections will maintain this coherence and flatten the frequency-dependent phase. Using \texttt{caltype=\textquotesingle sbd\textquotesingle} in this instance will introduce phase offsets among spectral windows that reflect the multi-band delay.

For antenna position corrections (\texttt{caltype=\textquotesingle antpos\textquotesingle}), the antenna position offsets are specified in the ITRF frame. If the \texttt{antenna} field is left empty, \texttt{gencal} will try to look up the appropriate antenna position offsets at the time of the observation from the VLA baseline webpage \url{http://www.vla.nrao.edu/astro/archive/baselines/}. For VLA position corrections in the VLA-centric frame, use \texttt{caltype=\textquotesingle antposvla\textquotesingle}, and \texttt{gencal} will rotate them to ITRF before storing them in the output caltable.

The sign and scale convention for \texttt{gencal} corrections (indeed for all CASA caltables) is such that the specified parameters (and as stored in caltables) are the factors that \textit{corrupt} ideal data to yield the observed data. Thus, when applied to correct the data, their effective inverse will automatically be taken. I.e., amplitude factors will be divided into the data on correction. Phase-like parameters adopt the convention that the complex factor for the second antenna in the baseline is conjugated, and then both antenna factors are divided into the data on correction. (These conventions differ from AIPS in that \textit{multiplying} correction factors are stored in AIPS calibration tables; however, the phase convention ends up being the same since AIPS conjugates the complex factor for the \textit{first} antenna in the baseline.)

The following series of examples illustrate the use of \texttt{gencal}.

For the dataset \texttt{\'data.ms\textquoteright}, the following sequence of \texttt{gencal} runs introduces, into a single caltable (\texttt{\textquotesingle test.G\textquotesingle}), (1) an antenna-based amplitude scale correction of 3.0 for all polarizations, antennas, and spectral windows, (2) phase corrections for all spectral windows and polarizations of 45 and 120 degrees to antennas EA03 and EA04, respectively, (3) phase corrections for all spectral windows of 63 and -34 in R (only) for antennas EA05 and EA06, respectively, and (4) phase corrections for all spectral windows of 14, -23, -130, and 145 degrees for antenna/polarizations EA09/R, EA09/L, EA10/R, and EA10/L, respectively:

\begin{verbatim}
gencal(vis='data.ms',caltable='test.G',caltype='amp', \ 
     spw='',antenna='',pol='', \ 
     parameter=[3])
gencal(vis='data.ms',caltable='test.G',caltype='ph', \ 
     spw='',antenna='EA03,EA04',pol='', \ 
     parameter=[45,120])
\end{verbatim}
In the following example, delay corrections in both polarizations will be adjusted for antenna EA09 by 14 nsec in spw 2 and -130 nsec in spw 3, and for antenna EA10 by -23 nsec in spw 2 and 145 nsec in spw 3:

```
gencal(vis='test.ms', caltable='test.sbd', caltype='sbd', spw='2,3', antenna='EA09,EA10', pol='', parameter=[14,-23,-130,145])
```

In the following example, antenna position corrections in meters (in ITRF) for antenna EA09 (dBx=0.01, dBy=0.02, dBz=0.03) and for antenna EA10 (dBx=-0.03, dBy=-0.01, dBz=-0.02) are introduced. Note that three parameters are required for each antenna. The antenna offsets can be obtained for the 'Jansky VLA/old VLA Baseline Corrections' web page: [http://www.vla.nrao.edu/astro/archive/baselines](http://www.vla.nrao.edu/astro/archive/baselines). The table given on this webpage has a format like:

```
; 2010 BASELINE CORRECTIONS IN METERS
;ANT
;MOVED OBSDATE Put_In_ MC(IAT) ANT PAD Bx By Bz
;
JAN27 FEB12 FEB21 01:57 11 E04 0.0000 0.0000 0.0000
JAN27 FEB12 FEB21 01:57 26 W03 -0.0170 0.0204 0.0041
MAR24 MAR25 MAR26 18:28 17 W07 -0.0061 -0.0069 -0.0055
APR21 MAY02 MAY04 23:25 12 E08 -0.0072 0.0045 -0.0017
```

If your observations fall in between the 'Antenna Moved' and 'Put_In' dates of a given antenna, you may choose to apply the offsets in that table; the 'Put_In' time stamp marks the date where the more accurate solution was introduced in the data stream directly and no correction is required anymore. In `gencal` the offsets will be inserted as:

```
gencal(vis='test.ms', caltable='test.antpos', caltype='antpos', antenna='EA09,EA10', parameter=[0.01,0.02,0.03,-0.03,-0.01,-0.02])
```

In the following example, antenna position corrections (in the traditional VLA-centric frame) will be introduced in meters for antenna EA09 (dBx=0.01, dBy=0.02, dBz=0.03) and for antenna EA10 (dBx=-0.03, dBy=-0.01, dBz=-0.02) These offsets will be rotated to the ITRF frame before storing them in the caltable.
gencal(vis='test.ms', caltable='test.antposvla', caltype='antposvla', \ 
    antenna='EA09,EA10', \ 
    parameter=[0.01,0.02,0.03, -0.03,-0.01,-0.02])

gencal is also the task to generate gaincurve, antenna efficiency, and opacity tables. The first two items can be determined together with caltype='gceff' and the latter with caltype='opac'. These tables are treated just like any other calibration table and will be carried through the calibration steps. This method replaces the older method where 'gaincurve' and 'opacity' keywords were present in calibration tasks such as gaincal, bandpass, or applycal.

4.3.7 Applying Jansky VLA switched power or ALMA Tsys using gencal

Noise diodes in the Jansky VLA antennas can be used to pre-calibrate the data. The diodes follow an ON-OFF cycle and the power for both states is measured and recorded. This is called the 'VLA switched power' calibration. To apply the switched power data, one needs to create a calibration table with gencal using caltype='evlagain', like

gencal(vis='test.ms', caltable='VLAswitchedpower.cal', caltype='evlagain')

For ALMA the calibration of system temperature is done via hot loads and the data recorded similar to the VLA in the measurement set (ALMA will provide measurement sets where these data are available. To derive the calibration table from it, use caltype='tsys':

gencal(vis='test.ms', caltable='ALMAtsys.cal', caltype='tsys')

This calibration tables created for ALMA or VLA are then carried along all further calibration steps in the gaintable parameter.

4.3.8 Generate a gain table based on Water Vapor Radiometer data wvrgcal

# wvrgcal :: Generate a gain table based on Water Vapour Radiometer data
vis    = ''       # Name of input visibility file
caltab = ''       # Name of output gain calibration table
toffset = 0       # Time offset (sec) between interferometric and WVR data
segsource = True  # Do a new coefficient calculation for each source
tie = []          # Prioritise tieing the phase of these sources as well as possible (requires segsource=True)
_sourceflag = []  # Flag the WVR data for these source(s) as bad and do not produce corrections for it (requires segsource=True)
disperse = False  # Apply correction for dispersion
wvrflag = ['']    # Flag the WVR data for these antenna(s) as bad and replace its data with interpolated values
statfield = ''    # Compute the statistics (Phase RMS, Disc) on this field only
statsource = ''   # Compute the statistics (Phase RMS, Disc) on this source only
The task `wvrgcal` generates a gain table based on Water Vapor Radiometer (WVR) data and is used for ALMA data reduction. It is an interface to the executable “wvrgcal” which is part of the CASA 4.3 distribution and can also be called from outside CASA. The wvrgcal software is based on the libair and libbnmin libraries which were developed by Bojan Nikolic at the University of Cambridge as part of EU FP6 ALMA Enhancement program.

CASA 4.3 contains version 2.0 of wvrgcal. Source code of the stand-alone package and links to documentation up to version 1.2.1 can be found online[1]. In particular, there are three ALMA memos (number 587, 588, and 593 (submitted)) which describe the algorithms implemented in the software. They can be found at at the ALMA Memo Series[2].

With `wvrgcal` version 2.0, maintenance of the tool has now fullt transitioned to ESO. Newly added features include full support for flags, and the new parameters `spw`, `wvrspw`, `maxdistm`, `mingoodfrac`, and `usefieldtab`.

Briefly, wvrgcal follows a Bayesian approach to calculate the coefficients that convert the outputs of the ALMA 183 GHz water-vapor radiometers (mounted on each antenna) into estimates of path fluctuations which can then be used to correct the observed interferometric visibilities.

The CASA task interface to wvrgcal follows closely the interface of the shell executable at the same time staying within the CASA task parameter conventions.

In ALMA data, the WVR measurements belonging to a given observation are contained in the ASDM for that observation. After conversion to an MS using `importasdm`, the WVR information can be found in separate spectral windows (as of November 2014, it is still spectral window id 0 only). This spectral window must be present in the MS for wvrgcal to work.

The various features of wvrgcal are then controlled by a number of task parameters (see the list above). They have default values which will work for ALMA data. An example for a typical wvrgcal call can be found in the ALMA CASA guide for the NGC 3256 analysis:

---


Here, \texttt{vis} is the name of input visibility file (which as mentioned above also contains the WVR data in spectral window 0) and \texttt{caltable} is the name of the output gain calibration table. 

\texttt{toffset} is the known time offset in seconds between the WVR measurements and the visibility integrations they are valid for. For ALMA, this offset is presently -1 s (which is also the default value).

The parameter \texttt{segsource} (segregate source) controls whether separate coefficients are calculated for each source. The default value True is the recommended one for ALMA. When \texttt{segsource} is True, the subparameter \texttt{tie} is available. It permits to form groups of sources for which common coefficients are calculated as well as possible. The \texttt{tie} parameter ensures best possible phase transfer between a group of sources. In general it is recommended to tie together all of the sources in a single Science Goal (in ALMA speak) and their phase calibrator(s). The recommended maximum angular distance up to which two sources can be tied is 15°.

The parameter \texttt{statsource} controls for which sources statistics are calculated and displayed in the logger. This has no influence on the generated calibration table.

\texttt{wvrgcal} respects the flags in the Main and ANTENNA table of the MS. The parameter \texttt{mingood-frac} lets the user set a requirement on the minimum fraction of good measurements for accepting the WVR data from an antenna. If antennas are flagged, their WVR solution is interpolated from the three nearest neighbouring antennas. This process can be controlled with the new parameters \texttt{maxdistm} and \texttt{minnumants}. The former sets the maximum distance an antenna used for interpolation may have from the flagged one. And \texttt{minnumants} sets how many near antennas there have to be for interpolation to take place.

For more details on the WVR Phase correction, see also the the ALMA Memo “Quality Control of WVR Phase Correction Based on Differences Between WVR Channels” by B. Nikolic, R. C. Bolton & J. S. Richer\(^3\) see also ALMA memo #593\(^4\).

### 4.3.8.1 Statistical parameters shown in the logger output of \texttt{wvrgcal}

\texttt{wvrgcal} writes out a variety of information to the logger, including various statistical measures of the performance. This allows the user to judge whether WVR correction is appropriate for the ms, to check whether any antennas have problematic WVR values, and to examine the predicted performance of the WVR correction when applied.

For each set of correction coefficients which are calculated (the number of coefficient sets are controlled by the parameters \texttt{nsol}, \texttt{segsource} and \texttt{tie}), the \texttt{wvrgcal} output to the logger first of all shows the time sample, the individual temperatures of each of the four WVR channels and the elevation of the source in question at that time.

\(^3\)http://www.casa.nrao.edu/Memos/memoqachannels.pdf

\(^4\)https://science.nrao.edu/facilities/alma/aboutALMA/Technology/ALMA_Memo_Series/alma593/abs593
For each of these coefficient sets, it then gives the evidence of the bayesian parameter estimation, the calculated precipitable water vapour (PWV) and its error in mm, and the correction coefficients found for each WVR channel (dTdL).

The output then shows the statistical information about the observation. First of all it gives the start and end times for the parts of the observation used to calculate these statistics (controlled by segsource). It then shows a break down for each of the antennas in the data set. This gives the antenna name and number; whether or not it has a WVR (column WVR); whether or not it has been flagged (column FLAG); the RMS of the path length variation with time towards that antenna (column RMS); and the discrepancy between the RMS path length calculated separately for different WVR channels (column Disc.). These values allow the user to see if an individual WVR appears to have been suffering from problems during the observation, and to flag that antenna using wvrflag if necessary.

This discrepancy value, Disc., can in addition be used as a simple diagnostic tool to evaluate whether or not the WVR correction caltable created by wvrgcal should be applied. In the event of the WVR observations being contaminated by strong cloud emission in the atmosphere, the attempt by wvrgcal to fit the water vapour line may not be successful, and applying the produced calibration table can in extreme cases reduce the quality of the data. However, these weather conditions should identified by a high value in the discrepancy column produced when running wvrgcal.

Although there have not currently been enough cases checked to give definitive limits, the available data sets as of summer 2012 suggest that discrepancy values of greater than a 1000 microns usually indicate strong cloud contamination of the WVR data, and the output calibration table should probably not be applied. If the values are between 100 and 1000 microns, then the user should manually examine the phases before and after applying the caltable to decide if WVR correction is appropriate.

After the antenna-by-antenna statistics, the output then displays some estimates of the performance of the wvrgcal correction. These are the thermal contribution from the water vapour to the path fluctuations per antenna (in microns), the largest path fluctuation found on a baseline (in microns), and the expected error on the path length calculated for each baseline due to the error in the coefficients (in microns).

4.3.8.2 Antenna position calculation

The information about antenna pointing direction is by default taken from the POINTING table. Should this table not be present for some reason, the user can instead switch to determining the antenna positions from the phase directions in the FIELD table (under the assumption that all antennas were pointing ideally). The switch is performed by setting the parameter usefieldtab to True.
4.3.8.3 Spectral window selection

By default, \texttt{wvrgcal} has traditionally put solutions for all spectral windows of the MS into the output calibration table. Since usually only the spectral windows are of interest in which the science target and the calibrators were observed, it is not necessary to store solutions for other spectral windows.

The spectral windows for which solutions are stored can be selected with the parameter \texttt{spw}, e.g., \texttt{spw = [17, 19, 21, 23]} will make \texttt{wvrgcal} write only solutions for spectral windows 17, 19, 21, and 23.

Should ALMA in the future use other spectral windows than id 0 to store the WVR information, then the input WVR spectral window can be selected using the parameter \texttt{wvrspw}. The syntax is the same as for parameter \texttt{spw}.

4.3.9 Ionospheric corrections

CASA 4.3 introduces initial support for on-axis ionospheric corrections, using time- and direction-dependent total electron content (TEC) information obtained from the internet. The correction includes the dispersive delay ($\propto \nu^{-1}$) delay and Faraday rotation ($\propto \nu^{-2}$) terms. These corrections are most relevant at observing frequencies less than $\sim$5 GHz. When relevant, the ionosphere correction table should be generated at the beginning of a reduction along with other calibration priors (antenna position errors, gain curve, opacity, etc.), and carried through all subsequent calibration steps. Formally, the idea is that the ionospheric effects (as a function of time and on-axis direction) will be nominally accounted for by this calibration table, and thus not spuriously leak into gain and bandpass solves, etc. In practice, the quality of the ionospheric correction is limited by the relatively sparse sampling (in time and direction) of the available TEC information. Especially active ionospheric conditions may not be corrected very well. Also, direction-dependent ionosphere corrections are not yet supported. (Various improvements are under study for future releases.)

To generate the ionosphere correction table, first import a helper function from the casapy recipes repository:

```
from recipes import tec_maps
```

Then, generate a TEC surface image:

```
tec_maps.create(vis='mydata.ms', doplot=T, imname='tec.im')
```

This function goes to the web to obtain TEC information for the observing date and location, and generates a time-dependent CASA image containing this information. The quality of the information improves with time after the observing date, becoming optimal 1-2 weeks later. This image can be viewed as a movie in the CASA \texttt{viewer}. If \texttt{doplot=T}, the function will also produce a plot of the TEC as a function of time in a vertical direction over the observatory.

Finally, to generate the ionosphere correction caltable, pass the TEC image into \texttt{gencal}, using \texttt{calmode='tecim'}:
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```
gencal(vis='mydata.ms',caltable='tec.cal',calmode='tecim',infile='tec.im')
```

This iterates through the dataset and samples the zenith angle-dependent projected line-of-sight TEC for all times in the observation, storing the result in a standard CASA caltable. Plotting this caltable will show how the TEC varies between observing directions for different fields and times, in particular how it changes as zenith angle changes, and including the nominal difference between science targets and calibrators.

This caltable should then be used as a prior in all subsequent calibration solves, and included in the final applycal.

A few warnings:

- The TEC information obtained from the web is relatively poorly sampled in time and direction, and so will not always described the details of the ionospheric corruption, especially during active periods.
- For instrumental polarization calibration, it is recommended that an unpolarized calibrator be used; polarized calibrators may not yield as accurate a solution since the ionospheric corrections are not yet used properly in the source polarization portion of the solve.

Special thanks are due to Jason Kooi (UIowa) for his contributions to ionospheric corrections in CASA.

4.3.10 Other a priori Calibrations and Corrections

Other a priori calibrations will be added to the calibrator (cb) tool in the near future. These will include instrumental line-length corrections, etc. Where appropriate, solving capabilities for these effects will also be added.

4.4 Solving for Calibration — Bandpass, Gain, Polarization

The gaincal, bandpass, polcal, and blcal tasks actually solve for the unknown calibration parameters from the visibility data obtained on calibrator sources, placing the results in a calibration table. They take as input an MS, and a number of parameters that specify any prior calibration or previous calibration tables to pre-apply before computing the solution. These are placed in the proper sequence of the Measurement Equation automatically.

We first discuss the parameters that are in common between many of the calibration tasks. Then we describe each solver in turn.

4.4.1 Common Calibration Solver Parameters

There are a number of parameters that are in common between the calibration “solver” tasks. These also appear in some of the other calibration manipulation and application tasks.
4.4.1.1 Parameters for Specification: vis and caltable

The input measurement set and output table are controlled by the following parameters:

```python
vis       = ''   # Name of input visibility file
caltable  = ''   # Name of output calibration table
```

The MS name is input in `vis`. If it is highlighted red in the inputs (§1.4.5.4) then it does not exist, and the task will not execute. Check the name and path in this case.

The output table name is placed in `caltable`. Be sure to give a unique name to the output table, or be careful. If the table exists, then what happens next will depend on the task and the values of other parameters (e.g. §4.4.1.6). The task may not execute giving a warning that the table already exists, or will go ahead and overwrite the solutions in that table, or append them. Be careful.

4.4.1.2 Selection: field, spw, selectdata, intent, and observation

Selection is controlled by the parameters:

```python
field     = ''   # field names or index of calibrators: ''==>all
spw       = ''   # spectral window:channels: ''==>all
intent    = ''   # Select observing intent
selectdata = False # Other data selection parameters
```

Field and spectral window selection are so often used, that we have made these standard parameters `field` and `spw` respectively. `intent` is the scan intent that was specified when the observations were set up. They typically describe what was intended with a specific scan, i.e. a flux or phase calibration, a bandpass, a pointing, an observation of your target, or something else or a combination. The format for the scan intents of your observations are listed in the logger when you run `listobs`. Minimum matching with wildcards will work, like "*BANDPASS*". This is especially useful when multiple intents are attached to scans. Finally, `observation` is an identifier to distinguish between different observing runs, mainly used for ALMA.

The `selectdata` parameter expands as usual, uncovering other selection sub-parameters:

```python
selectdata = True   # data selection parameters
field      = ''     # field names or field index
               # numbers (blank for all)
spw        = ''     # spectral windows:channels (blank for all)
timerange  = ''     # time range (blank for all)
urvrange   = ''     # uv range (blank for all)
antenna    = ''     # antenna/baselines (blank for all)
scan       = ''     # scan numbers (blank for all)
correlation= ''     # correlations (blank for all)
array      = ''     # (sub)array numbers (blank for all)
observation= ''     # Select by observation ID(s)
msselect  = ''     # MS selection (blank for all)
```
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Note that if selectdata=False these parameters are not used when the task is executed, even if set underneath.

The most common selectdata parameter to use is uvrange, which can be used to exclude longer baselines if the calibrator is resolved, or short baselines of the calibrator contains extended flux not accounted for in the model (e.g. §4.3.5.1).

See §2.3 for more on the selection parameters.

4.4.1.3 Prior Calibration and Correction: parang

These parameters control the on-the-fly application of various calibration or effect-based corrections prior to the solving process.

The parang parameter turns on the application of the antenna-based parallactic angle correction (‘P’) in the measurement equation. This is necessary for polarization calibration and imaging, or for cases where the parallactic angles are different for geographically spaced antennas and it is desired that the ordinary gain calibration not absorb the inter-antenna parallactic angle phase. When dealing with only the parallel-hand data (e.g. RR, LL, XX, YY), and an unpolarized calibrator model for a co-located array (e.g. the VLA or ALMA), you can set parang=False and save some computational effort. Otherwise, set parang=True to apply this correction.

See §4.3 for more on Prior Calibration, including how to invoke gaincurve and opacity correction using gencal.

4.4.1.4 Previous Calibration: gaintable, gainfield, interp and spwmap

Calibration tables that have already been determined can also be applied before solving for the new table:

docallib = False # Use traditional cal apply parameters
gaintable = []  # Gain calibration table(s) to apply on the fly
gainfield = []  # Select a subset of calibrators from gaintable(s)
interp = []     # Interpolation mode (in time) to use for each gaintable
spwmap = []     # Spectral windows combinations to form for gaintable(s)

This is controlled by the gaintable parameter, which takes a string or list of strings giving one or more calibration tables to pre-apply. For example,

```
gaintable = ['ngc5921.bcal','ngc5921.gcal']
```

specifies two tables, in this case bandpass and gain calibration tables respectively.

The other parameters key off gaintable, taking single values or lists, with an entry for each table in gaintable. The order is given by that in gaintable.

The gainfield parameter specifies which fields from the respective gaintable to select for apply. This is a list, with each entry a string or list of strings. The default ’’ for an entry means to use all in that table. For example,
gaintable = ['ngc5921.bcal', 'ngc5921.gcal']
gainfield = ['1331+305', ['1331+305', '1445+099']]

or using indices

gainfield = ['0', ['0', '1']]

to specify the field '1331+305' from the table 'ngc5921.bcal' and fields '1331+305' and '1445+099' from the second table 'ngc5921.gcal'. We could also have wildcarded the selection, e.g.

gainfield = ['0', '*']

taking all fields from the second table. And of course we could have used the default

gainfield = ['0', '']
or even

gainfield = ['0']

which is to take all for the second table in gaintable. In addition, gainfield can be specified by

gainfield = ['nearest']

which selects the calibrator that is the spatially closest (in sky coordinates) to each of the selected MS fields specified in the field parameter. This is particularly useful for running applycal with a number of different sources to be calibrated in a single run.

The interp parameter chooses the interpolation scheme to be used when pre-applying the solution in the tables. Interpolation in both time and frequency (for channel-dependent calibrations) are supported. The choices are currently 'nearest' and 'linear', and 'nearest', 'linear', cubic, and spline for frequency-dependent interpolation. Frequency-dependent interpolation is only relevant for channel-dependent calibration tables (like bandpasses) that are undersampled in frequency relative to the data.

- 'nearest' just picks the entry nearest in time or freq to the visibility in question;

- 'linear' interpolation calibrates each datum with calibration phases and amplitudes linearly interpolated from neighboring values. In the case of phase, this mode will assume that phase never jumps more than 180° between neighboring points, and so undersampled cycle-slips will not be corrected for. Solutions will not be extrapolated arbitrarily in time or frequency for data before the first solution or after the last solution; such data will be calibrated using 'nearest' to avoid unreasonable extrapolations.

- 'cubic' interpolation forms a 3rd-order polynomial that passes through the nearest 4 calibration samples (separately in phase and amplitude
- 'spline' interpolation forms a cubic spline that passes through the nearest 4 calibration samples (separately in phase and amplitude)

For each gaintable, specify the interpolation style in quotes, with the frequency-dependent interpolation style specified after a comma, if relevant.

If the uncalibrated phase is changing rapidly, a 'nearest' interpolation is not desirable. Usually, interp='linear' is the best choice. For example,

```plaintext
gaintable=['gain','bandpass']
interp = [ 'nearest', 'linear,spline' ]
```

uses nearest “interpolation” on the first table, and linear (in time) and spline (in freq) on the second.

The spwmap parameter sets the spectral window combinations to form for the gaintable(s). This is a list, or a list of lists, of integers giving the spw IDs to map. There is one list for each table in gaintable, with an entry for each ID in the MS. For example,

```plaintext
spwmap=[0,0,1,1] # apply from spw=0 to 0,1 and 1 to 2,3
```

for an MS with spw=0,1,2,3. For multiple gaintable, use lists of lists, e.g.

```plaintext
spwmap=[[0,0,1,1], [0,1,0,1]] # 2nd table spw=0 to 0,2 and 1 to 1,3
```

### 4.4.1.5 Solving: solint, combine, preavg, refant, minblperant, minsnr

The parameters controlling common aspects of the solution are:

```plaintext
solint = 'inf' # Solution interval: egs. 'inf', '60s' (see help)
combine = 'scan' # Data axes which to combine for solve (obs, scan, spw, and/or field)
preavg = -1.0 # Pre-averaging interval (sec) (rarely needed)
refant = '' # Reference antenna name(s)
minblperant = 4 # Minimum baselines _per antenna_ required for solve
minsnr = 3.0 # Reject solutions below this SNR
```

The time and frequency (if relevant) solution interval is given by solint. Optionally a frequency interval for each solution can be added after a comma, e.g. solint='60s,300Hz'. Time units are in seconds unless specified differently. Frequency units can be either channels or Hz and only make sense for bandpass of frequency dependent polarization calibration. The special values 'inf' and -1 specify an “infinite” solution interval encompassing the entire dataset, while 'int' or zero specify a solution every integration. You can use time quanta in the string, e.g. solint='1min' and solint='60s' both specify solution intervals of one minute. Note that 'm' is a unit of distance (meters); 'min' must be used to specify minutes. The solint parameter interacts with combine to determine whether the solutions cross scan or field boundaries.
The parameter controlling the scope of the solution is **combine**. For the default **combine=','** solutions will break at obsId, scan, field, and spw boundaries. Specification of any of these in **combine** will extend the solutions over the boundaries (up to the **solint**). For example, **combine='spw'** will combine spectral windows together for solving, while **combine='scan'** will cross scans, and **combine='obs,scan'** will use data across different observation IDs and scans (usually, obsIds consist of many scans, so it is not meaningful to combine obsIds without also combining scans). Thus, to do scan-based solutions (single solution for each scan), set

```python
    solint = 'inf'
    combine = ''
```

while

```python
    solint = 'inf'
    combine = 'scan'
```

will make a single solution for the entire dataset (for a given field and spw).

```python
    solint = 'inf,30ch'
```

will calculate a bandpass solution for each scan, averaging over 30 channels.

You can specify multiple choices for combination:

```python
    combine = 'scan,spw'
```

for example.

The reference antenna is specified by the **refant** parameter. A list of antennas can be provided to this parameter and if the first antenna is not present in the data, the next antenna in the list will be used, etc. It is useful to “lock” the solutions with time, effectively rotating (after solving) the phase of the gain solutions for all antennas such that the reference antenna’s phase is constant at zero. If the selected antenna drops out, another antenna will be selected for ongoing consistency in time (at its “current” value) until the refant returns, usually at a new value (not zero), which will be kept fixed thenceforth. You can also run without a reference antenna, but in this case the solutions will formally float with time; in practice, the first antenna will be approximately constant near zero phase. It is usually prudent to select an antenna in the center of the array that is known to be particularly stable, as any gain jumps or wanders in the **refant** will be transferred to the other antenna solutions. Also, it is best to choose a reference antenna that never drops out.

Setting a **preavg** time (only needed in **polcal**) will let you average data over periods shorter than the solution interval first before solving on longer timescales.

The minimum signal-to-noise ratio allowed for an acceptable solution is specified in the **minsnr** parameter. Default is **minsnr=3**. The **minblperant** parameter sets the minimum number of baselines to other antennas that must be preset for each antenna to be included in a solution. This enables control of the constraints that a solution will require for each antenna.
4.4.1.6 Action: append and solnorm

The following parameters control some things that happen after solutions are obtained:

solnorm = False  # Normalize solution amplitudes post-solve.
append = False   # Append solutions to (existing) table.

The solnorm parameter toggles on the option to normalize the solution after the solutions are obtained. The exact effect of this depends upon the type of solution. Not all tasks use this parameter.

One should be aware when using solnorm that if this is done in the last stage of a chain of calibration, then the part of the calibration that is “normalized” away will be lost. It is best to use this in early stages (for example in a first bandpass calibration) so that later stages (such as final gain calibration) can absorb the lost normalization scaling. It is not strictly necessary to use solnorm=True at all, but is sometimes helpful if you want to have a normalized bandpass for example.

The append parameter, if set to True, will append the solutions from this run to existing solutions in caltable. Of course, this only matters if the table already exists. If append=False and caltable exists, it will overwrite.

4.4.2 Spectral Bandpass Calibration (bandpass)

For channelized data, it is usually desirable to solve for the gain variations in frequency as well as in time. Variation in frequency arises as a result of non-uniform filter passbands or other frequency-dependent effects in signal transmission. It is usually the case that these frequency-dependent effects vary on timescales much longer than the time-dependent effects handled by the gain types 'G' and 'T'. Thus, it makes sense to solve for them as a separate term: 'B', using the bandpass task.

The inputs to bandpass are:

```python
# bandpass :: Calculates a bandpass calibration solution
vis = ''       # Name of input visibility file
caltable = ''  # Name of output gain calibration table
field = ''     # Select field using field id(s) or field name(s)
spw = ''       # Select spectral window/channels
intent = ''    # Select observing intent
selectdata = True # Other data selection parameters
timerange = ''  # Select data based on time range
uvrange = ''    # Select data within uvrange (default units meters)
antenna = ''    # Select data based on antenna/baseline
scan = ''      # Scan number range
observation = '' # Select by observation ID(s)
msselect = ''  # Optional complex data selection (ignore for now)
```
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solint = 'inf'  # Solution interval in time[,freq]
combine = 'scan'  # Data axes which to combine
# for solve (obs, scan, spw, and/or field)
refant = ''  # Reference antenna name(s)
minblperant = 4  # Minimum baselines _per _antenna_ required for solve
minsnr = 3.0  # Reject solutions below this
# SNR (only applies for bandtype = B)
solnorm = False  # Normalize average solution amplitudes to 1.0
bandtype = 'B'  # Type of bandpass solution (B or BPOLY)
fillgaps = 0  # Fill flagged solution channels by interpolation
smodel = []  # Point source Stokes parameters for source model.
append = False  # Append solutions to the (existing) table
docallib = False  # Use callib or traditional cal apply parameters
gaintable = []  # Gain calibration table(s) to apply on the fly
gainfield = []  # Select a subset of calibrators from gaintable(s)
interp = []  # Interpolation mode (in
# time) to use for each gaintable
spwmap = []  # Spectral windows
# combinations to form for gaintable(s)
parang = False  # Apply parallactic angle correction

Many of these parameters are in common with the other calibration tasks and are described above in §[4.4.1].

The bandtype parameter selects the type of solution used for the bandpass. The choices are 'B' and 'BPOLY'. The former solves for a complex gain in each channel in the selected part of the MS. See §[4.4.2.2] for more on 'B'. The latter uses a polynomial as a function of channel to fit the bandpass, and expands further to reveal a number of sub-parameters See §[4.4.2.3] for more on 'BPOLY'.

It is usually best to solve for the bandpass in channel data before solving for the gain as a function of time. However, if the gains of the bandpass calibrator observations are fluctuating over the timerange of those observations, then it can be helpful to first solve for the gains of that source with gaincal, and input these to bandpass via gaintable. See more below on this strategy.

We now describe the issue of bandpass normalization, followed by a description of the options bandtype='B' and bandtype='BPOLY'.

4.4.2.1 Bandpass Normalization

The solnorm parameter (§[4.4.1.6]) deserves more explanation in the context of the bandpass. Most users are used to seeing a normalized bandpass, where the mean amplitude is unity and fiducial phase is zero. The toggle solnorm=True allows this. However, the parts of the bandpass solution normalized away will be still left in the data, and thus you should not use solnorm=True if the bandpass calibration is the end of your calibration sequence (e.g. you have already done all the
gain calibration you want to). Note that setting solnorm=True will NOT rescale any previous calibration tables that the user may have supplied in gaintable.

You can safely use solnorm=True if you do the bandpass first (perhaps after a throw-away initial gain calibration) as we suggest above in §4.2, as later gain calibration stages will deal with this remaining calibration term. This does have the benefit of isolating the overall (channel independent) gains to the following gaincal stage. It is also recommended for the case where you have multiple scans on possibly different bandpass calibrators. It may also be preferred when applying the bandpass before doing gaincal and then fluxscale (§4.4.4), as significant variation of bandpass among antennas could otherwise enter the gain solution and make (probably subtle) adjustments to the flux scale.

We finally note that solnorm=False at the bandpass step in the calibration chain will still in the end produce the correct results. It only means that there will be a part of what we usually think of the gain calibration inside the bandpass solution, particularly if bandpass is run as the first step.

### 4.4.2.2 B solutions

Calibration type ’B’ differs from ’G’ only in that it is determined for each channel in each spectral window. It is possible to solve for it as a function of time, but it is most efficient to keep the ’B’ solving timescale as long as possible, and use ’G’ or ’T’ for frequency-independent rapid time-scale variations.

The ’B’ solutions are limited by the signal-to-noise ratio available per channel, which may be quite small. It is therefore important that the data be coherent over the time-range of the ’B’ solutions. As a result, ’B’ solutions are almost always preceded by an initial ’G’ or ’T’ solve using gaincal (§4.4.3). In turn, if the ’B’ solution improves the frequency domain coherence significantly, a ’G’ or ’T’ solution following it will be better than the original.

For example, to solve for a ’B’ bandpass using a single short scan on the calibrator, then

```python
default(‘bandpass’)  
vis = ‘n5921.ms’  
caltab = ‘n5921.bcal’  
gaintable = ’’  
gainfield = ’’  
interp = ’’  
field = ‘0’  
spw = ‘’  
selectdata = False  
bandtype = ‘B’  
solint = ‘inf’  
refant = ‘15’  
bandpass()  
```
On the other hand, we might have a number of scans on the bandpass calibrator spread over time, but we want a single bandpass solution. In this case, we could solve for and then pre-apply an initial gain calibration, and let the bandpass solution cross scans:

```python
gaintable = 'n5921.init.gcal'  # Our previously determined G table
gainfield = '0'
interp = 'linear'               # Do linear interpolation
solint = 'inf'                  # One interval over dataset
combine = 'scan'                # Solution crosses scans
```

Note that we obtained a bandpass solution for all channels in the MS. If explicit channel selection is desired, for example some channels are useless and can be avoided entirely (e.g. edge channels or those dominated by Gibbs ringing), then `spw` can be set to select only these channels, e.g.

```python
spw = '0:4~59'                  # channels 4-59 of spw 0
```

This is not so critical for ‘B’ solutions as for ‘BPOLY’, as each channel is solved for independently, and poor solutions at edges can be ignored.

If you have multiple time solutions, then these will be applied using whatever time interpolation scheme is specified in later tasks.

The `combine` parameter (§ 4.4.1.5) can be used to combine data across spectral windows, scans, and fields.

### 4.4.2.3 BPOLY solutions

For some observations, it may be the case that the SNR per channel is insufficient to obtain a usable per-channel ‘B’ solution. In this case it is desirable to solve instead for a best-fit functional form for each antenna using the `bandtype='BPOLY'` solver. The ‘BPOLY’ solver naturally enough fits (Chebychev) polynomials to the amplitude and phase of the calibrator visibilities as a function of frequency. Unlike ordinary ‘B’, a single common ‘BPOLY’ solution will be determined for all spectral windows specified (or implicit) in the selection. As such, it is usually most meaningful to select individual spectral windows for ‘BPOLY’ solves, unless groups of adjacent spectral windows are known a priori to share a single continuous bandpass response over their combined frequency range (e.g., PdBI data).

The ‘BPOLY’ solver requires a number of unique sub-parameters:

```python
bandtype     = 'BPOLY'  # Type of bandpass solution (B or BPOLY)
degamp       = 3      # Polynomial degree for BPOLY amplitude solution
degphase     = 3      # Polynomial degree for BPOLY phase solution
visnorm      = False  # Normalize data prior to BPOLY solution
maskcenter   = 0      # Number of channels in BPOLY to avoid in center of band
maskedge     = 0      # Percent of channels in BPOLY to avoid at each band edge
```
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The `degamp` and `degphase` parameters indicate the polynomial degree desired for the amplitude and phase solutions. The `maskcenter` parameter is used to indicate the number of channels in the center of the band to avoid passing to the solution (e.g., to avoid Gibbs ringing in central channels for PdBI data). The `maskedge` drops beginning and end channels. The `visnorm` parameter turns on normalization before the solution is obtained (rather than after for `solnorm`).

The `combine` parameter (§ 4.4.1.5) can be used to combine data across spectral windows, scans, and fields.

Note that `bandpass` will allow you to use multiple fields, and can determine a single solution for all specified fields using `combine='field'`. If you want to use more than one field in the solution it is prudent to use an initial `gaincal` using proper flux densities for all sources (not just 1Jy) and use this table as an input to `bandpass` because in general the phase towards two (widely separated) sources will not be sufficiently similar to combine them, and you want the same amplitude scale. If you do not include amplitude in the initial `gaincal`, you probably want to set `visnorm=True` also to take out the amplitude normalization change. Note also in the case of multiple fields, that the `BPOLY` solution will be labeled with the field ID of the first field used in the `BPOLY` solution, so if for example you point `plotcal` at the name or ID of one of the other fields used in the solution, `plotcal` does not plot.

For example, to solve for a `BPOLY` (5th order in amplitude, 7th order in phase), using data from field 2, with G corrections pre-applied:

```python
bandpass(vis='data.ms', # input data set
caltab='cal.BPOLY', #
spw='0:2~56', # Use channels 3-57 (avoid end channels)
field='0', # Select bandpass calibrator (field 0)
bandtype='BPOLY', # Select bandpass polynomials
degamp=5, # 5th order amp
degphase=7, # 7th order phase
gaintable='cal.G', # Pre-apply gain solutions derived previously
refant='14') #
```

### 4.4.2.4 What if the bandpass calibrator has a significant slope?

The bandpass calibrator can have a spectral slope that will change the spectral properties of the solutions. If the slope is significant, the best way is to model the slope and store that model in the bandpass calibrator MS. To do so, go through the normal steps of `bandpass` and the `gaincal` runs on the bandpass and flux calibrators, followed by `setjy` of the flux calibrator. The next step would be to use `fluxscale` on the bandpass calibrator to derive the slope of it. `fluxscale` can store this information in a python dictionary which is subsequently fed into a second `setjy` run, this time using the bandpass calibrator as the source and the derived slope (the python dictionary) as input. This step will create a source model with the correct overall spectral slope for the bandpass. Finally, rerun `bandpass` and all other calibration steps again, making use of the newly created internal bandpass model.
4.4.3 Complex Gain Calibration (gaincal)

The fundamental calibration to be done on your interferometer data is to calibrate the antenna-based gains as a function of time. Some of these calibrations are known beforehand (“a priori”) and others must be determined from observations of calibrators, or from observations of the target itself (“self-calibration”).

It is best to have determined a (constant or slowly-varying) “bandpass” from the frequency channels by solving for the bandpass (see above). Thus, the bandpass calibration table would be input to gaincal via the gaintable parameter (see below).

The gaincal task has the following inputs:

```plaintext
# gaincal :: Determine temporal gains from calibrator observations
vis = '' # Name of input visibility file
caltab = '' # Name of output gain calibration table
field = '' # Select field using field id(s) or field name(s)
spw = '' # Select spectral window/channels
intent = '' # Select observing intent
selectdata = True # Other data selection parameters
timerange = '' # Select data based on time range
uvrange = '' # Select data within uvrange (default units meters)
antenna = '' # Select data based on antenna/baseline
scan = '' # Scan number range
observation = '' # Select by observation ID(s)
msselect = '' # Optional complex data selection (ignore for now)

solint = 'inf' # Solution interval: egs. 'inf', '60s' (see help)
combine = '' # Data axes which to combine
preavg = -1.0 # Pre-averaging interval (sec) (rarely needed)
refant = '' # Reference antenna name(s)
minblperant = 4 # Minimum baselines _per antenna_ required for solve
minsnr = 3.0 # Reject solutions below this SNR
solnorm = False # Normalize average solution
# amplitudes to 1.0 (G, T only)
gaintype = 'G' # Type of gain solution (G,T,GSPLINE,K,KCROSS)
smodel = [] # Point source Stokes parameters for source model.
calmode = 'ap' # Type of solution: ('ap', 'p', 'a')
append = False # Append solutions to the (existing) table
docallib = False # Use callib or traditional cal apply parameters
gaintable = [] # Gain calibration table(s) to apply on the fly
gainfield = [] # Select a subset of calibrators from gaintable(s)
interp = [] # Temporal interpolation for
each gaintable (=linear)
spwmap = [] # Spectral windows
# combinations to form for gaintable(s)
parang = False # Apply parallactic angle correction on the fly
```

It is best to have determined a (constant or slowly-varying) “bandpass” from the frequency channels by solving for the bandpass (see above). Thus, the **bandpass** calibration table would be input to gaincal via the gaintable parameter (see below).
Data selection is done through the standard `field`, `spw`, `intent`, and `selectdata` expandable sub-parameters (see §2.3). The bulk of the other parameters are the standard solver parameters. See §4.4.1 above for a description of these.

The `gaintype` parameter selects the type of gain solution to compute. The choices are 'T', 'G', and 'GSPLINE'. The 'G' and 'T' options solve for independent complex gains in each solution interval (classic AIPS style), with 'T' enforcing a single polarization-independent gain for each co-polar correlation (e.g. RR and LL, or XX and YY) and 'G' having independent gains for these. See §4.4.3.1 for a more detailed description of 'G' solutions, and §4.4.3.2 for more on 'T'. The 'GSPLINE' fits cubic splines to the gain as a function of time. See §4.4.3.3 for more on this option.

### 4.4.3.1 Polarization-dependent Gain (G)

Systematic time-dependent complex gain errors are almost always the dominant calibration effect, and a solution for them is almost always necessary before proceeding with any other calibration. Traditionally, this calibration type has been a catch-all for a variety of similar effects, including: the relative amplitude and phase gain for each antenna, phase and amplitude drifts in the electronics of each antenna, amplitude response as a function of elevation (gain curve), and tropospheric amplitude and phase effects. In CASA, it is possible to handle many of these effects separately, as available information and circumstances warrant, but it is still possible to solve for the net effect using calibration type G.

Generally speaking, type G can represent any per-spectral window multiplicative polarization- and time-dependent complex gain effect downstream of the polarizers. (Polarization- and time-independent effects upstream of the polarizers may also be treated implicitly with G.) Multi-channel data (per spectral window) will be averaged in frequency before solving (use calibration type B to solve for frequency-dependent effects within each spectral window).

To solve for G on, say, fields 1 & 2, on a 90s timescale, and do so relative to gaincurve corrections:

```bash
gaincal('data.ms',
    caltable='cal.G', # Write solutions to disk file 'cal.G'
    field='0,1', # Restrict field selection
    solint=90.0, # Solve for phase and amp on a 90s timescale
    gaintable=['cal.gc'] # a gain curve table from gencal
    refant='3') #

plotcal('cal.G','amp') # Inspect solutions
```

These G solution will be referenced to antenna 4. Choose a well-behaved antenna that is located near the center of the array and is ever-present for the reference antenna. For non-polarization datasets, reference antennas need not be specified although you can if you want. If no reference antenna is specified, an effective phase reference that is an average over the data will be calculated and used. For data that requires polarization calibration, you must choose a reference antenna that has a constant phase difference between the right and left polarizations (e.g. no phase jumps or drifts). If no reference antenna (or a poor one) is specified, the phase reference may have jumps
in the R–L phase, and the resulting polarization angle response will vary during the observation, thus corrupting the polarization imaging.

To apply this solution, along with the gain curve correction, to the calibrators (fields 0,1) and the target source (field 2):

```python
applycal('data.ms',
    field='0,1,2', # Restrict field selection (cals + src)
    gaintable=['cal.gc','cal.G']) # Apply gc and G solutions to correct data
```

The calibrated data is written to the **CORRECTED_DATA** column, with **calwt=True** by default. This parameter can also be a list of Boolean values for which each entry then controls the calculation of weights based on each individual input calibration table. **calwt=False** will recompute the weights form the **SIGMA** column, thus resetting the weights to their original value.

**Alert**: Current (as of February 2014) Jansky VLA data has no calibrated weights (unless they are computed from switched power calibration). To avoid trouble, **calwt=False** should be set for those data sets. Older, pre-upgrade VLA data should still be calibrated with **calwt=True**.

### 4.4.3.2 Polarization-independent Gain (T)

At high frequencies, it is often the case that the most rapid time-dependent gain errors are introduced by the troposphere, and are polarization-independent. It is therefore unnecessary to solve for separate time-dependent solutions for both polarizations, as is the case for 'G'. Calibration type 'T' is available to calibrate such tropospheric effects, differing from 'G' only in that a single common solution for both polarizations is determined. In cases where only one polarization is observed, type 'T' is adequate to describe the time-dependent complex multiplicative gain calibration.

In the following example, we assume we have a 'G' solution obtained on a longish timescale (longer than a few minutes, say), and we want a residual 'T' solution to track the polarization-independent variations on a very short timescale:

```python
gaincal('data.ms',
    caltable='cal.T', # Specify output table name
    gaintype='T', # Solve for T
    field='0,1', # Restrict data selection to calibrators
    solint=3.0, # Obtain solutions on a 3s timescale
    gaintable='cal120.G') # Pre-apply prior G solution
```

For dual-polarization observations, it will always be necessary to obtain a 'G' solution to account for differences and drifts between the polarizations (which traverse different electronics), but solutions for rapidly varying polarization-independent effects such as those introduced by the troposphere will be optimized by using 'T'. Note that 'T' can be used in this way for self-calibration purposes, too.
4.4.3.3 GSPLINE solutions

At high radio frequencies, where tropospheric phase fluctuates rapidly, it is often the case that there is insufficient signal-to-noise ratio to obtain robust ‘G’ or ‘T’ solutions on timescales short enough to track the variation. In this case it is desirable to solve for a best-fit functional form for each antenna using the ‘GSPLINE’ solver. This fits a time-series of cubic B-splines to the phase and/or amplitude of the calibrator visibilities.

The combine parameter (§4.4.1.5) can be used to combine data across spectral windows, scans, and fields. Note that if you want to use combine='field', then all fields used to obtain a ‘GSPLINE’ amplitude solution must have models with accurate relative flux densities. Use of incorrect relative flux densities will introduce spurious variations in the ‘GSPLINE’ amplitude solution.

The ‘GSPLINE’ solver requires a number of unique additional parameters, compared to ordinary ‘G’ and ‘T’ solving. The sub-parameters are:

- gaintype = 'GSPLINE' # Type of solution (G, T, or GSPLINE)
- splinetime = 3600.0 # Spline (smooth) timescale (sec), default=1 hours
- npointaver = 3 # Points to average for phase wrap (okay)
- phasewrap = 180 # Wrap phase when greater than this (okay)

The duration of each spline segment is controlled by splinetime. The actual splinetime will be adjusted such that an integral number of equal-length spline segments will fit within the overall range of data.

Phase splines require that cycle ambiguities be resolved prior to the fit; this operation is controlled by npointaver and phasewrap. The npointaver parameter controls how many contiguous points in the time-series are used to predict the cycle ambiguity of the next point in the time-series, and phasewrap sets the threshold phase jump (in degrees) that would indicate a cycle slip. Large values of npointaver improve the SNR of the cycle estimate, but tend to frustrate ambiguity detection if the phase rates are large. The phasewrap parameter may be adjusted to influence when cycles are detected. Generally speaking, large values (> 180°) are useful when SNR is high and phase rates are low. Smaller values for phasewrap can force cycle slip detection when low SNR conspires to obscure the jump, but the algorithm becomes significantly less robust. More robust algorithms for phase-tracking are under development (including fringe-fitting).

For example, to solve for ‘GSPLINE’ phase and amplitudes, with splines of duration 600 seconds,

```
 gain('data.ms',
    caltable='cal.spline.ap',
    gaintype='GSPLINE' # Solve for GSPLINE
    calmode='ap' # Solve for amp & phase
    field='0,1', # Restrict data selection to calibrators
    splinetime=600.) # Set spline timescale to 10min
```

**ALERT**: The ‘GSPLINE’ solutions cannot yet be used in fluxscale. You should do at least some ‘G’ amplitude solutions to establish the flux scale, then do ‘GSPLINE’ in phase before or after to fix up the short timescale variations. Note that the “phase tracking” algorithm in ‘GSPLINE’ needs some improvement.
4.4.3.4 Antenna Delays — 'K' solutions

gaintype='K' solves for simple antenna-based delays via Fourier transforms of the spectra on baselines to the reference antenna. This is not a global fringe fit but will be useful for deriving delays from data of reasonable snr. If combine includes 'spw', multi-band delays solved jointly from all selected spectral windows will be determined, and will be identified with the first spectral window id in the output caltable. When applying a multi-band delay table, spwmap is required to distribute the solutions to all spectral windows.

After solving for delays, a subsequent bandpass is recommended to describe higher-order channel-dependent variation in the phase (and amplitude).

4.4.3.5 Cross-Hand Delays — 'KCROSS' solutions

gaintype='KCROSS' solves for a global cross-hand delay. Use parang=T and apply prior gain and bandpass solutions. Alert: Multi-band delays are not yet supported for KCROSS solutions.

4.4.4 Establishing the Flux Density Scale (fluxscale)

The 'G' or 'T' solutions obtained from calibrators for which the flux density was unknown and assumed to be 1 Jansky are correct in a time- and antenna- relative sense, but are mis-scaled by a factor equal to the inverse of the square root of the true flux density. This scaling can be corrected by enforcing the constraint that mean gain amplitudes determined from calibrators of unknown flux density should be the same as determined from those with known flux densities. The fluxscale task exists for this purpose.

The inputs for fluxscale are:

```python
# fluxscale :: Bootstrap the flux density scale from standard calibrators
vis = '' # Name of input visibility file (MS)
caltab = '' # Name of input calibration table
fluxtable = '' # Name of output, flux-scaled calibration table
reference = [''] # Reference field name(s) (transfer flux scale FROM)
transfer = [''] # Transfer field name(s) (transfer flux scale TO), '' -> all
listfile = '' # Name of listfile that contains the fit information. Default is (no file).
append = False # Append solutions?
refspwmap = [-1] # Scale across spectral window boundaries. See help
# fluxscale
gainthreshold = -1.0 # Threshold (% deviation from the median) on gain amplitudes to be used in the flux scale calculation
antenna = '' # antennas to include/exclude
incremental = False # incremental caltable
fitorder = 1 # order of spectral fitting
display = False # display some statistics of flux scaling
```
Before running *fluxscale*, one must have first run *setjy* for the reference sources and run a gaincal that includes reference and transfer fields. After running *fluxscale* the output fluxtable caltable will have been scaled such that the correct scaling will be applied to the transfer sources.

For example, given a 'G' table, e.g. 'cal.G', containing solutions for a flux density calibrator (in this case '3C286') and for one or more gain calibrator sources with unknown flux densities (in this example '0234+285' and '0323+022'):

```python
fluxscale(vis='data.ms',
    caltable='cal.G', # Select input table
    fluxtable= 'cal.Gflx', # Write scaled solutions to cal.Gflx
    reference='3C286', # 3C286 = flux calibrator
    transfer='0234+258, 0323+022') # Select calibrators to scale
```

The output table, 'cal.Gflx', contains either the scaling factors alone (incremental=T) to be used alongside with the input gain table 'cal.G', or a scaled version of the gain table (incremental=F), that replaces it for the execution of applycal.

Note that the assertion that the gain solutions are independent of the calibrator includes the assumption that the gain amplitudes are strictly not systematically time-dependent in any way. While synthesis antennas are designed as much as possible to achieve this goal, in practice, a number of effects conspire to frustrate it. When relevant, it is advisable to pre-apply gaincurve and opacity corrections when solving for the 'G' solutions that will be flux-scaled (see §4.3 and §4.4.1.3). When the 'G' solutions are essentially constant for each calibrator separately, the fluxscale operation is likely to be robust.

*fluxscale* will report the fluxes of each spw for each source. In addition, it will attempt a fit across the spws of each source and report a spectral index and curvature ($S \propto (\nu/\nu_0)^{\alpha+\beta\log(\nu/\nu_0)}$). This information can be subsequently used to build up a model for the spectral slope of a calibrator with the *setjy* task if required.

The *fluxscale* task can be executed on either 'G' or 'T' solutions, but it should only be used on one of these types if solutions exist for both and one was solved relative to the other (use fluxscale only on the first of the two).

**ALERT:** The 'GSPLINE' option is not yet supported in *fluxscale* (see §4.4.3.3).

If the reference and transfer fields were observed in different spectral windows, the refspwmap parameter may be used to achieve the scaling calculation across spectral window boundaries.

The refspwmap parameter functions similarly to the standard spwmap parameter (§4.4.1.4), and takes a list of indices indicating the spectral window mapping for the reference fields, such that refspwmap[i]=j means that reference field amplitudes from spectral window j will be used for spectral window i.

**Note:** You should be careful when you have a dataset with spectral windows with different bandwidths, and you have observed the calibrators differently in the different spw. The flux-scaling will probably be different in windows with different bandwidths.

For example,
fluxscale(vis='data.ms',
caltablename='cal.G',
fluxtable='cal.Gflx',
reference='3C286',
transfer='0234+258,0323+022',
refspwmap=[0,0,0])

will use spw=0 to scale the others, while in

fluxscale(vis='data.ms',
caltablename='cal.G',
fluxtable='cal.Gflx',
reference='3C286',
transfer='0234+285, 0323+022',
refspwmap=[0,0,1,1])

the reference amplitudes from spectral window 0 will be used for spectral windows 0 and 1 and reference amplitudes from spectral window 2 will be used for spectral windows 2 and 3.

### 4.4.4.1 Using Resolved Calibrators

If the flux density calibrator is resolved, the assumption that it is a point source will cause solutions on outlying antennas to be biased in amplitude. In turn, the fluxscale step will be biased on these antennas as well. In general, it is best to use model for the calibrator, but if such a model is not available, it is important to limit the solution on the flux density calibrator to only the subset of antennas that have baselines short enough that the point-source assumption is valid. This can be done by using antenna and uvrange selection when solving for the flux density calibrator. For example, if antennas 1 through 8 are the antennas among which the baselines are short enough that the point-source assumption is valid, and we want to be sure to limit the solutions to the use of baselines shorter than 15000 wavelengths, then we can assemble properly scaled solutions for the other calibrator as follows (note: specifying both an antenna and a uvrange constraint prevents inclusion of antennas with only a small number of baselines within the specified uvrange from being included in the solution; such antennas will have poorly constrained solutions):

As an example, we first solve for gain solutions for the flux density calibrator (3C286 observed in field 0) using a subset of antennas

gaincal(vis='data.ms',
caltablename='cal.G',
field='0',
selectdata=True,
 antenna='0~7',
uvrange='0~15klambda',
solint=90)

Now solve for other calibrator (0234+285 in field 1) using all antennas (implicitly) and append these solutions to the same table
gaincal(vis='data.ms',
    caltable='cal.G',         # write solutions to cal.G
    field='1',
    solint=90,
    append=T)                  # Set up to write to the same table

Finally, run fluxscale to adjust scaling

fluxscale(vis='data.ms',
          caltable='cal.G',  # Input table with unscaled cal solutions
          fluxtable='cal.Gflx',  # Write scaled solutions to cal.Gflx
          reference='3C286',   # Use 3c286 as ref with limited uvrange
          transfer='0234+285')  # Transfer scaling to 0234+285

The fluxscale calculation will be performed using only the antennas common to both fields, but
the result will be applied to all antennas on the transfer field. Note that one can nominally get by
only with the uvrange selection, but you may find that you get strange effects from some antennas
only having visibilities to a subset of the baselines and thus causing problems in the solving.

4.4.5 Instrumental Polarization Calibration (D,X)

Full support for instrumental polarization calibration for the circular feed basis (e.g., VLA) is
provided in CASA. Support for the linear feed basis (e.g., ALMA) is now practical (as of v4.0)
and is also described below. The linear feed basis treatment will continue to be expanded and
streamlined for the v4.3 release.

The inputs to polcal are:

```plaintext
# polcal :: Determine instrumental polarization calibrations
vis = ''    # Name of input visibility file
caltab = '' # Name of output gain calibration table
field = ''  # Select field using field id(s) or field name(s)
spw = ''   # Select spectral window/channels
intent = '' # Select observing intent
selectdata = True  # Other data selection parameters
timerange = ''  # Select data based on time range
uvrange = ''   # Select data within uvrange (default units meters)
antenna = ''   # Select data based on antenna/baseline
scan = ''     # Scan number range
observation = ''  # Select by observation ID(s)
msselect = ''  # Optional complex data selection (ignore for now)
solint = 'inf'  # Solution interval
combine = 'obs,scan'  # Data axes which to combine
    # for solve (obs, scan, spw, and/or field)
preavg = 300.0  # Pre-averaging interval (sec)
refant = ''   # Reference antenna name(s)
minbaselants = 4 # Minimum baselines _per antenna_ required for solve
```
minsnr = 3.0 # Reject solutions below this SNR
poltype = 'D+QU' # Type of instrumental polarization solution (see help)
# Type of instrumental polarization solution (see help)
smodel = [] # Point source Stokes parameters for source model.
append = False # Append solutions to the (existing) table
docallib = False # Use calib or traditional cal apply parameters
gaintable = [] # Gain calibration table(s) to apply
gainfield = [] # Select a subset of calibrators from gaintable(s)
interp = [] # Interpolation mode (in time) to use for each gaintable
spwmap = [] # Spectral windows combinations to form for gaintable(s)

The polcal task uses many of the standard calibration parameters as described above in §4.4.1.

The key parameter controlling polcal is poltype. The choices are:

'D' — Solve for instrumental polarization (leakage D-terms), using the transform of an IQU model; requires no parallactic angle coverage, but if the source polarization is non-zero, the gain calibration must have the correct R-L phase registration. (Note: this is unlikely, so just use 'D+X' to let the position angle registration float.) This will produce a calibration table of type D.

'D+X' — Solve for instrumental polarization D-terms and the polarization position angle correction, using the transform of an IQU model; this mode requires at least 2 distinct parallactic angles to separate the net instrumental polarization from the PA. This will produce a calibration table of type 'D'. ALERT: no table of type 'X' will be produced, so you must follow this by a run of polcal with polmode='X' (see below).

'D+QU' — Solve for instrumental polarization and source Q + iU; requires at least 3 distinct parallactic angles to separate the net instrumental polarization from the source Q and U. Effectively sets the polarization PA to the value if the R-L phase difference were 0°. This will produce a calibration table of type 'D'.

'X' — Solve only for the position angle correction; best to use this after getting the D-terms from one of the above modes. Requires the observation of a calibrator with known Q + iU (or at least known U/Q). This will produce a calibration table of type 'X'.

'Dflls' — A specialized mode for instrumental polarization solving for the linear feed basis. This will probably be consolidated with other options in a future release.

There are channelized solution modes for the above options. For example, substitute 'Df' for 'D' in the 'D*' modes described above to get a channelized D-term solution; substitute 'Xf' for 'X' to get channelized position angle correction.

ALERT: polcal will obtain a separate D-term solution for each field supplied to it. This limitation will be relaxed in the future, enabling more sensitive solutions.
4.4.5.1 Heuristics and Strategies for Polarization Calibration

**ALERT:** This section concentrates on polarization calibration for the circular feed basis. It will be generalized to include the linear feed basis for the v4.3 release. See §4.4.5.4 for the currently supported processing steps for the linear feed basis.

Fundamentally, with good ordinary gain (and bandpass, if relevant) calibration already in hand, good polarization calibration must deliver both the instrumental polarization and position angle calibration. An unpolarized source can deliver only the first of these, but does not require parallactic angle coverage. A polarized source can only deliver the position angle calibration also if its polarization is known a priori. Sources that are polarized, but with unknown polarization, must always be observed with sufficient parallactic angle coverage, where ”sufficient” is determined by SNR and the details of the solving mode.

These principles are stated assuming the instrumental polarization solution is solved using the ”linear approximation” where cross-terms in more than a single product of the instrumental or source polarizations are ignored in the Measurement Equation (see §E). A more general non-linearized solution, with sufficient SNR, may enable some relaxation of the requirements indicated here, and modes supporting such an approach are currently under development.

For instrumental polarization calibration, there are 3 types of calibrator choice:

### CASA Polarization Calibration Modes

<table>
<thead>
<tr>
<th>Cal Polarization</th>
<th>Parallactic Angles</th>
<th>model</th>
<th>polmode</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>unpolarized</td>
<td>any</td>
<td>set $Q = U = 0$</td>
<td>'D' or 'Df'</td>
<td>D-terms only</td>
</tr>
<tr>
<td>known non-zero</td>
<td>2+ scans</td>
<td>set $Q,U$</td>
<td>'D+X' or 'Df+X'</td>
<td>D-terms and PA</td>
</tr>
<tr>
<td>unknown</td>
<td>2+ scans</td>
<td>ignored</td>
<td>'D+QU' or 'Df+QU'</td>
<td>D-terms and source</td>
</tr>
</tbody>
</table>

Note that the parallactic angle ranges spanned by the scans in the modes that require this should be large enough to give good separation between the components of the solution. In practice, 60° is a good target.

Each of these solutions should be followed with a 'X' solution on a source with known polarization position angle (and correct $Q + iU$ in the model). **ALERT:** polmode='D+X' will soon be enhanced to deliver this automatically.

The polcal task will solve for the 'D' or 'X' terms using the model visibilities that are in the model attached to the MS. Calibration of the parallel hands must have already been carried out using gaincal and/or bandpass in order to align the phases over time and frequency. This calibration must be supplied through the gaintable parameters, but any cal-tables to be used in polcal must agree (e.g. have been derived from) the data in the DATA column and the FT of the model. Thus, for example, one would not use the cal-table produced by fluxscale as the rescaled amplitudes would no longer agree with the contents of the model.

Be careful when using resolved calibrators for polarization calibration. A particular problem is if the structure in Q and U is offset from that in I. Use of a point model, or a resolved model for I but point models for Q and U, can lead to errors in the 'X' calibration. Use of a uvrange will
help here. The use of a full-Stokes model with the correct polarization is the only way to ensure a correct calibration if these offsets are large.

4.4.5.2 A Note on channelized polarization calibration

When your data has more than one channel per spectral window, it is important to note that the calibrator polarization estimate currently assumes the source polarization signal is coherent across each spectral window. In this case, it is important to be sure there is no large cross-hand delay still present in your data. Unless the online system has accounted for cross-hand delays (typically intended, but not always achieved), the gain and bandpass calibration will only correct for parallel-hand delay residuals since the two polarizations are referenced independently. Good gain and bandpass calibration will typically leave a single cross-hand delay (and phase) residual from the reference antenna. Plots of cross-hand phases as a function of frequency for a strongly polarized source (i.e., that dominates the instrumental polarization) will show the cross-hand delay as a phase slope with frequency. This slope will be the same magnitude on all baselines, but with different sign in the two cross-hand correlations. This cross-hand delay can be estimated using the `gaintype='KCROSS'` mode of `gaincal` (in this case, using the strongly polarized source 3C286):

```python
default('gaincal')
vis = 'polcal_20080224.cband.all.ms'
caltab = 'polcal.xdelcal'
field = '3C286'
spw = '',
solint = 'inf'
combine = 'scan'
refant = 'VA15'
smodel = [1.0,0.11,0.0,0.0]
gaintype = 'KCROSS'
gaintable = ['polcal.gcal','polcal.bcal']
gaincal()
```

Note that `smodel` is used to specify that 3C286 is polarized; it is not important to specify this polarization stokes parameters correctly, as only the delay will be solved for (not any absolute position angle or amplitude scaling). The resulting solution should be carried forward and applied along with the gain (.gcal) and bandpass (.bcal) solutions in subsequent polarization calibration steps.

4.4.5.3 A Polarization Calibration Example - Circular Feed Basis (e.g., VLA $\nu > 1$ GHz)

In the following example, we do a standard 'D+QU' solution on the bright source BLLac ($2202+422$) which has been tracked through a range in parallactic angle:

```python
default('polcal')
vis = 'polcal_20080224.cband.all.ms'
```
caltable = 'polcal.pcal'
field = '2202+422'
spw = ''
solint = 'inf'
combine = 'scan'
preavg = 300.0
refant = 'VA15'
minsnr = 3
poltype = 'D+QU'
gaintable = ['polcal.gcal','polcal.bcal','polcal.xdelcal']
gainfield = ['']
polcal()

This assumes setjy and gaincal have already been run. Note that the original gain-calibration table is used in gaintable so that what is in the model is in agreement with what is in the gaintable, rather than using the table resulting from fluxscale.

Now, we need to set the R-L phase using a scan on 3C48 (0137+331):

default('polcal')
vis = 'polcal_20080224.cband.all.ms'
caltable = 'polcal.polx'
field = '0137+331'
refant = 'VA15'
minsnr = 3
poltype = 'X'
smodel = [1.0,-0.0348,-0.0217,0.0] # the fractional Stokes for 3C48
gaintable = ['polcal.gcal','polcal.bcal','polcal.xdelcal','polcal.pcal']
polcal()

Note that the fractional polarization of 3C48 has been properly specified in smodel here.

If, on the other hand, we had a scan on an unpolarized bright source, for example 3C84 (0319+415), we could use this to calibrate the leakages:

default('polcal')
vis = 'polcal_20080224.cband.all.ms'
caltable = 'polcal.pcal'
field = '0319+415'
refant = 'VA15'
poltype = 'D'
gaintable = ['polcal.gcal','polcal.bcal','polcal.xdelcal']
polcal()

We would then do the 'X' calibration as before (but using this D-table in gaintable).

4.4.5.4 A Polarization Calibration Example - Linear Feed Basis (e.g., ALMA, VLA $\nu < 1$ GHz)

CASA v4.0.0 introduces supports for instrumental polarization calibration for the linear feed basis at a level that is now practical for the general user. Some details remain to be implemented with
full flexibility, and much of what follows will be streamlined for the v4.1 release.

Calibrating the instrumental polarization for the linear feed basis is somewhat more complicated than the circular feed basis because the polarization effects (source and instrument) appear in all four correlations at first or zeroth order (whereas for circular feeds, the polarization information only enters the parallel hand correlations at second order). As a result, e.g., the time-dependent gain calibration will be distorted by any non-zero source polarization, and some degree of iteration will be required to isolate the gain calibration if the source polarization is not initially known. These complications can actually be used to advantage in solving for the instrumental calibration; in can be shown, for example, that a significantly linearly polarized calibrator enables a better instrumental polarization solution than an unpolarized calibrator.

In the following example, we show the processing steps for calibrating the instrumental polarization using a strongly (> 5%) polarized point-source calibrator (which is also the time-dependent gain calibrator) that has been observed over a range of parallactic angle (a single scan is not sufficient). We assume that we have calibrated the gain, bandpass, and cross-hand delay as described above, and that the gain calibration (polcal.gcal) was obtained assuming the calibrator was unpolarized.

First, we import some utility functions from the CASA recipes area:

from recipes.almapolhelpers import *

Since the gain calibrator was assumed unpolarized, the time-dependent gain solutions contain information about the source polarization. This can be seen by plotting the amp vs. time for this table using poln='/'. The antenna-based polarization amplitude ratios will reveal the sinusoidal (in parallactic angle) of the source polarization. Run a utility method (qufromgain()) to extract the apparent source polarization estimates for each spw:

qu=qufromgain('polcal.gcal')

The source polarization reported for all spws should be reasonably consistent. This estimate is not as good as can be obtained from the cross-hands (see below) since it relies on the gain amplitude polarization ratio being stable which may not be precisely true. However, this estimate will be useful in resolving an ambiguity that occurs in the cross-hand estimates.

Next we estimate both the XY-phase offset and source polarization from the cross-hands. The XY-phase offset is a spectral phase-only bandpass relating the X and Y systems of the reference antenna. The cross-hand delay solved for above represents a systematic component (linear phase in frequency). If the XY-phase is solved for in a channel-dependent manner (as below), it is strictly not necessary to have solved for the cross-hand delay above, but it does not hurt (at it allows reasonably coherent channel averages for data examination). The source polarization occurs in the cross-hands as a sinusoidal function of parallactic angle that is common to both cross-hands on all baselines (for a point-source). If the XY-phase bandpass is uniformly zero, then the source linear polarization function will occur entirely in the real part of the cross-hand visibilities. Non-zero XY-phase has the effect of rotating the source linear polarization signature partially into the imaginary part, where circular (and instrumental) polarization occur (cf. the circular feed basis where the cross-hand
phase merely rotates the position angle of linear polarization. The following solve averages all baselines together and first solves for a channelized XY-phase (the slope of the source polarization function in the complex plane), then corrects the slope and solves for a channel-averaged source polarization. This calibration is obtained using `gaintype='XYf+QU'` in `gaincal`:

```python
default('gaincal')
vis = 'polcal_linfeed.ms'
caltab = 'polcal.xy0amb' # possibly with 180deg ambiguity
field = '1' # the calibrator
solint = 'inf'
combine = 'scan'
preev = 200.0 # minimal parang change
smodel = [1,0,1,0] # non-zero U assumed
gaintype = 'XYf+QU'
gaintable = ['polcal.gcal','polcal.bcal','polcal.xdelcal']
gaincal()
```

Note that we imply non-zero Stokes U in `smodel`; this is to enforce the assumption of non-zero source polarization signature in the cross-hands in the ratio of data and model. This solve will report the center-channel XY-phase and apparent Q,U for each spw. The Q,U results should be recognizable in comparison to that reported by `qufromgain()` above. However, since the XY-phase has a 180 degree ambiguity (you can rotate the source polarization signature to lie entirely in the visibility real part by rotating clockwise or counter-clockwise), some or all spw QU estimates may have the wrong sign. We correct this using the `xyamb()` utility method, using the `qu` obtained from `qufromgain()` above (which is not ambiguous):

```python
S=xyamb(xy='polcal.xy0amb',qu=qu,xyout='polcal.xy0')
```

The python variable `S` now contains the mean source model (Stokes I = 1; fractional Q,U; V=0) that can be used in a revision of the gain calibration and instrumental polarization calibration.

Next we revise the gain calibration using the full polarization source model:

```python
default('gaincal')
vis = 'polcal_linfeed.ms'
caltab = 'polcal.gcal1'
field = '1'
solint = 'int' # or whatever was used previously
smodel = S # obtained from xyamb
gaintype = 'G'
gaintable = ['polcal.bcal']
parang = T # so source poln properly rotated
gaincal()
```

Note that `parang=T` so that the supplied source linear polarization is properly rotated in the parallel-hand visibility model. This new gain solution can be plotted with `poln='/'` as above to
show that the source polarization is no longer distorting it. Also, if \texttt{qufromgain} is run on this new gain table, the reported source polarization should be statistically indistinguishable from zero.

Finally, we can now solve for the instrumental polarization:

```python
default('polcal')
vis = 'polcal_linfeed.ms'
caltab = 'polcal.dcal'
field = '1'
solint = 'inf'
combine = 'scan'
preavg = 200
poltype = 'Dflls' # freq-dep LLS solver
refant = '' # no reference antenna
smodel = S
gaintable = ['polcal.gcal1','polcal.bcal','polcal.xdelcal','polcal.xy0']
polcal()
```

Note that no reference antenna is used since this solve will produce an absolute instrumental polarization solution that is registered to the assumed source polarization (S) and prior calibrations. Applying a refant (referring all instrumental polarization terms to a reference antenna’s X feed, which would then be assumed perfect) would, in fact, discard valid information about the imperfections in the reference antenna’s X feed. (Had we used an unpolarized calibrator, we would not have a valid xy-phase solution, nor would we have had access to the absolute instrumental polarization solution demonstrated here.)

A few points:

- Since the gain, bandpass, and XY-phase calibrations were obtained prior to the instrumental polarization solution and maybe distorted by it, it is generally desirable to resolve for them using the instrumental polarization solution. In effect, this means iterating the sequence of calibration steps using all of the best of the available information at each stage, including the source polarization (and \texttt{parang=T}). This is a generalization of traditional self-calibration. For the CASA v4.1 release, we expect to provide utility methods for iteration.

- If the source linear polarization fraction and position angle is known \textit{a priori}, the processing steps outlined above can be amended to use that source polarization assertion in the gain and instrumental calibration solves. The \texttt{qufromgain()} method is not needed (but can be used to verify assumptions), the \texttt{gaincal(...,gaintype='XYf+QU',...)} should not be altered (parallactic angle coverage is still required!), and the \texttt{xyamb()} run should use the \textit{a priori} polarization for \texttt{qu}. If there is likely to be a large systematic offset in the mean feed position angle, iteration of the gain, bandpass, and instrumental polarization terms is required to properly isolate the calibration effects.

- Note that the above process does not explicitly include a position angle calibration. In effect, the estimated source polarization sets the mean feed position angle as the reference position angle, and this is usually within a degree or so of optimal. If your mean X feed position angle
is not \(\sim 0\) degrees, and your MS does not account for the offset in its FEED subtable, be careful in your interpretation of the final position angle. Currently, the circular feed-specific position angle calibration modes of \texttt{polcal} (\texttt{poltype='X'} or \texttt{'Xf'}) will not properly handle the linear feed basis; this will be fixed in the CASA v4.1 release.

A full processing example for linear feed basis polarimetry is under development and will be distributed with an upcoming CASA release.

### 4.4.6 Baseline-based Calibration (\texttt{blcal})

You can use the \texttt{blcal} task to solve for baseline-dependent (non-closing) errors. **WARNING:** this is in general a very dangerous thing to do, since baseline-dependent errors once introduced are difficult to remove. You must be sure you have an excellent model for the source (better than the magnitude of the baseline-dependent errors).

The inputs are (note that \texttt{blcal} does not yet use the \texttt{docallib} parameter):

```
# blcal :: Calculate a baseline-based calibration solution (gain or bandpass)
vis = ' ' # Name of input visibility file
caltab = ' ' # Name of output gain calibration table
field = ' ' # Select field using field id(s) or field name(s)
spw = ' ' # Select spectral window/channels
intent = ' ' # Select observing intent
selectdata = False # Other data selection parameters
solint = 'inf' # Solution interval
combine = 'scan' # Data axes which to combine for solve (scan, spw, # and/or field)
freqdep = False # Solve for frequency dependent solutions
calmode = 'ap' # Type of solution" ('ap', 'p', 'a')
solnorm = False # Normalize average solution amplitudes to 1.0
gaintable = [''] # Gain calibration table(s) to apply on the fly
gainfield = [''] # Select a subset of # calibrators from gaintable(s)
interp = [''] # Interpolation mode (in # time) to use for each gaintable
spwmap = [] # Spectral windows combinations to form for # gaintable(s)
gaincurve = False # Apply internal VLA antenna # gain curve correction
opacity = [] # Opacity correction to apply (neper), per spw parang = False # Apply parallactic angle correction
```

The \texttt{freqdep} parameter controls whether \texttt{blcal} solves for “gain” (\texttt{freqdep=False}) or “bandpass” (\texttt{freqdep=True}) style non-closing calibration.

Other parameters are the same as in other calibration tasks. These common calibration parameters are described in \S 4.4.1.
4.5 Plotting and Manipulating Calibration Tables

At some point, the user should examine (plotting or listing) the calibration solutions. Calibration tables can also be manipulated in various ways, such as by interpolating between times (and sources), smoothing of solutions, and accumulating various separate calibrations into a single table.

4.5.1 Plotting Calibration Solutions (plotcal)

The `plotcal` task is available for examining solutions of all of the basic solvable types (G, T, B, D, M, MF, K). The inputs are:

```
# plotcal :: An all-purpose plotter for calibration results:

caltablaee = '' # Name of input calibration table
xaxis = '' # Value to plot along x axis (time,chan,amp,phase,real,imag,snr)
yaxis = '' # Value to plot along y axis (amp,phase,real,imag,snr)
poln = '' # Polarization to plot (RL,R,L,XY,X,Y,/)  
field = '' # Field names or index: '='all, '3C286,P1321*', '0~3'
antenna = '' # Antenna selection. E.g., antenna='3^5'
spw = '' # Spectral window: '='all, '0,1' means spw 0 and 1

timerange = '' # Time selection '='all
subplot = 111 # Panel number on display screen (yxn)
overplot = False # Overplot solutions on existing display

clearpanel = 'Auto' # Specify if old plots are cleared or not
iteration = '' # Iterate on antenna,time,spw,field

plotrange = [] # plot axes ranges: [xmin,xmax,ymin,ymax]

showflags = False # If true, show flags
plotsymbol = '.' # pylab plot symbol
plotcolor = 'blue' # initial plotting color
markersize = 5.0 # size of plot symbols
fontsize = 10.0 # size of label font
showgui = True # Show plot on gui
figfile = '' # ''= no plot hardcopy, otherwise supply name
```

**ALERT:** Currently, `plotcal` needs to know the MS from which `caltablaee` was derived to get indexing information. It does this using the name stored inside the table, which does not include the full path, but assumes the MS is in the `cwd`. Thus if you are using a MS in a directory other than the current one, it will not find it. You need to change directories using `cd` in IPython (or `os.chdir()` inside a script) to the MS location.

The controls for the `plotcal` window are the same as for `plotxy` (see §3.3.2.1).

The `xaxis` and `yaxis` plot options available are:

- `'amp'` — amplitude,
- `'phase'` — phase,
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- 'real' – the real part,
- 'imag' — the imaginary part,
- 'snr' – the signal-to-noise ratio,

of the calibration solutions that are in the caltable. The xaxis choices also include 'time' and 'channel' which will be used as the sensible defaults (if xaxis='') for gain and bandpass solutions respectively.

The poln parameter determines what polarization or combination of polarization is being plotted. The poln='RL' plots both R and L polarizations on the same plot. The respective XY options do equivalent things. The poln='/' option plots amplitude ratios or phase differences between whatever polarizations are in the MS (R and L or X and Y).

The field, spw, and antenna selection parameters are available to obtain plots of subsets of solutions. The syntax for selection is given in §2.3.

The subplot parameter is particularly helpful in making multi-panel plots. The format is subplot=yxn where yxn is an integer with digit y representing the number of plots in the y-axis, digit x the number of panels along the x-axis, and digit n giving the location of the plot in the panel array (where n = 1, ..., xy, in order upper left to right, then down). See §3.3.2.8 for more details on this option.

The iteration parameter allows you to select an identifier to iterate over when producing multi-panel plots. The choices for iteration are: 'antenna', 'time', 'spw', 'field'. For example, if per-antenna solution plots are desired, use iteration='antenna'. You can then use subplot to specify the number of plots to appear on each page. In this case, set the n to 1 for subplot=yxn. Use the Next button on the plotcal window to advance to the next set of plots. Note that if there is more than one timestamp in a 'B' table, the user will be queried to interactively advance the plot to each timestamp, or if multiplot=True, the antennas plots will be cycled through for each timestamp in turn. Note that iteration can take more than one iteration choice (as a single string containing a comma-separated list of the options). ALERT: the iteration order is fixed (independent of the order specified in the iteration string), for example:

iteration = 'antenna, time, field'
iteration = 'time, antenna, field'

will both iterate over each field (fastest) then time (next) and antenna (slowest). The order is:

iteration = 'antenna, time, field, spw'

from the slowest (outer loop) to fastest (inner loop).

The markersize and fontsize parameters are especially helpful in making the dot and label sizes appropriate for the plot being made. The screen shots in this section used this feature to make the plots more readable in the cookbook. Adjusting the fontsize can be tricky on multi-panel plots, as the labels can run together if too large. You can also help yourself by manually resizing the Plotter window to get better aspect ratios on the plots.

ALERT: Unfortunately, plotcal has many of the same problems that plotxy does, as they use similar code underneath. An overhaul is underway, so stay tuned.
4.5.1.1 Examples for plotcal

For example, to plot amplitude or phase as a function of time for 'G' solutions (after rescaling by fluxscale can look like

```python
default('plotcal')
fontsize = 14.0  # Make labels larger
markersize = 10.0  # Make dots bigger

caltab = 'ngc5921.usecase.fluxscale'
yaxis = 'amp'
subplot = 211
plotcal()

yaxis = 'phase'
subplot = 212
plotcal()
```

The results are shown in Figure 4.4. This makes use of the subplot option to make multi-panel displays.

Similarly, to plot amplitude or phase as a function of channel for 'B' solutions for NGC5921:

```python
default('plotcal')
fontsize = 14.0  # Make labels larger
markersize = 10.0  # Make dots bigger

caltab = 'ngc5921.usecase.bcal'
antenna = '1'
yaxis = 'amp'
subplot = 311
plotcal()

yaxis = 'phase'
subplot = 312
plotcal()

yaxis = 'snr'
subplot = 313
plotcal()
```

The results are shown in Figure 4.5. This stacks three panels with amplitude, phase, and signal-to-noise ratio. We have picked antenna='1' to show.

For example, to show 6 plots per page of 'B' amplitudes on a 3 × 2 grid:

```python
default('plotcal')
fontsize = 12.0  # Make labels just large enough
markersize = 10.0  # Make dots bigger
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Figure 4.4: Display of the amplitude (upper) and phase (lower) gain solutions for all antennas and polarizations in the *ngc5921* post-fluxscale table.

```
caltab = 'ngc5921.usecase.bcal'
yaxis = 'amp'
subplot = 231
iteration = 'antenna'

plotcal()
```

See Figure 4.6 for this example. This uses the *iteration* parameter.

### 4.5.2 Listing calibration solutions with (**listcal**)  

The **listcal** task will list the solutions in a specified calibration table. The inputs are:
Figure 4.5: Display of the amplitude (upper), phase (middle), and signal-to-noise ratio (lower) of the bandpass 'B' solutions for antenna='0' and both polarizations for ngc5921. Note the falloff of the SNR at the band edges in the lower panel.

# listcal :: List data set summary in the logger:

vis = ''   # Name of input visibility file (MS)
caltab = '' # Input calibration table to list
field = ''  # Select data based on field name or index
antenna = '' # Select data based on antenna name or index
spw = ''    # Spectral window, channel to list
listfile = '' # Disk file to write, else to terminal
pagerows = 50 # Rows listed per page

An example listing is:

Listing CalTable: jupiter6cm.usecase.split.ms.smoothcal2 (G Jones)
Figure 4.6: Display of the amplitude of the bandpass 'B' solutions. Iteration over antennas was turned on using iteration='antenna'. The first page is shown. The user would use the Next button to advance to the next set of antennas.

SpwId = 0, channel = 0.

Time    Field      Ant  Amp  Phase  Amp  Phase
-------- --------  ----  ------  ----  ----
1999/04/16/14:10:43.5 'JUPITER' '1'  1.016  -11.5  1.016  -9.2
      '2'  1.013  -5.3  0.993  -3.1
      '3'  0.993  -0.8  0.990  -5.1
      '4'  0.997  -10.7  0.999  -8.3
      '5'  0.985  -2.7  0.988  -4.0
      '6'  1.005  -8.4  1.009  -5.3
      '7'  0.894  -8.7  0.897  -6.8
      '8'  1.001  -0.1  0.992  -0.7
      '9'  0.989  -12.4  0.992  -13.5
     '10'  1.000F  -4.2F  1.000F  -3.2F
4.5.3 Calibration table statistics (calstat)

The calstat task will print the statistics of solutions in a specified calibration table.

The inputs are:

```python
# calstat :: Displays statistical information on a calibration table
caltablen = '' # Name of input calibration table
axis = 'amp' # Which values to use
datacolumn = 'gain' # Which data column to use
useflags = True # Take flagging into account? # (not implemented)
```

For example:

```python
CASA <3>: calstat('ngc5921.demo.gcal',axis='amp',datacolumn='gain')
Out[3]:
{'GAIN': {'max': 1.6031942367553711,
'mean': 1.4448433067117419,
'medabsdevmed': 0.0086394548416137695,
'median': 1.5732669830322266,
'min': 0.99916577339172363,
'npts': 280.0,
'quartile': 0.020265340805053711,
'rms': 1.4650156497955322,
'stddev': 0.24271160321065546,
'sum': 404.55612587928772,
'sumsq': 600.9579999685287,
}
```
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{'var': 0.05890892233308665}]

CASA <4>: calstat('ngc5921.demo.gcal',axis='phase',datacolumn='gain')

Out[4]:

{'GAIN': {'max': 0.091214209794998169,
  'mean': -0.015221830284565011,
  'medabsdevmed': 0.012778861448168755,
  'median': -0.012778861448168755,
  'min': -0.15903720259666443,
  'npts': 280.0,
  'quartile': 0.02537553571164608,
  'rms': 0.031241731718182564,
  'stddev': 0.027331476552707856,
  'sum': -4.2621124796782031,
  'sumsq': 0.27329283416317834,
  'var': 0.00074700961055121926}}

The statistics can be captured as return variables from the task:

CASA <7>: mystat = calstat('ngc5921.demo.gcal',axis='amp',datacolumn='gain')

CASA <8>: print 'Gain Amp = ',mystat['GAIN']['mean']+'

Gain Amp = 1.44484330671 +/- 0.242711603211

ALERT: This task is still under development and currently offers no selection (e.g. by antenna) for the statistics.

4.5.4 Calibration Smoothing (smoothcal)

The smoothcal task will smooth calibration solutions (most usefully G or T) over a longer time interval to reduce noise and outliers. The inputs are:

# smoothcal :: Smooth calibration solution(s) derived from one or more sources:

vis = '' # Name of input visibility file
tablein = '' # Input calibration table
caltbtable = '' # Output calibration table
field = '' # Field name list
smoothtype = 'median' # Smoothing filter to use
smoothtime = 60.0 # Smoothing time (sec)

Note that if no caltable is specified as output, smoothcal will overwrite the input tablein calibration table.

The smoothing will use the smoothtime and smoothtype parameters to determine the new data points which will replace the previous points on the same time sampling grid as for the tablein solutions. The currently supported smoothtype options:
• 'mean' — use the mean of the points within the window defined by smoothtime (a “boxcar” average),

• 'median' — use the median of the points within the window defined by smoothtime (most useful when many points lie in the interval).

Note that smoothtime defines the width of the time window that is used for the smoothing.

ALERT: Note that smoothcal currently smooths by field and spw, and thus you cannot smooth solutions from different sources or bands together into one solution.

Figure 4.7: The 'amp' of gain solutions for NGC4826 before (top) and after (bottom) smoothing with a 7200 sec smoothtime and smoothtype='mean'. Note that the first solution is in a different spw and on a different source, and is not smoothed together with the subsequent solutions.

An example using the smoothcal task to smooth an existing table:

smoothcal('n4826_16apr.ms',

tablein='n4826_16apr.gcal',
caltablen='n4826_16apr.smoothcal',
smoothtime=7200.,
smoothtype='mean')

# Plot up before and after tables
plotcal('n4826_16apr.gcal','','amp',antenna='1',subplot=211)
plotcal('n4826_16apr.smoothcal','','amp',antenna='1',subplot=212)

This example uses 2 hours (7200 sec) for the smoothing time and smoothtype='mean'. The plotcal results are shown in Figure 4.7.

### 4.5.5 Calibration Interpolation and Accumulation (accum)

**ALERT:** The accum task is generally no longer recommended for most calibration scenarios. Please write to the NRAO CASA helpdesk if you need support using accum.

The accum task is used to interpolate calibration solutions onto a different time grid, and to accumulate incremental calibrations into a cumulative calibration table. The manual accumulation of calibration is rarely required and can usually be achieved implicitly simply by running applycal with all the calibration tables given as a list in the gaintable parameter (and using gainfield, spwmap, and interp appropriately. However, sometimes it is desirable to see the interpolated calibration prior to application, and this section describes how this can be done.

Its inputs are:

```python
# accum :: Accumulate incremental calibration solutions
vis = ''  # Name of input visibility file
tablein = ''  # Input (cumulative) calibration table; use '' on first run
accumtime = 1.0  # Timescale on which to create cumulative table
incrtable = ''  # Input incremental calibration table to add
caltablen = ''  # Output (cumulative) calibration table
field = ''  # List of field names to process from tablein.
calfield = ''  # List of field names to use from incrtable.
interp = 'linear'  # Interpolation mode to use for resampling incrtable solutions
spwmap = [-1]  # Spectral window combinations to apply
```

The mapping implied here is

```
tablein + incrtable => caltable
```

(mathematically the cal solutions are multiplied as complex numbers as per the Measurement Equation). The tablein is optional (see below). You must specify an incrtable and a caltable.

The tablein parameter is used to specify the existing cumulative calibration table to which an incremental table is to be applied. Initially, no such table exists, and if tablein='' then accumulate will generate one from scratch (on-the-fly), using the timescale (in seconds) specified by
the sub-parameter **accumtime**. These nominal solutions will be unit-amplitude, zero-phase calibration, ready to be adjusted by accumulation according to the settings of other parameters. When **accumtime** is negative (the default), the table name specified in **tablein** must exist and will be used. If **tablein** is specified, then the entries in that table will be used.

The **incrtable** parameter is used to specify the incremental table that should be applied to **tablein**. The calibration type of **incrtable** sets the type assumed in the operation, so **tablein** (if specified) must be of the same type. If it is not, **accum** will exit with an error message. (Certain combinations of types and subtypes will be supported by **accum** in the future.)

The **caltable** parameter is used to specify the name of the output table to write. If un-specified (""), then **tablein** will be overwritten. Use this feature with care, since an error here will require building up the cumulative table from the most recent distinct version (if any).

The **field** parameter specifies those field names in **tablein** to which the incremental solution should be applied. The solutions for other fields will be passed to **caltable** unaltered. If the cumulative table was created from scratch in this run of accumulate, then the solutions for these other fields will be unit-amplitude, zero-phase, as described above.

The **calfield** parameter is used to specify the fields to select from **incrtable** to use when applying to **tablein**. Together, use of **field** and **calfield** permit completely flexible combinations of calibration accumulation with respect to fields. Multiple runs of **accum** can be used to generate a single table with many combinations. In future, a 'self' mode will be enabled that will simplify the accumulation of field-specific solutions.

The **spwmap** parameter gives the mapping of the spectral windows in the **incrtable** onto those in **tablein** and **caltable**. The syntax is described in §4.4.1.4.

The **interp** parameter controls the method used for interpolation. The options are (currently): 'nearest' and 'linear' for time-dependent interpolation, and 'nearest', 'linear', cubic, and spline for (optional) frequency-dependent interpolation. These are described in §4.4.1.4. For most purposes, the 'linear' option should suffice.

We now describe the two uses of **accum**.

### 4.5.5.1 Interpolation using (accum)

**ALERT:** The **accum** task is generally no longer recommended for most calibration scenarios. Please write to the NRAO CASA helpdesk if you need support using **accum**.

Calibration solutions (most notably $G$ or $T$) can be interpolated onto the timestamps of the science target observations using **accum**.

The following example uses **accum** to interpolate an existing table onto a new time grid:

```plaintext
accum(vis='n4826_16apr.ms',
    tablein='',
    accumtime=20.0,
    incrtable='n4826_16apr.gcal',
```
caltable='n4826_16apr.20s.gcal',
interp='linear',
spwmap=[0,1,1,1,1])

plotcal('n4826_16apr.gcal','','phase',antenna='1',subplot=211)
plotcal('n4826_16apr.20s.gcal','','phase',antenna='1',subplot=212)

See Figure 4.8 for the plotcal results. The data used in this example is BIMA data (single polarization YY) where the calibrators were observed in single continuum spectral windows (spw='0,1') and the target NGC4826 was observed in 64-channel line windows (spw='2,3,4,5'). Thus, it is necessary to use spwmap=[0,1,1,1,1] to map the bandpass calibrator in spw='0' onto itself, and the phase calibrator in spw='1' onto the target source in spw='2,3,4,5'.

Figure 4.8: The 'phase' of gain solutions for NGC4826 before (top) and after (bottom) 'linear' interpolation onto a 20 sec accumtime grid. The first scan was 3C273 in spw='0' while the calibrator scans on 1331+305 were in spw='1'. The use of spwmap was necessary to transfer the interpolation correctly onto the NGC4826 scans.
4.5.5.2 Incremental Calibration using (accum)

It is occasionally desirable to solve for and apply calibration incrementally. This is the case when a calibration table of a certain type already exists (from a previous solve), a solution of the same type and incremental relative to the first is required, and it is not possible or convenient to recover the cumulative solution by a single solve.

Much of the time, it is, in fact, possible to recover the cumulative solution. This is because the equation describing the solution for the incremental solution (using the original solution), and that describing the solution for their product are fundamentally the same equation—the cumulative solution, if unique, must always be the same no matter what initial solution is. One circumstance where an incremental solution is necessary is the case of phase-only self-calibration relative to a full amplitude and phase calibration already obtained (from a different field).

For example, a phase-only 'G' self-calibration on a target source may be desired to tweak the full amplitude and phase 'G' calibration already obtained from a calibrator. The initial calibration (from the calibrator) contains amplitude information, and so must be carried forward, yet the phase-only solution itself cannot (by definition) recover this information, as a full amplitude and phase self-calibration would. In this case, the initial solution must be applied while solving for the phase-only solution, then the two solutions combined to form a cumulative calibration embodying the net effect of both. In terms of the Measurement Equation, the net calibration is the product of the initial and incremental solutions.

Cumulative calibration tables also provide a means of generating carefully interpolated calibration, on variable user-defined timescales, that can be examined prior to application to the data with applycal. The solutions for different fields and/or spectral windows can be interpolated in different ways, with all solutions stored in the same table.

The only difference between incremental and cumulative calibration tables is that incremental tables are generated directly from the calibration solving tasks (gaincal, bandpass, etc.), and cumulative tables are generated from other cumulative and incremental tables via accum. In all other respects (internal format, application to data with applycal, plotting with plotcal, etc.), they are the same, and therefore interchangeable. Thus, accumulate and cumulative calibration tables need only be used when circumstances require it.

The accum task represents a generalization on the classic AIPS CLCAL (see sidebox) model of cumulative calibration in that its application is not limited to accumulation of 'G' solutions. In principle, any basic calibration type can be accumulated (onto itself), as long as the result of the accumulation (matrix product) is of the same type. This is true of all the basic types, except 'D'. Accumulation is currently supported for 'B', 'G', and 'T', and, in future, 'F' (ionospheric Faraday rotation), delay-rate, and perhaps others. Accumulation of certain specialized types (e.g., 'GSPLINE', 'TOPAC', etc.) onto the basic types will be supported in the near future. The treatment of various calibration from ancillary data (e.g., system temperatures, weather data, WVR, etc.), as they become available, will also make use of accumulate to achieve the net calibration.

Other Packages:
The analog of accum in classic AIPS is the use of CLCAL to combine a series of (incremental) SN calibration tables to form successive (cumulative) CL calibration tables. AIPS SN/CL tables are the analog of 'G' tables in CASA.
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Note that accumulation only makes sense if treatment of a uniquely incremental solution is required (as described above), or if a careful interpolation or sampling of a solution is desired. In all other cases, re-solving for the type in question will suffice to form the net calibration of that type. For example, the product of an existing ’G’ solution and an amplitude and phase ’G’ self-cal (solved with the existing solution applied), is equivalent to full amplitude and phase ’G’ self-cal (with no prior solution applied), as long as the timescale of this solution is at least as short as that of the existing solution.

One obvious application is to calibrate the amplitudes and phases on different timescales during self-calibration. Here is an example:

```python
# Add clean model
ft(vis='jupiter6cm.usecase.split.ms',
    model='jupiter6cm.usecase.clean1.model')

# Phase only self-cal on 10s timescales
gaincal(vis='jupiter6cm.usecase.split.ms',
    caltable='jupiter6cm.usecase.phasecal1',
    gaintype='G',
    calmode='p',
    refant='6',
    solint=10.0,
    minsnr=1.0)

# Plot up solution phase and SNR
plotcal('jupiter6cm.usecase.phasecal1','','phase',antenna='1',subplot=211)
plotcal('jupiter6cm.usecase.phasecal1','','snr',antenna='1',subplot=212)

# Amplitude and phase self-cal on scans
gaincal(vis='jupiter6cm.usecase.split.ms',
    caltable='jupiter6cm.usecase.scancal1',
    gaintable='jupiter6cm.usecase.phasecal1',
    gaintype='G',
    calmode='ap',
    refant='6',
    solint='inf',
    minsnr=1.0)

# Plot up solution amp and SNR
plotcal('jupiter6cm.usecase.scancal1','','amp',antenna='1',subplot=211)
plotcal('jupiter6cm.usecase.scancal1','','snr',antenna='1',subplot=212)

# Now accumulate these - they will be on the 10s grid
accum(vis='jupiter6cm.usecase.split.ms',
    tablein='jupiter6cm.usecase.phasecal1',
    incrtable='jupiter6cm.usecase.scancal1',
    caltable='jupiter6cm.usecase.selfcal1',
    interp='linear')

# Plot this up
plotcal('jupiter6cm.usecase.selfcal1','','amp',antenna='1',subplot=211)
```
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plotcal('jupiter6cm.usecase.selfcal1','','phase','antenna='1',subplot=212)

The final plot is shown in Figure 4.9

![Figure 4.9: The final 'amp' (top) and 'phase' (bottom) of the self-calibration gain solutions for Jupiter. An initial phase calibration on 10s solint was followed by an incremental gain solution on each scan. These were accumulated into the cumulative solution shown here.](image)

**ALERT:** Only interpolation is offered in `accum`, no smoothing (as in `smoothcal`).

### 4.6 Application of Calibration to the Data

After the calibration solutions are computed and written to one or more calibration tables, one then needs to apply them to the data.
4.6.1 Application of Calibration (applycal)

After all relevant calibration types have been determined, they must be applied to the target source(s) before splitting off to a new MS or before imaging. This is currently done by explicitly taking the data in the DATA column in the MAIN table of the MS, applying the relevant calibration tables, and creating the CORRECTED_DATA scratch column. The original DATA column is untouched.

The applycal task does this. The inputs are:

```plaintext
# applycal :: Apply calibrations solutions(s) to data
vis = '' # Name of input visibility file
field = '' # Select field using field id(s) or field name(s)
spw = '' # Select spectral window/channels
intent = '' # Select observing intent
selectdata = True # Other data selection parameters
timerange = '' # Select data based on time range
uvrange = '' # Select data within uvrange (default units meters)
antenna = '' # Select data based on antenna/baseline
scan = '' # Scan number range
observation = '' # Select by observation ID(s)
msselect = '' # Optional complex data selection (ignore for now)
docallib = False # Use callib or traditional cal apply parameters
gaintable = [] # Select a subset of calibrators from gaintable(s)
interp = [] # Interp type in time[,freq], per gaintable. default=linear,linear
spwmap = [] # Spectral windows combinations to form for gaintable(s)
calwt = [True] # Calibrate data weights per gaintable.
parang = False # Apply parallactic angle correction
applymode = '' # Calibration mode:
flagbackup = True # Automatically back up the state of flags before the run?
```

As in other tasks, setting selectdata=True will open up the other selection sub-parameters (see §[2.3]). In addition, you can also select data based on the scan intents that were set during the observations (find them through listobs). Many of the other parameters are the common calibration parameters that are described in §[4.4.1].

The single non-standard parameter is the calwt option to toggle the ability to scale the visibility weights by the inverse of the products of the scale factors applied to the amplitude of the antenna gains (for the pair of antennas of a given visibility). This should in almost all cases be set to its default (True). The weights should reflect the inverse noise variance of the visibility, and errors in amplitude are usually also in the weights.

**Alert:** Current (as of February 2014) Jansky VLA data has no calibrated weights to the data (unless they are created from switched power). To avoid trouble, calwt=False should be set for
those data sets. Older, pre-Jansky VLA data should still be calibrated with `calwt=True`.

For `applycal`, the list of final cumulative tables is given in `gaintable`. In this case you will have run `accum` if you have done incremental calibration for any of the types, such as `G`. You can also feed `gaintable` the full sets and rely on use of `gainfield`, `interp` and `spwmap` to do the correct interpolation and transfer. In particular, for frequency interpolation, the interpolation methods ending in `PD`, `nearestPD` and `linearPD` also scale the phase by the frequency ratio between the measured and interpolated values. It is often more convenient to go through accumulation of each type with `accum` as described above (see §4.5.5.2), as this makes it easier to keep track of the sequence of incremental calibration as it is solved and applied. You can also do any required smoothing of tables using `smoothcal` (§4.5.4), as this is not yet available in `accum` or `applycal`.

`applycal` has different `applymodes`: `calflag` will apply all flags from a calibration table to the data and apply the calibration itself to the remaining visibilities. `trial` will only report on the calibration table flags but not manipulate the data, `flagonly` applies the flags but not the calibration itself, and `calonly` will apply the calibration and but not the solution table flags. Data that would `calflag` would flag are thus passed through uncalibrated. This option can be useful when `applycal` is executed in consecutive steps, one calibration table at a time. Portions of the data that were not calibrated in the first run can then be calibrated in a second run with a different calibration table. This option should be used with care such that no uncalibrated data remains in the final data product.

`applycal` will flag all data that have no calibration solution. Flags will distribute into all of your scratch columns, i.e. it will affect your uncalibrated visibilities, too. To be able to restore the flags to the state before `applycal` is starting its duty, the task will make a backup of your current flags by default (`flagbackup=True`). Restore them with `flagmanager`, if you are not happy with the `applycal` results.

If you are not doing polarization calibration or imaging, then you can set `parang=False` to make the calculations faster. If you are applying polarization calibration, or wish to make polarization images, then set `parang=True` so that the parallactic angle rotation is applied to the appropriate correlations. Currently, you must do this in `applycal` as this cannot be done on-the-fly in `clean` or `mosaic`. See §4.4.1.3 for more on `parang`.

For example, to apply the final bandpass and flux-scaled gain calibration tables solutions to the NGC5921 data:

```python
default('applycal')
vis='ngc5921.usecase.ms'

# We want to correct the calibrators using themselves
# and transfer from 1445+099 to itself and the target N5921

# Start with the fluxscale/gain and bandpass tables
gaintable=['ngc5921.usecase.fluxscale','ngc5921.usecase.bcal']

# pick the 1445+099 (field 1) out of the gain table for transfer
# use all of the bandpass table
```
gainfield = ['1', '*']

# interpolation using linear for gain, nearest for bandpass
interp = ['linear', 'nearest']

# only one spw, do not need mapping
spwmap = []

# all channels, no other selection
spw = ''
selectdata = False

gaincurve = False
opacity = 0.0

# select the fields for 1445+099 and N5921 (fields 1 and 2)
field = '1,2'
applycal()

# Now for completeness apply 1331+305 (field 0) to itself
field = '0'

gainfield = ['0', '*']
applycal()

# The CORRECTED_DATA column now contains the calibrated visibilities

In another example, we apply the final cumulative self-calibration of the Jupiter continuum data obtained in the example of §4.5.5.2:

applycal(vis='jupiter6cm.usecase.split.ms',
         gaintable='jupiter6cm.usecase.selfcal1',
         selectdata=False)

Again, it is important to remember the relative nature of each calibration term. A term solved for in the presence of others is, in effect, residual to the others, and so must be used in combination with them (or new versions of them) in subsequent processing. At the same time, it is important to avoid isolating the same calibration effects in more than one term, e.g., by solving for both \'G\' and \'T\' separately (without applying the other), and then using them together.

It is always a good idea to examine the corrected data after calibration (using plotxy to compare the raw (\'data\') and corrected (\'corrected\') visibilities), as we describe next.

### 4.6.2 Examine the Calibrated Data

Once the source data is calibrated using applycal, you should examine the $uv$ data and flag anything that looks bad. If you find source data that has not been flanked by calibration scans,
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delete it (it will not be calibrated).

For example, to look at the calibrated Jupiter data in the last example given in the previous section:

\[
\text{plotxy('jupiter6cm.usecase.split.ms','uvdist','amp','corrected', selectdata=True,correlation='RR LL',fontsize = 14.0)}
\]

will show the CORRECTED_DATA column. See Figure 4.10.

Figure 4.10: The final 'amp' versus 'uvdist' plot of the self-calibrated Jupiter data, as shown in plotxy. The 'RR LL' correlations are selected. No outliers that need flagging are seen.

See §3.3 for a description of how to display and edit data using plotms or plotxy, and §7.5 for use of the viewer to visualize and edit a Measurement Set.
4.6.3 Resetting the Calibration Models (delmod and clearcal)

Whenever calibration tasks are run, the models associated with the MS will be overwritten. Sometimes, however, one would like to completely remove the model and the task delmod can perform this functionality:

```python
# delmod :: Deletes model representations in the MS
vis = '' # Name of input visibility file (MS)
otf = True # Delete the on-the-fly model data keywords
field = '' # Select field using field id(s) or field name(s)
scr = False # Delete the MODEL_DATA scr col (if it exists)
```

To do so, the parameter otf should be set to True. delmod can also be used if for any reason a MODEL.column was created and should be removed to avoid confusion between the on-the-fly model and the MODEL column (the MODEL_DATA column was required in CASA 3.3 and earlier). This can be achieved with scr=T.

delmod generally replaces the functionality of the older clearcal task. If one still decides to use the MODEL_DATA columns, however, clearcal is still useful and will reset both the MODEL_DATA and CORRECTED_DATA columns to unity:

```python
CASA <11>: inp clearcal
# clearcal :: Re-initializes the calibration for a visibility data set
vis = '' # Name of input visibility file (MS)
field = '' # Select field using field id(s) or field name(s)
spw = '' # Select spectral window/channel.
intent = '' # Select observing intent
addmodel = False # Add MODEL_DATA scratch column
```

with field, spw, and intent being data selection parameters. addmodel can be used to opt in/out of formation of the MODEL_DATA column.

With the introduction of the on-the-fly calculation of the MODEL visibilities, and the fact that applycal overwrites any previously existing CORRECTED_DATA column, clearcal is not required anymore unless usescratch=True is chosen in calibration tasks, and it is also not recommended to use clearcal to create the scratch columns at the beginning of data calibration; all benefits from the on-the-fly model would be made obsolete.

4.7 Other Calibration and UV-Plane Analysis Options

4.7.1 Splitting out Calibrated uv data (split)

The split task will apply calibration and output a new sub-MS containing a specified list of sources (usually a single source). The inputs are:
# split :: Create a visibility subset from an existing visibility set:
vis = ''  # Name of input measurement set
outputvis = ''  # Name of output measurement set
datacolumn = 'corrected'  # Which data column(s) to split out
field = ''  # Select field using field id(s) or field name(s)
spw = ''  # Select spectral window/channels
width = 1  # Number of channels to average to form one output channel
antenna = ''  # Select data based on antenna/baseline
timebin = '0s'  # Value for timeaveraging
timerange = ''  # Select data based on time range
scan = ''  # Select data based on scan numbers
array = ''  # Select (sub)array by array ID number(s)
vrang = ''  # Select data based on uv distance range

Usually you will run split with datacolumn='corrected' as previous operations (e.g. applycal) will have placed the calibrated data in the CORRECTED_DATA column of the MS. This will produce a new MS with this corrected data in its DATA column. The modes available in datacolumn are:

- 'data', 'model', 'corrected',  # produce MS with single DATA column
- 'data,model', 'data,corrected', 'model,corrected',  # pairs of columns
- 'all'  # all columns 'data,model,corrected'

We recommend sticking to the simple single-column modes (e.g. 'data' or 'corrected') or 'all' if all columns are in the MS. Further processing may get confused by mismatched pairs of columns.

For example, to split out 46 channels (5-50) from spw 1 of our NGC5921 calibrated dataset:

```python
split(vis='ngc5921.usecase.ms',
     outputvis='ngc5921.split.ms',
     field='2',  # Output NGC5921 data (field 2)
     spw='0:5~50',  # Select 46 chans from spw 0
datacolumn='corrected')  # Take the calibrated data column
```

### 4.7.1.1 Averaging in split

Time and channel averaging are available using the timebin and width parameters.

The `timebin` parameter gives the averaging interval. It takes a quantity, e.g.

```
    timebin = '30s'
```

and will combine scans during averaging.

When time averaging, the `ignorables` subparameter can be used to specify that the bins should not be split by changes in SCAN_NUMBER, (sub)ARRAY_ID, and/or STATE_ID.

The `width` parameter defines the number of channels to average to form a given output channel. This can be specified globally for all `spw`, e.g.
width = 5

or specified per spw, e.g.

width = [2,3]

to average 2 channels of 1st spectral window selected and 3 in the second one.

**ALERT:** When averaging channels split will produce negative channel widths (as reported by listobs) if frequency goes down with increasing channel number, whether or not the input channel widths are negative. The bandwidths and channel resolutions will still be positive.

### 4.7.2 Recalculation of uvw values (**fixvis**)

Sometimes the u,v,w coordinates of a measurement set are not recorded correctly by the correlator. In those cases, it may be necessary to recalculate them based on the antenna positions. **fixvis** will perform this task.

```
# fixvis :: Recalculates (u, v, w) and/or changes Phase Center
vis           = ''   # Name of the input visibility set.
outputvis     = ''   # Name of the output visibility set. (Can be the same as vis.)
field         = ''   # Fields to operate on. = all.
refcode       = ''   # reference frame to convert UVW coordinates to
reuse         = True # base UVW calculation on the old values?
phasedir      = ''   # use this direction as phase center
```

A useful feature of **fixvis** is that it can also change the phase center of a measurement set. This can be done with absolute coordinates or using offsets. An example is:

```
fixvis(vis='Moon.ms',outputvis='Moon-fixed.ms',field='Moon', phasedir='J2000 9h25m00s 05d12m00s')
```

that will recalculate the u,v,w coordinates relative to the new phase center for the field 'Moon'.

### 4.7.3 Hanning smoothing of uv data (**hanningsmooth**)

For strong spectral line sources (like RFI sources), the Gibbs phenomenon may cause ringing across the frequency channels of an observation. This is called the Gibbs phenomenon and a proven remedy is the Hanning smoothing algorithm. Hanning smoothing is a running mean across the spectral axis with a triangle as a smoothing kernel. The central channel is weighted by 0.5 and the two adjacent channels by 0.25 to preserve the flux. Hanning smoothing significantly reduces Gibbs ringing but there’s no gain without a penalty and here it is the loss of a factor of two in spectral resolution.

In CASA, the **hanningsmooth** task will apply Hanning smoothing to a spectral line uv data measurement set. The inputs are:
hanningsmooth will operate on the input measurement set if no outputvis file name is provided. This option will keep the disk usage of large datasets under control. But one should be aware that the data is overwritten. If outputvis is provided, the task will copy the input MS to a new file with that name and operate there. The datacolumn parameter determines which of the data columns is to be Hanning smoothed: 'all', 'corrected' or 'data'. 'all' refers to both, the CORRECTED_DATA and the DATA column. If 'corrected' is specified but does not exist in the MS, hannahingsmooth will create this column for your convenience.

**ALERT:** We intend to make the kernel size a user supplied parameter. In the longer term we intend to offer other varieties of spectral smoothing as well.

### 4.7.4 MStransform (mstransform)

MStransform is an experimental task. It combines the functionality of split, cvel, hanningsmooth, to split an MS, combine/separate/regrid spws and do channel and time averaging, all in a single step.

Up to date documentation for mstransform can be found in the inline help of mstransform or at this web address:

http://www.eso.org/~scastro/ALMA/casa/MST/MSTransformDocs/MSTransformDocs.html

### 4.7.5 Model subtraction from uv data (uvsub)

The uvsub task will subtract the Fourier transform of the associated model of the MS (added to the MS with the tasks ft or setjy) from that in the CORRECTED_DATA column in the input MS and store the result in that same CORRECTED_DATA column.

The reverse operation is achieved by specifying reverse = True: in that case uvsub will add the value of the Fourier transform of the associated model to that in the CORRECTED_DATA column in the input MS and store the result in that same CORRECTED_DATA column.

The inputs are:

```python
# uvsub :: Subtract/add model from/to the corrected visibility data.
vis = '' # Name of input visibility file (MS)
reverse = False # reverse the operation (add rather than subtract)
```
For example:

uvsub('ngc5921.split.ms')

**ALERT:** Currently, *uvsub* operates on the **CORRECTED_DATA** column in the MS *vis*. Eventually we will provide the option to write out a new MS.

### 4.7.6 UV-Plane Continuum Subtraction (*uvcontsub*)

At this point, consider whether you are likely to need continuum subtraction. If there is significant continuum emission present in what is intended as a spectral line observation, continuum subtraction may be desirable. You can estimate and subtract continuum emission in the *uv*-plane prior to imaging or wait and subtract an estimate of it in the image-plane. Note that neither method is ideal, and the choice depends primarily upon the distribution and strength of the continuum emission. Subtraction in the *uv*-plane is desirable if continuum emission dominates the source, since deconvolution of the line emission will be more robust if it not subject to the deconvolution errors of the brighter continuum. There is also a performance benefit since the continuum is nearly the same in each channel of the observation, and it is desirable to avoid repeating its deconvolution in each channel. However, doing the continuum estimation in the *uv*-plane has the serious drawback that interpolating visibilities between channels is only a good approximation for emission from near the phase center. Thus, *uvcontsub* will do an increasingly poor job for emission distributed further from the phase center. If the continuum emission is relatively weak, it is usually adequate to subtract it in the image plane; this is described in the Image Analysis section of this cookbook. Here, we describe how to do continuum subtraction in the *uv*-plane.

The *uv*-plane continuum subtraction is performed by the *uvcontsub* task. First, determine which channels in your data cube do not have line emission, perhaps by forming a preliminary image as described in the next chapter. This image will also help you decide whether or not you need to come back and do *uv*-plane continuum subtraction at all.

The inputs to *uvcontsub* are:

```python
# uvcontsub :: Continuum fitting and subtraction in the uv plane
vis = '' # Name of input MS. Output goes to vis + ".contsub"
    # (will be overwritten if already exists)
field = '' # Select field(s) using id(s) or name(s)
fitspw = '' # Spectral window:channel selection for fitting the
    # continuum
combine = '' # Data axes to combine for the continuum estimation
    # (none, or spw and/or scan)
solint = 'int' # Continuum fit timescale (int recommended!)
fitorder = 0 # Polynomial order for the fits
spw = '' # Spectral window selection for output
want_cont = False # Create vis + ".cont" to hold the continuum estimate.
```

For each baseline, and over the timescale specified in *solint*, *uvcontsub* will provide a polynomial fit to the real and imaginary parts of the (continuum-only) channels specified in *fitspw* (using
the standard \texttt{spw} selection syntax), and then subtract this model from all channels specified in \texttt{spw}, or from all channels in spectral windows of \texttt{fitspw} if \texttt{spw=''} . By setting the subparameter \texttt{excludechannels=True}, the channel selection in \texttt{fitspw} will be inverted. In that case one can select the line channels themselves and/or corrupted channels that are not used in the continuum fit to the data. \texttt{fitspw} can also take frequency ranges, e.g.

\texttt{fitspw='*:113.767\textasciitilde114.528GHz;114.744\textasciitilde115.447GHz'}

where \texttt{'*'} indicates to go across all spws.

Typically, low orders for the polynomial work best, like \(0^{\text{th}}\) (a constant), or \(1^{\text{st}}\) order (a linear fit). Use higher orders with caution and check your results carefully.

Usually, one should set \texttt{solint='int'} which does no averaging and fits each integration. However, if the continuum emission comes from a small region around the phase center and \texttt{fitorder = 0}, then you can set \texttt{solint} larger (as long as it is shorter than the timescale for changes in the visibility function of the continuum). If your scans are short enough you can also use scan averaging with \texttt{combine='scan'} and \texttt{solint='inf'}. Be warned, setting \texttt{solint} too large will introduce “time smearing” in the estimated continuum and thus not properly subtract emission not at the phase center. Increasing \texttt{solint} speeds up the calculation but it does not improve the overall result quality of \texttt{uvcontsub} - although the continuum estimates of each baseline may be noisy (just like each visibility in a continuum MS may be noisy), it is better to use the ensemble of individual fits than to average the ensemble before fitting. Note that \texttt{plotms} can do time and baseline averaging on the fly to help you examine noisy data.

So, the recommended procedure is as follows:

- Finish calibration as described in the previous chapter.
- Use the invert or clean task on the split result to form an exploratory image that is useful for determining the line-free channels.
- Use \texttt{uvcontsub} with as low fit orders as possible to estimate and subtract the continuum from \texttt{vis}, and write the continuum-subtracted dataset to \texttt{vis + '.contsub'}. 
- Use \texttt{clean} with \texttt{vis + '.contsub'} to make an image cube of the line emission.
- If a continuum image is desired, clean the line-free channels of the original MS with \texttt{mode='mfs'} and \texttt{spw=fitspw}. Note that using the line free channels directly is preferred over the imaging the 'continuum' model fitted by \texttt{uvcontsub}. The fitting procedure will also fit noise and artifacts which produce a nice line cube when subtracted, but the model may not represent the true underlying continuum.

For example, we perform \texttt{uv}-plane continuum subtraction on our NGC5921 dataset:

\begin{verbatim}
# Want to use channels 4-6 and 50-59 for continuum
uvcontsub(vis='ngc5921.usecase.ms,
\end{verbatim}
4.7.7 Spectral regridding of the MS (cvel)

Although not strictly a calibration operation, spectral regridding of a MS is available to aid in calibration operations (e.g. continuum subtraction) and preparation for imaging. For this purpose, the cvel task has been developed.

The inputs are:

```python
# cvel :: regrid an MS to a new spectral window / channel structure or frame
vis = ''  # Name of input measurement set
outputvis = ''  # Name of output measurement set
passall = False  # Pass through (write to output MS) non-selected data with no change
field = ''  # Select field using field id(s) or field name(s)
spw = ''  # Select spectral window/channels
selectdata = True  # Other data selection parameters
timerange = ''  # Range of time to select from data
array = ''  # (sub)array indices
antenna = ''  # Select data based on antenna/baseline
scan = ''  # scan number range
mode = 'channel'  # Regridding mode
nchan = -1  # Number of channels in output spw (-1=all)
start = 0  # first input channel to use
width = 1  # Number of input channels to average
interpolation = 'linear'  # Spectral interpolation method
phasecenter = ''  # Image phase center: position or field index
restfreq = ''  # rest frequency (see help)
outframe = ''  # Output frame (not case-sensitive, ''=keep input frame)
veltype = 'radio'  # velocity definition
hanning = False  # If true, Hanning smooth data before regridding to remove Gibbs ringing.
```

The key parameters for the operation of cvel are the regridding mode, the output reference outframe, veltype, restfreq (which may be a list of rest frequencies to match the different spws) and the standard selection parameters (in particular spw and field).

The syntax for mode options (‘channel’, ‘velocity’, ‘frequency’, ‘channel_b’) has been made compatible with the respective modes of clean (§ 5.2.5). The combination of selected spw and mode will determine the output channels and spw(s):
The simplest use of `cvel` is to shift a single spectral window into an output frame without regridding. This is done with `mode='channel'`. For example:

```python
cvel(vis='test_w3oh_nohann.ms',
      outputvis='test_w3oh_nohann_chanbary.ms',
      mode='channel', nchan=-1, start=0, width=1,
      interpolation='linear',
      phasecenter='',
      spw='',
      restfreq='1665.4018MHz',
      outframe='BARY')
```

does this for an observation of the OH line.

There is also a special `mode='channel_b'` that does not force a linear output frequency grid, e.g. for irregularly spaced/overlapping spectral windows, but is nominally faster. This is not equivalent to a `clean` output gridding mode, although `clean` will work on this spectral lattice.

Mode `channel` is intended to not interpolate between channels. It will perform binning if needed. For most scientific applications we recommend using the `mode='velocity'` and `mode='frequency'` options, as it is easiest to determine what the resulting channelization will be. For example:

```python
cvel(vis='test_w3oh_nohann.ms',
      outputvis='test_w3oh_nohann_cvellsrk.ms',
      mode='velocity', nchan=45, start='-35.0km/s', width='-0.55km/s',
      interpolation='linear',
      phasecenter='',
      spw='',
      restfreq='1665.4018MHz',
      outframe='LSRK')
```
cvel(vis='test_w3oh_nohann.ms',
    outputvis = 'test_w3oh_nohann_cvelbary.ms',
    mode='velocity',nchan=45,start='35.0km/s',width='-0.55km/s',
    interpolation='linear',
    phasecenter='',
    spw='',
    restfreq='1665.4018MHz',
    outframe='BARY')

will transform a MS into the LSRK and BARYcenter frames respectively.

The sign of the width parameter determines whether the channels run along increasing or decreasing values of frequency or velocity (i.e. if the cube is reversed or not).

The intent of cvel regridding is to transform channel labels and the visibilities to a spectral reference frame which is appropriate for the science analysis, e.g. from TOPO to LSRK, e.g. to correct for Doppler shifts throughout the time of the observation. Naturally, this will change the shape of the spectral features to some extent. According to the Nyquist theorem you should oversample a spectrum with twice the numbers of channels to retain the shape. Based on some tests, however, we recommend to observe with at least 3-4 times the number of channels for each significant spectral feature (like 3-4 channels per linewidth). This will minimize regridding artifacts in cvel.

If cvel has already established the grid that is desired for the imaging, clean should be run with the default channel mode (> width=1) or with exactly the same frequency/velocity parameters as was used in cvel. This will avoid additional regridding in clean. Hanning smoothing is optionally offered in cvel, but tests have shown that already the regridding process itself, if it involved a transformation from TOPO to a non-terrestrial reference frame, implies some smoothing (due to channel interpolation) such that Hanning smoothing may not be necessary.

The interpolation method fftshift calculates the transformed visibilities by applying a FFT, then a phase ramp, and then an inverse FFT. It will also perform pre-averaging, if necessary (this will increase the S/N). Note that if you want to use this interpolation method, your frequency grid needs to be equidistant, i.e. it only works in mode velocity with veltype=radio, in mode frequency, and in mode channel (in the latter only if the input grid is itself equidistant in frequency). Note also that, as opposed to all other interpolation methods, this method will apply a constant (frequency-independent) shift in frequency which is not fully correct in the case of large fractional bandwidth of the given spectral window.

4.7.8 UV-Plane Model Fitting (uvmodelfit)

It is often desirable to fit simple analytic source component models directly to visibility data. Such fitting has its origins in early interferometry, especially VLBI, where arrays consisted of only a few antennas and the calibration and deconvolution problems were poorly constrained. These methods overcame the calibration uncertainties by fitting the models to calibration-independent closure quantities and the deconvolution problem by drastically limiting the number of free parameters
required to describe the visibilities. Today, even with larger and better calibrated arrays, it is still desirable to use visibility model fitting in order to extract geometric properties such as the positions and sizes of discrete components in radio sources. Fits for physically meaningful component shapes such as disks, rings, and optically thin spheres, though idealized, enable connecting source geometry directly to the physics of the emission regions.

Visibility model fitting is carried out by the uvmodelfit task. The inputs are:

```python
# uvmodelfit :: Fit a single component source model to the uv data:

vis = '' # Name of input visibility file
field = '' # field name or index
spw = '' # spectral window
selectdata = False # Activate data selection details
niter = 5 # Number of fitting iterations to execute
comptype = 'P' # Component type (P=pt source,G=ell. gauss,D=ell. disk)
sourcepar = [1, 0, 0] # Starting guess (flux,xoff,yoff,bmajaxrat,bpa)
varypar = [] # Which parameters can vary in fit
outfile = '' # Optional output component list table
```

**ALERT:** This task currently only fits a single component.

The user specifies the number of non-linear solution iterations (niter), the component type (comptype), an initial guess for the component parameters (sourcepar), and optionally, a vector of Booleans selecting which component parameters should be allowed to vary (varypar), and a filename in which to store a CASA componentlist for use in other applications (file). Allowed comptypes are currently point 'P' or Gaussian 'G'.

The function returns a vector containing the resulting parameter list. This vector can be edited at the command line, and specified as input (sourcepar) for another round of fitting.

The sourcepar parameter is currently the only way to specify the starting parameters for the fit. For points, there are three parameters: I (total flux density), and relative direction (RA, Dec) offsets (in arcsec) from the observation’s phase center. For Gaussians, there are three additional parameters: the Gaussian’s semi-major axis width (arcsec), the aspect ratio, and position angle (degrees). It should be understood that the quality of the result is very sensitive to the starting parameters provided by the user. If this first guess is not sufficiently close to the global \( \chi^2 \) minimum, the algorithm will happily converge to an incorrect local minimum. In fact, the \( \chi^2 \) surface, as a function of the component’s relative direction parameters, has a shape very much like the inverse of the absolute value of the dirty image of the field. Any peak in this image (positive or negative) corresponds to a local \( \chi^2 \) minimum that could conceivably capture the fit. It is the user’s responsibility to ensure that the correct minimum does the capturing.

Currently, uvmodelfit relies on the likelihood that the source is very near the phase center (within a beamwidth) and/or the user’s savvy in specifying the starting parameters. This fairly serious constraint will soon be relieved somewhat by enabling a rudimentary form of uv-plane weighting to increase the likelihood that the starting guess is on a slope in the correct \( \chi^2 \) valley.

Improvements in the works for visibility model fitting include:
• User-specifiable uv-plane weighting
• Additional component shapes, including elliptical disks, rings, and optically thin spheroids.
• Optional calibration pre-application
• Multiple components. The handling of more than one component depends mostly on efficient means of managing the list itself (not easy in command line options), which are currently under development.
• Combined component and calibration fitting.

Example (see Figure 4.11):

```python
# Note: It's best to channel average the data if many channels
# before running a modelfit

split('ngc5921.ms', '1445+099_avg.ms',
     datacolumn='corrected', field='1445*', width='63')

# Initial guess is that it's close to the phase center
# and has a flux of 2.0 (a priori we know it's 2.47)

uvmodelfit('1445+099_avg.ms',
           niter=5,  # Do 5 iterations
           comptype='P',  # P=Point source, G=Gaussian, D=Disk
           sourcepar=[2.0, 1.1, 1.1],  # Source parameters for a point source
           spw='0',
           outfile='gcal.cl')  # Output component list file

# Output looks like:
There are 19656 - 3 = 19653 degrees of freedom.
iter=0: reduced chi2=0.0418509: I=2, dir=[0.1, 0.1] arcsec
iter=1: reduced chi2=0.003382: I=2.48614, dir=[0.00325324, 0.00228963] arcsec
iter=2: reduced chi2=0.00338012: I=2.48614, dir=[0.00325324, 0.00228963] arcsec
iter=3: reduced chi2=0.00338012: I=2.48614, dir=[0.00325324, 0.00228963] arcsec
iter=4: reduced chi2=0.00338012: I=2.48614, dir=[0.00325324, 0.00228963] arcsec
iter=5: reduced chi2=0.00338012: I=2.48614, dir=[0.00325324, 0.00228963] arcsec

If data weights are arbitrarily scaled, the following formal errors will be underestimated by at least a factor sqrt(reduced chi2). If the fit is systematically poor, the errors are much worse.
I = 2.48614 +/- 0.0176859
x = 0.00325324 +/- 0.163019 arcsec
y = -0.00228963 +/- 0.174458 arcsec
Writing componentlist to file: /home/sandrock/smyers/Testing/Patch2/N5921/gcal.cl

# Fourier transform the component list to a model of the MS
ft('1445+099_avg.ms', complist='gcal.cl')

# Plot data versus uv distance
plotxy('1445+099_avg.ms', xaxis='uvdist', datacolumn='corrected')

# Specify green circles for model data (overplotted)
plotxy('1445+099_avg.ms', xaxis='uvdist', datacolumn='model',
overplot=True, plotsymbol='go')

Figure 4.11: Use of plotxy to display corrected data (red and blue points) and uv model fit data (green circles).

4.7.9 Reweighing visibilities based on their scatter (\texttt{statwgt})

\textbf{Alert:} \texttt{statwgt} is still an experimental task. Please check the results carefully and report any problems to the NRAO CASA helpdesk.

In most cases, the data that comes from the telescopes have the correct absolute or relative weights associated (absolute weights will be supplied once the VLA switched power application becomes standard; for ALMA the Tsys application is already in place). However, there are data sets where one would like to adjust the weights based on the scatter of the visibilities (typically as a function
of time, antenna, and/or baseline). This calculation is performed by the task `statwt` that updates the WEIGHT and SIGMA columns of the measurement set. `statwt` inputs are:

```
# statwt :: Reweight visibilities according to their scatter
vis = '' # Name of measurement set
dorms = False # Use rms instead of stddev?
byantenna = False # Estimate the noise per antenna -not
    # implemented (vs. per baseline)
fitspw = '' # The signal-free spectral window: channels
    # to estimate the scatter from
fitcorr = '' # The signal-free correlation(s) to estimate
    # the scatter from (not implemented)
combine = '' # Let estimates span changes in spw, corr,
    # scan and/or state
timebin = '0s' # Bin length for estimates (not implemented)
minsamp = 2 # Minimum number of unflagged visibilities
    # for estimating the scatter
field = '' # Select field using ID(s) or name(s)
spw = '' # Select spectral window/channels
antenna = '' # Select data based on antenna/baseline
timerange = '' # Select data by time range
scan = '' # Select data by scan numbers
intent = '' # Select data by scan intents
array = '' # Select (sub)array(s) by array ID number
correlation = '' # Select correlations to reweight
observation = '' # Select by observation ID(s)
datacolumn = 'corrected' # Which data column to calculate the scatter
    # from
```

`statwt` should only be run after all calibration steps have been performed. The parameter `dorms` switches from a scatter standard deviation to a root mean square scatter estimator. `datacolumn` specifies the column on which the task operates and the usual data selection parameters apply. Channels with strong RFI or a spectral line should be avoided for the calculation and good channel range should be specified via `fitspw`. In its current implementation, `statwt` uses data samples of an integration time interval but eventually wider sample intervals can be specified by the `timebin` parameter. Those samples are contained within a scan, spw, and polarization product but using the `combine` can relax this restriction. `minsamp` sets the minimum number of unflagged visibilities used for the calculation.

### 4.7.10 Change the signs of visibility phases (`conjugatevis`)

`conjugatevis` is an easy task to flip the signs of the phases of visibilities, thus creating the complex conjugate numbers. The inputs are like:

```
# conjugatevis :: Change the sign of the phases in all visibility columns.
vis = '' # Name of input visibility file.
spwlist = '' # Spectral window selection
outputvis = '' # Name of output visibility file
overwrite = False # Overwrite the outputvis if it exists.
```

The task works on all scratch columns.
4.7.11 Manipulation of Ephemeris Objects

When an astronomical object has a proper motion, in particular objects in our solar system, a static (RA, Dec) position in the FIELD table of the MeasurementSet will not accurately describe the time-dependent position. Prior to CASA 4.2, there was no support for ephemeris objects other than the built-in reference frames for the Sun, the Moon, and the planets out to PLUTO. With CASA 4.2, several new features were introduced which help the user to attach an arbitrary ephemeris to a given field and work with the object from calibration to imaging.

4.7.11.1 Ephemeris tables

The CASA format for ephemeris tables has been defined was introduced in the early development stages of CASA in connection with the Measures module. The `me` tool permits position calculations based on ephemerides in this format. Two examples for such tables can be found in the distribution directory in subdirectory `data/ephemerides`: `VGEO` is an ephemeris of Venus in the geocentric reference frame while `VTOP` is an ephemeris for the same object in the `TOPO` reference frame for the observatory location of the VLA. With the introduction of solar system source models (Butler) in the `setjy` task, a nearly complete set of ephemerides for the larger bodies in our solar system had to be made available. These are stored in nearly the same format as the above examples `VGEO` and `VTOP` (but with a few enhancements) in directory `data/ephemerides/JPL-Horizons`. If your object’s ephemeris is among those stored in `data/ephemerides/JPL-Horizons`, you can simply copy the ephemeris from there. Otherwise, you can request the ephemeris from the JPL-Horizons using the CASA commands (for example)

```python
import recipes.ephemerides.request as jplreq
jplreq.request_from_JPL(objnam='Mars', startdate='2012-01-01', enddate='2013-12-31',
date_incr='0.1 d', get_axis_orientation=False,
get_axis_ang_orientation=True,
get_sub_long=True, use_apparent=False, get_sep=False,
return_address='YOUR_EMAIL_ADDRESS',
mailserver='YOUR_MAIL_SERVER_ADDRESS')
```

where you need to fill in the parameters `objnam`, `startdate`, `enddate`, `date_incr` (the time interval between individual ephemeris table entries), `return_address` (your email address where you want to receive the ephemeris), and `mailserver` (the smtp server through which you want to send the request email). The other parameters should be set as shown. Within a short time, you should receive the requested ephemeris as an email from NASA’s JPL-Horizons system. Save the email into a file with the “save as” function of your mail client. See the next section on how to attach it to your dataset.

4.7.11.2 Using `fixplanets` to attach ephemerides to a field of a MeasurementSet

In order to set the ephemeris of a given field in a MeasurementSet, you can use the task `fixplanets` as in the following example:

```
fixplanets(vis='uid___A002_X1c6e54_X223.ms',
field='Titan', fixuvw=True, direction='mytitanephemeris')

where you need to set the parameters vis to the name of your MS and the parameter field to the name of the field to which you want to attach the ephemeris. The parameter direction must be set to the name of your ephemeris table. Accepted formats are (a) the CASA format (as in VGE0 or the ephemerides in data/ephemerides/JPL-Horizons as described above) and (b) the JPL-Horizons mail format which you obtain by saving an ephemeris email you received from JPL-Horizons. The parameter fixuvw should be set to True in order to trigger a recalculation of the UVW coordinates in your MS based on the new ephemeris. The task fixplanets can also be used for other field direction modifications. Please refer to the help text of the task.

Note that among the ephemerides in the directory data /ephemerides/JPL-Horizons/ you should only use those ending in '.J2000.tab'. They are the ones in J2000 coordinates.

4.7.11.3 Use of the ephemeris after attachment

Once you have attached the ephemeris to a field of an MS, it will automatically be handled in tasks like split and concat which need to hand on the ephemeris to their output MSs. In particular concat recognizes when fields of the MSs to be concatenated use the same ephemeris and merges these fields if the time coverage of the provided ephemeris in the first MS also covers the observation time of the second MS. The ephemeris of the field in the first MS will then be used for the merged field. In order to inspect the ephemeris attached to a field, you can find it inside the FIELD subdirectory of your MS. The optional column EPHMERIS_ID in the FIELD table points to the running number of the ephemeris table. A value of -1 indicates that no ephemeris is attached. Note that in case an ephemeris is attached to a field, the direction column entries for that field in the FIELD table will be interpreted as an offset to the ephemeris direction and are therefore set to (0.,0.) by default. This offset feature is used in mosaic observations where several fields share the same ephemeris with different offsets. The TIME column in the FIELD table should be set to the beginning of the observation for that field and serves as the nominal time for ephemeris queries.

4.7.11.4 Spectral frame transformation to the rest frame of the ephemeris object in task cvel

The ephemerides contain radial velocity information. The task cvel can be used to transform spectral windows into the rest frame of the ephemeris object by setting the parameter outframe to “SOURCE” as in the following example:

\[
cvel(vis='europa.ms', 
outputvis='cvel_europa.ms', outframe='SOURCE', mode = 'velocity', 
width = '0.3km/s', restfreq = '354.50547GHz')
\]

This will make cvel perform a transformation to the GEO reference frame followed by an additional Doppler correction for the radial velocity given by the ephemeris for the each field. (Typically,
this should happen after calibration and after splitting out the spectral widows and the target of interest). The result is an MS with a single combined spectral window in reference frame REST. From this frame, further transformations to other reference frames are not possible.

4.7.11.5 Ephemerides in ALMA datasets

The ALMA Science Data Model (the raw data format for ALMA data) now foresees an ephemeris table. However, this feature is not yet used in ALMA Cycle 1. Once this table will be filled at the observatory, the task importasdm will automatically translate it into an ephemeris table in CASA format and attach it to the respective fields.

4.8 Examples of Calibration

The data reduction tutorials on casaguides.nrao.edu provide walkthroughs for high and low frequency, spectral line and polarization calibration techniques.
Chapter 5

Synthesis Imaging

This chapter describes how to make and deconvolve images starting from calibrated interferometric data, possibly supplemented with single-dish data or an image made from single-dish data. This data must be available in CASA (see §2 on importing data). See §4 for information on calibrating synthesis data. In the following sections, the user will learn how to make various types of images from synthesis data, reconstruct images of the sky using the available deconvolution techniques, include single-dish information in the imaging process, and to prepare to use the results of imaging for improvement of the calibration process (“self-calibration”).

5.1 Imaging Tasks Overview

The current imaging and deconvolution tasks are:

- **clean** — calculate a deconvolved image with a selected clean algorithm, including mosaicing, or make a dirty image (§5.3),

- **feather** — combine a single dish and synthesis image in the Fourier plane (§5.5),

- **deconvolve** — image-plane only deconvolution based on the dirty image and beam, using one of several algorithms (§5.8),

- **pclean** — an experimental task for clean to work in a parallelized way for multi-node and core computing systems (§5.10)

There are also tasks that help you set up the imaging or interface imaging with calibration:

- **boxit** - create “cleanbox” deconvolution regions automatically from an image (§5.6.1),
• **ft** - add a source model to the MS (§ 5.7).

The full “tool kit” that allows expert-level imaging must still be used if you do not find enough functionality within the tasks above.

Information on other useful tasks and parameter setting can be found in:

- **listobs** — list what’s in a MS (§ 2.2.7),
- **split** — Write out new MS containing calibrated data from a subset of the original MS (§ section:cal.split),
- **cvel** — regrid a spectral MS onto a new frequency channel system (§ 4.7.7),
- data selection — general data selection syntax (§ 2.3),
- **viewer** — image display including region statistics and image cube slice and profile capabilities (§ 7).

### 5.2 Common Imaging Task Parameters

We now describe some parameters that are common to the imaging tasks. These should behave the same way in any imaging task that they are found in. These are in alphabetical order.

**ALERT:** **clean** tries to use up to four cores on the computer that it is running on. If this is not desired, the environment variable **OMP_NUM_THREAD** can be set to a lower value.

#### 5.2.1 Parameter **cell**

The **cell** parameter defines the pixel size in the x and y axes for the output image. If given as floats or integers, this is the cell size in arc seconds, e.g.

```plaintext
cell=[0.5,0.5]
```

make 0.5″ pixels. You can also give the cell size in **quantities**, e.g.

```plaintext
cell=['1arcmin', '1arcmin']
```

If a single value is given, then square pixels of that size are assumed.
5.2.2 Parameter field

The field parameter selects the field indexes or names to be used in imaging. Unless you are making a mosaic, this is usually a single index or name:

```python
field = '0'  # First field (index 0)
field = '1331+305'  # 3c286
field = '*'  # all fields in dataset
```

The syntax for field selection is given in §2.3.2.

5.2.3 Parameter imagename

The value of the imagename parameter is used as the root name of the output image. Depending on the particular task and the options chosen, one or more images with names built from that root will be created. For example, the clean task run with imagename='ngc5921 a series of output images will be created with the names `ngc5921.clean`, `ngc5921.residual`, `ngc5921.model`, etc.

If an image with that name already exists, it will in general be overwritten. Beware using names of existing images however. If the clean is run using an imagename where `<imagename>.residual` and `<imagename>.model` already exist then clean will continue starting from these (effectively restarting from the end of the previous clean). Thus, if multiple runs of clean are run consecutively with the same imagename, then the cleaning is incremental (as in the difmap package).

The output image may also have a different beam per plane. For datasets with very large fractional bandwidth, clean will use a different PSF for each channel when the PSF changes by more than half a pixel as a function of frequency. To smooth to a common resolution, one can either use the parameter ressmooth (§5.2.6) to smooth to the smallest common possible beam, restoringbeam for an arbitrary, larger beam, (§5.3.11), or the task imsmooth (§6.17) after cleaning. Data analysis tasks such as immoments in CASA support changing beams per plane.

5.2.4 Parameter imsize

The image size in numbers of pixels on the x and y axes is set by imsize. For example,

```python
imsize = [640, 640]
```

makes a square image 640 pixels on a side. If a single value is given, then a square image of that dimension is made. The underlying algorithms work best for certain image sizes. If you pick a size where that algorithm will be particularly slow, the logger will send a warning message, suggesting the nearest optimal values. In general, the best performance is obtained with image sizes that are even and factorizable to 2,3,5,7 only. An easy rule of thumb would be $2^n \times 10$ where $n$ is an integer number, like 160, 320, 640, 1280, 2560, etc.
5.2.5 Parameter mode

The mode parameter defines how the frequency channels in the synthesis MS are mapped onto the image. The allowed values are: mfs, channel, velocity, frequency. The mode parameter is expandable, with some options uncovering a number of sub-parameters, depending upon its value.

5.2.5.1 Mode mfs

```
mode = 'mfs' # Spectral gridding type (mfs, channel, velocity, frequency)
nterms = 1 # Number of terms used to model the sky frequency dependence (Note: nterms>1 is under development)
reffreq = '' # Reference frequency for MFS (relevant only if nterms > 1), '' defaults to central data-frequency
```

The default mode='mfs' emulates multi-frequency synthesis in that each visibility-channel datum \( k \) with baseline vector \( \mathbf{B}_k \) at wavelength \( \lambda_k \) is gridded into the uv-plane at \( \mathbf{u}_k = \mathbf{B}_k / \lambda_k \). The result is one or more images (depending on nterms), regardless of how many channels are in the input dataset. The first image plane is at the frequency given by the midpoint between the highest and lowest frequency channels in the input spw(s). Currently, there is no way to choose the center frequency of the output image plane independently.

WideBand imaging (mfs with nterms > 1) is now available in CASA. This algorithm models the wide-band sky brightness as a linear combination of Gaussian-like functions whose amplitudes follow a Taylor-polynomial in frequency. The output images are a set of Taylor-coefficient images, from which spectral index and curvature maps are derived. The reffreq parameter sets the reference frequency \( \nu_0 \) about which the Taylor expansion is done. The Taylor expansion is a polynomial in frequency:

\[
I_{\nu}^{\text{sky}} = \sum_t I_t^{\text{sky}} \left( \frac{\nu - \nu_0}{\nu_0} \right)^t
\]

(5.1)

\( I_t^{\text{sky}} \) an image of the \( t^{\text{th}} \) coefficient of the Taylor-polynomial expansion.

When Eq. 5.1 is applied on a source with a spectral index

\[
I_{\nu}^{\text{sky}} = I_{\nu_0}^{\text{sky}} \left( \frac{\nu}{\nu_0} \right)^{\alpha + \beta \log(\nu/\nu_0)}
\]

(5.2)

The Taylor terms \( I_t^{\text{sky}} \) can be used to constrain the sky brightness, \( \alpha \), and \( \beta \) through

\[
I_{\nu_0}^{\text{sky}} = I_0^{\text{sky}}
\]

(5.3)
\[ \alpha = \frac{I^\text{sky}}{I^\text{sky}_0} = \frac{I^\text{sky}_1}{I^\text{sky}_0} \]  
\[ \beta = \frac{I^\text{sky}_2}{I^\text{sky}_0} - \frac{\alpha(\alpha - 1)}{2} = \frac{I^\text{sky}_2}{I^\text{sky}_0} - \frac{\alpha(\alpha - 1)}{2} \]  
\[ (5.4) \]

\[ (5.5) \]


Alert: The MS-MFS (multiscale-multifrequency) algorithm in the current release is new and is still being developed/tested/debugged. Its basic operation has been tested on wide-band JVLA data for Stokes I imaging.

Explanation of the Parameters:
\textbf{nterms}: The number of terms in the Taylor polynomial used to model the frequency structure. \texttt{nterms} > 1 triggers MS-MFS. \texttt{nterms}= 1 triggers standard point-source clean or multi-scale-clean. Note: The choice of \texttt{nterms} follows the same rules used while fitting a polynomial to a 1D set of noisy data points. To prevent overfitting, the order of the polynomial needs to depend on the available signal-to-noise in the data. A very rough rule-of-thumb is as follows: For high SNR data (single channel SNR $> 100$), and fields dominated by point-sources with spectral indices around $-1.0$ across a 2:1 bandwidth, choose \texttt{nterms}= 3 or 4. For lower SNR data ($5 < \text{SNR} < 100$), flatter spectra, or when there is significant extended emission, \texttt{nterms}= 2 is a much safer option. For very low SNR data (SNR $< 5$), choose \texttt{nterms} = 1).

\textbf{reffreq}: The reference frequency used to compute Taylor functions \( [(\text{freq} - \text{reffreq})/\text{reffreq}]^i \). If left blank (\texttt{reffreq}=”), it defaults to the middle frequency of the selected data. Note: For the current release, the use of \texttt{reffreq}=” is recommended.

\textbf{multiscale}: The MS-MFS algorithm always uses scale sizes set via the \texttt{multiscale} parameter. For point-source deconvolution, set \texttt{multiscale}=[0] (also the default). Note: Unlike standard mscclean (\texttt{multiscale} = [0, 6, 10, ...] with \texttt{nterms}=1), with higher \texttt{nterms} the largest specified scale size must lie within the sampled range of the interferometer. If not, there can be an ambiguity in the spectral reconstruction at very large spatial scales.

\textbf{gridmode}: Wideband W-Projection is supported, and can be triggered via \texttt{gridmode}='widefield'.

\textbf{modelimage}: Supply a list of Taylor-coefficient images, to start the deconvolution from. If only one image is specified, it will be used as the model for the ‘tt0’ image.

\textbf{Output images}: [\texttt{xxx.image.tt0}, \texttt{xxx.image.tt1},... ] : Images of Taylor coefficients that describe the frequency-structure. The “tt0” image is the total-intensity image at the reference frequency, and is equivalent to ”\texttt{xxx.image}” obtained via standard imaging.

[\texttt{xxx.image.alpha, xxx.image.beta}] : Spectral index and spectral curvature at the reference-frequency. These are computed from \texttt{tt0}, \texttt{tt1}, \texttt{tt2} only for regions of the image where there is sufficient signal-to-noise to compute them. These regions are chosen via a threshold on the intensity image (\texttt{tt0}) computed as \texttt{MAX(userthreshold*5 , peakresidual/10 )}. This threshold is reported in the logger. Elsewhere, the values are currently set to zero.
[xxx.image.alpha.error] contains the errors of the spectral index solutions.

The following is a list of differences between MS-MFS (nterms > 1) and standard imaging, in the current CASA release.

1. Iterations always proceed as cs-clean major/minor cycles, and uses the full psf during minor cycle iterations. There are currently no user-controls on the cyclespeedup, and the flux-limit per major cycle is chosen as 10% of the peak residual. In future releases, this will be made more adaptive/controllable.

2. Currently, the following options are not supported for nterms > 1: psfmode, pbcorr, minpb, imagermode='mosaic', gridmode='aprojection', cyclespeedup, and allowed are one of Stokes I, Q, U, V, RR, LL, XX, YY at a time. More options and combinations are currently under development and testing. Under 'Using CASA' → 'Other Documentation' → 'Imaging Algorithms in CASA' you can find the latest implementations.

5.2.5.2 Mode channel

**ALERT:** Note that mode='channel' is intended as a shortcut to produce a cube based on the input MS channelization. It will be in the frame of the input MS. We recommend that users instead use the ‘velocity’ and ‘frequency’ modes which will produce cubes in other frames with more control of the cube spacing. These modes have defaults that will work from the MS spacing, reproducing the action of mode='channel'.

If mode='channel' is chosen, then an image cube will be created. This is an expandable parameter, with dependent parameters:

```plaintext
mode = 'channel'       # Spectral image definition(mfs, 
                      # channel, velocity,frequency)
 nchan = -1             # Number of channels (planes) in output image
 start = 0              # first input channel to use
 width = 1              # Number of input channels to average
 interpolation = 'nearest'  # Spectral interpolation(nearest, linear, cubic)
```

The default nchan=-1 will automatically produce a cube with the number of channels needed to span the (regridded) spectral windows of the MS. If multiple MSs are used, the spectral frames of these need to be identical, e.g. LSRK. **ALERT:** This often results in extra blank channels at the beginning and end of the image cube, so it is usually more precise to specify nchan and start to get what you want. For best results, we also recommend 'nearest' interpolation for the mode=channel.

The channelization of the resulting image is determined by the channelization in the MS of vis of the first spw specified (the “reference spw”). The actual channels to be gridded and used in

---

1Note that when TOPO is used, it refers to a time stamp at a given observation date. If more than one observation in TOPO is specified, this may lead to vastly erroneous values. Any conversion from TOPO to other frames such as BARY and LSRK should be performed for each individual observation, prior to clean or concatenation.
the clean are selected via the spw parameter as usual. The resulting image cube will have nchan channels spaced evenly in frequency. The first output channel will be located at the frequency of channel start in the (first) reference spw (independent of what channels are selected using spw). If width > 1, then input MS channels with centers within a frequency range given by \((\text{width} + 1)/2\) times the reference spw spacing will be gridded together (as in mode = 'mfs' above) into the channels of the output image cube. The output channel spacing is thus given by width channels in the reference spw of the MS.

The interpolation sub-parameter (§ 5.2.5.5) sets how channels are gridded into the image cube planes. For 'nearest', the channels in spw beyond the first are mapped into the nearest output image channel within half a channel (if any). Otherwise, the chosen interpolation scheme will be used. Image channels that lie outside the MS frequency range or have no data mapped to them will be blank in the output image, but will be in the cube.

Example:

```python
mode = 'channel'
nchan = 46
start = 5
width = 1
```

which will produce a 46-channel cube starting with channel 5 of the MS with the same channel width as the MS. Note: the start channel is in reference to the channels in the MS, not the subset selected by spw.

5.2.5.3 Mode frequency

For mode='frequency', an output image cube is created with nchan channels spaced evenly in frequency.

```python
mode = 'frequency'  # Spectral image definition
nchan = -1  # Number of channels (planes) in output image
start = ''  # Frequency of first image channel:
width = ''  # Image channel frequency width:
interpolation = 'linear'  # Spectral interpolation
outframe = ''  # velocity frame of output image
```

The frequency of the first output channel is given by start and spacing by width. Output channels have width also given by width. The sign of width determines whether the output channels ascend or descend in frequency. Data from the input MS with centers that lie within one-half an input channel overlap of the frequency range of ±width/2 centered on the output channels are gridded together.

The defaults are designed to safely choose output cube channels to span the input MS(s). The default nchan=-1 will choose the number of channels needed to span the frequencies of the channels.
in the MS. The defaults \texttt{start='} and \texttt{width=}’ will use the channel frequency and width of the first channel of the first specified spectral window selected in \texttt{spw}. \textbf{ALERT:} As in “channel” mode, this is currently the first channel of the first \texttt{spw}, not the first channel selected from that \texttt{spw}.

The \texttt{interpolation} sub-parameter (§5.2.5.5) sets how channels are gridded into the image cube planes.

Using the NGC5921 dataset as an example:

\begin{verbatim}
mode = 'frequency'
  nchan = 21
  start = '1412.830MHz'
  width = '50kHz'
  outframe = 'LSRK'
\end{verbatim}

would produce a 21-channel output cube with 50 kHz wide channels rather than the default channelization of the MS (24.4 kHz).

\subsection*{5.2.5.4 Mode velocity}

If mode='velocity' is chosen, then an output image cube with \texttt{nchan} channels will be created, with channels spaced evenly in velocity. Parameters are:

\begin{verbatim}
mode = 'velocity' # Spectral image definition(mfs,
    # channel, velocity,frequency)
  nchan = -1 # Number of channels (planes) in output image
  start = '' # Velocity of first image channel:
    # e.g '0.0km/s'(''=default)
  width = '' # Image channel velocity width: e.g
    # '-1.0km/s'(''=default)
  interpolation = 'linear' # Spectral interpolation(nearest,
    # linear, cubic)
  outframe = '' # velocity reference frame of output
    # image; '' =input
  veltype = 'radio' # velocity definition
\end{verbatim}

Note that velocities are calculated with respect to the rest frequency in the MS or specified through the \texttt{restfreq} parameter (§5.2.8).

The velocity of the first output channel is given by \texttt{start} and spacing by \texttt{width}. Averaging is as in \texttt{mode='frequency'}. The \texttt{interpolation} sub-parameter (§5.2.5.5) sets how channels are gridded into the image cube planes.

The defaults are designed to safely choose output cube channels to span the input MS(s). The default \texttt{nchan=-1} will choose the number of channels needed to span the velocities of the channels in the MS. The defaults \texttt{start=''} and \texttt{width=''} will use the channel velocity and width of the first channel of the first specified spectral window selected in \texttt{spw}. \textbf{ALERT:} As in “channel” mode, this is currently the first channel of the first \texttt{spw}, not the first channel selected from that \texttt{spw}.

Again, using the NGC5921 dataset as an example:
mode = 'velocity'
nchan = 21
start = '1383.0 km/s'
width = '10 km/s'
outframe = 'LSRK'

Note that in this case the velocity axis runs forward, as opposed to the default channelization for 'channel' or 'frequency'.

5.2.5.5 Sub-parameter interpolation

The interpolation sub-parameter controls how spectral channels in the MS are gridded into the output image cube. This is available in all modes except 'mfs'. The options are: 'nearest', 'linear', 'cubic'.

For 'nearest', the channels in spw beyond the first are mapped into the nearest output image channel within half a channel (if any).

For 'linear', the channels are gridded into the planes using weights given by a linear function of the frequency of the MS channel versus the plane. Each input channel will be mapped to 1 or 2 output planes. For most users, this is the best choice.

For 'cubic', the channels are gridded using a cubic interpolation function.

'Linear' and 'cubic' interpolation methods require that there are two datapoints that sandwich your new, regridded bin. This can introduce edge effects like in the first or last channel or adjacent to flagged channels where data is only available on one side of the spectrum. interpolation='nearest' will avoid such edge effects but may not work so well for data with spws that overlap. For mode='velocity' or 'frequency', 'linear' interpolation usually works best and for mode='channel' the 'nearest' interpolation method is superior. But this could be different for your dataset and you should carefully check your results.

5.2.6 Parameter resmooth

For large cubes, the psf will change as a function of frequency. clean will produce cubes with different synthesized beams per plane. All CASA analysis tasks can deal with such cubes. If one would like a common psf for all planes, typically the smallest possible beam, one can invoke the resmooth Boolean parameter. Alternatively, the cube can be convolved to the smallest common beam in a separate step vis immosaic (see Sect. 6.17).

5.2.7 Parameter phasecenter

The phasecenter parameter indicates which of the field IDs should be used to define the phase center of the mosaic image, or what that phase center is in RA and Dec. The default action is to use the first one given in the field list.

For example:
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\begin{alltt}
\verb|phasecenter='5'| # field 5 in multi-src ms
\verb|phasecenter='J2000 19h30m00 -40d00m00'| # specify position
\end{alltt}

Note that the format for angles prefers to use \texttt{hm} for RA/time units and \texttt{dm} for Dec/Angle units as separators. The colon :: separator is interpreted as RA/time even if its used for the Dec, so be careful not to copy/paste from other sources.

5.2.8 Parameter \texttt{restfreq}

The value of the \texttt{restfreq} parameter, if set, will over-ride the rest frequency in the header of the first input MS to define the velocity frame of the output image.

\textbf{ALERT:} The \texttt{restfreq} parameter takes the options of transitions and frequencies as in the corresponding \texttt{plotxy} parameter (§3.3.2.12), but the frame information is controlled under the \texttt{mode} parameter (§5.2.5).

For example:

\begin{verbatim}
restfreq='115.2712GHz',
\end{verbatim}

will set the rest frequency to that of the CO 1-0 line.

\textbf{ALERT:} Setting \texttt{restfreq} explicitly here in \texttt{clean} is good practice, and may be necessary if your MS has been concatenated from different files for different spectral windows (§2.2.12).

5.2.9 Parameter \texttt{spw}

The \texttt{spw} parameter selects the spectral windows that will be used to form the image, and possibly a subset of channels within these windows.

The \texttt{spw} parameter is a string with an integer, list of integers, or a range, e.g.

\begin{verbatim}
spw = '1' # select spw 1
spw = '0,1,2,3' # select spw 0,1,2,3
spw = '0~3' # same thing using ranges
\end{verbatim}

You can select channels in the same string with a \texttt{:} separator, for example

\begin{verbatim}
spw = '1:10~30' # select channels 10-30 of spw 1
spw = '0:5~55,3:5,6,7' # chans 5-55 of spw 0 and 5,6,7 of spw 3
\end{verbatim}

This uses the standard syntax for \texttt{spw} selection is given in §2.3.3. See that section for more options.

Note that the order in which multiple \texttt{spw}s are given is important for \texttt{mode = 'channel'}, as this defines the origin for the channelization of the resulting image.
5.2.10 Parameter stokes

The stokes parameter specifies the Stokes parameters for the resulting images. Note that forming Stokes Q and U images requires the presence of cross-hand polarizations (e.g. RL and LR for circularly polarized systems such as the VLA) in the data. Stokes V requires both parallel hands (RR and LL) for circularly polarized systems or the cross-hands (XY and YX) for linearly polarized systems such as ALMA and ATCA.

This parameter is specified as a string of up to four letters and can indicate stokes parameters themselves, Right/Left hand polarization products, or linear polarization products (X/Y). For example,

```
stokes = 'I'  # Intensity only
stokes = 'IQU'  # Intensity and linear polarization
stokes = 'IV'  # Intensity and circular polarization
stokes = 'IQUV'  # All Stokes imaging
stokes = 'RR'  # Right hand polarization only
stokes = 'XXYY'  # Both linear polarizations
```

are common choices (see the inline help of clean for a full range of possible options). The output image will have planes (along the “polarization axis”) corresponding to the chosen Stokes parameters.

If as input to deconvolution tasks such as clean, the stokes parameter includes polarization planes other than I, then choosing psfmode='hogbom' (§ 5.3.1.2) or psfmode='clarkstokes' (§ 5.3.1.3) will clean (search for components) each plane sequentially, while psfmode='clark' (§ 5.3.1.1) will deconvolve jointly.

**Alert:** As of Release 3.2, clean expects that all input polarizations are present. E.g. if you have RR and LL dual polarization data and you flagged parts of RR but not LL, clean will ignore both polarizations in slice. It is possible to split out a polarization product with split and image separately. But you will not be able to combine these part-flagged data in the uv-domain. We will remove that restriction in a future CASA release.

5.2.11 Parameter uvtaper

This controls the radial weighting of visibilities in the uv-plane (see § 5.2.12 below) through the multiplication of the visibilities by the Fourier transform of an elliptical Gaussian. This is itself a Gaussian, and thus the visibilities are “tapered” with weights decreasing as a function of uv-radius.

The uvtaper parameter expands the menu upon setting uvtaper=True to reveal the following sub-parameters:

```
uvtaper = True  # Apply additional uv tapering of visibilities.
outertaper = []  # uv-taper on outer baselines in uv-plane
innertaper = []  # uv-taper in center of uv-plane (not implemented)
```
The sub-parameters specify the size and optionally shape and orientation of this Gaussian in the uv-plane or optionally the sky plane. The \texttt{outertaper} refers to a Gaussian centered on the origin of the uv-plane.

Some examples:

\begin{verbatim}
outertaper=[]                        # no outer taper applied
outertaper=['5klambda']            # circular uv taper FWHM=5 kilo-lambda
outertaper=['5klambda','3klambda','45.0deg'] # elliptical Gaussian
outertaper=['10arcsec']            # on-sky FWHM 10"
outertaper=['300.0']               # 300m in aperture plane
\end{verbatim}

Note that if no units are given on the taper, then the default units are assumed to be meters in aperture plane.

\textbf{ALERT:} The \texttt{innertaper} option is not yet implemented.

\subsection*{5.2.12 Parameter weighting}

In order to image your data, we must have a map from the visibilities to the image. Part of that map, which is effectively a convolution, is the weights by which each visibility is multiplied before gridding. The first factor in the weighting is the “noise” in that visibility, represented by the data weights in the MS (which is calibrated along with the visibility data). The weighting function can also depend upon the uv locus of that visibility (e.g. a “taper” to change resolution). This is actually controlled by the \texttt{uvtaper} parameter (see \S\ref{5.2.11}). The weighting matrix also includes the convolution kernel that distributes that visibility onto the uv-plane during gridding before Fourier transforming to make the image of the sky. This depends upon the density of visibilities in the uv-plane (e.g. “natural”, “uniform”, “robust” weighting).

The user has control over all of these.

\textbf{ALERT:} You can find a weighting description in the online User Reference Manual at:

\url{http://casa.nrao.edu/docs/casaref/imager.weight.html}

The \texttt{weighting} parameter expands the menu to include various sub-parameters depending upon the mode chosen:

\subsubsection*{5.2.12.1 ‘natural’ weighting}

For \texttt{weighting='natural'}, visibilities are weighted only by the data weights, which are calculated during filling and calibration and should be equal to the inverse noise variance on that visibility.
Imaging weight $w_i$ of sample $i$ is given by

$$w_i = \omega_i = \frac{1}{\sigma_i^2}$$

(5.6)

where the data weight $\omega_i$ is determined from $\sigma_i$ is the rms noise on visibility $i$. When data is gridded into the same uv-cell for imaging, the weights are summed, and thus a higher uv density results in higher imaging weights. No sub-parameters are linked to this mode choice. It is the default imaging weight mode, and it should produce “optimum” image with the lowest noise (highest signal-to-noise ratio). Note that this generally produces images with the poorest angular resolution, since the density of visibilities falls radially in the uv-plane.

### 5.2.12.2 'uniform' weighting

For `weighting = 'uniform'`, the data weights are calculated as in 'natural' weighting. The data is then gridded to a number of cells in the uv-plane, and after all data is gridded the uv-cells are re-weighted to have “uniform” imaging weights. This pumps up the influence on the image of data with low weights (they are multiplied up to be the same as for the highest weighted data), which sharpens resolution and reduces the sidelobe level in the field-of-view, but increases the rms image noise. No sub-parameters are linked to this mode choice.

For uniform weighting, we first grid the inverse variance $\omega_i$ for all selected data onto a grid with uv cell-size given by $2/\text{FOV}$ where $\text{FOV}$ is the specified field of view (defaults to the image field of view). This forms the gridded weights $W_k$. The weight of the $i$-th sample is then:

$$w_i = \frac{\omega_i}{W_k}.$$  

(5.7)

### 5.2.12.3 'superuniform' weighting

The `weighting = 'superuniform'` mode is similar to the 'uniform' weighting mode but there is now an additional `npixels` sub-parameter that specifies a change to the number of cells on a side (with respect to uniform weighting) to define a uv-plane patch for the weighting renormalization. If `npixels=0` you get uniform weighting.

### 5.2.12.4 'radial' weighting

The `weighting = 'radial'` mode is a seldom-used option that increases the weight by the radius in the uv-plane, i.e.

$$w_i = \omega_i \cdot \sqrt{u_i^2 + v_i^2}. $$

(5.8)

Technically, I would call that an inverse uv-taper since it depends on uv-coordinates and not on the data per-se. Its effect is to reduce the rms sidelobes for an east-west synthesis array. This option has limited utility.
5.2.12.5 'briggs' weighting

The \texttt{weighting = 'briggs'} mode is an implementation of the flexible weighting scheme developed by Dan Briggs in his PhD thesis. See:

\url{http://www.aoc.nrao.edu/dissertations/dbriggs/}

This choice brings up the sub-parameters:

\begin{verbatim}
weighting = 'briggs' # Weighting to apply to visibilities
robust = 0.0 # Briggs robustness parameter
npixels = 0 # number of pixels to determine uv-cell size 0=> field of view
\end{verbatim}

The actual weighting scheme used is:

\[ w_i = \frac{\omega_i}{1 + W_k f^2} \]  \hspace{1cm} (5.9)

where \( W_k \) is defined as in uniform and superuniform weighting, and

\[ f^2 = \frac{(5 \times 10^{-R})^2}{\sum_i W_i^2 / \sum_i \omega_i} \]  \hspace{1cm} (5.10)

and \( R \) is the robust parameter.

The key parameter is the \texttt{robust} parameter, which sets \( R \) in the Briggs equations. The scaling of \( R \) is such that \( R = 0 \) gives a good trade-off between resolution and sensitivity. The \texttt{robust} \( R \) takes value between \(-2.0\) (close to uniform weighting) to \(2.0\) (close to natural).

Superuniform weighting can be combined with Briggs weighting using the \texttt{npixels} sub-parameter. This works as in 'superuniform' weighting (§ 5.2.12.3).

5.2.12.6 'briggsabs' weighting

For \texttt{weighting='briggsabs'}, a slightly different Briggs weighting is used, with

\[ w_i = \frac{\omega_i}{W_k R^2 + 2\sigma_R^2} \]  \hspace{1cm} (5.11)

where \( R \) is the robust parameter and \( \sigma_R \) is the noise parameter.

This choice brings up the sub-parameters:

\begin{verbatim}
weighting = 'briggsabs' # Weighting to apply to visibilities
robust = 0.0 # Briggs robustness parameter
noise = '0.0Jy' # noise parameter for briggs weighting when rmode='abs'
npixels = 0 # number of pixels to determine uv-cell size 0=> field of view
\end{verbatim}

Otherwise, this works as \texttt{weighting='briggs'} above (§ 5.2.12.5).
5.2.13 Parameter vis

The value of the vis parameter is either the name of a single MS, or a list of strings containing the names of multiple MSs, that should be processed to produce the image. The MS referred to by the first name in the list (if more than one) is used to determine properties of the image such as channelization and rest frequency.

For example,

\[
\text{vis} = \text{'ngc5921.ms'}
\]

set a single input MS, while

\[
\text{vis} = [\text{'ngc5921_day1.ms', 'ngc5921_day2.ms', 'ngc5921_day3.ms'}]
\]

points to three separate measurement sets that will be gridded together to form the image. This means that you do not have to concatenate datasets, for example from different configurations, before imaging.

For the multiple MS case, all selection commands like field, spw, etc. are lists that refer to the list of input MSs, like

\[
\text{spw}=[\text{'1:2~9','0:10~22','<2'}]
\]

\[
\text{field}=[\text{'0','ngc5921','12'}]
\]

will use the first entry of each selection criterion and apply it to the first dataset (spw='1:2~9' and field='0' to 'ngc5921_day1.ms'), the second selection criterion to the second dataset etc.

5.2.14 Primary beams in imaging

The CASA imaging task and tools use primary beams based on models for each observatory's antenna types. In addition to different antenna diameters, different functions may be used.

The voltage patterns are based on the following antenna primary beams, based on the TELESCOPE_NAME keyword in the OBSERVATION table:

VLA — Airy disk fitted to measurement. Note that a R/L beam squint is also included with feed dependent angle;

ALMA — Airy disk for 12m dish with a blockage of 1m;

ATA — Airy disk for 6m dish;

ATCA — polynomial fitted to measurement of main lobe;

BIMA, HATCREEK — Gaussian with halfwidth of \(\lambda/2D\);
CARMA — Airy patterns for the BIMA or OVRO dish sizes as appropriate;

GBT — polynomial fitted to measurement of main lobe;

GMRT — VLA Airy disk scaled to 45.0m;

IRAMPDB — Airy disk for dish of 15m with a blockage of 1m;

NRAO12M — VLA beam scaled to 12m;

OVRO — VLA Airy disk scaled to 10.4m;

SMA — Spheroidal function fit to FWHM;

WSRT — polynomial fitted to measurement of main lobe;

If the telescope name is unknown, or is CARMA or ALMA, then the DISH DIAMETER in the ANTENNA table is used with a scaled VLA pattern.

In mosaicking mode, clean will use frequency-dependent primary beams. It also appears that Airy or spheroidal beams are best behaved for mosaics (see § 5.3.15).

### 5.3 Deconvolution using CLEAN (clean)

To create an image and then deconvolve it with the CLEAN algorithm, use the clean task. This task will work for single-field data, or for multi-field mosaics (§5.3.15), in both narrow and wide-field imaging modes.

**Alert:** For large fractional bandwidths the psf in clean may vary considerably with frequency in data cubes. To accommodate this fact we have introduced a per-plane psf (dirty beam) when the change is larger than half the size of a pixel. Analysis tasks in CASA can deal with such beam variation. If a single beam size is requested, imsmooth can be invoked on the clean products to smooth to a common, uniform beam for all channels.

**Toolkit Note:** MEM is not included in clean, but is available in the toolkit.

Clean will use the CORRECTED_DATA column from your measurement set if it exists. If that column is not available, it will use DATA. The clean task utilizes many of the common imaging parameters. These are described above in § 5.2. There are also a number of parameters specific to clean. These are listed and described below.

The default inputs to clean are:

```
# clean :: Deconvolve an image with selected algorithm
vis = ''   # name of input visibility file
imagename = ['']   # Pre-name of output images
cleanfile = ''   # Text file with image names, sizes, centers
field = ''   # Field Name
spw = ''   # Spectral windows:channels: '' is all
```
selectdata = False # Other data selection parameters
mode = 'mfs' # Type of selection (mfs, channel, velocity, frequency)
nterms = 1 # Number of taylor terms to use
# for modeling the sky frequency dependence
reffreq = '' # Reference frequency for MFS
# (relevant only if nterms > 1)
gridmode = '' # The kind gridding kernel to be
# used for FFT-based transforms
niter = 500 # Maximum number of iterations
gain = 0.1 # Loop gain for cleaning
threshold = '0.0mJy' # Flux level to stop cleaning. Must include units
psfmode = 'clark' # method of PSF calculation to use during minor cycles
imagermode = '' # Use csclean or mosaic. If '', use psfmode
multiscale = [] # deconvolution scales (pixels);
# [] = default standard clean
interactive = False # use interactive clean (with GUI viewer)
mask = [] # cleanbox(es), mask image(s),
# and/or region(s) used in cleaning
imsiz e = [256, 256] # x and y image size in pixels,
# symmetric for single value
 cell = ['1.0arcsec', '1.0arcsec'] # x and y cell size. default unit arcsec
phasescenter = '' # Image phase center: position or field index
restfreq = '' # rest frequency to assign to image (see help)
stokes = 'I' # Stokes params to image (eg I,IV, QU, IQUV)
weighting = 'natural' # Weighting of uv (natural, uniform, briggs, ...)
atuve taper = False # Apply additional uv tapering of visibilities.
modelimage = '' # Name of model image(s) to initialize cleaning
restoringbeam = [''] # Output Gaussian restoring beam for CLEAN image
pbcor = False # Output primary beam-corrected image
minpb = 0.1 # Minimum PB level to use
usescratch = False # True if to save model
# visibilities in MODEL_DATA column

The clean task will produce a number of output images based on the root name given in imagename. These include:

<imagename>.clean.image # the restored image
<imagename>.clean.flux # the effective response (e.g. for pbcor)
<imagename>.clean.flux.pbcov erage # the PB coverage (ftmachine='mosaic' only)
<imagename>.clean.model # the model image
<imagename>.clean.residual # the residual image
<imagename>.clean.psf # the synthesized (dirty) beam

The mode, psfmode, imagermode, and weighting parameters open up other sub-parameters. These are detailed in the common imaging task parameters section (§5.2). The gridmode parameter (§5.3.13) is available to select more advanced imaging options such as widefield imaging and beam squint correction.

A typical setup for clean on the NGC5921 dataset, after setting parameter values, might look like:

vis = 'ngc5921.usecase.ms.contsub' # Name of input visibility file
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imagename = 'ngc5921.usecase.clean'  # Pre-name of output images
field = '0'  # Field Name
spw = ''  # Spectral windows:channels: '' is all
selectdata = False  # Other data selection parameters
mode = 'channel'  # Type of selection (mfs, channel, velocity, frequency)
nchan = 46  # Number of channels (planes) in output image
start = 5  # first input channel to use
width = 1  # Number of input channels to average
interpolation = 'linear'  # Spectral interpolation (nearest, linear, cubic)

gridmode = ''  # The kind gridding kernel to be used for FFT-based transforms
niter = 6000  # Maximum number of iterations
gain = 0.1  # Loop gain for cleaning
threshold = 8.0  # Flux level to stop cleaning. Must include units
psfmode = 'clark'  # method of PSF calculation to use during minor cycles
imagermode = ''  # Use csplane or mosaic, or image-plane only if ''
multiscale = []  # set deconvolution scales (pixels)
interactive = False  # use interactive clean (with GUI viewer)
mask = [108, 108, 148, 148]  # cleanbox(es), mask image(s), and/or region(s)
imsize = [256, 256]  # x and y image size in pixels
cell = [15.0, 15.0]  # x and y cell size. default unit arcsec
phasecenter = ''  # Image phase center: position or field index
restfreq = ''  # rest frequency to assign to image (see help)
stokes = 'I'  # Stokes params to image (eg I,IV, QU,IQUV)
weighting = 'briggs'  # Weighting to apply to visibilities
robust = 0.5  # Briggs robustness parameter

uv taper = False  # Apply additional uv tapering of visibilities.
modelimage = ''  # Name of model image(s) to initialize cleaning
restoringbeam = ['']  # Output Gaussian restoring beam for CLEAN image
pbcor = False  # Output primary beam-corrected image
minpb = 0.1  # Minimum PB level to use

An example of the `clean` task to create a continuum image from many channels is given below:

clean(vis='ggtau.1mm.split.ms',  # Use data in ggtau.1mm.split.ms
    imagename='ggtau.1mm',  # Name output images 'ggtau.1mm.*' on disk
    psfmode='clark',  # Use the Clark CLEAN algorithm
    imagermode='''',  # Do not mosaic or use csplane
    mask='''',  # Do not use clean box or mask
    niter=500, gain=0.1,  # Iterate 500 times using gain of 0.1
    mode='mfs',  # multi-frequency synthesis (combine channels)
    spw='0~2:2~57',  # Combine channels from 3 spectral windows
    field='0',
    stokes='I',  # Image stokes I polarization
    weighting='briggs',  # Use Briggs robust weighting
    rmode='norm', robust=0.5,  # with robustness parameter of 0.5
    cell=[0.1,0.1],  # Using 0.1 arcsec pixels
imsize=[256,256])  # Set image size = 256x256 pixels

This example will clean the entire inner quarter of the primary beam. However, if you want to limit the region over which you allow the algorithm to find clean components then you can make a deconvolution region (or mask). To use a deconvolution region, box, or mask, set the mask parameter (§5.3.6).

For example, you can set up a simple 'cleanbox' region. To do this, make a first cut at the image and clean the inner quarter. Then use the viewer to look at the image and get an idea of where the emission is located. You can use the viewer adjustment panel to view the image in pixel coordinates and read out the pixel locations of your cursor.

Then, you can use those pixel read-outs you just go to define a clean box region with the CASA region format described in Chapter D. For example, say you have a continuum source near the center of your image between the pixel coordinates [80,80] and [120,120], you may use the rectangular region:

\[
\text{mask} = 'box[[80\text{pix},80\text{pix}],[120\text{pix},120\text{pix}]]'
\]

For more complicated and multiple clean regions, it will be best to use the viewer to create them interactively or to create a region file (Chapter D) and use that file as an input like:

\[
\text{mask} = 'myregions.txt'
\]

The following are the clean specific parameters and their allowed values, followed by a description of carrying out interactive cleaning.

### 5.3.1 Parameter psfmode

The psfmode parameter chooses the “algorithm” that will be used to calculate the synthesized beam for use during the minor cycles in the image plane. The value types are strings. Allowed choices are 'clark' (default) and 'hogbom'.

#### 5.3.1.1 The clark algorithm

In the 'clark' algorithm, the cleaning is split into minor and major cycles. In the minor cycles only the brightest points are cleaned, using a subset of the point spread function. In the major cycle, the points thus found are subtracted correctly by using an FFT-based convolution. This algorithm is reasonably fast. Also, for polarization imaging, Clark searches for the peak in \(I^2 + Q^2 + U^2 + V^2\).
5.3.1.2 The hogbom algorithm

The hogbom algorithm is the “Classic” image-plane CLEAN, where model pixels are found iteratively by searching for the peak. Each point is subtracted from the full residual image using the shifted and scaled point spread function. In general, this is not a good choice for most imaging problems (clark or csclean are preferred) as it does not calculate the residuals accurately. But in some cases, with poor uv-coverage and/or a PSF with bad sidelobes, the Hogbom algorithm will do better as it uses a smaller beam patch. For polarization cleaning, Hogbom searches for clean peak in $I$, $Q$, $U$, and $V$ independently.

5.3.1.3 The clarkstokes algorithm

In the 'clarkstokes' algorithm, the Clark psf (§5.3.1.1) is used, but for polarization imaging the Stokes planes are cleaned sequentially for components instead of jointly as in 'clark'. This means that this is the same as 'clark' for Stokes I imaging only. This option can also be combined with imagermode='csclean' (§5.3.4).

5.3.2 The multiscale parameter

To activate multi-scale mode, specify a non-blank list of scales in the multiscale parameter. A good rule of thumb for starters is $[0, 2 \times \text{beam}, 5 \times \text{beam}]$, and maybe adding larger scales up to the maximum scale the interferometer can image. E.g. for a 2 arcsecond beam

$$\text{multiscale} = [0, 6, 10, 30]$$ # Four scales including point sources

These are given in numbers of pixels, and specify FWHM of the Gaussians used to compute the filtered images.

Setting the multiscale parameter to a non-empty list opens up the sub-parameter:

```plaintext
multiscale = [0, 6, 10, 30] # Four scales including point sources
negcomponent = -1 # Stop cleaning if the largest scale finds this number of neg components
smallscalebias = 0.6 # a bias to give more weight # toward smaller scales
```

The negcomponent sub-parameter is here to set the point at which the clean terminates because of negative components. For negcomponent > 0, component search will cease when this number of
negative components are found at the largest scale. If \texttt{negcomponent} = -1 then component search will continue even if the largest component is negative.

Increasing \texttt{smallscalebias} gives more weight to small scales. A value of 1.0 weighs the largest scale to zero and a value < 0.2 weighs all scales nearly equally. The default of 0.6 is usually a good number as it corresponds to a weighting that approximates the normalization of each component by its area. Depending on the image, however, it may be necessary to tweak the \texttt{smallscalebias} for a better convergence of the algorithm. Note that currently, this parameter is ignored by the MS-MFS algorithm. It will be available in a future release.

Multi-scale cleaning is also not as sensitive to the loop gain as regular cleaning algorithms. A loop gain of 0.3 may still work fine and will considerably speed up the processing time. Increasing the \texttt{cyclefactor} by a few (e.g. 5) may provide better stability in the solution, in particular when the data exhibit a severely non-Gaussian dirty beam.

The CASA multi-scale algorithm uses “Multi-scale CLEAN” to deconvolve using delta-functions and circular Gaussians as the basis functions for the model, instead of just delta-functions or pixels as in the other clean algorithms. This algorithm is still in the experimental stage, mostly because we are working on better algorithms for setting the scales for the Gaussians. The sizes of the Gaussians are set using the \texttt{scales} sub-parameter.

We are working on defining a better algorithm for scale setting. In the toolkit, there is a \texttt{nscale} argument which sets scales

\[ \theta_i = \theta_{bmin} 10^{(i-N/2)/2} \]  

(5.12)

where \( N = \text{nscales} \) and \( \theta_{bmin} \) is the fitted FWHM of the minor axis of the CLEAN beam.

### 5.3.3 Parameter \texttt{gain}

The \texttt{gain} parameter sets the fraction of the flux density in the residual image that is removed and placed into the clean model at each minor cycle iteration. The default value is \texttt{gain} = 0.1 and is suitable for a wide-range of imaging problems. Setting it to a smaller gain per cycle, such as \texttt{gain} = 0.05, can sometimes help when cleaning images with lots of diffuse emission. Larger values, up to \texttt{gain}=1, are probably too aggressive and are not recommended.

### 5.3.4 Parameter \texttt{imagermode}

This choose the mode of operation of \texttt{clean}, either as single-field deconvolution using image-plane major and minor cycles only (\texttt{imagermode=''}), single-field deconvolution using Cotton-Schwab (CS) residual visibilities for major cycles (\texttt{imagermode='csclean'}), or multi-field mosaics using CS major cycles (\texttt{imagermode='mosaic'}).

The default \texttt{imagermode='csclean'} choice specifies the Cotton-Schwab algorithm. This opens up the sub-parameters

\begin{verbatim}
  imagermode = 'csclean'  # Options: 'csclean' or
  # 'mosaic', '', uses psfmode
\end{verbatim}
Figure 5.1: Close-up of the top of the interactive clean window. Note the boxes at the left (where the iterations, cycles, and threshold can be changed), the buttons that control add/erase, the application of mask to channels, and whether to stop, complete, or continue cleaning, and the row of Mouse-button tool assignment icons.

```
cyclefactor = 1.5  # Controls how often major
                 # cycles are done. (e.g. 5 for
                 # frequently)
cyclespeedup = -1  # Cycle threshold doubles in
                 # this number of iterations
```

These options are explained below. In the CS mode, cleaning is split into minor and major cycles. For each field, a minor cycle is performed using the PSF algorithm specified in psfmode (§ 5.3.1). At major-cycle breakpoints, the points thus found are subtracted from the original visibilities. A fast variant does a convolution using a FFT. This will be faster for large numbers of visibilities. If you want to be extra careful, double the image size from that used for the Clark clean and set a mask to clean only the inner quarter or less (this is not done by default). This is probably the best choice for high-fidelity deconvolution of images without lots of large-scale structure.

Note that when using the Cotton-Schwab algorithm with a threshold (§ 5.3.12), there may be strange behavior when you hit the threshold with a major cycle. In particular, it may be above threshold again at the start of the next major cycle. This is particularly noticeable when cleaning a cube, where different channels will hit the threshold at different times.

In the empty mode (imagermode=''), the major and minor clean cycles work off of the gridded FFT dirty image, with residuals updated using the PSF calculation algorithm set by the psfmode parameter (§ 5.3.1). This method is not recommended for high dynamic range or high fidelity imaging applications, but can be significantly faster than CS clean (the default). Note that for this option only, if mask='', (no mask or box set) then it will clean the inner quarter of the image by default.
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**ALERT:** You will see a warning message in the logger, similar to this:

Zero Pixels selected with a Flux limit of 0.000551377 and a maximum Residual of 0.00751239

whenever it find 0 pixels above the threshold. This is normal, and not a problem, if you’ve specified a non-zero threshold. On the other hand, if you get this warning with the threshold set to the default of '0Jy', then you should look carefully at your inputs or your data, since this usually means that the masking is bad.

The option `imagermode='mosaic'` is for multi-field mosaics. This choice opens up the sub-parameters

- `imagermode` = 'mosaic'  # Use csclean or mosaic. If '', use psfmode
- `mosweight` = False     # Individually weight the fields of the mosaic
- `ftmachine` = 'mosaic'  # Gridding method for the image
- `scaletype` = 'SAULT'   # Controls scaling of pixels in the image plane.
- `cyclefactor` = 1.5     # change depth in between of csclean cycle
- `cyclespeedup` = -1     # Cycle threshold doubles in this number of iteration

These options are explained below.

### 5.3.4.1 Sub-parameter cyclefactor

This sub-parameter is activated for `imagermode='csclean'` and 'mosaic'.

The `cyclefactor` parameter allows the user to change the threshold at which the deconvolution cycle will stop and then degrid and subtract the model from the visibilities to form the residual. This is with respect to the breaks between minor and major cycles that the `clean` part would normally force. Larger values force a major cycle more often.

This parameter in effect controls the threshold used by CLEAN to test whether a major cycle break and reconciliation occurs:

\[
\text{cycle threshold} = \text{cyclefactor} \times \text{max sidelobe} \times \text{max residual}
\]

If mosaic or csclean diverges on your data, try a larger `cyclefactor`. A larger value typically increases the robustness of your deconvolution. The price, however, will be a slower algorithm. On the other hand, if you find that the cleaning is slow due to taking too many major cycle breaks, then reduce `cyclefactor`.

Note that currently the `cycle_threshold` will saturate at a maximum value of 0.80 even when you set `cyclefactor` to a very high value or you have very high PSF sidelobes. This means that with a `gain` = 0.1 you will get 3 minor cycles per major cycle when hitting the limit.
Some rules of thumb:

If you have data taken with a small number of antennas, for example from ALMA in the commissioning and early-science phase, then you will have high sidelobes in the PSF. In this case, you will have to reduce \texttt{cyclefactor} considerably, likely into the range 0.25 to 0.5, if you want efficient cleaning of simple source structures (e.g. point sources). You can use the \texttt{viewer} to look at your PSF image and see what the maximum sidelobe level is and judge accordingly.

However, if your uv-coverage results in a poor PSF and you have complex source structure, then you should reconcile often (a \texttt{cyclefactor} of 4 or 5). For reasonable PSFs, use \texttt{cyclefactor} in the range 1.5 to 2.0. For good PSFs, or for faster cleaning at the expense of some fidelity, we recommend trying a lower value, e.g. \texttt{cyclefactor} = 0.25, which at least in some of our mosaicing tests led to a speedup of around a factor of two with little change in residuals.

### 5.3.4.2 Sub-parameter cyclespeedup

This sub-parameter is activated for \texttt{imagermode='csclean'} and \texttt{'mosaic'}. The \texttt{cyclespeedup} parameter allows the user to let \texttt{clean} raise the threshold at which a major cycle is forced if it is not converging to that threshold. To do this, set \texttt{cyclespeedup} to an integer number of iterations at which if the threshold is not reached, the threshold will be doubled. See \texttt{cyclefactor} above for more details. By default this is turned off (\texttt{cyclespeedup} = -1). In our tests, a value like \texttt{cyclespeedup} = 50 has been used successfully.

### 5.3.4.3 Sub-parameter ftmachine

This sub-parameter is activated for \texttt{imagermode='mosaic'}. \textbf{Note:} The actual “ftmachine” used may be overridden by choices made to other parameters, such as \texttt{gridmode}.

The \texttt{ftmachine} parameter controls the gridding method and kernel to be used to make the image. A string value type is expected. Choices are: \texttt{‘ft’}, \texttt{‘sd’}, \texttt{‘both’}, or \texttt{‘mosaic’} (the default).

The \texttt{‘ft’} option uses the standard gridding kernel (as used in \texttt{clean}). The \texttt{‘sd’} option forces gridding as in single-dish data.

For combining single-dish and interferometer MS in the imaging, the \texttt{‘both’} option will allow \texttt{clean} to choose the \texttt{‘ft’} or \texttt{‘sd’} machines as appropriate for the data.

The \texttt{‘mosaic’} option (the default) uses the Fourier transform of the frequency-dependent primary beam (the aperture cross-correlation function in the uv-plane) as the gridding kernel. This allows the data from the multiple fields to be gridded down to a single uv-plane, with a significant speed-up in performance in most (non-memory limited) cases. The effect of this extra convolution is an addi-
tional multiplication (apodization) by the primary beam in the image plane. This can be corrected for, but does result in an image with optimal signal to noise ratio across it.

The primary beams used in CASA are described in §5.2.14.

**ALERT:** Note that making a non-square image (e.g. using unequal values in imsize) for ftmachine='mosaic' will grid the data into a uv-plane with correspondingly non-square cells. This has not been extensively tested, and may results in undesired image artifacts. We recommend that the user make square mosaic images when possible, but in principle non-square images should work.

### 5.3.4.4 Sub-parameter mosweight

If mosweight=False (default) then the data will be weighted for optimal signal to noise ratio across the mosaic image. This should be used for most cases.

If mosweight=True then individual mosaic field relative weights will be readjusted on a per visibility basis (much like uniform gridding weights). This may give better performance in cases where one or a few fields in the mosaic have drastically different weights and/or integration time, and it is desired that the noise be more “uniform” across the mosaic image. Use this with care, we have not explored its use fully.

### 5.3.4.5 Sub-parameter scaletype

The scaletype parameter controls weighting of pixels in the image plane. This sub-parameter is activated for imagermode='mosaic'.

The default scaletype='PBCOR' scales the image to have the correct flux scale across it (out to the beam level cutoff minpb). This means that the noise level will vary across the image, being increased by the inverse of the weighted primary beam responses that are used to rescale the fluxes. This option should be used with care, particularly if your data has very different exposure times (and hence intrinsic noise levels) between the mosaic fields.

If scaletype='SAULT' then the image will be scaled so as to have constant noise across it. This means that the point source response function varies across the image attenuated by the weighted primary beam(s). However, this response is output in the .flux image and can be later used to correct this.

Note that this scaling as a function of position in the image occurs after the weighting of mosaic fields specified by mosweight and implied by the gridding weights (ftmachine and weighting).

### 5.3.4.6 The threshold revisited

For mosaics, the specification of the threshold is not straightforward, as it is in the single field case. This is because the different fields can be observed to different depths, and get different weights in
the mosaic. We now provide internal rescaling (based on `scaletype`) so `clean` does its component
search on a properly weighted and scaled version of the sky.

For `ftmachine='ft'`, the minor cycles of the deconvolution are performed on an image that has
been weighted to have constant noise, as in 'SAULT' weighting (see §[5.3.4.5]). This is equivalent
to making a dirty mosaic by coadding dirty images made from the individual pointings with a
sum of the mosaic contributions to a given pixel weighted by so as to give constant noise across
the image. This means that the flux scale can vary across the mosaic depending on the effective
noise (higher weighted regions have lower noise, and thus will have higher “fluxes” in the 'SAULT'
map). Effectively, the flux scale that threshold applies to is that at the center of the highest-
weighted mosaic field, with higher-noise regions down-scaled accordingly. Compared to the true
sky, this image has a factor of the PB, plus a scaling map (returned in the `.flux` image). You will
preferentially find components in the low-noise regions near mosaic centers.

When `ftmachine='mosaic'` and `scaletype='SAULT'`, the deconvolution is also performed on a
“constant noise image”, as detailed above for 'ft'.

**ALERT:** The intrinsic image made using `ftmachine='mosaic'` is equivalent to a dirty mosaic that
is formed by coadding dirty images made from the individual fields after apodizing each by the
PB function. Thus compared to the true sky, this has a factor of the PB² in it. You would thus
preferentially find components in the centers of the mosaic fields (even more so than in the 'ft'
mosaics). We now rescale this image internally at major-cycle (and interactive) boundaries based
on `scaletype`, and do not have a way to clean on the raw unscaled dirty image (as was done in
previous released versions).

### 5.3.5 Parameter `interactive`

If `interactive=True` is set, then an interactive window will appear at various “cycle” stages while
you clean, so you can set and change mask regions. These breakpoints are controlled by the
`npercycle` sub-parameter which sets the number of iterations of clean before stopping.

```plaintext
interactive    =  True  # use interactive clean (with GUI viewer)
npercycle     =   100  # Number of iterations before interactive prompt
```

Note that `npercycle` is currently the only way to control the breakpoints in interactive clean.

For spectral cube imaging, it is often easier to deal with each channel in turn, rather than cleaning
all channels in each cycle. We therefore provide the `chaniter=True` option under 'mode', where it
will clean a channel fully before moving to the next one. You will set masks for each channel.

See the example of interactive cleaning in §[5.3.14]

### 5.3.6 Parameter `mask`

The `mask` parameter takes a list of elements, each of which can be a list of coordinates specifying a
box, or a string pointing to the name of a cleanbox file, mask image, or region file. These are used
by CLEAN to define a region to search for components.
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Note that for imagermode=''' (§5.3.4) the default with mask=''' is to restrict clean to the inner quarter of the image.

5.3.6.1 Setting clean boxes

mask can be a list of CASA regions. For example,

    mask = 'box[[80pix, 80pix],[120pix,120pix]],circle[[150pix,150pix],10pix]'

defines a box and a circle. They will be applied to all channels. To define different regions for different channel ranges, it will be best to use interactive mode in clean, the viewer (note that the viewer still created old format regions - they are still supported in CASA 3.3) or to create a CASA region file that contain the different regions. Chapter D describes the syntax of CASA regions. They can be specified by;

    mask = 'regionfile.rgn, regionfile2.rgn'

5.3.6.2 Using clean mask images

You can give the mask parameter a string containing the name of a mask image to be used for CLEAN to search for components. You can use interactive=True to create such a mask for your image (§5.3.5).

5.3.7 Parameter minpb

The minpb parameter sets the level down to which the primary beam (or more correctly the voltage patterns in the array) can go and have a given pixel included in the image. This is important as it defines where the edge of the visible image or mosaic is. The default is 0.1 or equivalent to the 10% response level. If there is a lot of emission near the edge, then set this lower if you want to be able to clean it out.

NOTE: The minpb parameter is the level in the “primary beam” (PB) at which the cut is made. If you are using ftmachine='mosaic' (§5.3.4.3), this will show up in the .flux.pbcov image (new in version 2.4.0). See the discussion of threshold (§5.3.4.6) for related issues.

5.3.8 Parameter modelimage

The modelimage parameter specifies the name(s) of one or more input starting image(s) to use to calculate the first residual before cleaning. These are used in addition to any image with a name defaulting from the imagename root (e.g. on a restart). The output model will contain this model plus clean components found during deconvolution.

If the units of the image are Jy/pixel, then this is treated as a model image.
If the units of the image are $\text{Jy/beam}$ or $\text{Jy per solid angle}$, then this is treated as a “single-dish” image and rescaled by the resolution (in the ‘beam’ image header keyword). Inclusion of the SD image here is superior to feathering it in later. See §5.5 for more information on feathering.

### 5.3.9 Parameter $niter$

The $niter$ parameter sets the maximum total number of minor-cycle CLEAN iterations to be performed during this run of $\text{clean}$. If restarting from a previous state, it will carry on from where it was. Note that the $\text{threshold}$ parameter can cause the CLEAN to be terminated before the requested number of iterations is reached.

### 5.3.10 Parameter $pbcor$

The $pbcor$ parameter controls whether the final $\text{.image}$ is scaled to correct for the Primary Beam of the array or not.

If $pbcor=False$ (the default), then no such scaling is done and the image is in whatever “raw” scaling used by the $\text{imagermode}$ algorithm underneath. For single-field cleaning with $\text{imagermode}=''$ or ‘$\text{csclean}$’, this is the standard constant-noise image. If $\text{imagermode}='\text{mosaic}'$, then this is the ‘SAULT’ scaled image (regardless of what $\text{scaletype}$ is set to).

If $pbcor=True$, the at the end of deconvolution and imaging the “raw” image is rescaled by dividing by the noise and PB correction image. This is what is output by $\text{clean}$ as the $\text{.flux}$ image.

Note that regardless of what you set $pbcor$ to, you can recover the other option using $\text{immath}$ ($§6.6$) to either multiply or divide by the $\text{.flux}$ image.

### 5.3.11 Parameter $\text{restoringbeam}$

The $\text{restoringbeam}$ parameter allows the user to set a specific Gaussian restoring beam to make the final restored $\text{.image}$ from the final $\text{.model}$ and residuals.

If $\text{restoringbeam}=''$ (the default), then the restoring beam is calculated by fitting to the PSF (e.g. the $\text{.psf}$ image). For a mosaic, this is at the center of the field closest to the $\text{phasecenter}$.

The restoring beam can also be used to establish a single beam for large fractional bandwidths. If the PSF changes more than half a pixel across all channels in a cube, the PSF itself will be stored in the form of a cube, changing size from channel to channel. A specified restoring beam will output all planes at the same resolution and thus collapse to a single PSF (note that this can also be done in hindsight using $\text{imsmooth}$).

To specify a restoring beam, provide $\text{restoringbeam}$ a list of $[\text{bmaj}, \text{bmin}, \text{bpa}]$ which are the parameters of an elliptical Gaussian. The default units are in arc-seconds for $\text{bmaj}$, $\text{bmin}$ components and degrees for the $\text{bpa}$ component.

For example,
restoringbeam=['10arcsec'] # circular Gaussian FWHM 10"
restoringbeam=['10.0','5.0','45.0deg'] # 10"x5" at PA=45 degrees

5.3.12 Parameter threshold

The threshold parameter instructs clean to terminate when the maximum absolute residual reaches this level or below. Note that it may not reach this residual level due to the value of the niter parameter which may cause it to terminate early.

If threshold is given a floating-point number, then this is the threshold in milli-Jansky.

You can also supply a flux density quanta to threshold, e.g.

```
threshold = '8.0mJy'
threshold = '0.008Jy'
```

(these do the same thing).

5.3.13 Parameter gridmode

The gridmode parameter is now provided to access more advanced deconvolution capabilities. The default gridmode='' is recommended for most cases.

The gridmode='widefield' option allows imaging in the wide-field regime where the W-term is not negligible. The CASA implementation allows both standard uv-plane faceting as well as the W-Projection algorithm\(^2\) or a combination of the two. Its sub-parameters are:

```
gridmode = 'widefield' # Gridding kernel for FFT-based
# transforms, default='' None
wprojplanes = 1 # Number of w-projection planes for convolution
facets = 1 # Number of facets along each axis (main image only)
```

The wprojplanes parameter sets the number of pre-computed w-planes used for the W-Projection algorithm (wprojplanes=1 disables w-projection). The facets parameter sets the number of facets used. W-Projection, if used, is done for each facet. See §5.3.18 below for more on wide-field imaging.

gridmode='aprojection': A-Projection is an algorithm to account for the effects of the antenna primary beam (PB) during imaging. The time-dependent effects of the PB are projected-out during the imaging phase and the PB is included in the prediction phase of the iterative image deconvolution (see Bhatnagar, Cornwell, Golap & Uson 2008, A&A, 487, 419) for more details. Please also refer to this publication in your papers if this algorithm is used for imaging. The narrow-band A-Projection can be used by setting the gridmode='aprojection' in the clean task. This opens up the following new parameters:

gridmode = 'aprojection'  # Gridding kernel for FFT-based
# transforms, default='None

cfcache = 'cfcache.dir'  # Convolution function cache directory

rotpainc = 5.0  # Parallactic angle increment
# (degrees) for OTF A-term rotation

painc = 360.0  # Parallactic angle increment
# (degrees) for computing A-term

cfcache is used to cache functions required in the A-Projection algorithm. The PB is rotated
on-the-fly when a change of greater than rotpainc is detected. Alternatively, PB is re-computed
if the P.A. changes by greater than painc.

Note that this code is still in the development and testing stage and should be used on shared-risk
basis. Note also that the cost of imaging will be higher when using A-Projection. Therefore make
a careful evaluation of whether you need to invoke it.

5.3.14 Interactive Cleaning — Example

If interactive=True is set, then an interactive window will appear at various “cycle” stages while
you clean, so you can set and change mask regions. These breakpoints are controlled by the
npercycle sub-parameter which sets the number of iterations of clean before stopping.

The window controls are fairly self-explanatory. It is basically a form of the viewer. A close-up of
the controls are shown in Figure 5.1, and an example can be found in Figures 5.2–5.4. You assign
one of the drawing functions (rectangle or polygon, default is rectangle) to the right-mouse button
(usually), then use it to mark out regions on the image. Zoom in if necessary (standard with the
left-mouse button assignment). Double-click inside the marked region to add it to the mask. If you
want to reduce the mask, click the Erase radio button (rather than Add), then mark and select as
normal. When finished setting or changing your mask, click the green clockwise arrow “Continue
Cleaning” Next Action button. If you want to finish your clean with no more changes to the mask,
hit the blue right arrow “Apply mask edits and proceed with non-interactive clean” button. If you
want to terminate the clean, click the red X “Stop deconvolving now” button.

While stopped in an interactive step, you can change a number of control parameters in the boxes
provided at the left of the menu bar. The main use of this is to control how many iterations
before the next breakpoint (initially set to npercycle), how many cycles before completion (initially
equal to niter/npercycle), and to change the threshold for ending cleaning. Typically, the user
would start with a relatively small number of iterations (50 or 100) to clean the bright emission
in tight mask regions, and then increase this as you get deeper and the masking covers more of
the emission region. For extended sources, you may end up needing to clean a large number of
components (10000 or more) and thus it is useful to set niter to a large number to begin with
— you can always terminate the clean interactively when you think it is done. Note that if you
change iterations you may also want to change cycles or your clean may terminate before you
expect it to.

For strangely shaped emission regions, you may find using the polygon region marking tool (the
second from the right in the button assignment toolbar) the most useful.
Figure 5.2: Screen-shots of the interactive clean window during deconvolution of the VLA 6m Jupiter dataset. We start from the calibrated data, but before any self-calibration. In the initial stage (left), the window pops up and you can see it dominated by a bright source in the center. Next (right), we zoom in and draw a box around this emission. We have also at this stage dismissed the tape deck and Position Tracking parts of the display (§7.2) as they are not used here. We have also changed the iterations to 30 for this boxed clean. We will now hit the Next Action Continue Cleaning button (the green clockwise arrow) to start cleaning.

The sequence of cleaning starting with the “raw” externally calibrated data is shown in Figures 5.2 – 5.4.

The final result of all this cleaning for Jupiter is shown in Figure 5.5. The viewer (§7) was used to overplot the polarized intensity contours and linear polarization vectors calculated using immath (§6.6) on the total intensity. See the following chapters on how to make the most of your imaging results.

For spectral cube images you can use the tapedeck to move through the channels. You also use the panel with radio buttons for choosing whether the mask you draw applies to the Displayed Plane or to All Channels. See Figure 5.6 for an example. Note that currently the Displayed Plane option is set by default. This toggle is unimportant for single-channel images or mode='mfs'.

**Advanced Tip:** Note that while in interactive clean, you are using the viewer. Thus, you have the ability to open and register other images in order to help you set up the clean mask. For
Figure 5.3: We continue in our interactive cleaning of Jupiter from where Figure 5.2 left off. In the first (left) panel, we have cleaned 30 iterations in the region previously marked, and are zoomed in again ready to extend the mask to pick up the newly revealed emission. Next (right), we have used the Polygon tool to redraw the mask around the emission, and are ready to **Continue Cleaning** for another 100 iterations.

Example, if you have a previously cleaned image of a complex source or mosaic that you wish to use to guide the placement of boxes or polygons, just use the Open button or menu item to bring in that image, which will be visible and registered on top of your dirty residual image that you are cleaning on. You can then draw masks as usual, which will be stored in the mask layer as before. Note you can blink between the new and dirty image, change the colormap and/or contrast, and carry out other standard viewer operations. See § 7 for more on the use of the viewer.

**ALERT:** Currently, interactive spectral line cleaning is done globally over the cube, with halts for interaction after searching all channels for the requested npercycle total iterations. It is more convenient for the user to treat the channels in order, cleaning each in turn before moving on. This will be implemented in an upcoming update.
Figure 5.4: We continue in our interactive cleaning of Jupiter from where Figure 5.3 left off. In the first (left) panel, it has cleaned deeper, and we come back and zoom in to see that our current mask is good and we should clean further. We change `npercycle` to 500 (from 100) in the box at upper right of the window. In the final panel (right), we see the results after this clean. The residuals are such that we should terminate the clean using the red X button and use our model for self-calibration.

5.3.15 Mosaic imaging

The `clean` task contains the capability to image multiple pointing centers together into a single “mosaic” image. This ability is controlled by setting `imagermode='mosaic'` (§ 5.3.4).

The key parameter that controls how `clean` produces the mosaic is the `ftmachine` sub-parameter (§ 5.3.4.3). For `ftmachine='ft'`, `clean` will perform a weighted combination of the images produced by transforming each mosaic pointing separately. This can be slow, as the individual sub-images must be recombined in the image plane. **NOTE:** this option is preferred for data taken with sub-optimal mosaic sampling (e.g. fields too far apart, on a sparse irregular pattern, etc.).

The primary beams used in CASA are described in § 5.2.14.

If `ftmachine='mosaic'`, then the data are gridded onto a single uv-plane which is then transformed to produce the single output image. This is accomplished by using a gridding kernel that approximates the transform of the primary beam pattern. Note that for this mode the
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Figure 5.5: After clean and self-calibration using the intensity image, we arrive at the final polarization image of Jupiter. Shown in the viewer superimposed on the intensity raster is the linear polarization intensity (green contours) and linear polarization B-vectors (vectors). The color of the contours and the sampling and rotation by 90 degrees of the vectors was set in the Display Options panel. A LEL expression was used in the Load Data panel to mask the vectors on the polarized intensity.

<imagername>.flux image includes this convolution kernel in its effective weighted response pattern (needed to “primary-beam correct” the output image). For this mode only, an additional image <imagername>.flux.pbcoverage is produced that is the primary-beam coverage only used to compute the minpb cutoff (§5.3.7).

ALERT: In order to avoid aliasing artifacts for ftmachine=’mosaic’ in the mosaic image, due to the discrete sampling of the mosaic pattern on the sky, you should make an image in which the desired unmasked part of the image (above minpb) lies within the inner quarter. In other words, make an image twice as big as necessary to encompass the mosaic.

It is also important to choose an appropriate phascenter for your output mosaic image (§ section:im.pars.phascenter). The phase center should not be at the edge of an image with pointings
Figure 5.6: Screen-shot of the interactive clean window during deconvolution of the NGC5921 spectral line dataset. Note where we have selected the mask to apply to the Displayed Plane rather than All Channels. We have just used the Polygon tool to draw a mask region around the emission in this channel, which will apply to this channel only.

around it. In that case, FFT aliasing may creep into the image.

An example of a simple mosaic clean call is shown below:

clean(vis='n4826_tboth.ms',
      imagename='tmosaic',
      mode='channel',
      ...)
5.3.16 Heterogeneous imaging

The `clean` task and underlying tools can handle cases where there are multiple dish sizes, and thus voltage patterns and primary beams, in the array. This is effected by using the dish sizes stored in the ANTIENNA sub-table of the MS. Depending on how the data was written and imported into CASA, the user may have to manually edit this table to insert the correct dish sizes (e.g. using `browsetable` or the `tb` table tool).

5.3.17 Polarization imaging

The `clean` task handles full and partial Stokes polarization imaging through the setting of the `stokes` parameter (§5.2.10). The subsequent deconvolution of the polarization planes of the image and the search for clean components is controlled by the `psfmode` parameter (§5.3.1). If the `stokes` parameter includes polarization planes other than I, then choosing `psfmode='hogbom'` (§5.3.1.2) or `psfmode='clarkstokes'` (§5.3.1.3) will clean (search for components) each plane sequentially, while `psfmode='clark'` (§5.3.1.1) will deconvolve jointly.

The interactive `clean` example given above (§5.3.14) shows a case of polarization imaging.

5.3.18 Wide-field imaging and deconvolution in `clean`

When imaging sufficiently large angular regions, the sky can no longer be treated as a two-dimensional plane and the use of the standard `clean` task will produce distortions around sources that become increasingly severe with increasing distance from the phase center. In this case, one must use a “wide-field” imaging algorithm such as w-projection or faceting.

When is wide-field imaging needed? The number of required facets $N$ depends on the maximum baseline $B_{\text{max}}$, the dish diameter $D$ and the wavelength $\lambda$ as:

$$N = \frac{B_{\text{max}} \lambda}{D^2} \quad (5.13)$$
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Table 5.1: Combinations of observing band (wavelength,) and antenna array configurations that require w-projection.

<table>
<thead>
<tr>
<th>Receiver Band</th>
<th>Wavelength [cm]</th>
<th>Array configurations</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>430</td>
<td>A/B/C/D</td>
</tr>
<tr>
<td>L</td>
<td>20</td>
<td>A/B/C</td>
</tr>
<tr>
<td>S</td>
<td>10</td>
<td>A/B</td>
</tr>
<tr>
<td>C</td>
<td>5</td>
<td>A</td>
</tr>
<tr>
<td>X</td>
<td>3</td>
<td>A</td>
</tr>
<tr>
<td>Ku</td>
<td>2</td>
<td>A</td>
</tr>
<tr>
<td>K</td>
<td>1.4</td>
<td>–</td>
</tr>
<tr>
<td>Ka</td>
<td>0.9</td>
<td>–</td>
</tr>
<tr>
<td>Q</td>
<td>0.7</td>
<td>–</td>
</tr>
</tbody>
</table>

and w-projection is required when \( N > 1 \). (For details, see “Synthesis Imaging in Radio Astronomy II”, ed. Taylor, G., Carilli, C., Perley, R. 1999). With 25 m diameter JVLA dishes (which implies that imaging is requested out to the primary beam FWHM), w-projection is required for array configurations as listed in Table 5.1.

The relevant inputs for clean for wide-field imaging are:

```plaintext
gridmode = 'widefield'  # The kind gridding kernel to be used for FFT-based transforms
wprojplanes = 1         # Number of w-projection planes for convolution
facets = 1              # Number of facets along each axis (main image only)
```

Most of the clean parameters behave as described previously.

Wide-field imaging can be carried out using two major modes: First, the w-projection mode as chosen with ftmachine deals with the w-term (the phase associated with the sky/array curvature) internally. Secondly, the image can be broken into many facets, each small enough so that the w-term is not significant. These two basic methods can be combined, as discussed below in § 5.3.18.4.

5.3.18.1 Outlier fields

When using wide-field imaging, the position and image size of any independent images must be specified. Those positions will be used to add additional cleaning components to strong sources that may reside in that area and influence the central image.

There are two options to specify the outlier fields:

- **Direct listing of fields** The outlier field directions are provided via their centers (phasecenter parameter), and their sizes as a second entry in the imsize parameter, e.g. 128 pixels in the example below. clean will derive two additional images and their names are to be provided in the imagename field that will then be a list of the main field name plus the outlier field names:
vis = 'wfield.ms'  # name of input visibility file
imagename=['n5921','outlier1','outlier2']  # Pre-name of output images
outlierfile = ''  # Text file with image names, sizes, centers
mask = [[['image_setup.rgn']],[''],['']]
imsiz = [[2048,2048],[128,128],[128,128]]  # Image size in pixels (nx,ny)
cell = '1.0arcsec'  # The image cell size in arcseconds [x,y].
phaseten = ['','J2000 13h27m20.98 43d26m28.00', 'J2000 13h30m52.158 43d23m08.00']

**Outlier file** For many outlier fields, it may be easier to setup the main interface to clean for the main field only and list outlier fields in an additional outlierfile:

imagename='n5921'
outlierfile = 'outliers.txt'
imsiz=[1024,1024]
phaseten = ''

*outliers.txt* provides all outlier fields with a syntax that is similar to the direct input, but separated by field. Below is an example for an outlierfile:

```plaintext
# content of outliers.txt
#
# outlier field1
imagename='outlier1'
imsiz=[512,512]
phaseten = 'J2000 13h30m52.15 43d23m08.00'
mask='box[[245pix,245pix],[265pix,265pix]]'
#
# outlier field2
imagename='outlier2'
imsiz=[512,512]
phaseten = 'J2000 13h24m08.16 43d09m48.0'
```

The syntax rules for the outlier files are:

- each field must begin with imagename followed by
- imsize and phaseten must be given
- optionally a mask can be provided. The mask parameter follows the CASA region file convention (Chapter D) or can be a mask file or LEL string.

The older AIPS-style convention (and box definition) that was used in CASA 3.2 and earlier is still supported in CASA 3.3 but will be deprecated for CASA 3.4 and higher.
5.3.18.2 Setting up w-projection

The w-projection mode is controlled using wprojplanes sub-parameter, e.g.

```
gridmode = 'widefield'  # The kind gridding kernel to be used for FFT-based transforms
wprojplanes = 64       # Number of w-projection planes for convolution
facets = 1             # Number of facets along each axis (main image only)
```

will construct 64 w-projection planes.

The w-projection algorithm is much faster than using faceting, but it does consume a lot of memory. On most 32-bit machines with 1 or 2 Mbytes of memory, images larger than about 4000 × 4000 cannot be made.

5.3.18.3 Setting up faceting

Faceting will break the image into many small parts. This is invoked using facets:

```
gridmode = 'widefield'  # The kind gridding kernel to be used for FFT-based transforms
wprojplanes = 1        # Number of w-projection planes for convolution
facets = 7             # Number of facets along each axis (main image only)
```

In this example the image is broken into 49 (7 × 7) facets.

A reasonable value of facets is such that the image width of each facet does not need the w-term correction. The computation method with pure faceting is slow, so that w-projection is recommended.

5.3.18.4 Combination of w-projection and faceting

You can also use a combination of w-projection and faceting:

```
gridmode = 'widefield'  # The kind gridding kernel to be used for FFT-based transforms
wprojplanes = 32       # Number of w-projection planes for convolution
facets = 3             # Number of facets along each axis (main image only)
```

This hybrid method allows for a smaller number of wprojplanes in order to try to conserve memory if the image size approached the memory limit of the computer. However, there is a large penalty in execution time.
5.4 Primary Beam Correction (impbcor, widebandpbcor)

The primary beam correction can be applied during the imaging with clean. It is also possible to correct after imaging using the task impbcor for 'regular' data sets, or widebandpbcor for those that used the Taylor-term expansion function in clean (nterms>1). pbcor has the following inputs:

```python
# impbcor :: Construct a primary beam corrected image from an image
# and a primary beam pattern.
imagename = '' # Name of the input image
pbimage = '' # Name of the primary beam
# image which must exist or
# array of values for the pb response. Default ""
outfile = '' # Output image name. If empty, no image is written.
# Default ""
box = '' # One or more boxes to use
# for fit region(s). Default is
# to use the entire directional plane.
region = '' # The region to correct. Default is entire image. If
# both box and region are specified, box is used and
# region is not.
chans = '' # The frequency planes to correct. Default is all
# frequencies.
stokes = 'I' # The correlations to correct. Default is all.
mask = [] # Boolean LEL expression or mask region. Default is
# none.
mode = 'velocity' # Divide or multiply the image by the primary beam
# image. Minimal match supported. Default "divide"
cutoff = -1.0 # PB cutoff. If mode is "d", all values less than this
# will be masked. If "m", all values greater will be
# masked. Less than 0, no cutoff. Default no cutoff
wantreturn = False # Return an image tool
# referencing the corrected image?
```

The main inputs are the input image and the image of a primary beam (usually your “image.flux” output image from clean) in the pbimage parameter. The mode parameter will typically be 'divide' but it is also possible to multiply with the beam pattern.

widebandpbcor has the following options

```python
# widebandpbcor :: Wideband PB-correction on the output of the MS-MFS algorithm
vis = '' # Name of measurement set.
imagename = '' # Name-prefix of multi-termimages to operate on.
nterms = 2 # Number of taylor terms to use
threshold = '' # Intensity above which to
# re-calculate spectral index
action = 'pbcor' # PB-correction (pbcor) or
# only calc spectral-index (calcalpha)
```
action='pbcor' computes Taylor-coefficient images that represent the primary beam spectrum and applies them to the input Taylor coefficient images. The action='calcalpha' will recalculate spectral index maps based on the primary beam correction polynomials.

5.5 Combined Single Dish and Interferometric Imaging (feather)

The term “feathering” is used in radio imaging to describe how to combine or “feather” two images together by forming a weighted sum of their Fourier transforms in the (gridded) uv-plane. Intermediate size scales are down-weighted to give interferometer resolution while preserving single-dish total flux density.

The feathering technique does the following:

1. The single-dish and interferometer images are Fourier transformed.

2. The beam from the single-dish image is Fourier transformed ($FTSDB(u, v)$), (alternatively, one can specify some smaller portion of the single dish aperture, which corresponds to a wider beam).

3. The Fourier transform of the interferometer image is multiplied by ($1 - FTSDB(u, v)$). This basically down weights the shorter spacing data from the interferometer image.

4. The Fourier transform of the single-dish image is scaled by the volume ratio of the interferometer restoring beam to the single dish beam.

5. The results from 3 and 4 are added and Fourier transformed back to the image plane.

The term feathering derives from the tapering or down-weighting of the data in this technique; the overlapping, shorter spacing data from the deconvolved interferometer image is weighted down compared to the single dish image while the overlapping, longer spacing data from the single-dish are weighted down compared to the interferometer image.

The tapering uses the transform of the low resolution point spread function. This can be specified as an input image or the appropriate telescope beam for the single-dish. The point spread function for a single dish image may also be calculated using clean.

Other Packages:
The feather task is analogous to the AIPS IMERG task and the MIRIAD immerge task with option 'feather'.
Advice: Note that if you are feathering large images, be advised to have the number of pixels along the X and Y axes to be composite numbers and definitely not prime numbers. In general FFTs work much faster on even and composite numbers. You may use subimage function of the image tool to trim the number of pixels to something desirable.

The inputs for feather are:

```plaintext
# feather :: Combine two images using their Fourier transforms
imagename = '' # Name of output feathered image
highres = '' # Name of high resolution (interferometer) image
lowres = '' # Name of low resolution (single dish) image
sdfactor = 1.0 # Scale factor to apply to Single Dish image
effdishdiam = -1.0 # New effective SingleDish diameter to use in m
lowpassfiltersd = False # Filter out the high spatial frequencies of the SD

The single-dish data cube is specified by the lowres and the interferometric data cube by the highres keyword. The combined, feathered output cube name is given by the imagename parameter. sdfactor can be used to adjust the flux calibration of the images. Since single-dish processing typically involves the fit of a baseline level, it might be the one with the most uncertain calibration and sdfactor will multiply with the single-dish image values for any needed correction.

The weighting functions for the data are usually the Fourier transform of the Single Dish beam FFT(PB_{SD}) for the Single dish data, and the inverse, 1-FFT(PB_{SD}) for the interferometric data. It is possible, however, to change the weighting functions by pretending that the SD is smaller in size via the effdishdiameter parameter. This tapers the high spatial frequencies of the SD data and adds more weight to the interferometric data. The lowpassfiltersd can take out artifacts at very high spatial frequencies that are often present but non-physical in SD data.

Note that the only inputs are for images and feather will attempt to regrid the images to a common shape, i.e. pixel size, pixel numbers, and spectral channels. feather does not do any deconvolution but combines presumably deconvolved images after the fact. This implies that the short spacings extrapolated by the deconvolution process will be those that are down-weighted the most. The single dish image must have a well-defined beam shape and the correct flux units for a model image (Jy/beam instead of Jy/pixel) so use the tasks imhead and immath first to convert if needed.

Starting with a cleaned synthesis image and a low resolution image from a single dish telescope, the following example shows how they can be feathered:

```plaintext
feather(imagename='feather.im', # Create an image called feather.im
        highres='synth.im', # The synthesis image is called synth.im
        lowres='single_dish.im' # The SD image is called single_dish.im
)
```

### 5.5.1 Visual Interface for feather (casafeather)

CASA also provides a visual interface to the feather task. The interface is run from a command line outside CASA by typing casafeather in a shell. Fig. 5.7 shows an example. As a first step,
one needs to specify a high and a low resolution image, typically an interferometric and a single dish map. Note that the single dish map needs to be in units of Jy beam\(^{-1}\). An output image is usually specified, too, and an additional image, such as a non-deconvolved (dirty) interferometric image can be specified, too. On the main GUI, press “Feather” to start the feathering process, which includes regridding the low resolution image to the high resolution image.

Figure 5.7: Visual “casafeather” interface to the feather task.

“casafeather” has the ability to show two major rows of displays (see Fig. 5.7) that can be turned on or off. A good visualization is usually obtained by making both axes logarithmic. This can be specified in the “Customize menu”, the toothed wheel symbol at the top of the panel. The two rows of displays are: 1) “Original Data Slice”: Cuts through the u and v directions of the Fourier transformed input images. A vertical line shows the location of the effective dish diameter(s). 2) “Feathered Data Slice”: The same cuts, but scaled by the “low resolution scale factor” and
weighted by the weighting functions (see \S5.5). In this display, the weighting functions themselves are shown, too.

At the top of the display **effdshdiameter** for u and v and **sdfactor** can be provided in the “Effective Dish Diameter” and “Low Resolution Scale Factor” input boxes.

The data can be visualized in different forms. The data type to be displayed can be selected in the “Color Options” menu. The data can be the unmodified, original data, or data that have been convolved with the high or low resolution beams. One can also select to display data that were weighted and scaled by the functions discussed above.

The data can also be displayed in the form of a “scatter plot” (Fig.5.8). This allows one to check for differences in flux. In particular, the scaling parameter should be adjusted such that the flux of the Low-resolution data, convolved with the High beam, weighted and scaled, is the same as the Dirty data, convolved with the Low beam, weighted (use the High resolution data instead of the Dirty data if the latter are not available). If that can be achieved, the flux adjustments should be roughly correct. The “scatter plot” can display any two data sets on the two axes, selected from the “Color Preferences” menu.

![Figure 5.8: The scatter plot in casafeather.](image)

The “Customize” button at the top (toothed wheel), allows one to set the display parameters as seen in Fig.5.9. Options are to show the slice plot, the scatter plot, or the legend. One can also select between logarithmic and linear axes, and whether the x-axis for the slices are in the u, or v, or both directions, or, alternatively a radial average in the uv-plane can be used. For data cubes one can also select a particular velocity plane, or to average the data across all velocity channels.

### 5.6 Making Deconvolution Masks or Box Regions

For most careful imaging, you will want to restrict the region over which you allow CLEAN components to be found. To do this, you can create a ‘deconvolution region’ or ‘mask’ image using the **boxit** or the **viewer**. Note that **clean** can take simple boxes or box files as direct input to its **mask** parameter, so these tasks are most useful when direct input to clean (or use of interactive clean) will not suffice.
There are two ways to construct region files or mask images for use in deconvolution. The boxit task will find a set of box regions based upon an input image and control parameters.

5.6.1 Making Deconvolution Regions from an Image (boxit)

The boxit task creates “cleanbox” deconvolution regions automatically from an image. It searches the image to find “islands”: all contiguous sets of pixels above the given threshold. The extreme x- and y-pixels of the island are used to determine the corners of a rectangular box that covers each island. The set of boxes are written out into a single region file with extension .rgn. Boxit works on single-plane images as well as multi-channel images: in the latter case, the thresholding and boxing is done separately in each plane of the image. The output region file from boxit can be used as the mask input parameter for the clean task (§ 5.3).

The parameter inputs for boxit are:

```plaintext
# boxit :: Box regions in image above given threshold value.
imagename = '' # Name of image to threshold
regionfile = '' # Output region file
threshold = '0.0mJy' # Threshold value. Must include units.
minsize = 2 # Minimum number of pixels for a boxable island
diag = False # Count diagonal connections?
boxstretch = 1 # Increase box sizes by this many pixels beyond thresholded pixels.
overwrite = False # Overwrite existing region file?
```
The `regionfile` parameter specifies the root name of the region file. It will automatically be given `.rgn` as the file extension. The `minsize` parameter specifies the smallest island that qualifies to be boxed. It refers to the total number of pixels in the island. To include pixels connected only on the diagonal as being part of the same island, set the `diag` parameter to `True`. The `boxstretch` parameter increases the size of the boxes beyond the extent of the island, and can range from -1 to 5. For a value of 1 (the default), the box is stretched by one pixel in each outward direction; therefore, each side of the box lengthens by two pixels. Finally, the parameter `overwrite` specifies whether an existing region file can be overwritten.

**ALERT:** The `boxit` task is a prototype under active development and coded in Python. Eventually we will add functionality to deal with the creation of non-rectangular regions and with multi-plane masks, as well as efficiency improvements.

### 5.7 Insert an Image Model (ft)

The `ft` task will add a source model (units should be Jy/pixel) or a clean component list to a Measurement Set. This is especially useful if you have a resolved calibrator and you want to start with a model of the source before you derive accurate gain solutions. This is also helpful for self-calibration (see §5.9 below).

The inputs for `ft` are:

```plaintext
# ft :: Insert a source model a visibility set:
vis = ''  # Name of input visibility file (MS)
field = ''  # Field selection
spw = 'all'  # Spw selection
model = ''  # Name of input model image(s)
nterms = 1  # Number of terms used to model the sky frequency dependence
complist = ''  # Name of component list
incremental = False  # Add to the existing model visibility?
usescratch = False  # If True predicted visibility is stored in MODEL_DATA column
```

An example on how to do this:

```plaintext
ft(vis='n75.ms',  # Start with the visibility dataset n75.ms
    field='1328',  # Select field name '1328+307' (minimum match)
    model='1328.model.image')  # Name of the model image you have already
```

This example will add the source model '1328.model.imag' to all entries that match the field name '1328'. If the parameter `usescratch` is set to 'True', `ft` will Fourier transform the source model and fill the `MODEL_DATA` column with the data. This, however, is only needed in special applications and `usescratch=F` is the default.

Alternatively, one can add a clean component list to be used as a model to the MS. The following procedure is an example:
# for a point source with no spectral index
cl.addcomponent(flux=0.39, fluxunit='Jy', shape='point', dir='J2000 19h33m09s 15d01m20s')

# for a Gaussian with a spectral index
cl.addcomponent(flux=1.25, fluxunit='mJy', polarization='Stokes',
dir='J2000 19h30m00s 15d00m00s', shape='gaussian', majoraxis='10arcsec',
minoraxis='6arcsec', positionangle='0deg', freq='1.25GHz',
spectrumtype='spectral index', index=-0.8)
### you can add more components if you wish by calling addcomponent repeatedly with different params

## save it to disk
cl.rename('my_component.cl')
cl.close()

## write the model into the measurement set ('myms')
ft(vis='myms', complist='my_component.cl')

5.8 Image-plane deconvolution (deconvolve)

If you have only an image (obtained from some telescope) and an image of its point spread function, then
you can attempt a simple image-plane deconvolution. Note that for interferometer data, full uv-plane
deconvolution using clean or similar algorithm is superior!

The default inputs for deconvolve are:

# deconvolve :: Deconvolving a point spread function from an image

imagename = '' # Name of image to deconvolve
model = '' # Name of output image to which deconvolved components are stored
psf = '' # Name of psf or gaussian parameters if psf is assumed gaussian
alg = 'clark' # Deconvolution algorithm to use
niter = 10 # number of iteration to use in deconvolution process
gain = 0.1 # CLEAN gain parameter
threshold = '0.0Jy' # level below which sources will not be deconvolved
mask = '' # Name of image that has mask to limit region of deconvolution

The algorithm (alg) options are: 'clark', 'hogbom', 'multiscale' or 'mem'. The 'multiscale'
and 'mem' options will open the usual set of sub-parameters for these methods.

5.9 Self-Calibration

Once you have a model image or set of model components reconstructed from your data using one
of the deconvolution techniques described above, you can use it to refine your calibration. This is
called self-calibration as it uses the data to determine its own calibration (rather than observations
of special calibration sources).
In principle, self-calibration is no different than the calibration process we described earlier (§ 4). In effect, you alternate between calibration and imaging cycles, refining the calibration and the model as you go. The trick is you have to be careful, as defects in early stages of the calibration can get into the model, and thus prevent the calibration from improving. In practice, it is best to not clean very deeply early on, so that the CLEAN model contains correct components only.

One important thing to keep in mind is that the self-calibration relies upon having the most recent source model inside the MS. This is indeed the case if you follow the imaging (using clean) directly by the self-calibration. If you have done something strange in between and have lost or overwritten source model (for example done some extra cleaning that you do not want to keep), then use the ft task (see § 5.7 above), which adds a source model image or clean component lists to an MS.

Likewise, during self-calibration (once you have a new calibration solution) the imaging part relies upon having the CORRECTED_DATA column contain the self-calibrated data. This is done with the applycal task (§ 4.6.1).

The clearcal command can be used during the self-calibration if you need to clear the CORRECTED_DATA column and revert to the original DATA. If you need to restore the CORRECTED_DATA to any previous stage in the self-calibration, use applycal again with the appropriate calibration tables.

**ALERT:** In later patches we will change the tasks so that users need not worry what is contained in the MS scratch columns and how to fill them. CASA will handle that underneath for you!

For now, we refer the user back to the calibration chapter for a reminder on how to run the calibration tasks.

### 5.10 Parallel Cleaning (pclean)

pclean is an experimental task to allow clean speedup by using more than a single computing node.

The inputs are somewhat different from clean but we will merge the two tasks in the future:

```plaintext
# pclean :: Invert and deconvolve images with parallel engines
vis = '' # Name of input visibility file
imagename = '' # Pre-name of output images
imsize = [256, 256] # Image size in pixels (nx,ny), symmetric for single value

cell = ['1.0arcsec', '1.0arcsec'] # The image cell size in arcseconds.
phasedcenter = '' # Image center: direction or field index
stokes = '' # Stokes params to image (eg I,IV,IQ,IQUV)
mask = '' # mask image
field = '' # Field Name or id
spw = '' # Spectral windows e.g. '0~3', '' is all
ftmachine = 'ft' # Fourier Transform Engine ('ft', 'sd', 'mosaic'
# or 'wproject')
alg = 'clark' # Deconvolution algorithm ('clark', 'hogbom',
# 'multiscale')
```
cyclefactor = 1.5 # Control number of major cycle, threshold of
# cycle=residualPeak*psfSidelobe*cyclefactor
niter = 500 # Maximum number of iterations
gain = 0.1 # Gain to use in deconvolution
threshold = '0.0Jy' # Flux level to stop cleaning, must include
# units: '1.0mJy'
weighting = 'natural' # Type of weighting
mode = 'continuum' # Clean mode ('continuum', 'cube')
interactive = False # Interactive clean
overwrite = False # Overwrite an existing model image
uv taper = False # Apply additional uv tapering of visibilities
timerange = '' # Range of time to select from data
uvrange = '' # Select data within uvrange
antenna = '' # Select data based on antenna/baseline
scan = '' # Scan number range
observation = '' # Observation ID range
pbcor = False # Correct for the primary beam post deconvolution
minpb = 0.2 # Fractional of peak of pb coverage where to stop
# the pb correction
clusterdef = '' # File that contains cluster definition

In pclean, the parameter alg controls whether the 'clark', 'hogbom', or 'multiscale' cleaning
algorithms are used. 'ft', 'sd', 'mosaic', and 'wproject' are specified via ftmachine, similar to
imagermode in clean. Important is the clusterdef parameter. It specifies a file with all computer
names, the number of CPU cores and temporary directories that can be used for pclean. It is
formatted like:

##############################################
hal9000, 10, /home/ptest
sal9000, 12, /home/ptest
nearstar, 6, /home/ptest
##############################################

It is advisable to leave one or to cores unused by pclean on computers that run the task. This will
allow other, sometimes vital, processes to continue.

IMPORTANT: The user has to have password-less ssh access3 to all the computers
used in a cluster definition file and the working directories have to be cross-mounted
by all the computers under the same name.

pclean attempts so slice the data in time bins for continuum imaging and in channel bins for
spectral imaging and sends sub-processes to the individual nodes. Since they all report back to the
main terminal while they are executed, the logging may look a bit messy. After the clean processes
on the nodes have concluded, the data will be put back together to create single image files.

3see e.g. http://www.linuxproblem.org/art_9.html
5.11 Examples of Imaging

The data reduction tutorials on casaguides.nrao.edu provide walkthroughs for high and low frequency, spectral line and polarization imaging techniques.
Chapter 6

Image Analysis

Once data has been calibrated (and imaged in the case of synthesis data), the resulting image or image cube must be displayed or analyzed in order to extract quantitative information, such as statistics or moment images. In addition, there need to be facilities for the coordinate conversion of images for direct comparison.

The image analysis tasks are:

- **imhead** — summarize and manipulate the “header” information in a CASA image (§6.2)
- **imsubimage** — Create a (sub)image from a region of the image (§6.3)
- **imcontsub** — perform continuum subtraction on a spectral-line image cube (§6.4)
- **imfit** — image plane Gaussian component fitting (§6.5)
- **immath** — perform mathematical operations on or between images (§6.6)
- **immoments** — compute the moments of an image cube (§6.7)
- **impv** — generate a position-velocity diagram along a slit (§6.8)
- **imstat** — calculate statistics on an image or part of an image (§6.9)

**Inside the Toolkit:**
Image analysis is handled in the ia tool. Many functions exist there, including region statistics and image math. See §6.25 below for more information.
• **imval** — extract the data and mask values from a pixel or region of an image (§ 6.10)

• **imtrans** — reorder the axes of an image or cube (§ 6.11)

• **imcollapse** — collapse image along one or more axes by aggregating pixel values along that axis (§ 6.12)

• **imregrid** — regrid an image onto the coordinate system of another image (§ 6.13)

• **imreframe** — change the frame in which the image reports its spectral values (§ 6.14)

• **imrebin** — rebin an image (§ 6.15)

• **specsmooth** — 1-dimensional smooth images in the spectral and angular directions (§ 6.16)

• **imsmooth** — 2-dimensional smooth images in the spectral and angular directions (§ 6.17)

• **specfit** — fit 1-dimensional Gaussians, polynomial, and/or Lorentzians models to an image or image region (§ 6.18)

• **rmfit** — Calculation of rotation measures (§ 6.19)

• **spxfit** — Calculation of Spectral Indices and higher order polynomials (§ 6.20)

• **makemask** — image mask handling (§ 6.21)

• **slsearch** — query a subset of the Splatalogue spectral line catalog (§ 6.22)

• **splattotable** — convert a file exported from Splatalogue to a CASA table (§ 6.23)

• **importfits** — import a FITS image into a CASA *image* format table (§ 6.24.2)

• **exportfits** — write out an image in FITS format (§ 6.24.1)

There are other tasks which are useful during image analysis. These include:

• **viewer** — there are useful region statistics and image cube slice and profile capabilities in the viewer (§ 7)

We also give some examples of using the CASA Toolkit to aid in image analysis (§ 6.25).
6.1 Common Image Analysis Task Parameters

We now describe some sets of parameters are are common to the image analysis. These should
behave the same way in any of the tasks described in this section that they are found in.

**ALERT:** As of CASA 4.0.0 we introduced changed clean such that for large fractional bandwidths,
if the PSF changes more than half a pixel, a different beam will be used for each image plane. These
cubes are supported in all image analysis tasks.

6.1.1 Input Image (imagename)

The input image typically is an image cube. Most analysis tasks and tools also accept complex
valued images.

6.1.2 Region Selection (box)

Direction (e.g. RA, Dec) areal selection in the image analysis tasks is controlled by the box
parameter or through the region parameter (§ 6.1.6). Note that one should either specify a
region (recommended) or any of box/chans/stokes. Specifying both at the same time is not unique
anymore and can lead to unwanted selections. In the future we may remove the box/chans/stokes
selection (for CASA 3.3 we keep both selection methods for backward compatibility).

The box parameter selects spatial rectangular areas:

```
box = '' # Select one or more box regions
#
# string containing blcx,blcy,trcx,trcy
#
# A box selection in the directional portion of an image.
# The directional portion of an image are the axes for right
# ascension and declination, for example. Boxes are specified
# by their bottom-left corner (blc) and top-right corner (trc)
# as follows: blcx, blcy, trcx, trcy;
# ONLY pixel values acceptable at this time.
# Default: none (all);
# Example: box='0,0,50,50'
```

To get help on box, see the in-line help

```
help(par.box)
```
6.1.3 Plane Selection (chans, stokes)

The channel, frequency, or velocity plane(s) of the image is chosen using the chans parameter:

```plaintext
chans = '' # Select the channel(spectral) range
```

# string containing channel range

# immath, imstat, and imcontsub - takes a string listing
# of channel numbers, velocity, and/or frequency
# numbers, much like the spw parameter
# Only channel numbers acceptable at this time.
# Default: none (all);
# Example: chans='3~20'
# chans="0,3,4,8"
# chans="3~20,50,51"

chans can also be set in the CASA region format to allow settings ins frequency and velocity, e.g.

```plaintext
chans=("range=[-50km/s,50km/s], restfreq=100GHz, frame=LSRK")
```

this example would even define a new velocity system independent of the one in the image itself. If the rest frequency and velocity frame within the image are being used, the latter two entries are not needed. The parentheses are needed when the call is in a single command.

A frequency selection looks as follows:

```plaintext
chans=("range=[100GHz,100.125GHz]")
```

The polarization plane(s) of the image is chosen with the stokes parameter:

```plaintext
stokes = '' # Stokes params to image (I,IV,IQU,IQUV)
```

# string containing Stokes selections

# Stokes parameters to image, may or may not be separated
# by commas but best if you use commas.
# Default: none (all); Example: stokes='IQUV';
# Example:stokes='I,Q'
# Options: 'I','Q','U','V',
# 'RR','RL','LR','LL',
# 'XX','YY',

To get help on these parameters, see the in-line help

```plaintext
help(par.chans)
help(par.stokes)
```

Sometimes, as in the immoments task, the channel/plane selection is generalized to work on more than one axis type. In this case, the planes parameter is used. This behaves like chans in syntax.
6.1.4 Lattice Expressions (expr)

Lattice expressions are strings that describe operations on a set of input images to form an output image. These strings use the Lattice Expression Language (LEL). LEL syntax is described in detail in AIPS++ Note 223

http://aips2.nrao.edu/docs/notes/223/223.html

**ALERT:** This document was written in the context of glish-based AIPS++ and is not yet updated to CASA syntax (see below).

The `expr` string contains the LEL expression:

```plaintext
expr = '' # Mathematical expression using images

# string containing LEL expression

# A mathematical expression, with image file names.
# image file names must be enclosed in double quotes ("")
# Default: none
# Example: expr='min("image2.im")+(2*max("image1.im"))'

# Available functions in the expr and mask parameters:
# pi(), e(), sin(), sinh(), asinh(), cos(), cosh(), tan(), tanh(),
# atan(), exp(), log(), log10(), pow(), sqrt(), complex(), conj()
# real(), imag(), abs(), arg(), phase(), amplitude(), min(), max()
# round(), isgn(), floor(), ceil(), rebin(), spectralindex(), pa(),
# iif(), indexin(), replace(), ...
```

For examples using LEL `expr`, see §6.6.1 below. Note that in `immath`, shortcut names have been given to the images provided by the user in `imagename` that can be used in the LEL expression, for the above example:

```plaintext
imagename=['image2.im','image1.im']
expr='min(IM0)+(2*max(IM1))'
```

**ALERT:** LEL expressions use 0-based indices. Also, the functions must be lowercase (in almost all cases we know about).

6.1.5 Masks (mask)

A mask can be used to define whether part of an image is used or not. There are different options for masks:

- an image cube with Boolean True/False values
- an image cube with zero and non-zero values
• an LEL string for a condition.

Using image cubes is useful to mask on a pixel by pixel basis, where False and zeros mark masked pixels. Both versions can be converted into each other `makemask` (§ 6.21). Some analysis tasks show an optional `stretch` parameter which is useful, e.g., to expand a single plane mask to an entire cube along the spectral axis.

An LEL string (see § 6.1.4 above) can be an on-the-fly (OTF) mask expression or refer to an image pixel mask.

```python
mask = ''  # Mask to be applied to the images

# string containing LEL expression

# Name of mask applied to each image in the calculation
# Default '' means no mask;
# Example: mask='ngc5921.clean.cleanbox.mask">0.5'
# mask='mask(ngc5921.clean.cleanbox.mask)'
```

Note that the mask file supplied in the `mask` parameter must have the same shape, same number of axes and same axes length, as the images supplied in the `expr` parameter, with one exception. The mask may be missing some of the axes — if this is the case then the mask will be expanded along these axes to become the same shape.

For examples using `mask`, see § 6.6.2 below.

### 6.1.6 Regions (region)

The `region` parameter points to a CASA region which can be directly specified or listed in a `ImageRegion` file. An `ImageRegion` file can be created with the CASA viewer’s region manager (§ 7.4.3). Or directly using the CASA region syntax (Chapter D). Typically `ImageRegion` files will have the suffix ‘.crtf’ for CASA Region Text Format.

**Alert:** When both the `region` parameter and any of box/chans/stokes are specified simultaneously, the task may perform unwanted selections. Only specify one of these (sets of) parameters. We recommend the use of CASA regions and may remove the box/chans/stokes selection in later releases.

For example:

```python
region='circle[[18h12m24s, ~23d11m00s], 2.3arcsec]'
```

or

```python
region='myimage.im.crtf'
```
for to specify a region file.

For the most part, the region parameter in tasks only accepts strings (e.g. file names, region shape descriptions) while the region parameter in ia tool methods only accepts python region dictionaries (e.g. produced using the rg tool).

### 6.2 Image Header Manipulation (imhead)

To summarize and change keywords and values in the “header” of your image, use the **imhead** task. Its inputs are:

```python
# imhead :: List, get and put image header parameters
imagename = '' # Name of the input image
mode = 'summary' # imhead options: add, del,
                  # get, history, list, put, summary
verbose = False # Give a full listing of
                # beams or just a short summary?
                # Only used when the image has multiple beams
                # and mode="summary".
```

The **mode** parameter controls the operation of **imhead**.

Setting **mode='summary'** will print out a summary of the image properties and the header to the logger.

Setting **mode='list'** prints out a list of the header keywords and values to the terminal.

The **mode='get'** allows the user to retrieve the current value for a specified keyword **hdkey**:

```python
mode = 'get' # imhead options: list, summary, get, put
            hdkey = '' # The FITS keyword
```

Note that to catch this value, you need to assign it to a Python variable. See § [1.4.3](#) for more on return values.

The **mode='put'** allows the user to replace the current value for a given keyword **hditem** with that specified in **hdvalue**. There are two sub-parameters that are opened by this option:

```python
mode = 'put' # imhead options: list, summary, get, put
            hdkey = '' # The FITS keyword
            hdvalue = '' # Value of hdkey
            hdtype = '' # Data type of the header keyword.
            hdcomment = '' # Comment associated with the header keyword
```

**WARNING:** Be careful when using **mode='put'**. This task does no checking on whether the values you specify (e.g. for the axes types) are valid, and you can render your image invalid. Make sure you know what you are doing when using this option!
6.2.1 Examples for imhead

Here is an example – we can print the summary to the logger:

CASA <51>: imhead('ngc5921.demo.cleanimg.image',mode='summary')

prints in the logger:

##### Begin Task: imhead #####
Image name : ngc5921.demo.cleanimg.image
Object name : N5921_2
Image type : PagedImage
Image quantity : Intensity
Pixel mask(s) : None
Region(s) : None
Image units : Jy/beam
Restoring Beam : 52.3782 arcsec, 45.7319 arcsec, -165.572 deg
Direction reference : J2000
Spectral reference : LSRK
Velocity type : RADIO
Rest frequency : 1.42041e+09 Hz
Pointing center : 15:22:00.000000 +05.04.00.000000
Telescope : VLA
Observer : TEST
Date observation : 1995/04/13/00:00:00
Telescope position: [-1.60119e+06m, -5.04198e+06m, 3.55488e+06m] (ITRF)

Axis Coord Type Name Proj Shape Tile Coord value at pixel Coord incr Units
-----------------------------------------------------------------------------------------------
0 0 Direction Right Ascension SIN 256 64 15:22:00.000 128.00 -1.500000e+01 arcsec
1 0 Direction Declination SIN 256 64 +05.04.00.000 128.00 1.500000e+01 arcsec
2 1 Stokes Stokes 1 1 I
3 2 Spectral Frequency 46 8 1.41279e+09 0.00 2.4414062e+04 Hz
Velocity 1607.99 0.00 -5.152860e+00 km/s

##### End Task: imhead

If the beam size per plane differs, the beam information will be displayed for the channel with the smallest beam, the one with the largest beam, and the channel closest to the median beam size. E.g.,

Restoring Beams
Pol Type Chan Freq Vel
I Max 0 9.680e+08 0 39.59 arcsec x 22.77 arcsec pa=-70.57 deg
I Min 511 1.990e+09 -316516 20.36 arcsec x 12.05 arcsec pa=-65.67 deg
I Median 255 1.478e+09 -157949 27.11 arcsec x 15.54 arcsec pa=-70.36 deg

If verbose=T the beam information for every plane will be provided.

If you choose mode='list', you get the summary in the logger and a listing of keywords and values to the terminal:
CASA <52>: imhead('ngc5921.demo.cleanimg.image',mode='list')
  Out[52]:
  {'beammajor': 52.378242492675781,
   'beamminor': 45.731891632080078,
   'beampa': -165.5721435546875,
   'bunit': 'Jy/beam',
   'cdelt1': '-7.27220521664e-05',
   'cdelt2': '7.27220521664e-05',
   'cdelt3': '1.0',
   'cdelt4': '24414.0625',
   'crpix1': 128.0,
   'crpix2': 128.0,
   'crpix3': 0.0,
   'crpix4': 0.0,
   'crval1': '4.02298392585',
   'crval2': '0.0884300154344',
   'crval3': 'I',
   'crval4': '1412787144.08',
   'ctype1': 'Right Ascension',
   'ctype2': 'Declination',
   'ctype3': 'Stokes',
   'ctype4': 'Frequency',
   'cunit1': 'rad',
   'cunit2': 'rad',
   'cunit3': '',
   'cunit4': 'Hz',
   'datamax': ' Not Known ',
   'datamin': -0.010392956435680389,
   'date-obs': '1995/04/13/00:00:00',
   'equinox': 'J2000',
   'imtype': 'Intensity',
   'masks': ' Not Known ',
   'maxpixpos': array([134, 134, 0, 38], dtype=int32),
   'maxpos': '15:21:53.976, +05.05.29.998, I, 1.41371e+09Hz',
   'minpixpos': array([117, 0, 0, 21], dtype=int32),
   'minpos': '15:22:11.035, +04.31.59.966, I, 1.4133e+09Hz',
   'object': 'N5921_2',
   'observer': 'TEST',
   'projection': 'SIN',
   'reffreftype': 'LSRK',
   'restfreq': [1420405752.0],
   'telescope': 'VLA'}

Note that this list is a return value and can be captured in a variable:

    mylist = imhead('ngc5921.demo.cleanimg.image',mode='list')

The values for these keywords can be queried using mode='get'. At this point you should capture the return value:

CASA <53>: mybmaj = imhead('ngc5921.demo.cleanimg.image',mode='get',hdkey='beammajor')

CASA <54>: mybmaj
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Out[54]: {'unit': 'arcsec', 'value': 52.378242492699997}

CASA <55>: myobserver = imhead('ngc5921.demo.cleaning.image', mode='get', hdkey='observer')

CASA <56>: print myobserver
{'value': 'TEST', 'unit': ''}

You can set the values for these keywords using mode='put'. For example:

CASA <57>: imhead('ngc5921.demo.cleaning.image', mode='put', hdkey='observer', hdvalue='CASA')
Out[57]: 'CASA'

CASA <58>: imhead('ngc5921.demo.cleaning.image', mode='get', hdkey='observer')
Out[58]: {'unit': '', 'value': 'CASA'}

6.3 Extracting sub-images (imsubimage)

The task `imsubimage` provides a way to extract a smaller data cube from a bigger one. The inputs are:

```python
# imsubimage :: Create a (sub)image from a region of the image
imagename = '' # Input image name. Default is unset.
outfile = '' # Output image name. Default is unset.
region = '' # Region of interest. Default is whole image.
mask = '' # Mask to use. See help par.mask. Default is none.
dropdeg = False # Drop degenerate axes
overwrite = False # Overwrite (unprompted) pre-existing output file?
verbose = True # Post additional informative messages to the logger
stretch = False # Stretch the mask if necessary and possible?
wantreturn = True # If True, return an image analysis tool associated with the subimage. If False,
# return nothing.
```

The `region` keyword defines the size of the smaller cube and is specified via the CASA region CRTF syntax. E.g.

```python
region='box [ [ 100pix , 130pix] , [120pix, 150pix ] ]'
```

will extract the portion of the image that is between pixel coordinates (100,130) and (12,150). `dropdeg=T` is useful to remove axes in the data cube that are degenerate, i.e. axes with a single plane only. A single Stokes I axis is a common example.
6.4 Continuum Subtraction on an Image Cube (imcontsub)

One method to separate line and continuum emission in an image cube is to specify a number of line-free channels in that cube, make a linear fit to the visibilities in those channels, and subtract the fit from the whole cube. Note that the task uvcontsub serves a similar purpose; see §4.7.6 for a synopsis of the pros and cons of either method.

The imcontsub task will subtract a polynomial baseline fit to the specified channels from an image cube.

The default inputs are:

```plaintext
# imcontsub :: Continuum subtraction on images
imagename = '' # Name of the input image
linefile = '' # Output line image file name
contfile = '' # Output continuum image file name
fitorder = 0 # Polynomial order for the continuum estimation
region = '' # Image region or name to process see viewer
box = '' # Select one or more box regions
chans = '' # Select the channel(spectral) range
stokes = '' # Stokes params to image (I,IV,IQU,IQUV)
```

Area selection using box and region is detailed in §6.1.2 and §6.1.6 respectively.

Image cube plane selection using chans and stokes are described in §6.1.3.

**ALERT:** imcontsub has issues when the image does not contain a spectral or stokes axis. Errors are generated when run on an image missing one or both of these axes. You will need to use the Toolkit (e.g. the ia.adddegaxes method) to add degenerate missing axes to the image.

6.4.1 Examples for imcontsub

For example, we first make a clean image without the uv-plane continuum subtraction:

```plaintext
# First, run clearcal to clear the uvcontsub results from the
# corrected column
clearcal('ngc5921.demo.src.split.ms')

# Now clean, keeping all the channels except first and last
default('clean')
vis = 'ngc5921.demo.src.split.ms'
imagename = 'ngc5921.demo.nouvcontsub'
mode = 'channel'
nchan = 61
start = 1
width = 1
imsize = [256,256]
psfmode = 'clark'
imagermode = ''
```
cell = [15.,15.]
niter = 6000
threshold='8.0mJy'
weighting = 'briggs'
robust = 0.5
interactive=False
clean()

# It will have made the image:
# ----------------------------------
# ngc5921.demo.nouvcontsub.image

# You can view this image
viewer('ngc5921.demo.nouvcontsub.image')

You can clearly see continuum sources in the image which were removed previously in the script by the use of uvcontsub. Let’s see if imcontsub can work as well.

Using the viewer, it looks like channels 0 through 4 and 50 through 60 are line-free. Then:

default('imcontsub')
imagename = 'ngc5921.demo.nouvcontsub.image'
linefile = 'ngc5921.demo.nouvcontsub.lineimage'
contfile = 'ngc5921.demo.nouvcontsub.contimage'
fitorder = 1
chans = '0~4,50~60'
stokes = 'I'
imcontsub()

This did not do too badly!

### 6.5 Image-plane Component Fitting (imfit)

The inputs are:

```python
# imfit :: Fit one or more elliptical Gaussian components on an image region(s)
imagename = '' # Name of the input image
box = '' # Specify one or more box regions for the fit.
region = '' # Region. See help par.region for specs.
chans = '' # Spectral channels on which to perform fit. See "help par.chans" for examples.
stokes = '' # Stokes parameter to fit. If blank, first stokes plane is used.
mask = '' # Mask to use. See help par.mask. Default is none.
includepix = [] # Range of pixel values to include for fitting.
excludepix = [] # Range of pixel values to exclude for fitting.
residual = '' # Name of output residual image.
```
model = '' # Name of output model image.
estimates = '' # Name of file containing initial estimates of component
# parameters.
logfile = '' # Name of file to write fit results.
newestimates = '' # File to write fit results which can be used as initial
# estimates for next run.
complist = '' # Name of output component list table.
dooff = False # Also fit a zero level offset? Default is False
rms = -1 # RMS to use in calculation of uncertainties. Numeric or
# valid quantity (record or string). If numeric, it is
# given units of the input image. If quantity, units must
# conform to image units. If not positive, the rms of the
# residual image, in the region of the fit, is used.
oisefwhm = '' # Noise correlation beam FWHM. If numeric value,
# interpreted as pixel widths. If quantity (dictionary,
# string), it must have angular units.

This task will return (as a Python dictionary) the results of the fit, but the results can also be
written into a component list table or a logfile.

Note that to fit more than a single component, you must provide starting estimates for each
component via the estimates file. See `help imfit` for more details on this. An noise estimate
will be calculated automatically or can be provided through the rms and noisefwhm keywords.

### 6.5.1 Examples for imfit

The following are some examples using the B1608+656 Tutorial

as an example.

# First fit only a single component at a time
# This is OK since the components are well-separated and not blended
# Box around component A
xfit_A_res = imfit('b1608.demo.clean2.image',box='121,121,136,136',
newestimates='b1608.demo.clean2.newestimate')

# Now extract the fit part of the return value
xfit_A = xfit_A_res['results']['component0']
# xfit_A
# Out[7]:
#{'flux': {'error': array([ 6.73398035e-05, 0.00000000e+00, 0.00000000e+00,
# 0.00000000e+00]),
# 'polarisation': 'Stokes',
# 'unit': 'Jy',
# 'value': array([ 0.01753742, 0. , 0. , 0. ])},
# 'label': ''},
# 'shape': {'direction': {'error': {'latitude': {'unit': 'arcsec', 'value': 0.00041154866279462775}, 'longitude': {'unit': 'arcsec', 'value': 0.00046695916589535109}}, 'm0': {'unit': 'rad', 'value': -2.0541102061078207}, 'm1': {'unit': 'rad', 'value': 1.1439131060384089}, 'refer': 'J2000', 'type': 'direction'}, 'majoraxis': {'unit': 'arcsec', 'value': 0.29100166137741568}, 'majoraxiserror': {'unit': 'arcsec', 'value': 0.0011186420613222663}, 'minoraxis': {'unit': 'arcsec', 'value': 0.24738110059830495}, 'minoraxiserror': {'unit': 'arcsec', 'value': 0.0013431999725066338}, 'positionangle': {'unit': 'deg', 'value': 19.369249322401796}, 'positionangleerror': {'unit': 'rad', 'value': 0.016663189295782171}, 'type': 'Gaussian'}, 'spectrum': {'frequency': {'m0': {'unit': 'GHz', 'value': 1.0}, 'refer': 'LSRK', 'type': 'frequency'}, 'type': 'Constant'}}

# Now the other components
xfit_B_res = imfit('b1608.demo.clean2.image', box='108,114,120,126', newestimates='b1608.demo.clean2.newestimate', append=True)
xfit_B = xfit_B_res['results']['component0']

xfit_C_res = imfit('b1608.demo.clean2.image', box='108,84,120,96')
xfit_C = xfit_C_res['results']['component0']

xfit_D_res = imfit('b1608.demo.clean2.image', box='144,98,157,110')
xfit_D = xfit_D_res['results']['component0']

print """ print 'Imfit Results:' print """"
print "A Flux = %6.4f Bmaj = %6.4f" % (xfit_A['flux']['value'][0], xfit_A['shape']['majoraxis']['value'])
print "B Flux = %6.4f Bmaj = %6.4f" % (xfit_B['flux']['value'][0], xfit_B['shape']['majoraxis']['value'])
print "C Flux = %6.4f Bmaj = %6.4f" % (xfit_C['flux']['value'][0], xfit_C['shape']['majoraxis']['value'])
print "D Flux = %6.4f Bmaj = %6.4f" % (xfit_D['flux']['value'][0], xfit_D['shape']['majoraxis']['value'])
print ""

Now try fitting four components together. For this we will have to provide an estimate file. We will use the clean beam for the estimate of the component sizes:

estfile=open('b1608.demo.clean2.estimate','w')
print >>estfile,'# peak, x, y, bmaj, bmin, bpa'
print >>estfile,'0.017, 128, 129, 0.293arcsec, 0.238arcsec, 21.7deg'
print >>estfile,'0.008, 113, 120, 0.293arcsec, 0.238arcsec, 21.7deg'
print >>estfile,'0.008, 113, 90, 0.293arcsec, 0.238arcsec, 21.7deg'
print >>estfile,'0.002, 151, 104, 0.293arcsec, 0.238arcsec, 21.7deg'
estfile.close()

Then, this can be used in imfit:

```python
xfit_all_res = imfit('b1608.demo.clean2.image',
                     estimates='b1608.demo.clean2.estimate',
                     logfile='b1608.demo.clean2.imfitall.log',
                     newestimates='b1608.demo.clean2.newestimate',
                     box='121,121,136,136,108,114,120,126,108,84,120,96,144,98,157,110')
```

# Now extract the fit part of the return values
xfit_allA = xfit_all_res['results']['component0']
xfit_allB = xfit_all_res['results']['component1']
xfit_allC = xfit_all_res['results']['component2']
xfit_allD = xfit_all_res['results']['component3']

These results are almost identical to those from the individual fits. You can see a nicer printout of the fit results in the logfile.

## 6.6 Mathematical Operations on an Image (immath)

The inputs are:

```python
# immath :: Perform math operations on images
 imagename = '' # a list of input images
 mode = 'evalexpr' # mode for math operation (evalexpr, spix, pola, poli)
 expr = '' # Mathematical expression using images
 varnames = '' # a list of variable names to use with the image files

 outfile = 'immath_results.im' # File where the output is saved
 mask = '' # Mask to be applied to the images
 region = '' # File path which contains an Image Region
 box = '' # Select one or more box regions in the input images
 chans = '' # Select the channel(spectral) range
 stokes = 'I' # Stokes params to image (I,IV,IQU,IQUV)
```

In all cases, `outfile` must be supplied with the name of the new output file to create.

The `mode` parameter selects what `immath` is to do.

The default `mode='evalexpr'` lets the user specify a mathematical operation to carry out on one or more input images. The sub-parameter `expr` contains the Lattice Expression Language (LEL) string describing the image operations based on the images in the `imagename` parameter. See §6.1.4 for more on LEL strings and the `expr` parameter.

Mask specification is done using the `mask` parameter. This can optionally contain an on-the-fly mask expression (in LEL) or point to an image with a pixel mask. See §6.1.5 for more on the use
of the `mask` parameter. See also §6.1.4 for more on LEL strings. Sometimes, one would like to use a flat image (e.g. a moment image) mask to be applied to an entire cube. The `stretch=True` subparameter in `mask` allows one to expand the mask to all planes of the cube.

Region selection is carried out through the `region` and `box` parameters. See §6.1.2 and §6.1.6 for more on area selection.

Image plane selection is controlled by `chans` and `stokes`. See §6.1.3 for details on plane selection.

For `mode='evalexpr'`, the standard usage for specifying images to be used in the LEL expression is to provide them as a list in the `imagename` parameter, and then access there in the LEL expression by the names `IM0`, `IM1`, .... For example,

```python
immath(imagename=['image1.im','image2.im'],expr='IM0-IM1',outfile='ImageDiff.im')
```

would subtract the second image given from the first.

For the special modes `spix`, `pola`, `poli`, the required images for the given operation are to be provided in `imagename` (sometimes in a particular order). **V3.0 ALERT:** For `mode='pola'` you MUST call as a function as in the example below (§6.6.1.2), giving the parameters as arguments, or `immath` will fail.

Detailed examples are given below.

### 6.6.1 Examples for `immath`

In the following, we show a few examples of `immath`. Note that the image names in the `expr` are assumed to refer to existing image files in the current working directory.

#### 6.6.1.1 Simple math

Select a single plane (channel 22) of the 3-D cube and subtract it from the original image:

```python
immath(imagename='ngc5921.demo.cleanimg.image',
       expr='IM0',chans='22',
       outfile='ngc5921.demo.chan22.image')
```

Double all values in our image:

```python
immath(imagename=['ngc5921.demo.chan22.image'],
       expr='IM0*2.0',
       outfile='ngc5921.demo.chan22double.image')
```

Square all values in our image:

```python
immath(imagename=['ngc5921.demo.chan22.image'],
       expr='IM0^2',
       outfile='ngc5921.demo.chan22squared.image')
```
Note that the units in the output image are still claimed to be “Jy/beam”, i.e. `immath` will not correctly scale the units in the image for non-linear cases like this. Beware.

You can do other mathematical operations on an image (e.g. trigonometric functions) as well as use scalars results from an image (e.g. max, min, median, mean, variance). You also have access to constants such as `e()` and `pi()` (which are doubles internally, while most images are floats). For example: Take the sine of an image:

```python
immath(imagename=['ngc5921.demo.chan22.image','ngc5921.demo.chan22squared.image'],
       expr='sin(float(pi())*IM0/sqrt(max(IM1)))',
       outfile='ngc5921.demo.chan22sine.image')
```

Note again that the units are again kept as they were.

Select a single plane (channel 22) of the 3-D cube and subtract it from the original image:

```python
immath(imagename='ngc5921.demo.cleanimg.image',
       expr='IM0',chans='22',
       outfile='ngc5921.demo.chan22.image')

immath(imagename=['ngc5921.demo.cleanimg.image','ngc5921.demo.chan22.image'],
       expr='IM0-IM1',
       outfile='ngc5921.demo.sub22.image')
```

Note that in this example the 2-D plane gets extended in the third dimension and the 2-D values are applied to each plane in the 3-D cube.

Select and save the inner 1/4 of an image for channels 40,42,44 as well as channels 10 and below:

```python
default('immath')
imagename=['ngc5921.demo.cleaning.image']
expr='IM0'
region='box[[64pix,64pix],[192pix,192pix]]'
chans='<10;40,42,44'
outfile='ngc5921.demo.inner.image'
immath()
```

**ALERT:** Note that if chan selects more than one channel then the output image has a number of channels given by the span from the lowest and highest channel selected in chan. In the example above, it will have 45 channels. The ones not selected will be masked in the output cube. If we had set

```python
chans = '40,42,44'
```

then there would be 5 output channels corresponding to channels 40,41,42,43,44 of the MS with 41,43 masked. Also, the `chans='<10'` selects channels 0–9.

Note that the `chans` syntax allows the operators '<', '<=', '>', '>='. For example,
CHAPTER 6. IMAGE ANALYSIS

chans = '&lt;17,&gt;79'
chans = '&lt;=16,&gt;=80'

do the same thing.

Divide an image by another, with a threshold on one of the images:

\[
\text{immath(imagename=['ngc5921.demo.cleanimg.image','ngc5921.demo.chan22.image'],}
\text{expr='IM0/IM1[IM1>0.008]',}
\text{outfile='ngc5921.demo.div22.image')}
\]

6.6.1.2 Polarization manipulation

The following are some examples using the 3C129 Tutorial

[http://casa.nrao.edu/Doc/Scripts/3c129_tutorial.py](http://casa.nrao.edu/Doc/Scripts/3c129_tutorial.py)
as an example.

It is helpful to extract the Stokes planes from the cube into individual images:

\[
\text{default('immath')}
\text{imagename = '3C129BC.clean.image'}
\text{outfile='3C129BC.I'; expr='IM0'; stokes='I'; immath();}
\text{outfile='3C129BC.Q'; expr='IM0'; stokes='Q'; immath();}
\text{outfile='3C129BC.U'; expr='IM0'; stokes='U'; immath();}
\text{outfile='3C129BC.V'; expr='IM0'; stokes='V'; immath();}
\]

Extract linearly polarized intensity and polarization position angle images:

\[
\text{immath(stokes='', outfile='3C129BC.P', mode='poli',}
\text{imagename=['3C129BC.Q','3C129BC.U'], sigma='0.0mJy/beam');}
\text{immath(stokes='', outfile='3C129BC.X', mode='pola',}
\text{imagename=['3C129BC.Q','3C129BC.U'], sigma='0.0mJy/beam');}
\]

**V3.0 ALERT:** For mode='pola' you MUST call as a function as in this example (giving the parameters as arguments) or immath will fail.

Create a fractional linear polarization image:

\[
\text{default('immath')}
\text{imagename = ['3C129BC.I','3C129BC.Q','3C129BC.U']}
\text{outfile='3C129BC.fractional_linpol'
\text{expr='sqrt((IM1^2 + IM2^2)/IM0^2)'
\text{stokes=''
\text{immath()}
\]

Create a polarized intensity image:
 Toolkit Tricks: The following uses the toolkit (§ 6.25). You can make a complex linear polarization 
\((Q + iU)\) image using the imagepol tool:

```python
# See CASA User Reference Manual:
# http://casa.nrao.edu/docs/casaref/imagepol-Tool.html
#
# Make an imagepol tool and open the clean image
potool = casac.homefinder.find_home_by_name('imagepolHome')
po = potool.create()
po.open('3C129BC.clean.image')
# Use complexlinpol to make a Q+iU image
po.complexlinpol('3C129BC.cmplxlinpol')
po.close()
```

You can now display this in the viewer, in particular overlay this over the intensity raster with the 
intensity contours. When you load the image, use the LEL:

\['3C129BC.cmplxlinpol'\[3C129BC.P'>0.0001\]

which is entered into the LEL box at the bottom of the Load Data menu (§ 7.3.1).

### 6.6.2 Using masks in immath

The mask parameter is used inside immath to apply a mask to all the images used in expr before 
calculations are done (if you are curious, it uses the ia.subimage tool method to make virtual 
images that are then input in the LEL to the ia.imagecalc method).

For example, let’s assume that we have made a single channel image using clean

```python
default('clean')
vis = 'ngc5921.demo.src.split.ms.contsub'
imagename = 'ngc5921.demo.chan22.cleanimg'
mode = 'channel'
nchan = 1
start = 22
step = 1
field = ''
spw = ''
imsize = [256,256]
```
cell = [15.,15.]
psfalg = 'clark'
gain = 0.1
niter = 6000
threshold='8.0mJy'
weighting = 'briggs'
robust = 0.5

clean()

There is now a file 'ngc5921.demo.chan22.cleanimg.mask' that is an image with values 1.0 inside the cleanbox region and 0.0 outside.

We can use this to mask the clean image:

default('immath')
imagename = 'ngc5921.demo.chan22.cleanimg.image'
expr='IM0'
mask='"ngc5921.demo.chan22.cleanimg.mask">0.5'
outfile='ngc5921.demo.chan22.cleanimg.imasked'
immath()

**Toolbox Tricks:** Note that there are also *pixel masks* that can be contained in each image. These are Boolean masks, and are implicitly used in the calculation for each image in `expr`. If you want to use the mask in a different image not in `expr`, try it in `mask`:

```
# First make a pixel mask inside ngc5921.demo.chan22.cleanimg.mask
ia.open('ngc5921.demo.chan22.cleanimg.mask')
ia.calcmask('"ngc5921.demo.chan22.cleanimg.mask">0.5')
ia.summary()
ia.close()
# There is now a 'mask0' mask in this image as reported by the summary

# Now apply this pixel mask in immath
default('immath')
imagename='ngc5921.demo.chan22.cleanimg.image'
expr='IM0'
mask='mask(ngc5921.demo.chan22.cleanimg.mask)'
outfile='ngc5921.demo.chan22.cleanimg.imasked1'
immath()
```

Note that nominally the axes of the mask must be congruent to the axes of the images in `expr`. However, one exception is that the image in `mask` can have fewer axes (but not axes that exist but are of the wrong lengths). In this case `immath` will extend the missing axes to cover the range in the images in `expr`. Thus, you can apply a mask made from a single channel to a whole cube.
# drop degenerate stokes and freq axes from mask image
ia.open('ngc5921.demo_chan22_cleaning_mask')
im2 = ia.subimage(outfile='ngc5921.demo_chan22_cleaning_mymask', dropdeg=True)
im2.summary()
im2.close()
ia.close()
# mymask has only RA and Dec axes

# Now apply this mask to the whole cube
default('immath')
imagename='ngc5921.demo_cleaning_image'
expr='IM0'
mask='"ngc5921.demo_chan22_cleaning_mymask">0.5'
outfile='ngc5921.demo_cleaning_imasked'
immath()

For more on masks as used in LEL, see

http://aips2.nrao.edu/docs/notes/223/223.html

or in §6.1.5 above.

6.7 Computing the Moments of an Image Cube (immoments)

For spectral line datasets, the output of the imaging process is an image cube, with a frequency or velocity channel axis in addition to the two sky coordinate axes. This can be most easily thought of as a series of image planes stacked along the spectral dimension.

A useful product to compute is to collapse the cube into a moment image by taking a linear combination of the individual planes:

\[ M_m(x_i, y_i) = \sum_k w_m(x_i, y_i, v_k) I(x_i, y_i, v_k) \]  

(6.1)

for pixel \( i \) and channel \( k \) in the cube \( I \). There are a number of choices to form the \( m \) moment, usually approximating some polynomial expansion of the intensity distribution over velocity mean or sum, gradient, dispersion, skew, kurtosis, etc.). There are other possibilities (other than a weighted sum) for calculating the image, such as median filtering, finding minima or maxima along the spectral axis, or absolute mean deviations. And the axis along which to do these calculation need not be the spectral axis (i.e. do moments along Dec for a RA-Velocity image). We will treat all of these as generalized instances of a “moment” map.

The immoments task will compute basic moment images from a cube. The default inputs are:

# immoments :: Compute moments of an image cube:
imagename = ''   # Input image name
moments = [0]  # List of moments you would like to compute
axis = 'spectral'  # The moment axis: ra, dec, lat, long, spectral, or stokes
region = ''  # Image Region. Use viewer
box = ''  # Select one or more box regions
chans = ''  # Select the channel(spectral) range
stokes = ''  # Stokes params to image (I,IV,IQU,IQUV)
mask = ''  # mask used for selecting the area of the
# image to calculate the moments on
includepix = -1  # Range of pixel values to include
excludepix = -1  # Range of pixel values to exclude
outfile = ''  # Output image file name (or root for multiple moments)

This task will operate on the input file given by imagename and produce a new image or set of images based on the name given in outfile.

The moments parameter chooses which moments are calculated. The choices for the operation mode are:

- moments=-1 - mean value of the spectrum
- moments=0 - integrated value of the spectrum
- moments=1 - intensity weighted coordinate; traditionally used to get 'velocity fields'
- moments=2 - intensity weighted dispersion of the coordinate; traditionally used to get 'velocity dispersion'
- moments=3 - median of I
- moments=4 - median coordinate
- moments=5 - standard deviation about the mean of the spectrum
- moments=6 - root mean square of the spectrum
- moments=7 - absolute mean deviation of the spectrum
- moments=8 - maximum value of the spectrum
- moments=9 - coordinate of the maximum value of the spectrum
- moments=10 - minimum value of the spectrum
- moments=11 - coordinate of the minimum value of the spectrum

The meaning of these is described in the CASA Toolkit Manual, that describes the associated image.moments tool:

http://casa.nrao.edu/docs/CasaRef/image.moments.html

The axis parameter sets the axis along which the moment is “collapsed” or calculated. Choices are: ‘ra’, ‘dec’, ‘lat’, ‘long’, ‘spectral’, or ‘stokes’. A standard moment-0 or moment-1 image of a spectral cube would use the default choice ‘spectral’. One could make a position-velocity map by setting ‘ra’ or ‘dec’.

The includepix and excludepix parameters are used to set ranges for the inclusion and exclusion of pixels based on values. For example, includepix=[0.05,100.0] will include pixels with values from 50 mJy to 1000 Jy, and excludepix=[100.0,1000.0] will exclude pixels with values from 100 to 1000 Jy.
If a single moment is chosen, the `outfile` specifies the exact name of the output image. If multiple `moments` are chosen, then `outfile` will be used as the root of the output filenames, which will get different suffixes for each moment.

For image cubes that contain different beam sizes for each plane, `immoments` will smooth all planes to the largest beam size first, then collapse to the desired moment.

### 6.7.1 Hints for using (`immoments`)

In order to make an unbiased moment-0 image, do not put in any thresholding using `includepix` or `excludepix`. This is so that the (presumably) zero-mean noise fluctuations in off-line parts of the image cube will cancel out. If you image has large biases, like a pronounced clean bowl due to missing large-scale flux, then your moment-0 image will be biased also. It will be difficult to alleviate this with a threshold, but you can try.

To make a usable moment-1 (or higher) image, on the other hand, it is critical to set a reasonable threshold to exclude noise from being added to the moment maps. Something like a few times the rms noise level in the usable planes seems to work (put into `includepix` or `excludepix` as needed. Also use `chans` to ignore channels with bad data.

### 6.7.2 Examples using (`immoments`)

Below is an example for `immoments`:

```python
import immoments

default('immoments')
imagename = 'ngc5921.demo.cleanimg'
# Do first and second spectral moments
axis = 'spectral'
chans = ''
moments = [0, 1]
# Need to mask out noisy pixels, currently done
# using hard global limits
excludepix = [-100, 0.009]
outfile = 'ngc5921.demo.moments'

immoments()

# It will have made the images:
# --------------------------------------
# ngc5921.demo.moments.integrated
# ngc5921.demo.moments.weighted_coord
```

Other examples of NGC2403 (a moment zero image of a VLA line dataset) and NGC4826 (a moment one image of a BIMA CO line dataset) are shown in Figure 6.1.

**Alert:** We are working on improving the thresholding of planes beyond the global cutoffs in `includepix` and `excludepix`. 
6.8 Generating Position-Velocity Diagrams (impv)

CASA can generate position-velocity (pV) diagrams via the task impv or directly in the viewer (see §7.4.9). The viewer application calls the task:

```bash
# impv :: Construct a position-velocity image by choosing two points in the direction plane.
imagename = '' # Name of the input image
outfile = '' # Output image name. If empty, no image is written.
mode = 'coords' # If "coords", use start and end values. If "length", use
# center, length, and pa values.
width = 1 # Width of slice for averaging pixels perpendicular to the
# slice. Must be an odd positive integer or valid
# quantity. See help for details.
unit = 'arcsec' # Unit for the offset axis in the resulting image. Must be
# a unit of angular measure.
chans = '' # Channels to use. See "help par.chans" for examples.
# Channels must be contiguous. Default is to use all
# channels.
region = '' # Region selection. Default is entire image. No selection
# is permitted in the direction plane. See help
# par.region.

stokes = 'I' # Stokes planes to use. Planes must be contiguous. Default
# is to use all stokes.
mask = [] # Mask to use. See help par.mask. Default is none.
stretch = False # Stretch the mask if necessary and possible? See help
# par.stretch. Default False
```
PV diagrams are generated by placing a “slicing” a datacube through the RA/DEC planes. The “slit” can be defined either by start/end coordinates or by a length, center coordinate, and position angle. Averaged over the width of the 'slit' the image cube values are then stored in a new image with position and velocity as the two axes. The slit position is specified by a start and end pixel in the RA/DEC plane of the data cube. An angular unit can be set to define what is stored in the resulting pV image.

6.9 Computing image statistics (imstat)

The `imstat` task will calculate statistics on a region of an image, and return the results as a return value in a Python dictionary.

The inputs are:

```python
# imstat :: Displays statistical information from an image or image region
imagenname = '' # Name of the input image
axes = -1 # List of axes to evaluate statistics over. Default is all axes.
region = '' # Image Region or name. Use Viewer
box = '' # Select one or more box regions
chans = '' # Select the channel(spectral) range. See "help par.chans" for examples.
stokes = '' # Stokes params to image (I,IV,IQU,IQUV). Default "" => include all
listit = True # Print stats and bounding box to logger?
verbose = True # Print additional messages to logger?
mask = '' # Mask to use. See help par.mask. Default is none.
logfile = '' # Name of file to write fit results.
```

Area selection using `region` and `mask` is detailed in §6.1.6 and (§6.1.5) respectively.

Plane selection is controlled by `chans` and `stokes`. See §6.1.3 for details on plane selection.

The parameter `axes` will select the dimensions that the statistics is calculated over. Typical data cubes have axes like: RA axis 0, DEC axis 1, Velocity axis 2. So, e.g. `axes=[0,1]` would be the most common setting to calculate statistics per spectral channel.

A typical output of `imstat` on a cube with `axes=[0,1]` looks like:

```
No region specified. Using full positional plane.
Using all spectral channels.
Using polarizations ALL
Determining stats for image IRC10216_HC3N.cube_r0.5.image
Set region from supplied region record
Regions ---
-- bottom-left corner (pixel) [blc]: [0, 0, 0, 0]
-- top-right corner (pixel) [trc]: [299, 299, 0, 63]
-- bottom-left corner (world) [blcf]: 09:48:01.492, +13.15.40.658, I, 3.63994e+10Hz
-- top-right corner (world) [trcf]: 09:47:53.299, +13.17.40.258, I, 3.63915e+10Hz
```
No region specified. Using full positional plane. 
Using all spectral channels. 
Using polarizations ALL

Selected bounding box :
[0, 0, 0, 0] to [299, 299, 0, 63] (09:48:01.492, +13.15.40.658, I, 3.63994e+10Hz to 09:47:53.299, +13.17.40.258, I, 3.63915e+10Hz)

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<th>Npts</th>
<th>Sum</th>
<th>Mean</th>
<th>RMS</th>
<th>Std dev</th>
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<th>Maximum</th>
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<td>8</td>
<td>9.000000e+04</td>
<td>-3.706732e-01</td>
<td>-3.118591e-06</td>
<td>1.607191e-03</td>
<td>1.607194e-03</td>
<td>-8.871284e-03</td>
<td>6.591001e-03</td>
</tr>
</tbody>
</table>

where the header information provides the specifications of the data that were selected followed by the table with the frequency values of the lanes, the plane numbers, Npts the number of pixels per plane, and the Sum, Median, RMS, Standard deviations, Minimum, and Maximum of the pixel values for each plane. Similar output is provided when the data is averaged over different axes. The logger output can also be written into or appended to a log file for further processing elsewhere (logfile parameter).

### 6.9.1 Using the task return value

The contents of the return value of `imstat` are in a Python dictionary of key-value sets. For example,

```python
xstat = imstat()
```

will assign this to the Python variable `xstat`.

The keys for `xstat` are then:

```plaintext
KEYS
blc  # absolute PIXEL coordinate of the bottom left corner of
     # the bounding box surrounding the selected region
blcf # Same as blc, but uses WORLD coordinates instead of pixels
trc  # the absolute PIXEL coordinate of the top right corner
     # of the bounding box surrounding the selected region
trcf # Same as trc, but uses WORLD coordinates instead of pixels
flux # the integrated flux density if the beam is defined and
     # the brightness units are $Jy/beam$
Npts # the number of unmasked points used
max  # the maximum pixel value
min  # minimum pixel value
maxpos # absolute PIXEL coordinate of maximum pixel value
maxposf # Same as maxpos, but uses WORLD coordinates instead of pixels
minpos # absolute pixel coordinate of minimum pixel value
minposf # Same as minpos, but uses WORLD coordinates instead of pixels
sum  # the sum of the pixel values: $\sum I_i$
sumsq # the sum of the squares of the pixel values: $\sum I_i^2$
mean # the mean of pixel values:
```

```
\[
\text{ar(I)} = \sum I_i / n$
\]

\[
\sigma = \left(\sum I_i - \text{ar(I)}\right)^2 / (n-1)
\]

\[
\text{rms} = \sqrt{\sum I_i^2 / n}
\]

\[
\text{median} \quad \text{(if robust=T)}
\]

\[
\text{medabsdevmed} \quad \text{(if robust=T)}
\]

\[
\text{quartile} \quad \text{(if robust=T)}
\]

For example, an `imstat` call might be

```plaintext
default('imstat')

imagename = 'ngc5921.demo.cleanimg.image'  # The NGC5921 image cube
box = '108,108,148,148'  # 20 pixels around the center
chans = '21'  # channel 21

xstat = imstat()
```

In the terminal window, `imstat` reports:

```
Statistics on ngc5921.usecase.clean.image

Region ---
  -- bottom-left corner (pixel) [blc]: [108, 108, 0, 21]
  -- top-right corner (pixel) [trc]: [148, 148, 0, 21]
  -- bottom-left corner (world) [blcf]: 15:22:20.076, +04.58.59.981, I, 1.41332e+09Hz
  -- top-right corner (world) [trcf]: 15:21:39.919, +05.08.59.981, I, 1.41332e+09Hz

Values ---
  -- flux [flux]: 0.111799236126
  -- number of points [npts]: 1681.0
  -- maximum value [max]: 0.029451508075
  -- minimum value [min]: -0.00612453464419
  -- position of max value (pixel) [maxpos]: [124, 131, 0, 21]
  -- position of min value (pixel) [minpos]: [142, 110, 0, 21]
  -- position of max value (world) [maxposf]: 15:22:04.016, +05.04.44.999, I, 1.41332e+09Hz
  -- position of min value (world) [minposf]: 15:21:45.947, +04.59.29.990, I, 1.41332e+09Hz
  -- Sum of pixel values [sum]: 1.32267159822
  -- Sum of squared pixel values [sumsq]: 0.0284534543692

Statistics ---
  -- Mean of the pixel values [mean]: 0.000786836167885
  -- Standard deviation of the Mean [sigma]: 0.00403944306904
  -- Root mean square [rms]: 0.00411418313161
  -- Median of the pixel values [median]: 0.000137259965413
  -- Median of the deviations [medabsdevmed]: 0.00152346317191
```
The return value in xstat is

CASA <152>: xstat
Out[152]:
{'blc': array([108, 108, 0, 21]),
 'blcf': '15:22:04.016, +05.04.44.999, I, 1.41332e+09Hz',
 'flux': array([ 0.11179924]),
 'max': array([ 0.02945151]),
 'maxpos': array([124, 131, 0, 21]),
 'maxposf': '15:22:04.016, +05.04.44.999, I, 1.41332e+09Hz',
 'mean': array([ 0.00078684]),
 'medabsdevmed': array([ 0.00152346]),
 'median': array([ 0.00013726]),
 'min': array([-0.00612453]),
 'minpos': array([142, 110, 0, 21]),
 'minposf': '15:21:45.947, +04.59.29.990, I, 1.41332e+09Hz',
 'npts': array([ 1681.]),
 'quartile': array([ 0.00305395]),
 'rms': array([ 0.00411418]),
 'sigma': array([ 0.00403944]),
 'sum': array([ 1.3226716]),
 'sumsq': array([ 0.02845345]),
 'trc': array([148, 148, 0, 21]),
 'trcf': '15:21:39.919, +05.08.59.981, I, 1.41332e+09Hz')

**ALERT:** The return dictionary currently includes NumPy array values, which have to be accessed by an array index to get the array value. To access these dictionary elements, use the standard Python dictionary syntax, e.g.

    xstat['<key string>'][<array index>]

For example, to extract the standard deviation as a number

    mystddev = xstat['sigma'][0]
    print 'Sigma = ' + str(xstat['sigma'][0])

6.9.2 Examples for imstat

The following are some examples using the B1608+656 Tutorial

http://casa.nrao.edu/Doc/Scripts/b1608_demo.py

as an example.

To extract statistics for the final image:
xstat = imstat('b1608.demo.clean2.image')

# Printing out some of these
print 'Max = '+str(xstat['max'][0])
print 'Sigma = '+str(xstat['sigma'][0])

# results:
# Max = 0.016796965152
# Sigma = 0.00033631979385

In a box around the brightest component:

xstat_A = imstat('b1608.demo.clean2.image', box='124,125,132,133')

# Printing out some of these
print 'Comp A Max Flux = '+str(xstat_A['max'][0])
print 'Comp A Max X,Y = ('+str(xstat_A['maxpos'][0])+','+str(xstat_A['maxpos'][1])+')'

# results:
# Comp A Max Flux = 0.016796965152
# Comp A Max X,Y = (128,129)

6.10 Extracting data from an image (imval)

The imval task will extract the values of the data and mask from a specified region of an image and place in the task return value as a Python dictionary.

The inputs are:

# imval :: Get the data value(s) and/or mask value in an image.
imagename = '' # Name of the input image
region = '' # Image Region. Use viewer
box = '' # Select one or more box regions
chans = '' # Select the channel(spectral) range
stokes = '' # Stokes params to image (I,IV,IQU,IQUV)

Area selection using box and region is detailed in § 6.1.2 and § 6.1.6 respectively. By default, box='' will extract the image information at the reference pixel on the direction axes.

Plane selection is controlled by chans and stokes. See § 6.1.3 for details on plane selection. By default, chans='' and stokes='' will extract the image information in all channels and Stokes planes.

For instance,

xval = imval('myimage', box='144,144', stokes='I' )

will extract the Stokes I value or spectrum at pixel 144,144, while

xval = imval('myimage', box='134,134.154,154', stokes='I' )
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will extract a 21 by 21 pixel region.
Extractions are returned in NumPy arrays in the return value dictionary, plus some extra elements
describing the axes and selection:
CASA <2>: xval = imval(’ngc5921.demo.moments.integrated’)
CASA <3>: xval
Out[3]:
{’axes’: [[0, ’Right Ascension’],
[1, ’Declination’],
[3, ’Frequency’],
[2, ’Stokes’]],
’blc’: [128, 128, 0, 0],
’data’: array([ 0.89667124]),
’mask’: array([ True], dtype=bool),
’trc’: [128, 128, 0, 0],
’unit’: ’Jy/beam.km/s’}

extracts the reference pixel value in this 1-plane image. Note that the ’data’ and ’mask’ elements
are NumPy arrays, not Python lists.
To extract a spectrum from a cube:
CASA <8>: xval = imval(’ngc5921.demo.clean.image’,box=’125,125’)
CASA <9>: xval
Out[9]:
{’axes’: [[0, ’Right Ascension’],
[1, ’Declination’],
[3, ’Frequency’],
[2, ’Stokes’]],
’blc’: [125, 125, 0, 0],
’data’: array([ 8.45717848e-04,
1.93370355e-03,
1.53750915e-03,
2.88399984e-03,
2.38683447e-03,
2.89159478e-04,
3.16268904e-03,
9.93389636e-03,
1.88773088e-02,
3.01138610e-02,
3.14478502e-02,
4.03211266e-02,
3.82498614e-02,
3.06552909e-02,
2.80734301e-02,
1.72479432e-02,
1.20884273e-02,
6.13593217e-03,
9.04005766e-03,
1.71429547e-03,
5.22095338e-03,
2.49114982e-03,
5.30831399e-04,
4.80734324e-03,
1.19265869e-05,
1.29435991e-03,
3.75700940e-04,
2.34788167e-03,
2.72604497e-03,
1.78467855e-03,
9.74952069e-04,
2.24676146e-03,
1.82263291e-04,
1.98463408e-06,
2.02975096e-03,
9.65532148e-04,
1.68218743e-03,
2.92119570e-03,
1.29359076e-03,
-5.11484570e-04,
1.54162932e-03,
4.68662125e-04,
-8.50282842e-04, -7.91683051e-05,
2.95954203e-04,
-1.30133145e-03]),
’mask’: array([ True, True, True, True, True, True, True, True,
True, True, True, True, True, True, True, True, True,

True,


To extract a region from the plane of a cube:

CASA <13>: xval = imval('ngc5921.demo.clean.image', box='126,128,130,129', chans='23')

CASA <14>: xval
Out[14]:

={'axes': [[0, 'Right Ascension'],
        [1, 'Declination'],
        [3, 'Frequency'],
        [2, 'Stokes']],
 'blc': [126, 128, 0, 23],
'data': array([[ 0.00938627, 0.01487772],
               [ 0.00955847, 0.01688832],
               [ 0.00696965, 0.01501907],
               [ 0.00460964, 0.01220793],
               [ 0.00358087, 0.00990202]],
'mask': array([[ True, True],
               [ True, True],
               [ True, True],
               [ True, True]], dtype=bool),
 'trc': [130, 129, 0, 23],
'unit': 'Jy/beam'}

CASA <15>: print xval['data'][0][1]
0.0148777160794

In this example, a rectangular box was extracted, and you can see the order in the array and how to address specific elements.

### 6.11 Reordering the Axes of an Image Cube (imtrans)

Sometimes data cubes can be in axis orders that are not adequate for processing. The CASA task `imtrans` can change the ordering of the axis:

```python
# imtrans :: Reorder image axes
imname = '' # Name of the input image
outname = '' # Name of output CASA image.
order = '' # New zero-based axes order.
wantreturn = True # Return an image tool referencing the transposed image
```
The order parameter is the most important input here. It is a string of numbers that shows how axes 0, 1, 2, 3, ... are mapped onto the new cube (note that the first axis has the label 0, as typical in python). E.g. order='1032' will reorder the input axis 0 to be axis 1 in the output, input axis 1 to be output axis 0, input axis 2 to output axis 3 (the last axis) and input axis 3 to output axis 2. Alternatively, axes can be specified by their names. E.g., to reorder an image with right ascension, declination, and frequency and reverse the first two, order=['declination', 'right ascension', 'frequency'] will work. The axes names can be found typing (ia.coordsys().names()). Minimum match is supported, so that order=['d', 'f', 'r'] will produce the same results.

Axes can simultaneously be transposed and reversed. To reverse an axis, precede it by a "-". For example, order='10-32' will reverse the direction of the first and third axis of the input image (the zeroth and second axes in the output image).

Example:

Swap the stokes and spectral axes in an RA-Dec-Stokes-Frequency image

```python
imagename = "myim.im"
outfile = "outim.im"
order = "0132"
imtrans()
```

or

```python
outfile = "myim_2.im"
order = 132
imtrans()
```

or

```python
outfile = "myim_3.im"
order = ["r", "d", "f", "s"]
imtrans()
```

or

```python
outfile = "myim_4.im"
order = ["rig", "declin", "frequ", "stok"]
imtrans()
```

If the outfile parameter is empty, only a temporary image is created; no output image is written to disk. The temporary image can be captured in the returned value (assuming wantreturn is true).
6.12 Collapsing an Image Along an Axis (imcollapse)

imcollapse allows to apply an aggregation function along one or more axes of an image. Functions supported are 'max', 'mean', 'median', 'min', 'rms', 'stdev', 'sum', 'variance' (minimum match supported). The relevant axes will then collapse to a single value or plane (i.e. they will result in a degenerate axis). The functions are specified in the function parameter of the imcollapse inputs:

```python
# imcollapse :: Collapse image along one axis, aggregating pixel values along that axis.
# Name of the input image
imagename = ''
# Function used to compute aggregation of pixel values.
function = ''
# Zero-based axis number(s) or minimal match strings to collapse.
axes = [0]
# Name of output CASA image.
outfile = ''
# Optional direction plane box ("blcx, blcy, trcx, trcy").
box = ''
# Name of optional region file to use.
region = ''
# Optional zero-based contiguous frequency channel specification.
chans = ''
# Optional contiguous stokes planes specification.
stokes = ''
# Optional mask to use.
mask = ''
# Should an image analysis tool referencing the collapsed image be returned?
wantreturn = True
```

wantreturn=True returns an image analysis tool containing the newly created collapsed image.

Example:

myimage.im is a 512x512x128x4 (ra,dec,freq,stokes; i.e. in the 0-based system, frequency is labeled as axis 2) image and we want to collapse a subimage of it along its spectral axis avoiding the 8 edge channels at each end of the band, computing the mean value of the pixels (resulting image is 256x256x1x4 in size):

```
imcollapse(imagename="myimage.im", outfile="collapse_spec_mean.im", function="mean", axis=2, box="127,127,383,383", chans="8~119")
```

Note that imcollapse will not smooth to a common beam for all planes if they differ. If this is desired, run imsmooth before imcollapse.

6.13 Regridding an Image (imregrid)
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It is occasionally necessary to regrid an image onto a new coordinate system. The `imregrid` task will regrid one image onto the coordinate system of another, creating an output image. In this task, the user need only specify the names of the input, template, and output images.

The default inputs are:

```python
imregrid :: regrid an image onto a template image

imagename = '' # Name of the source image
template = 'get' # A dictionary, refcode, or name of an image that provides the output shape and coordinate system
output = '' # Name for the regridded image
asvelocity = True # Regrid spectral axis in velocity space rather than frequency space?
axes = [-1] # The pixel axes to regrid. -1 => all.
interpolation = 'linear' # The interpolation method. One of "nearest", "linear", "cubic"
decimate = 10 # Decimation factor for coordinate grid computation
replicate = False # Replicate image rather than regrid?
overwrite = False # Overwrite (unprompted) pre-existing output file?
```

The output image will have the data in `imagename` regrided onto the coordinate system provided by the `template` parameter. `template` is used universally for a range of ways to define the grid of the output image:

- **a template image**: specify an image name here and the input will be regridded to the same 3-dimensional coordinate system as the one in `template`. Values are filled in as blanks if they do not exist in the input. Note that the input and template images must have the same coordinate structure to begin with (like 3 or 4 axes, with the same ordering)

- **a coordinate system (reference code)**: to convert from one coordinate frame to another one, e.g. from B1950 to J2000, the `template` parameter can be used to specify the output coordinate system. These following recognized keywords are supported: 'J2000', 'B1950', 'B1950_VLA', 'GALACTIC', 'HADEC', 'AZEL', 'AZELSW', 'AZELNE', 'ECLIPTIC', 'MECLIPTIC', 'TECLIPTIC', 'SUPERGAL'

- `'get'`: This option returns a python dictionary in the `{‘csys’: csys_record, ‘shap’: shape}` format

- **a python dictionary**: In turn, such a dictionary can be used as a template to define the final grid

6.14 Redefining the Velocity System of an Image (imreframe)

`imreframe` can be used to change the velocity system of an image. It is not applying a regridding as a change from radio to optical conventions would require, but it will change the labels of the velocity axes.

Inside the Toolkit:

More complex coordinate system and image regridding operation can be carried out in the toolkit. The `coordsys (cs)` tool and the `ia.regrid` method are the relevant components.
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# imreframe :: Change the frame in which the image reports its spectral values
imagename = '' # Name of the input image
output = '' # Name of the output image; '' => modify input image
outframe = 'lsrk' # Spectral frame in which the frequency or velocity values will be reported by default
restfreq = '' # restfrequency to use for velocity values (e.g. "1.420GHz" for the HI line)

outframe defines the velocity frame (LSRK, BARY, etc., see §C.2) of the output image and a rest frequency should be specified to relabel the spectral axis in new velocity units.

6.15 Rebin an Image (imrebin)

The task imrebin allows one to rebin an image in any spatial or spectral direction:

# imrebin :: Rebin an image by the specified integer factors
imagename = '' # Name of the input image
outfile = '' # Output image name.
factor = [] # Binning factors for each axis
region = '' # The region to rebin. Default is entire image. Do not specify region and box/chans simultaneously.
    box = '' # Box in directional plane to rebin. Default is to use the entire directional plane.
    chans = '' # Channels to rebin. See "help par.chans" for examples.
    stokes = '' # The correlations to include in the output. Default is all. Stokes planes cannot be rebinned.

mask = '' # Mask to use. See help par.mask. Default is none.
dropdeg = False # Drop degenerate axes?
crop = True # Remove pixels from the end of an axis to be rebinned if there are not enough to form an integral bin?

where factor is a list of integers that provides the numbers of pixels to be binned for each axis. The crop parameters controls how pixels at the boundaries are treated if the bin values are not multiple integers of the image dimensions.

Example:

imrebin(imagename="my.im", outfile="rebinned.im", factor=[1,2,1,4], crop=T)

This leaves RA untouched, bins DEC by a factor of 2, leaves Stokes as is, and bins the spectral axis by a factor of 4. If the input image has a spectral axis with a length that is not a multiple of 4, the crop=T setting will discard the remaining 1-3 edge pixels.
6.16   1-dimensional Smoothing (specsmooth)

To gain higher signal-to-noise of data cubes, one can smooth the data along one dimension (for
2-dimensional smoothing, see $\text{imsmooth} \S 6.16$). Typically this is the spectral axis. Hanning and
Boxcar smoothing kernels are available in the task $\text{specsmooth}$:

\begin{verbatim}
# specsmooth :: Smooth an image region in one dimension
imagename = ''    # Name of the input image
outfile = ''      # Output image name.
region = ''       # Region selection. See help par.region for possible
                 # specifications. Default: Do not use a region.
     box = ''       # Rectangular box in direction coordinate blc, trc.
                 # Default: entire image ("").
mask = ''         # Mask to use. See help par.mask. Default is none..
axis = -1         # The profile axis. Default: use the spectral axis if one
                 # exists, axis 0 otherwise (<0).
function = 'hanning'    # Convolution function. hanning and boxcar are supported
                       # functions. Minimum match is supported.
dmethod = 'copy'    # Decimation method. "" means no decimation, "copy" and
                       # "mean" are also supported

The parameter $dmethod='copy'$ allows one to only keep every $w$'th channel, if the smoothing kernel
has a width of $w$. Leaving this parameter empty will return the same size cube as the input and
setting it to 'mean' will average planes using the kernel width.

6.17   2-dimensional Smoothing; Image Convolution (imsmooth)

A data cube can be smoothed across spatial dimensions with $\text{imsmooth}$. The inputs are:

\begin{verbatim}
# imsmooth :: Smooth an image or portion of an image
imagename = ''    # Name of the input image. Must be specified.
kernell = 'gauss'    # Type of kernel to use. Acceptable values are "b", "box",
                    # or "boxcar" for a boxcar kernel, "g", "gauss", or
                    # "gaussian" for a gaussian kernel, "c", "common", or
                    # "commonbeam" to use the common beam of an image with
                    # multiple beams as the gaussian to which to convolve all
                    # the planes, "i" or "image" to use an image as the
                    # kernel.
beam = ''         # Alternate way of describing a Gaussian. If specified,
                    # must be a dictionary with keys "major", "minor", and
                    # "pa" (or "positionangle"). Do not specify beam if
                    # specifying major, minor, and pa. Example: Example:
                    # {"major": "5arcsec", "minor": "2arcsec", "pa": "20deg"}.
\end{verbatim}
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targetres = False # If gaussian kernel, specified parameters are to be
# resolution of output image (True) or parameters of
# gaussian to convolve with input image (False).

major = '' # Major axis for the kernels. Standard quantity
# representation. Must be specified for kernel="boxcar".
# Example: "4arcsec".

minor = '' # Minor axis. Standard quantity representation. Must be
# specified for kernel="boxcar". Example: "2arcsec".

pa = '' # Position angle used only for gaussian kernel. Standard
# quantity representation. Example: "40deg".

region = '' # Region selection. See help par.region. Empty string means
# use box/chans/stokes if supplied, or else entire image.

box = '' # Rectangular region specification in directional plane. Do
# not specify region if you specify box.

chans = '' # Select the spectral channel range. See "help par.chans"
# for examples. Do not specify region if you specify
# chans.

stokes = 'I' # Stokes parameters to image (e.g., I,IV,IQU,IQUV). Do not
# specify region if you specify stokes.

mask = [] # Mask to use. See help par.mask. Default is none.

stretch = False # Stretch the mask if necessary and possible? See help
# par.stretch

outfile = '' # Output image name. Must be specified.

overwrite = False # Overwrite (unprompted) pre-existing output file?

where the cube/image imagename will be convolved with a kernel defined in the kernel keyword.
Kernels 'gauss' and 'boxcar' need the major and minor axes sizes as input, the Gaussian kernel
smoothing also requires a position angle. By default, the kernel size defines the kernel itself, i.e.
the data will be smoothed with this kernel. If the targetres parameter for Gaussian kernels is set
to 'True', major and minor axes will be those from the output resolution, and the kernel will be
adjusted for each plane to arrive at the final resolution.

The 'commonbeam' kernel is to be used when the beam shape is different as a function of frequency.
This option will then smooth all planes to a single beam, defined by the largest beam in the cube.

With the 'image' kernel, one can specify an image that will serve as the convolution kernel. A scale
factor can be applied, which defaults to flux conservation.

Examples:

1) smoothing to a Gaussian kernel 20" by 10"

imsmooth( imagename='my.image', kernel='gauss', major='20arcsec', minor='10arcsec', targetres=T)

2) Smoothing using pixel coordinates and a boxcar kernel.

imsmooth( imagename='new.image', major='20pix', minor='10pix', kernel='boxcar')
6.18 Spectral Line fitting with specfit

specfit is a powerful task to perform spectral line fits in data cubes. Three types of fitting functions are currently supported, polynomials, Gaussians, and Lorentzians. specfit can fit these functions in two ways: over data that were averaged across a region (multifit=False) or on a pixel by pixel basis (multifit=True).

```plaintext
# specfit :: Fit 1-dimensional Gaussians and/or polynomial models to an image or image region
imagename  = ''   # Name of the input image
box        = ''   # Rectangular box in direction coordinate
                # blc, trc. Default: entire image ("").
region     = ''   # Region of interest. See help par.region
                # for possible specifications. Default: Do
                # not use a region.
chans      = ''   # Channels to use. Channels must be
                # contiguous. Default: all channels ("").
stokes     = ''   # Stokes planes to use. Planes must be
                # contiguous. Default: all stokes ("").
axis       = -1   # The profile axis. Default: use the
                # spectral axis if one exists, axis 0
                # otherwise (<0).
mask       = ''   # Mask to use. See help par.mask. Default is
                # none..
poly       = -1   # Order of polynomial element. Default: do
                # not fit a polynomial (<0).
estimates  = ''   # Name of file containing initial estimates.
                # Default: No initial estimates ("").
ngauss     = 1    # Number of Gaussian elements. Default: 1.
pampest    = ''   # Initial estimate of PCF profile (gaussian
                # or lorentzian) amplitudes.
pcenterest = ''   # Initial estimate PCF profile centers, in
                # pixels.
pfwhmest   = ''   # Initial estimate PCF profile FWHMs, in
                # pixels.
pfix       = ''   # PCF profile parameters to fix during fit.
pfunc      = ''   # PCF singlet functions to fit. "gaussian"
                # or "lorentzian" (minimal match
                # supported). Unspecified means all
                # gaussians.
minpts     = 0    # Minimum number of unmasked points
                # necessary to attempt fit.
multifit   = True # If true, fit a profile along the desired
                # axis at each pixel in the specified
                # region. If false, average the non-fit
                # axis pixels and do a single fit to that
                # average profile. Default False.
amp        =''   # Name of amplitude solution image. Default:
                # do not write the image ("").
amperr     =''   # Name of amplitude solution error image.
```

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# Default: do not write the image ("").
center = '' # Name of center solution image. Default: do
# not write the image ("").

centererr = '' # Name of center solution error image.

fwhm = '' # Name of fwhm solution image. Default: do
# not write the image ("").

fwhmerr = '' # Name of fwhm solution error image.

integral = '' # Prefix of name of integral solution image.
# Name of image will have gaussian
# component number appended. Default: do
# not write the image ("").

integralerr = '' # Prefix of name of integral error solution
# image. Name of image will have gaussian
# component number appended. Default: do
# not write the image (""

model = '' # Name of model image. Default: do not write
# the model image ("

residual = '' # Name of residual image. Default: do not
# write the residual image ("

wantreturn = True # Should a record summarizing the results be
# returned?

logresults = True # Output results to logger?

gmcomps = 0 # Number of components in each gaussian
# multiplet to fit

gmampcon = '' # The amplitude ratio constraints for non-
# reference components to reference
# component in gaussian multiplets.

gmcentercon = '' # The center offset constraints (in pixels)
# for non-reference components to reference
# component in gaussian multiplets.

gmfwhmcon = '' # The FWHM ratio constraints for non-
# reference components to reference
# component in gaussian multiplets.

gmampest = [0.0] # Initial estimate of individual gaussian
# amplitudes in gaussian multiplets.

gmcenterest = [0.0] # Initial estimate of individual gaussian
# centers in gaussian multiplets, in
# pixels.

gmfwhmest = [0.0] # Initial estimate of individual gaussian
# FWHMss in gaussian multiplets, in pixels.

gmfix = '' # Parameters of individual gaussians in
# gaussian multiplets to fix during fit.

logfile = '' # File in which to log results. Default is
# not to write a logfile.

goodamprange = [0.0] # Acceptable amplitude solution range. [0.0]
# => all amplitude solutions are
# acceptable.

goodcenterrange = [0.0] # Acceptable center solution range in pixels
6.18.1 Polynomial Fits

Polynomials can be fit by specifying the polynomial order in poly. Negative orders will not fit any polynomials.

6.18.2 Lorentzian and Gaussian Fits

Gaussian and Lorentzian fits are very similar, they both require amplitude, center, and FWHM to be fully specified. All of the following discussion is thus valid for both functions. The parameter pfunc controls whether Gaussian or Lorentzian functions are to be used. Default is all Gaussians. pfunc=["L", "G", "G", "L"] would use Lorentzian, Gaussian, Gaussian, and Lorentzian components in the order they appear in the estimates file (see below).

6.18.2.1 One or more single Gaussian/Lorentzian

For Gaussian and Lorentzian fits, the task will allow multiple components and specfit will try to find the best solution. The parameter space, however, is usually not uniform and to avoid local minima in the goodness-of-fit space, one can provide initial start values for the fits. This can be done either through the parameters pampest, pccenterest, and pfwhmest for the amplitudes, center, and FWHM estimates in image coordinates. pfix can take parameters that specify fixed fit values. Any combination of the characters 'p' (peak), 'c' (center), and 'f' (fwhm) are permitted, e.g. "fc" will hold the fwhm and the center constant during the fit. Fixed parameters will have no errors associated with them in the solution. Alternatively, a file with initial values can be supplied by the estimates parameter (one Gaussian/Lorentzian parameter set per line). The file has the following format:

[peak intensity], [center], [fwhm], [optional fixed parameter string]

The first three values are required and must be numerical values. The peak intensity must be expressed in map units, while the center and fwhm must be specified in pixels. The fourth value is optional and if present, represents the parameter(s) that should be held constant during the fit (see above).

An example estimates file is:

# estimates file indicating that two Gaussians should be fit
# first guassian estimate, peak=40, center at pixel number 10.5,
# fwhm = 5.8 pixels, all parameters allowed to vary during
and the output of a typical execution, e.g.

```python
specfit(imagename='IRC10216_HC3N.cube_r0.5.image',
region='specfit.crtf', multifit=F, estimates='', ngauss=2)
```

('specfit.crtf' is a CASA regions file, see Section[D])

will be

```
Fit :
  RA : 09:47:57.49
  Dec : 13:16:46.46
  Stokes : I
  Pixel : [146.002, 164.499, 0.000, *]
  Attempted : YES
  Converged : YES
  Iterations : 28
  Results for component 0:
    Type : GAUSSIAN
    Peak : 5.76 +/- 0.45 mJy/beam
    Center : -15.96 +/- 0.32 km/s
              40.78 +/- 0.31 pixel
    FWHM : 7.70 +/- 0.77 km/s
           7.48 +/- 0.74 pixel
    Integral : 47.2 +/- 6.0 mJy/beam.km/s
  Results for component 1:
    Type : GAUSSIAN
    Peak : 4.37 +/- 0.33 mJy/beam
    Center : -33.51 +/- 0.58 km/s
              23.73 +/- 0.57 pixel
    FWHM : 15.1 +/- 1.5 km/s
           14.7 +/- 1.5 pixel
    Integral : 70.2 +/- 8.8 mJy/beam.km/s
```

If `wantreturn=True` (the default value), the task returns a python dictionary (here captured in a variable with the inventive name of 'fitresults'):

```python
fitresults=specfit(imagename='IRC10216_HC3N.cube_r0.5.image', region='specfit.rgn', multifit=F,
estimates='', ngauss=2)
```

The values can then be used by other python code for further processing.
6.18.2.2 Gaussian Multiplets

It is possible to fit a number of Gaussians together, as multiplets with restrictions. All restrictions are relative to a reference Gaussian (the zero’th component of each multiplet). \texttt{gncomps} specifies the number of Gaussians for each multiplet, and, in fact, a number of these multiplets can be fit simultaneously. \texttt{gncomps=[2,4,3]}, e.g. fits a 2-component Gaussian, a 4-component Gaussian, and a 3-component Gaussian all at once. The initial parameter estimates can be specified with the \texttt{gmampest}, \texttt{gmcenterest}, and \texttt{gmfwhmest} parameters and the estimates are simply listed in the sequence of \texttt{gncomps}. E.g. if \texttt{gncomps=[2,4,3]} is specified with multiplet G0 consisting of 2 Gaussians a, b, multiplet G1 of 4 Gaussians c, d, e, f, and multiplet G2 of three Gaussians g, h, i, the parameter list in \texttt{gm*est} would be like \texttt{gm*est=[a,b,c,d,e,f,g,h,i]}.

Restrictions can be specified via the \texttt{gmampcon} parameter for the amplitude ratio (non-reference to reference), \texttt{gcentercon} for the offset in pixels (to a reference), and \texttt{gmfwhmcon} for the FWHM ratio (non-reference to reference). A value of 0 will not constrain anything. The reference is always the zero’th component in each multiplet, in our example, Gaussians a, c, and g. They cannot be constrained. So \texttt{gncomps=[2, 4, 3], gmampcon=[0, 0.2, 0, 0.1, 4.5, 0]}, \texttt{gcentercon=[24.2, 45.6, 92.7, 0, -22.8, -33.5]}, and \texttt{gmfwhmcon=\"\"} would constrain Gaussians b relative to a with a 24.2 pixel offset, Gaussian d to c with a amplitude ratio of 0.2 and a 45.6 pixel offset, Gaussian e to c with a offset of 92.7 pixels, etc. Restrictions will overrule any estimates.

The parameters \texttt{goodamprange}, \texttt{goodcenterrange}, and \texttt{goodfwhmrange} can be used to limit the range of amplitude, center, and fwhm solutions for all Gaussians.

6.18.3 Pixel-by-pixel fits

As mentioned above, \texttt{specfit} can also fit spectral cubes on a pixel by pixel basis. In this case, one can choose to write none, any or all of the solution and error images for Gaussian/Lorentzian fits via the parameters \texttt{amp, amperr, center, centererr, fwhm, and fwhmerr}. The files written contain all the components along a “component number” axis. They can be inspected by using the “hidden axes” slider in the \texttt{viewer} (§7.4.1). Writing analogous images for polynomial coefficients is not yet supported although polynomial fits when \texttt{multifit=True} is supported. Best fit coefficients are written to the logger. Pixels for which fits were not attempted or did not converge will be masked as bad.

6.19 Calculation of Rotation Measures \texttt{rmfit}

\texttt{rmfit} is an image domain task that accepts an input cube image containing Stokes Q and U axes and will generate the rotation measure by performing a least square fit in the image domain to obtain the best fit to the equation $\chi = \chi_0 + RM \times \lambda^2$.

The inputs to \texttt{rmfit} are:
# rmfit :: Calculate rotation measure.

```python
# Name(s) of the input image(s). Must be specified.
imagename = ''

# Output rotation measure image name. If not specified, no image is written.
rm = ''

# Output rotation measure error image name. If not specified, no image is written.
rmerr = ''

# Output position angle (degrees) at zero wavelength image name. If not specified, no image is written.
pa0 = ''

# Output position angle (degrees) at zero wavelength error image name. If not specified, no image is written.
pa0err = ''

# Output number of turns image name. If not specified, no image is written.
nturns = ''

# Output reduced chi squared image name. If not specified, no image is written.
chisq = ''

# Estimate of the thermal noise. A value less than 0 means auto estimate.
sigma = ''

# Foreground rotation measure in rad/m/m to subtract.
rmfg = 0.0

# Maximum rotation measure in rad/m/m for which to solve. IMPORTANT TO SPECIFY.
rmax = 0.0

# Maximum input position angle error in degrees to allow in solution determination.
maxpaerr = 1e+30
```

This task generates the rotation measure image from stokes Q and U measurements at several different frequencies. You are required to specify the name of at least one image with a polarization axis containing stokes Q and U planes and with a frequency axis containing more than two pixels. The frequencies do not have to be equally spaced (ie the frequency coordinate can be a tabular coordinate). It will work out the position angle images for you. You may also specify multiple image names, in which case these images will first be concatenated along the spectral axis using `ia.imageconcat()`.

See also the `imagepol.fourierrotationmeasure` function for a new Fourier-based approach.

Rotation measure algorithms that work robustly are few. The main problem is in trying to account for the $n - \pi$ ambiguity (see Leahy et al, Astronomy & Astrophysics, 156, 234 or the MIRIAD manual\(^1\)).

The algorithm that this task uses is that of Leahy et al. in their Appendix A.1. But as in all these algorithms, the basic process is that for each spatial pixel, the position angle vs frequency data is fit to determine the rotation measure and the position angle at zero wavelength (and associated errors). An image containing the number of $n - \pi$ turns that were added to the data at each spatial pixel and for which the best fit was found can be written. The reduced $\chi^2$ image for the fits can also be written.

Note that no assessment of curvature (i.e. deviation from the simple linear position angle - $\lambda^2$ functional form) is made.

Any combination of output images can be written.

\(^1\)http://www.cfa.harvard.edu/sma/miriad/manuals/SMAuguide/smauserhtml/imrm.html
The parameter \( \text{sigma} \) gives the thermal noise in Stokes Q and U. By default it is determined automatically using the image data. But if it proves to be inaccurate (maybe not many signal-free pixels), it may be specified. This is used for calculating the error in the position angles (via propagation of Gaussian errors).

The argument \( \text{maxpaerr} \) specifies the maximum allowable error in the position angle that is acceptable. The default is an infinite value. From the standard propagation of errors, the error in the linearly polarized position angle is determined from the Stokes Q and U images (at each directional pixel for each frequency). If the position angle error for any pixel exceeds the specified value, the position angle at that pixel is omitted from the fit. The process generates an error for the fit and this is used to compute the errors in the output images.

Note that \( \text{maxpaerr} \) is not used to mask pixels in the output images.

The argument \( \text{rmfg} \) is used to specify a foreground RM value. For example, you may know the mean RM in some direction out of the Galaxy, then including this can improve the algorithm by reducing ambiguity.

The parameter \( \text{rmmax} \) specifies the maximum absolute RM value that should be solved for. This is quite an important parameter. If you leave it at the default, zero, no ambiguity handling will be used. So some a priori information should be supplied; this is the basic problem with rotation measure algorithms.

### 6.20 Calculation of Spectral Indices and Higher Order Polynomials (spxfit)

This application fits a power logarithmic polynomial or a logarithmic transformed polynomial to pixel values along a specified axis of an image or images. These functions are most often used for fitting the spectral index and higher order terms of a spectrum. A power logarithmic polynomial has the form

\[
y = \frac{c_0 x}{D(c_1 + c_2 \ln(x/D) + c_3 \ln(x/D)^2 + \ldots + c_n \ln(x/D)^n)}
\]  

and a logarithmic transformed polynomial is simply the result of this equation after taking the natural log of both sides so that it has the form

\[
\ln(y) = c_0 + c_1 \ln(x) + c_2 \ln(x/D)^2 + \ldots + c_n \ln(x/D)^n
\]  

Because the logarithm of the ordinate values must be taken before fitting a logarithmic transformed polynomial, all non-positive pixel values are effectively masked for the purposes of fitting.

The coefficients of the two forms are equal to each other except that \( c_0 \) in the second equation is equal to \( \ln(c_0) \) of the first. In the case of fitting a spectral index, which is traditionally represented as \( \alpha \), is equal to \( c_1 \).
In both cases, $D$ is a normalization constant used so that abscissa values are closer to unity when they are sent to the fitter. This generally improves the probability that the fit will converge. This parameter may be specified via the div parameter. A value of 0 (the default) indicates that the application should determine a reasonable value for $D$, which is determined via

$$D = 10\int (\log_{10}(\sqrt{\min(x)\cdot\max(x)}))$$

where $\min(x)$ and $\max(x)$ are the minimum and maximum abscissa values, respectively.

The inputs are:

```python
# spxfit :: Fit a 1-dimensional model to an image or image region for determination of spectral index.
# Name of the input image(s)
imagename = ''
# Rectangular box in direction coordinate blc, trc.
box = ''
# Region of interest. See help par.region for possible specifications. Default: Do not use a region.
region = ''
# Channels to use. Channels must be contiguous. See "help par.chans" for examples. Default: all channels ("").
chans = ''
# Stokes planes to use. Planes must be contiguous. Default: all stokes ("").
stokes = ''
# The profile axis. Default: use the spectral axis if one exists, axis 0 otherwise (<0).
axis = -1
# Mask to use. See help par.mask. Default is none.
mask = ''
# Minimum number of unmasked points necessary to attempt fit.
minpts = 1
# If true, fit a profile along the desired axis at each pixel in the specified region. If false, average the non-fit axis pixels and do a single fit to that average profile. Default False.
multifit = True
# Name of the spectral index function coefficient solution image to write.
spxsol = ''
# Name of the spectral index function coefficient error image to write.
spxerr = ''
# Name of model image. Default: do not write the model
model = ''
```
residual = '' # Name of residual image. Default: do not write the residual image ('').

spxtype = 'plp' # Type of function to fit. "plp" => power logarithmic polynomial, "ltp" => logarithmic transformed polynomial.

spxest = [] # Initial estimates for the spectral index function coefficients.

spxfix = [] # Fix the corresponding spectral index function coefficients during the fit. True=>hold fixed.

div = 0 # Divisor (numerical value or quantity) to use in the logarithmic terms of the plp or ltp function. 0 => calculate a useful value on the fly.

wantreturn = True # Should a record summarizing the results be returned?

logresults = True # Output results to logger?

logfile = '' # File in which to log results. Default is not to write a logfile.

sigma = -1 # Standard deviation array or image name(s).

outsigma = '' # Name of output image used for standard deviation. Ignored if sigma is empty.

for more than a single input image or cube, all images must have the same dimensions along all axes other than the fit axis. multifit will perform a per pixel fit, otherwise there will be a single value over the entire region.

6.21 Image Mask Handling makemask

makemask facilitates the handling of image masks in CASA. There are two basic mask formats: 1) one or more Boolean masks stored internally in the image file, and 2) images with zero and non-zero image values. makemask looks like:

# makemask :: Makes and manipulates image masks
mode = 'list' # Mask method (list, copy,expand,delete,setdefaultmask)
inpimage = '' # Name of input image.

To distinguish between Boolean internal masks and zero/non-zero images, makemask uses the syntax galaxy.image:mask0 for Boolean masks within an image, in this example the Boolean mask mask0
within the image `galaxy.image`. Without the colon separator, the image itself is assumed to be treated like a zero/non-zero mask.

`mode='list'` lists all the internal Boolean masks that are present in an image. The default masks can be set with mode `setdefaultmask` and they can be deleted with the mode `delete`. The default mask is used when an image is displayed in the `viewer` and is used in all analysis tasks.

`mode='copy'` lets a user to copy a Boolean mask from one image to another image, or to write out the Boolean mask as a zero/non-zero image. The latter format is very useful when different masks are combined or manipulated. All the image analysis tools, in particular `immath` are applicable for such zero/non-zero masks as they act like normal images. `makemask` will always attempt to regrid the input mask to the output image.

In addition `mode='copy'` can be used to merge masks and also accepts regions. E.g. to create a mask from a CASA region file (CRTF, see §6.1.6), the input would look like:

```python
# makemask :: Makes and manipulates image masks
mode = 'copy' # Mask method (list, copy,expand,delete,setdefaultmask)
inpimage = 'inputimage.im' # Name of input image.
inpmask = 'region.crtf' # mask(s) to be processed: image masks,T/F internal masks
output = 'imagemask.im' # Name of output mask (imagename or imagename:internal_maskname)
overwrite = False # overwrite output if exists?
```

`mode='expand'` furthermore expands a mask in the spectral domain. It regrids first then stretches the edge channels. E.g. a one plane continuum image would be stretched to all planes of a data cube.

### 6.22 Search for Spectral Line Rest Frequencies (slsearch)

The `slsearch` task allows the spectral line enthusiast to find their favorite spectral lines in subset of the Splatalogue spectral line catalog ([http://www.splatalogue.net](http://www.splatalogue.net)) which is distributed with CASA. In addition, one can export custom catalogs from Splatalogue and import them to CASA using the task `splattotable` (Sect. 6.23) or tool method `sl.splattotable()`. One can even import catalogs with lines not in Splatalogue using the same file format.

The inputs to `slsearch` are as follows:

```python
# slsearch :: Search a spectral line table.
tablename = '' # Input spectral line table name to search. If not specified, use the default table in the system.
outfile = '' # Results table name. Blank means do not write the table to disk.
freqrange = [84, 90] # Frequency range in GHz.
```
species = [''] # Species to search for.
reconly = False # List only NRAO recommended frequencies.
chemnames = [''] # Chemical names to search for.
qns = [''] # Resolved quantum numbers to search for.
rrlinclude = True # Include RRLs in the result set?
rrlonly = False # Include only RRLs in the result set?
i
ntensity = -1 # CDMS/JPL intensity range. -1 -> do not use an intensity range.
smu2 = -1 # S*mu*mu range in Debye**2. -1 -> do not use an S*mu*mu range.
loga = -1 # log(A) (Einstein coefficient) range. -1 -> do not use a loga range.
eu = -1 # Upper energy state range in Kelvin. -1 -> do not use an eu range.
el = -1 # Lower energy state range in Kelvin. -1 -> do not use an el range.
verbose = True # List result set to logger (and optionally logfile)?
logfile = '' # List result set to this logfile (only used if verbose=True).
append = True # If true, append to logfile if it already exists, if false overwrite logfile if it exists. Only used if verbose=True and logfile not blank.
wantreturn = True # If true, return the spectralline tool associated with the result set.

The table is provided in the tablename parameter but if it is blank (the default), the catalog which is included with CASA will be used. Searches can be made in a parameter space with large dimensionality:

- **freqrange** Frequency range in GHz.
- **species** Species to search for.
- **reconly** List only NRAO recommended frequencies.
- **chemnames** Chemical names to search for.
- **qns** Resolved quantum numbers to search for.
- **intensity** CDMS/JPL intensity range.
- **smu2** $S\mu^2$ range in Debye$^2$.
- **loga** $\log(A)$ (Einstein coefficient) range.
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- el  Lower energy state range in Kelvin.
- eu  Upper energy state range in Kelvin.
- rrlinclude  Include RRLs in the result set?
- rrlonly  Include only RRLs in the result set?

Notation is as found in the Splatalogue catalog.

Example:
Search for all lines of the species HOCN and HOCO+ in the 200-300GHz range:

```python
slsearch(outfile="myresults.tbl", freqrange = [200,300],
    species=['HOCN', 'HOCO+'])
```

The task can also return a python dictionary if assigned a variable like:

```python
myLines = slsearch(outfile="myresults.tbl", freqrange = [200,300],
    species=['HOCN', 'HOCO+'])
```

6.23 Convert Exported Splatalogue Catalogs to CASA Tables (splattotable)

In some cases the internal spectral line catalog may not contain the lines in which one is interested. In that case, one can export a catalog from Splatalogue \( http://www.splatalogue.net \) or even create their own "by hand" (be careful to get the format exactly right though!). CASA's task splattotable can then be used to create a CASA table that contains these lines and can be searched:

```python
--------> inp(splattotable)
# splattotable :: Convert a downloaded Splatalogue spectral line list to a casa table.
filenames  =  ["""]  # Files containing Splatalogue lists.
  table  =  ""  # Output table name.
wantreturn =  True  # Do you want the task to return a spectralline tool attached to the results table?
```

A search in Splatalogue will return a catalog that can be saved in a file (look for the "Export" section after the results on the search results page). The exported filename(s) should be entered in the filenames parameter of splattotable. The downloaded files must be in a specific format for this task to succeed. If you use the Splatalogue "Export CASA fields" feature, you should have no difficulties.

6.24 Image Import/Export to FITS

These tasks will allow you to write your CASA image to a FITS file that other packages can read, and to import existing FITS files into CASA as an image.
6.24.1 FITS Image Export (exportfits)

To export your images to fits format use the exportfits task. The inputs are:

```plaintext
# exportfits :: Convert a CASA image to a FITS file
imagename = '' # Name of input CASA image
fitsimage = '' # Name of output image FITS file
velocity = False # Use velocity (rather than frequency) as spectral axis
optical = False # Use the optical (rather than radio) velocity convention
bitpix = -32 # Bits per pixel
minpix = 0 # Minimum pixel value
maxpix = 0 # Maximum pixel value
overwrite = False # Overwrite pre-existing imagename
dropstokes = False # Drop the Stokes axis?
stokeslast = True # Put Stokes axis last in header?
```

The dropstokes or stokeslast parameter may be needed to make the FITS image compatible with an external application.

For example,

```python
exportfits('ngc5921.demo.cleanimg.image','ngc5921.demo.cleanimg.image.fits')
```

6.24.2 FITS Image Import (importfits)

You can also use the importfits task to import a FITS image into CASA image table format. Note, the CASA viewer can read fits images so you don’t need to do this if you just want to look at the image. The inputs for importfits are:

```plaintext
# importfits :: Convert an image FITS file into a CASA image
fitsimage = '' # Name of input image FITS file
imagename = '' # Name of output CASA image
whichrep = 0 # If fits image has multiple
whichhdu = 0 # If its file contains
zeroblanks = True # Set blanked pixels to zero (not NaN)
owrite = False # Overwrite pre-existing imagename
defaultaxes = False # Add the default 4D
defaultaxesvalues = [] # List of values to assign to
```

For example, we can read the above image back in

```python
importfits('ngc5921.demo.cleanimg.image.fits','ngc5921.demo.cleanimage')
```
6.25 Using the CASA Toolkit for Image Analysis

Although this cookbook is aimed at general users employing the tasks, we include here a more detailed description of doing image analysis in the CASA toolkit. This is because there are currently only a few tasks geared towards image analysis, as well as due to the breadth of possible manipulations that the toolkit allows that more sophisticated users will appreciate.

To see a list of the ia methods available, use the CASA help command:

```
CASA <1>: help ia
--------> help(ia)
Help on image object:

class image(__builtin__.object)
 | image object
 | Methods defined here:
 | __init__(...)
 | x.__init__(...) initializes x; see x.__class__.__doc__ for signature
 | __str__(...)
 | x.__str__() <==> str(x)
 | adddegaxes(...)
 | Add degenerate axes of the specified type to the image' :
 | outfile
 | direction = false
 | spectral = false
 | stokes
 | linear = false
 | tabular = false
 | overwrite = false
 | ----------------------------------------
 | addnoise(...) 
 |...
 | unlock(...) 
 | Release any lock on the image' :
 | ____________________________________________
 | ____________________________________________
 | Data and other attributes defined here:
 | __new__ = <built-in method __new__ of type object at 0x55d0f20>
 | T.__new__(S, ...) -> a new object with type S, a subtype of T
```

Inside the Toolkit:
The image analysis tool (ia) is the workhorse here. It appears in the User Reference Manual as the `image` tool. Other relevant tools for analysis and manipulation include `measures` (me), `quanta` (qa) and `coordsys` (cs).
or for a compact listing use <TAB> completion on ia., e.g.

CASA <5>: ia.
Display all 105 possibilities? (y or n)

A common use of the ia tool is to do region statistics on an image. The imhead task has
mode='stats' to do this quickly over the entire image cube. The tool can do this on specific
planes or sub-regions. Here is an example on how to use the ia tool to get on-source and off-source
statistics:

# The variable clnimage points to the clean image
# Pull the max and rms from the clean image
ia.open(clnimage)
on_statistics=ia.statistics()
thistest_immax=on_statistics['max'][0]
oldtest_immax = 1.07732224464
print ' Clean image ON-SRC max should be ',oldtest_immax
print ' Found : Max in image = ',thistest_immax
diff_immax = abs((oldtest_immax-thistest_immax)/oldtest_immax)
print ' Difference (fractional) = ',diff_immax

print ''
# Now do stats in the lower right corner of the image
box = ia.setboxregion([0.75,0.00],[1.00,0.25],frac=True)
off_statistics=ia.statistics(region=box)

thistest_imrms=off_statistics['rms'][0]

oldtest_imrms = 0.0010449
print ' Clean image OFF-SRC rms should be ',oldtest_imrms
print ' Found : rms in image = ',thistest_imrms

diff_imrms = abs((oldtest_imrms-thistest_imrms)/oldtest_imrms)
print ' Difference (fractional) = ',diff_imrms

print ''
print ' Final Clean image Dynamic Range = ',thistest_immax/thistest_imrms
print ''
print ' =============== '

ia.close()

Note: If you don’t close the file with, e.g., ia.close() the file will stay in a 'locked' state. Other processes won’t be able to access the file until the file is properly closed.

6.26 Examples of CASA Image Analysis

The data reduction tutorials on casaguides.nrao.edu provide walkthroughs for high and low frequency, spectral line and polarization analysis techniques.
Chapter 7

Visualization With The CASA Viewer

This chapter describes using the CASA Viewer to display data. The Viewer can be started as a stand-alone executable or by the viewer task inside a CASA shell. It can display both images and Measurement Sets. We are in the process of splitting the task-level access to the Viewer into two tasks: imview for images and msview for measurement sets. These tasks offer improved scriptability, giving command line access to many of the viewer features.

7.1 Starting the Viewer

Figure 7.1: The Viewer Display Panel (left) and the Data Manager (right) panel for a regular image or data cube.

Within the casapy environment, the viewer task can be used to start the CASA Viewer, displaying an image or MS. The inputs are:
CHAPTER 7. VISUALIZATION WITH THE CASA VIEWER

# viewer :: View an image or visibility data set.

infile = "" # (Optional) Name of file to visualize.
displaytype = 'raster' # (Optional) Type of visual rendering
# (raster, contour, vector or marker).
# lel if an lel expression is given
# for infile (advanced).

Examples of starting the CASA Viewer:

CASA <1>: viewer()

CASA <2>: viewer('ngc5921.demo.ms')

CASA <3>: viewer('ngc5921.demo.cleanimg.image')

CASA <4>: viewer('ngc5921.demo.cleanimg.image', 'contour')

CASA <5>: viewer('"ngc5921.demo.cleanimg.image"^2', 'lel')

The first command creates an empty Viewer Display Panel (§ 7.2) and a Load Data window (§ 7.3.1). The second starts the CASA Viewer and loads a Measurement Set. The third example starts the Viewer and opens an image data cube (see Figure 7.1).

Examples four and five make use of the second parameter (displaytype). Example four displays the image as a contour map rather than the default raster map. Example five uses 'Lattice (Image) Expression Language' to display the square of the image data.

Note that the Viewer can open FITS files, CASA image files, Measurement Sets, and saved viewer states. The Viewer determines the type of file being opened automatically.

For additional scripting options when opening the Viewer, see the discussion of the imview and msview tasks at the end of this Chapter (§ 7.7 and 7.8).

7.1.1 Running the CASA Viewer outside casapy

If you have CASA installed, then the CASA Viewer is available as a stand-alone application called casaviewer. From the operating system prompt, the following commands work the same as the casapy task commands given in the previous Section:

    casaviewer &
    casaviewer ms_filename &
    casaviewer image_filename &
    casaviewer image_filename contour &
    casaviewer '"image_filename"^2' lel &
7.2 The Viewer Display Panel

The CASA Viewer consists of a number of graphical user interface (GUI) windows. The main Viewer Display Panel (§7.2) is used for both image and Measurement Set viewing. It is shown in the left panels of Figures 7.1 and 7.2 and appears the same whether an image or Measurement Set is being displayed.

Figure 7.2: The Viewer Display Panel (left) and Data Display Options (right) panels with a Measurement Set open.

At the top of the Viewer Display Panel are drop down menus:

- **Data**
  - **Open** — open the Data Manager window (§7.3).
Register — select and de-select which of the loaded data file(s) should be displayed. The menu expands to the right showing all loaded data sets. Unchecking an image will cause it not to be displayed, but does not close it.

Close — close (unload) the selected data file. The menu expands to the right showing all loaded data.

Adjust Data Display — open the Data Display Options ('Adjust') window (§7.4.1).

Save as... — save/export data to a file

Print — print the displayed image

Save Panel State — to a 'restore' file (xml format)

Restore Panel State — from a restore file

Preferences — manually edit the viewer configuration

Close Panel — close this Viewer Display Panel. If this is the last display panel open, this will exit the Viewer.

Quit Viewer — close all display panels and exit

• Display Panel

New Panel — create a new, empty Viewer Display Panel

Panel Options — open the Viewer Canvas Manager window to edit margins, the number of panels, and the background (§7.4.1.5).

Save Panel State — save the current state of the viewer as a file that can later be reloaded.

Restore Panel State — restore the viewer to a state previously saved as a file.

Print — print displayed image

Close Panel — close this Viewer Display Panel. If this is the last display panel open, this will exit the Viewer.

• Tools

Spectral Profile — Open the Spectral Profile Browser window to look at intensity as a function of frequency for part of an image (§7.4.4.2)

Collapse Image — Open the Collapse/Moments window, which allows you to create new images from a data cube by integrating along the spectral axis (§7.4.6)

Histogram — Open the Histogram inspection window, which allows you to graphically examine the distribution of pixel values in a data cube (§7.4.7)

Fit — Open the two-d fitting window, which can be used to fit Gaussians to two dimensional intensity distributions (§7.4.8).

Interactive Clean — Open a window to look at ongoing interactive clean processes.

• View

Main Toolbar — show/hide the top row of icons (Figure 7.3, §7.2.1).
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– **Mouse Toolbar** — show/hide the second row of mouse-button action selection icons (Figure 7.4 §7.2.2).

– **Animator** — show/hide tapedeck control panel attachment to the main Viewer Display Panel (§7.2.3).

– **Position Tracking** — show/hide the position tracking attachment to the main Viewer Display Panel (§7.2.3).

– **Regions** — show/hide the region manager attachment to the main Viewer Display Panel (§7.2.3).

### 7.2.1 The Main Toolbar

![Main Toolbar](image)

Figure 7.3: The display panel’s **Main Toolbar** appears directly below the menus and contains ‘shortcut’ buttons for most of the frequently-used menu items.

Below the drop down menus is the **Main Toolbar** (Figure 7.3). This top row of icons offers fast access to these menu items:

- **folder** *(Data: Open shortcut)* — open the Data Manager window (§7.3)
- **wrench** *(Data: Adjust shortcut)* — open the Data Display Options ('Adjust') window (§7.4.1)
- **panels** *(Data: Register shortcut)* — select and de-select which of the loaded data file(s) should be displayed. The menu expands to the right showing all loaded data sets. Unchecking an image will cause it not to be displayed, but does not close it.
- **delete** *(Data: Close shortcut)* — close (unload) the selected data file. The menu expands to the right showing all loaded data.
- **save data** *(Data: Save as)* — save the current data to a file.
- **new panel** *(Display Panel: New Panel)* — create a new, empty Viewer Display Panel
- **panel wrench** *(Display Panel: Panel Options)* — open the Viewer Canvas Manager window to edit margins, the number of panels, and the background (§7.4.1.5).
- **save panel** *(Display Panel: Save Panel State)* — save panel state to a ‘restore’ file
- **restore panel** *(Display Panel: Restore Panel State)* — restore panel state from a restore file
- **spectral profile** *(Tools: Spectral Profile)* — Open the Spectral Profile Browser window to look at intensity as a function of frequency for part of an image (§7.4.4.2)
• **collapse/moments** *(Tools: Collapse Image)* — Open the Collapse/Moments window, which allows you to create new images from a data cube by integrating along the spectral axis (**7.4.6**).

• **histogram** *(Tools: Histogram)* — Open the Histogram inspection window, which allows you to graphically examine the distribution of pixel values in a data cube (**7.4.7**).

• **fitting** *(Tools: Fit)* — Open the two-d fitting window, which can be used to fit Gaussians to two dimensional intensity distributions (**7.4.8**).

• **print** *(Display Panel: Print)* — print the current display

• **magnifier box** — zoom out all the way

• **magnifier plus** — zoom in (by a factor of 2)

• **magnifier minus** — zoom out (by a factor of 2)

### 7.2.2 The Mouse Toolbar

![Mouse Tool Bar](image)

Figure 7.4: The ’Mouse Tool’ Bar allows you to assign how mouse buttons behave in the image display area. Initially, zooming, color adjustment, and rectangular regions are assigned to the left, middle and right mouse buttons. Click on a tool with a mouse button to assign that tool to that mouse button.

Below the Main Toolbar are eleven **Mouse Tool** buttons (Figure [7.4]). These allow you to assign what behavior the three mouse buttons have when clicked in the display area. Clicking a mouse tool icon will [re-]assign the mouse button that was clicked to that tool. Black and white squares beneath the icons show which mouse button is currently assigned to which tool.

The mouse tools available from the toolbar are:

(Notes that the ’escape’ key can be used to cancel any mouse tool operation that was begun but not completed, and to erase a region, point, or other tool showing in the display area.)

- **Zooming (magnifying glass icon):** To zoom into a selected area, press the Zoom tool’s mouse button (the left button by default) on one corner of the desired rectangle and drag to the desired opposite corner. Once the button is released, the zoom rectangle can still be moved or resized by dragging. To complete the zoom, double-click inside the selected rectangle. If you instead double-clicking outside the rectangle, you will zoom out.

- **Panning (hand icon):** Press the tool’s mouse button on a point you wish to move, drag it to the position where you want it moved, and release. Note: The arrow keys, Page Up, Page Down, Home and End keys can also be used to pan through your data any time you are zoomed in. (Click on the main display area first, to be sure the keyboard is ’focused’ there).
• **Stretch-shift colormap fiddling (crossed arrows):** This is usually the handiest color adjustment; it is assigned to the middle mouse button by default. Note that you can also adjust the color table quantitatively inside the Data Display Options window (§7.4.1).

• **Brightness-contrast colormap fiddling (light/dark sun):** Another tool to adjust the color stretch.

• **Positioning (plus):** This tool can place a point marker on the display to select a position. It is used to flag Measurement Set data or to select an image position for spectral profiles. Click on the desired position with the tool’s mouse button to place the point; once placed you can drag it to other locations. You can also place multiple points on the display (e.g. for different spectral profile positions) – remove them by hovering over and hitting ESC. Double-click is not needed for this tool. See §7.4.2 for more detail.

• **Rectangle, Ellipse and Polygon region drawing:** The rectangle region tool is assigned to the right mouse button by default. As with the zoom tool, a rectangle region is generated by dragging with the assigned mouse button; the selection is confirmed by double-clicking within the rectangle. An ellipse regions is created by dragging with the assigned mouse button. In addition to the elliptical region, also its surrounding rectangle is shown on the display. The selection is confirmed by double-clicking within the ellipse. Polygon regions are created by clicking the assigned mouse button at the desired vertices, clicking the final location twice to finish. Once created, a polygon can be moved by dragging from inside, or reshaped by dragging the handles at the vertices. See §7.4.3 for the uses of this tool.

• **Polyline drawing:** A polyline can be created by selecting this tool. It is manipulated similarly to the polygon region tool: create segments by clicking at the desired positions and then double-click to finish the line. [Uses for this tool are still to be implemented].

• **Distance tool:** After selecting the distance tool by assigning any mouse button to it, distances on the image can conveniently be measured by dragging the mouse with the assigned button pressed. The tool measures the distances along the world coordinate axes and along the hypotenuse. If the units in both axes are [deg], the distances are displayed in [arcsec].

• **Position-Velocity Diagram:** Use this mouse tool to drag out a position axis that can be used to generate a position velocity diagram in a new viewer panel from the region manager dock.

### 7.2.3 The Display Area

The main **Display Area** lies below the toolbars. This area shows the image or Measurement Set currently loaded. Clicking the mouse inside the display area allows region or position selection according to the settings in the mouse toolbar.

The Display Area may have up to three attached panels: the **Animator** panel, the **Position Tracking** panel, and the **Regions** panel. These may be displayed or hidden from the ”View” dropdown menu in the main Viewer Display Panel. If one of these is missing from your viewer,
check that it is checked "on" in that menu. The panels can also be turned off by clicked the "X" in the top right corner, in which case you will need to use the View menu to get them back.

By default, the three panels appear attached to the main Viewer Display Panel on the right side of the image. They may be dragged to new positions. Each of the three panels can be attached to the left, top, right, or bottom of the main Viewer Display Panel or they can be entirely undocked and left as free-floating panels.

NOTE: Depending on your window manager, windows without focus, including detached panels and tools like the Spectral Profile Browser may sometimes display odd behavior. As a general rule, giving the window focus by clicking on it will correct the issue. If you seem to "lose" a detached panel (like an Animator Panel), then click in the main window to get it back.

NOTE: With all three panels turned on (and especially with several images loaded), the main display panel can sometimes shrink to very small sizes as the panels grow. Try detaching the panels to get the main display panel back to a useful size.

A restart of the viewer will display all docks in the state of a previous viewer session, given that it was closed normally. In addition, the viewer docking can be changed under “Preferences” In the toolbar (Mac OS under the “CASA Viewer” tab on the toolbar, Linux: “Data”). Fig. 7.5 shows an example. Each item can be changed and the input box will only allow accepted input formats. A complete restart is required to apply the changes.

Figure 7.5: "Preferences" dialog to manually change the docking and size of the viewer panel.
7.2.3.1 The Animator Panel

The Animator Panel allows you to scroll through the channels of a data cube and to rotate among loaded images. The main features of the panel are the two “tape decks,” one labeled ”Channels” and one labeled ”Images” (note that you will only see the Images tape deck when multiple images are loaded.

The Channels tape deck scrolls between planes of an individual image. By default, the channel tape deck scrolls among frequency planes when R.A. and Declination are the displayed axes (in this case, frequency is the ”z axis”). From outside to inside, the buttons cause the display to jump all the way to the beginning/end of the z axis, cause the viewer to step one plane forward or backward along the z axis, or start a movie. The limits on the z axis can be set manually using the windows at the end of the scroll bar. The scroll bar can also be dragged or the user can jump the display to a manually entered plane by entering the plane into the text box.

When you have multiple images loaded, the Images tape deck cycles through which is image is being displayed. In the movie mode, it allows you continuously click between images. Functionally, the image tape deck works similarly to the channels tape deck, with the ability to step, jump, or continuously scroll through images.

NOTE: The check boxes next to the channel and images tabs enable or disable those panels. This doesn’t have much effect when the display has only a single panel, but with multiple panels (i.e., several maps at once in the main window) it changes the nature of the display. If the ”Images” box is checked then interleaved maps from different cubes are display. Otherwise a series of maps from a single cube are shown.

7.2.3.2 The Position Tracking Panel

The Position Tracking panel (below the images in Fig 7.1) shows the intensity, position (e.g., RA and Dec), Stokes, frequency (or velocity), and pixel location for the point currently under the cursor. A separate box appears for each registered image or Measurement Set and you can see the tracking information for each. Tracking can be ’frozen’ (and unfrozen again) by hitting the space bar when the viewer’s focus is on the main display area (to be sure that this is case first click on the main display area).
7.2.3.3 The Region Manager Panel

The Region Manager panel becomes active when regions are created. It has a large amount of functionality, from display region statistics and histograms to creating position-velocity cuts. We discuss these in §7.4.3. Like the Animator and Position Tracking panel, the Region Manager Panel can be moved relative to the main viewer display panel or entirely undocked.

7.2.4 Saving and Restoring the Display Panel State

You can save the display panel’s current state — meaning the panel settings and the data on display — or load a saved panel state from disk. To save the display panel state, select Save Panel State from the Display Panel drop-down menu or click the ”Save Display Panel State to File” icon on the main toolbar (an arrow pointing from a picture to a page, see Figure 7.3). It is advisable but not required to retain the file’s ’.rstr’ (”Restore”) extension.

You can restore the display panel to the saved state by loading the saved state from the Data Manager Panel, by selecting Restore Panel State from the Display Panel drop down menu, or by clicking the ”Restore Display Panel State” icon (just to the right of the ”Save Display Panel State” icon).

It is possible to restore panel states viewing Measurement Sets or image and panel states that have multiple layers, such as contour plots over raster images. You can also save LEL displays. You can also the save or restore the panel state with no data loaded, which is a convenient way to restore preferred initial settings such as overall panel size.

Data Locations: The viewer is fairly forgiving regarding data location when restore a saved panel state. It will find files located:

- in the original location recorded in the restore file
- in the current working directory (where you started the viewer)
- in the restore file’s directory
- in the original location relative to the restore file
This means that you can generally restore a saved panel state if you move that file together with data files. The exception to this rule is that the process is less forgiving if you save the display of an LEL expression. In this case the files must be in the locations specified in the original LEL expression. If a data file is not found, restore will attempt to proceed but results may not be ideal.

**Manually Editing Saved Display Panel States:** The saved "Restore" files are in ascii (xml) format, and manual edits are possible. However, these files are long and complex. Use caution, and back up restore files before editing. If you make a mistake, the viewer may not even recognize the file as a restore file. It is easier and safer to make changes on the display panel and then save the display panel state again.

### 7.3 The Data Manager Panel — Saving and Loading Data

![Data Manager Panel](image)

Figure 7.8: The load tab of the Data Manager panel. This appears if you open the viewer without any infile specified, if you use select Open from the Data drop down menu, or click the Open (Folder) icon. You can access the save image or save region tabs from this view or by selecting Save as... from the Data drop down menu. The load tab shows all files in the current directory that can be loaded into the viewer — images, MS, CASA region files, and Display Panel State files.

The Data Manager Panel is used to interactively load and save images, Measurement Sets, Display Panel States, and regions. An example of the loading tab in this panel is shown in Figure 7.8. This panel appears automatically if you open the viewer without specifying an input file or it can be accessed through the Data:Open menu or Open icon of the Viewer Display Panel.
7.3.1 Loading Data

The load tab of the Data Manager Panel allows you to interactively choose images or Measurement Sets to load into the viewer. The load tab automatically shows you the available images, Measurement Sets, and Display Panel States in the current directory that can be opened by the viewer. When you highlight an image in this view, the tab shows a brief summary of the image: pixel shape, extent of the image on the sky and in frequency/velocity, and restoring beam (if available).

Selecting a file will bring up information about that file in the panel on the right of the Data Manager Panel provide options for how to display the data. Images can be displayed as:

1. raster image
2. contour map
3. vector map
4. marker map

These options are each discussed in §7.4

slice: a subselection of a data cube can be loaded, the start and end pixel in each spatial, polarization, and spectral dimension can be selected.

LEL: Instead of only loading an image from disk, you may ask the viewer to evaluate a 'Lattice Expression Language' (LEL) expression (§6.1.4). This can be entered in the box provided after you click the "LEL" box. The images used in the LEL expression should have the same coordinates and extents.

Measurement Sets: A Measurement Set can only be displayed as a raster. For measurement sets, the load tab offers options for data selection. This will reduce loading and processing times for visibility flagging.

Regridding Images on Load: Optionally, you may regrid the velocity axis of an image on load to match the current coordinates grid in the Display Panel. In this case, the viewer will interpolate (using the selected interpolation scheme) the cube on disk to share the same velocity gridding as the loaded coordinates. This can be used, e.g., to overlay contour maps of different spectral lines or to make synchronized movies of multiple cubes. Note that the regridding depends on the rest frequency in the image, which is used to calculate the velocities used in regridding.

7.3.2 Registered vs. Open Datasets

When you load data as described above, it is first opened, and then registered on all existing Display Panels.

An open dataset has been prepared in memory from disk. All open datasets will have a tab in the Data Display Options window, whether currently registered or not.
When a data set is registered to a Display Panel its coordinates are aligned to the master coordinate image in the panel and it is ready for drawing. If multiple Display Panels are open then a data set may be registered on one Display Panel and not on another. Only those data sets registered on a particular Display Panel show up in its Position Tracking panel.

Why Register More Than One Image? It is useful to have more than one image registered on a panel if you are displaying a contour image over a raster image (§7.4.2.1), to 'blink' between images (see Animator in §7.2), or to compare images using the position tracking panel.

Unregistering Images: A data set can be registered or unregistered using the Image Manager in the Data drop down menu or the Image Manager icon (third from left). It will open the Image Manager window and the checkboxes can be used to register or unregister an image.

Closing vs. Unregistering: You can close a data set that is no longer needed using the Close option in the Data drop-down menu, the ”Close” icon (fourth from left), or right mouse button “Close” selection in the Image Manager (§7.3.3).

If you close a dataset, you must reload it from disk (or recreate it if it’s an LEL expression, regridded image, moment or something similar) to see it again. If you unregister a dataset, it will draw immediately if you re-register it, with its options as you have previously set them. In general, close unneeded datasets but unregister those that you intend to use again.

### 7.3.3 Image Manager

The Image Manager is used to define the master coordinate image, the sequence of images (e.g. for blinking), to register and unregister images, close images, change between raster, contour, vector, and marker displays, and to modify the properties of images. The panel can be invoked from the “Manage Images” tool, the third icon from the left (two overlapping squares).

An example is shown in Fig. 7.9. In this case, four images are loaded into the viewer. The sequence of images can be changed by dragging and dropping the images to new positions in the stack. The letter to the left indicates whether the image is a Raster, Contour, Vector, or Marker image. MC marks the coordinate master, in this case the second image. The checkboxes are to change the registration statuses. The Coordinate Master image can be defined by a right mouse click, and selection the corresponding option. The right mouse menu button also offers options for quick changes between contour and raster images and to close an image.

The Options button will open a drop down box (as shown in Fig. 7.9 for the first image), in which one can again change between image type, change to a different rest frequency, or open the “Display Options” panel for that specific image with all the adjustment options explained in (§7.4.1 or §7.4.2).

### 7.3.4 Saving Data or Regions

The viewer can create new images by carrying out velocity regridding, evaluating an LEL expression, or collapsing a data cube. You can save these images to disk using the Data Manager Panel. Select
Save as under the Data drop-down menu or click the Save as (disk) icon to bring up the Data Manager Panel set to the save tabs. This tab is shown in Figure 7.10.

From the Save Image tab of the Data Manager Panel, you can export images from the viewer to either a CASA image or FITS file on disk. Select the desired file name and click "save."
Manager also allows you to save your current regions to a file, either in the CASA or ds9 format. The left part lists all images that can be exported to disk. To save an image to a file, the user can either enter the new filename in the box labeled 'output name:' followed by the save-button (alternatively the 'Enter'-key), or choose a file name from the right hand side.

7.4 Viewing Images

There are several options for viewing an image. These are seen at the right of the Load Data - Viewer panel described in §7.3.1 and shown in Figure 7.1 after selecting an image. They are:

- **raster image** — a greyscale or color image,
- **contour map** — contours of intensity as a line plot,
- **vector map** — vectors (as in polarization) as a line plot,
- **marker map** — a line plot with symbols to mark positions.

The **raster image** is the default image display, and is what you get if you invoke the Viewer with an image file and no other options. In this case, you will need to use the Open menu to bring up the Load Data panel to choose a different display.

7.4.1 Viewing a Raster Map

A raster map of an image shows pixel intensities in a two-dimensional cross-section of gridded data with colors selected a colormap according to a scaling that can be specified by the user.

Starting the casaviewer with an image as a raster map will look something like the example in Figure 7.1.

Once loaded, the data display can be adjusted by the user through the Data Display Options panel, which appears when you choose the Data:Adjust menu or use the wrench icon from the Main Toolbar.

The Data Display Options window is shown in the right panel of Figure 7.1. It consists of a tab for each image or MS loaded, under which are a cascading series of expandable categories. For an image, these are:

- display axes
- hidden axes
- basic settings
- position tracking
- axis labels
• axis label properties
• beam ellipse
• color wedge

The **basic settings** category is expanded by default. To expand a category to show its options, click on it with the left mouse button.

### 7.4.1.1 Data Display Options — display and hidden axes

In this category the physical axes (i.e. Right Ascension, Declination, Velocity, Stokes) to be displayed can be selected and assigned to the x, y, and z axes of the display. The z axis will be the axis scrolled across by the channel bar in the Animator Panel.

If your image has a fourth axis (typically Stokes), then one of the axes will need to be hidden and not used in viewing. Which axis is hidden can be controlled by a slider within the **hidden axes** drop-down.

### 7.4.1.2 Data Display Options — basic settings

This roll-up is open by default showing some commonly-used parameters that alter the way the image is displayed. The most frequently used of these change how the intensity value of a pixel maps to a color on the screen. An example of this part of the panel is shown in Figure 7.11.

Figure 7.11: The **basic settings** category of the **Data Display Options** panel and the interactive tool for setting the mapping from intensity to color.

The options available are:
• basic settings: aspect ratio

This option controls the horizontal-vertical size ratio of data pixels on screen. fixed world (the default) means that the aspect ratio of the pixels is set according to the coordinate system of the image (i.e., true to the projected sky). fixed lattice means that data pixels will always be square on the screen. Selecting flexible allows the map to stretch independently in each direction to fill as much of the display area as possible.

• basic settings: pixel treatment

This option controls the precise alignment of the edge of the current 'zoom window' with the data lattice. edge (the default) means that whole data pixels are always drawn, even on the edges of the display. For most purposes, edge is recommended. center means that data pixels on the edge of the display are drawn only from their centers inwards. (Note that a data pixel’s center is considered its 'definitive' position, and corresponds to a whole number in 'data pixel' or 'lattice' coordinates).

• basic settings: resampling mode

This setting controls how the data are resampled to the resolution of the screen. nearest (the default) means that screen pixels are colored according to the intensity of the nearest data point, so that each data pixel is shown in a single color. bilinear applies a bilinear interpolation between data pixels to produce smoother looking images when data pixels are large on the screen. bicubic applies an even higher-order (and somewhat slower) interpolation.

• basic settings: data range

You can use the entry box provided to set the minimum and maximum data values mapped to the available range of colors as a list \([\text{min}, \text{max}]\). For very high dynamic range images, you will probably want to enter a \text{max} less than the data maximum in order to see detail in lower brightness-level pixels.

NOTE: By default you edit the scaling of a single image at once and can click between the tabs at the top of the Data Display Options window to manipulate different windows. By checking the Global Color Settings box at the bottom of this window, you will manipulate the scaling for all images at once. This can be very useful, for example, when attempting detailed comparison multiple reductions of the same data.

• basic settings: scaling power cycles

This option allows logarithmic scaling of data values to colormap cells, which can be very helpful in the case of very high dynamic range.

The color for a data value is determined as follows:

1. The value is clipped to lie within the data range \([\text{min}, \text{max}]\) specified above.
2. This clipped value is mapped to a new value depending on the selected scaling power cycles in the following way:
   - If the scaling power cycles is set to 0 (the default), the program considers a linear range from \([\text{min}, \text{max}]\) and scales this directly onto the set of available colors.
– For negative scaling values, the data value (after clipping on \([\text{min}, \text{max}]\)) is scaled linearly to lie between 0 and \(10^p\) (where \(p\) is the value chosen) and then program takes the logarithm of this values, yielding a value in the range 1 to \(p\). That value is scaled linearly to the set of available colors. Thus the data is treated as if it had \(p\) decades of range, with an equal number of colors assigned to each decade.

– For positive scaling values, the data value (after clipping on \([\text{min}, \text{max}]\)) is scaled linearly to lie between 0 and \(p\) (where \(p\) is the value chosen) and 10 is raised to this power, yielding a value in the range 1 to \(10^p\). That value is scaled linearly to the set of available colors.

3. The color corresponding to a number in final range is determined by the selected colormap and its ‘fiddling’ (shift/slope) and brightness/contrast settings (see Mouse Toolbar, above). Adding a color wedge to your image can help clarify the effect of the various color controls.

See Figure 7.12 for sample curves.

![Figure 7.12: Example curves for scaling power cycles.](image)

In practice, you will often manipulate the data range bringing the max down in high dynamic range images, raising the minimum to the near the noise level when using non-zero scaling cycles. It is also common to use negative power cycles when considering high dynamic range images — this lets you bring out the faint features around the bright peaks.

- **basic settings:** colormap

You can select from a variety of colormaps here. *Hot Metal, Rainbow* and *Greyscale* colormaps are the ones most commonly used.
7.4.1.3 Graphical Specification of the Intensity Scale

A histogram icon next to the data range in the Data Display opens the Image Color Mapping window, which allows visualization and graphical manipulation of the mapping of intensity to color. The window at the left shows a histogram of the data with a gray range showing the data range. You can use this window to select the data range graphically (with the mouse), manually (by typing into the empty windows), or as a percentile of the data. On the right, you can select the scaling power cycles and see a visualization of the transfer function mapping intensity ($x$-axis) to color ($y$-axis).

The functionality here follows the other histogram tools, with the Display tab used to change the histogram plotting parameters. It will often be useful to use a logarithmic scaling of the $y$-axis and filled histograms when manipulating the color table.

7.4.1.4 Data Display Options — other settings

Many of the other settings on the Data Options panel for raster images are self-explanatory such as those which affect beam ellipse drawing (only available if your image provides beam data), or the form of the axis labeling and position tracking information. You can also give your image a color wedge, a key to the current mapping from data values to colors.

7.4.1.5 Viewer Canvas Manager — Panels, Margins, and Backgrounds

The display area can also be manipulated from the Viewer Canvas Manager window. Use the wrench icon with a ‘P’ (or the ‘Display Panel’ menu) to show this window, which allows you to manipulate the infrastructure of the main display panel. You can set:

- **Margins** - specify the spacing for the left, right, top, and bottom margins
- **Number of panels** - specify the number of panels in $x$ and $y$ and the spacing between those panels.
- **Background Color** - white or black (more choices to come)

Figure 7.13 illustrates a multi-panel display along with the Viewer Canvas Manager settings which created it.

7.4.2 Viewing a Contour Map

Viewing a contour image is similar to viewing a raster map. A contour map shows lines of equal data value for the selected plane of gridded data (Figure 7.14). Contour maps are particularly useful for overlaying on raster images so that two different measurements of the same part of the sky can be shown simultaneously (§7.4.2.1).

Several basic settings options control the contour levels used:
The contours themselves are specified by a list in the box Relative Contour Levels. These are defined relative to the two other parameters:

- The Base Contour Level sets the zero level for the relative contour list corresponds to in units of intensity in the image.
- The Unit Contour Level sets what 1 in the relative contour list corresponds to in units of intensity in the image.

Additionally, you have the option to manipulate the thickness and color of the image and to have either positive or negative contours appear dashed.

For example, the following settings:

Relative Contour Levels = [0.2, 0.4, 0.6, 0.8]
Base Contour Level = 0.0
Unit Contour Level = <image max>

would map the maximum of the image to 1 in the relative contour levels and the base contour level to zero. So the contours will show 20%, 40%, 60%, and 80% of the peak.
Figure 7.14: The **Viewer Display Panel** (left) and **Data Display Options** panel (right) after choosing **contour map** from the **Load Data** panel. The image shown is for channel 11 of the NGC5921 cube, selected using the **Animator** tape deck, and zoomed in using the tool bar icon. Note the different options in the open **basic settings** category of the **Data Display Options** panel (as compared to **raster image** in Figure 7.1).

Another approach is to set the unit contour to 1, so that the contours are given in intensity units (usually Jy/beam). So this setup:

Relative Contour Levels = \([0.010, 0.0.020, 0.040, 0.080, 0.160, 0.320]\)
Base Contour Level = 0.0
Unit Contour Level = 1.0

would create contours starting at 10 mJy/beam and doubling every contour.

Another useful approach is to set contours in units of the rms noise level of the image, which can be worked out from a signal free region. Then a setup like this:

Relative Contour Levels = \([-3,3,5,10,15,20]\)
Base Contour Level = 0.0
Unit Contour Level = \(<\text{image rms}>\)

Would indicate significance in the image. The first two contours show emission at ± 3-sigma and so on.
You can get the image rms using the \texttt{imstat} task (§6.9) or using the Viewer statistics tool on a region of the image (§7.4.3.3).

Not all images are of intensity, for example a moment-1 image (§6.7) has units of velocity. In this case, absolute contours (like the last two examples) will work fine, but by default the Viewer will set fractional contours but referred to the min and max of the image:

\begin{itemize}
\item Relative Contour Levels = [0.2, 0.4, 0.6, 0.8]
\item Base Contour Level = <image min>
\item Unit Contour Level = <image max>
\end{itemize}

Here we have contours spaced evenly from min to max, and this is what you get by default if you load a non-intensity image (like the moment-1 image). See Figure 7.15 for an example of this.

### 7.4.2.1 Overlay Contours on a Raster Map

Contours of either a second data set or the same data set can be used for comparison or to enhance visualization of the data. The Data Options Panel will have multiple tabs (switch between them at the top of the window) that allow you to adjust each overlay individually.

\textbf{NOTE:} \textit{axis labeling} is controlled by the \textit{first-registered} image overlay that has labeling turned on (whether raster or contour), so make label adjustments within that tab.

To add a Contour overlay, open the \textbf{Load Data} panel (Use the Data menu or click on the folder icon), select the data set and click on \textit{contour map}. See Figure 7.15 for an example using NGC5921.

### 7.4.3 Regions and the Region Manager

CASA regions are following the CASA \texttt{crtf} standard as described in §D. CASA regions can be used in all applications, including \texttt{clean} and image analysis tasks (§6).

\textbf{NOTE:} A leading 'ann' (short for annotation) to a region definition indicates that it is for visual overlay purposes only.

\textbf{NOTE:} Whereas the region format is supported by all the data processing tasks, some aspects of the \texttt{viewer} implementation are still limited to rectangles, ellipses, and some markers. Full support for all region types is progressing with each CASA release.

Once one or more regions are created, the Region Manager Panel becomes active (see Figure 7.16). Like the Position Tracking and Animator Panels, this can be docked or detached from the main viewer display. It contains several tabs that can be used to adjust, analyze, and save or load regions.

\textbf{NOTE:} Moving the mouse into a region will bring it into focus for the Spectral Profile or Histogram tools.
Figure 7.15: The Viewer Display Panel (left) and Data Display Options panel (right) after overlaying a Contour Map of velocity on a Raster Image of intensity. The image shown is for the moments of the NGC5921 cube, zoomed in using the tool bar icon. The tab for the contour plot is open in the Data Display Options panel.

Figure 7.16: The Region Manager Panel, which becomes active once at least one region is created. Cycle through available regions using the slider bar at the bottom and use the various tabs to adjust, analyze, load, and save regions.
7.4.3.1 Region Creation, Selection, and Deletion

Within the display area, you can draw regions or select positions using the mouse. Regions can be created with the buttons marked as ‘R’ in the mouse tool bar (§7.2 §7.4.3.2). The viewer currently supports creation of rectangles, ellipses, polygons, and the point. As usual, a mouse button can be assigned to each button as indicated by the small black square in each button (marking the left, middle, or right mouse button §7.2 §7.4.3.2). An example is shown in Fig. 7.17.

Regions can be selected by SHIFT+click, de-selected by pressing SHIFT+click again. The bottom of the Region Manager Panel features a slider to switch between regions in the image. Regions can be removed by hovering over and pressing ESC or by pressing the buttons to the right side of the slider where the first button deletes all regions and the far right button deletes the region that is currently displayed in the panel.

Figure 7.17: Selecting an image region (done with SHIFT+click). The region can be resized by dragging the handles or deleted by hitting ESCAPE.

Once regions are selected, they will feature little, skeletal squares in the corners of their boundary boxes. Selected regions can be moved by dragging with the mouse button and manually resize by grabbing the corners as handles. If more than one region is selected, all selected regions move together.

The Rectangle Region drawing tool currently enables the full functionality of the various Region Manager tabs (see below) as well as:

- Region statistics reporting for images via double clicking (also shown in the stats tab of the Region Manager),
• Defining a region to be averaged for the spectral profile tool (accessed via the Tools:Spectral Profile drop down menu or "Open the Spectrum Profiler" icon),

• Flagging of Measurement Sets. Note that the Rectangle Region tool’s mouse button must also be double-clicked to confirm an MS flagging edit.

• Selecting Clean regions interactively (§ 5.3.5)

The Polygon Region and Ellipse Region drawing have the same uses, except that polygon region flagging of a Measurement Set is not supported.

7.4.3.2 Region Positioning

![Region Manager](Image)

Figure 7.18: The positioning tab in the Region Manager. Use it to manually adjust the location, width, and display style of the selected region.

With at least one region drawn, the region manager becomes active. Using the Properties tab, one can manually adjust the position, annotation, and display style of the region. The frames boxes set which planes of the image cube the region persists through (regions can have a depth associated with them and will only appear in the frames listed in this range). One can manually adjust the width and height and the center of the box in the chosen units. The 'selection' check box is an alternative way to the SHIFT+click to select a region. The 'annotation' checkbox will place the 'ann' string in front of the region ascii output – annotation regions are not be used for processing in, e.g. data analysis tasks. In the line and text tabs, one can set the style with which the region is displayed, the associated text, and the position and style of that text.

NOTE: Updating the position of a region will update the spectral profile shown if the Spectral Profile tool is open and the histogram if the Histogram tool is open. The views are linked. Dragging a region or adjusting it manually with the Position tab is a good way to explore an image.
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7.4.3.3 Region Statistics

One of the most useful features of defining a region is the ability to extract statistics characterizing
the intensity distribution inside the region. You can see these in the Statistics tab of the Region Manager Panel (see Figure 7.19). This displays statistics for the current region in the current plane of the current image. When more than a single region is drawn, you can select them one by one and the region panel will update the statistics to reflect the currently selected region. All values are updated on the fly when the region is dragged across the image.

A similar functionality can be achieved by double clicking inside of a region. This will send statistics information for this region in all registered images to the terminal, looking something like this:

```
(image) image
Stokes  Velocity  Frame  Doppler  Frequency
I -2.99447e+11km/s  LSRK  RADIO  3.63499e+10
BrightnessUnit  BeamArea  Npts  Sum  Flux
Jy/beam  36.2521  27547  1.087686e-01  3.000336e-03
Mean  Rms  Std dev  Minimum  Maximum
3.948473e-06  3.723835e-04  3.723693e-04  -1.045624e-03  9.968892e-03
```

This is an easy way to copy and paste the statistical data to a program outside of CASA for further use.

Taking the RMS of the signal-free portion of an image or cube is a good way to estimate the noise. Contrasting this number with the maximum of the image gives an estimate of the dynamic range of the image. The FluxDensity measurement gives a way to use the viewer to do very basic photometry.
7.4.3.4 Saving and Loading Regions

The File tab in the Region Manager allows one to save or load selected regions, either individually or en masse. You can choose between CASA and ds9 region format. The default is a CASA region file (saved with a *.crtf suffix, see §7.3). The DS9 format does not offer the full flexibility and cannot capture stokes and spectral axes. DS9 regions will only be usable as annotations in the viewer, they cannot be used for data processing in other CASA tasks. When saving regions, one can choose to save only the current region, all regions that were selected with SHIFT+click, or all regions that are visible on the screen.

NOTE: The load functionality for this tab will only become available once at least one region exists. To load a region when no regions exist, use the Data Manager window (§7.3).

7.4.3.5 The Region Fit

NOTE: This functionality is still under development. Its robustness and functionality will be improved in future version of CASA.

The Viewer can attempt to fit a two dimensional Gaussian to the emission distribution inside the currently selected region. To attempt the fit, go to the Fit tab of the Region Manager and click the gaussfit button in the bottom left of the panel. You can choose whether or not to fit a sky level (e.g., to account for a finite background, either astronomical, sky, or instrumental). After fitting the distribution, the Fit panel shows the results of the fit, the center, major and minor axis, and position angle of the Gaussian fit in pixels (I) and in world coordinates (W, RA and Dec). The detailed results of the fit will also appear in the terminal window, including a flag showing whether the fit converged.
7.4.3.6 The Region Histogram

![Figure 7.21: The histogram tab in the Region Manager. Right click to zoom. Hit SHIFT + Right Click to adjust the details of the histogram display.](image)

The Viewer will automatically derive a histogram of the pixel values inside the selected region. This can be viewed using the **Histogram** tab of the Region Manager Panel. This is a pared down version of the full Histogram Tool. You can manipulate the details of the histogram plot by clicking:

1. Use the **Right Click** to zoom - either to the full range, a selected percentile, or a range that you have graphically selected by dragging the mouse (may still be under development).

2. Hit **SHIFT + Right Click** to open the histogram options. This lets you toggle between a logarithmic and linear y-axis, choose between a line, outline, or filled histogram, and adjust the number of bins.

The histogram will update as you change the plane of the cube or shift between region.

7.4.4 The Spectral Profile Tool

The **Spectral Profile Tool** allows you examine the intensity as a function frequency or velocity. To start a new Spectral Profile window, click the **Spectral Profile** option from the **Tools** drop-down menu or click the "Spectral Profile" (red line graph) icon from the Main Toolbar (see Fig. 7.3). A new Spectral Profile window will appear.

**NOTE:** Make Sure That You Use the Radio Version! This section describes the "Radio" version of the profiler. To be sure that you have the radio version of the tool selected (this may not be
the default), click on the preferences icon (the gear fourth from the left) and make sure that the "Optical" option is not checked.

The Spectral Profile Tool consists of a toolbar (§7.4.4.1), a main display area (§7.4.4.2), and two associated tabs: Spectral-Line Fitting (§7.4.4.3) and Line Overlays (§7.4.4.4).

Interaction With the Main Display Panel: For the Spectral Profile tool to work, a region or point must be specified in the main Viewer Display window. Use the mouse tools to specify a point, rectangle, ellipse, or polygon region. Alternatively, load a region file. The Spectral Profile tool will show a spectrum extracted from the region most recently highlighted by the mouse in the main Viewer Display Panel. The method of extraction can be specified by the user using a drop down menu below the spectrum in the Spectral Profile window; the method of extraction is mean by default).

The Spectral Profile tool can also feed back to the Main Display Panel. By holding CTRL and right clicking in the spectrum, you will cause the Main Display Panel to jump to display the frequency channel corresponding to the spectral (x) coordinate of the region clicked in the Spectral Profile tool. Holding CTRL and dragging out a spectral range while holding the right mouse button will queue a movie scrolling through images across that spectral range. You can achieve the same effect with the two-ended-arrow icon towards the right of the toolbar in the Spectral Profile window.

In both tabs, it will be useful to do select regions of frequency or velocity. You can do this with the parallel lines-and-arrow icon (see below) or by holding shift, left clicking, and dragging out the
range of interest. A shaded gray region should appear indicating your selection.

7.4.4.1 Spectral Profile Toolbar

Figure 7.23: The toolbar for the Spectral Profile tool allows the user to save the spectrum, print or save the tool as an image, edit preferences (general, tool, legend), spectral smoothing, pan or zoom around the spectrum, select a range of interest, jump to a channel, or add a label.

Figure 7.23 shows the toolbar from the top portion of the Spectral Profile window. From left to right, the icons allow the user to:

- (disk) export the current profile to a FITS or ASCII file
- (printer) print the main window to a hard copy
- (writing desk) save the panel as an image (PNG, JPG, PDF, etc.)
- (gear) set plot preferences
- (color wheel) set color preferences for the plot
- (signpost) set legend preferences
- (triangle) set the spectral smoothing method and kernel width
- (arrows) pan the spectrum in the indicated direction  
  NOTE: The arrow keys also allow one to pan using the keyboard.
- (magnifying glass) zoom to the default zoom, in, and out  
  NOTE: the +/- keys allow one to zoom with the keyboard.
- (parallel lines+arrows) drag out a range of interest in the spectrum, for use with fitting or line overlays.
- (double-ended arrow) jump to a channel in the main viewer (single click) or define a range over which to play a movie in the viewer (with a drag).  
  NOTE: You can also jump to a channel with CTRL+Right Click and queue a movie by holding CTRL and dragging out a range while holding the right mouse button.
- notepad and pencil Add or edit a label on the plot. Click this icon to enter a mode where you can drag out a box to create a new annotation box or drag the corners of an existing one to resize it. You can edit the contents, color, and font of an existing annotation by right clicking on it and selecting "Edit Annotation" in the main Spectral Profile window.
Figure 7.24: Preferences options in the Spectral Profile Tool. From the toolbar, one can access dialogs to set overall viewer preferences, colors for plotting, how the plot legend is displayed, and spectral smoothing method and kernel width.

Figure 7.24 shows the setting dialogs accessible from the toolbar. This **Preferences** dialog opened by the gear icon allows the user to:

- Toggle automatic scaling the x- and y-ranges of the plot.
- Toggle the coordinate grid overlay in the background of the plot.
- Toggle whether registered images other than the current one appear as overlays on the plot.
- Toggle whether these profiles are plotted relative to the main profile (*in development*).
- Toggle the display of tooltips (*in development*).
- Toggle the plotting of a top axis.
- Toggle between a histogram and simple line style for the plot.
- Toggle between the radio and optical versions of the Spectral Profile tool. *Note: We discuss only the radio version here; this mainly impacts the Spectral Line Fitting and Collapse/Moments functionality.*
- Toggle the overplotting of a line showing the channel currently being displayed in the main Display Panel.

The **Color Curve Preferences** dialog opened by the color wheel icon allows the user to:

- Select the color of the line marking the current channel shown in the main Display Panel.
- Select the color used to overlay molecular lines from Splatalogue.
- Select the color to plot the initial Gaussian estimate used in spectral line fitting.
- Select the color used for the zoom rectangle.
• Set a queue of colors used to plot the various data sets registered in the Display Panel.
• Set a queue of colors to plot the set of Gaussian fits.
• Set a queue of colors to plot the synthesized curve.

Two sets of preset colors, "Traditional” or "Alternative”, are available and the user can define their own custom color palette.

The legend options opened by the signpost icon allow the user to toggle the plotting of a legend defining the curves shown in the main Spectral Profile window. Using a drop-down dialog, the legend can be placed in the top left corner of the plot, to the right of the plot, or below the plot. Toggling the color bar causes the color of the curve to be indicated either via a short bar or using the color of the text itself. Double click the names of the files or curves to edit the text shown for that curve by hand. A legend is only available if more than a single spectrum has been loaded.

The spectral smoothing option has two methods, Boxcar and Hanning with the selection of odd numbers for the smoothing kernel width in channels.

7.4.4.2 Main Spectral Profile Window

![Main Spectral Profile Window](image)

Figure 7.25: The main panel for the Spectral Profile tool. Buttons along the bottom row allow the axes to be set. Arrow keys pan and dragging out an area with the mouse zooms. Holding CTRL and right clicking in the spectrum will jump the main Viewer Display panel to display that frequency channel.

The main window shows the spectrum extracted from active region of the image in the main Display Panel. The spectra from the same region in any other registered images are also plotted if overlays are enabled. Menus along the bottom of the image allow the user to select how the spectrum is displayed. From left to right:

• The units for the bottom spectral axis.
• The units for the top spectral axis. **NOTE:** dual axes are enabled only if a single image is registered and the top axis option is enabled. In general, dual axes are not well-defined for mixed data sets. The exception is that open data cubes with matched frequency/spectral axes will allow dual axes.

• The units for the left intensity or flux axis *Note: fraction of peak allows easy comparison of data with disparate intensity scales.*

• The velocity reference frame used if a velocity axis is chosen for the top or bottom axis.

• The method used to extract spectrum from the region — a mean over all pixels in the region, a median, sum, or a sum converting units to get a flux density over the region (Jy).

• Toggle the calculation and overplotting of error bars calculated from scatter in the data (*rmse* refers to root mean square error).

In addition to these drop-down menus, the main Spectral Profile window allows the user to do the following using keyboard and mouse inputs:

• *jump the main Display Panel window to a specified channel (CTRL+Right click):* hold CTRL and right click in the spectrum. A marker will appear and the main Viewer Display Panel will jump to display that channel.

• *animate the main Display Panel in a movie across a frequency range (CTRL+Right click+drag):* hold CTRL, Right click, and drag. The main Viewer Display panel will respond by showing a movie scrolling across the selected spectral channels.

• *zoom the Spectral Profile (+/-, mouse drag):* Use the +/- keys to zoom in the same way as the toolbar buttons. Alternatively, press and dragging the left mouse button. A yellow box is drawn onto the panel. After releasing the mouse button, the plot will zoom to the selected range.

• *pan the Spectral Profile (arrows):* Use the arrow keys to pan the plot.

• *select a spectral range for analysis:* hold shift, left click, and drag. A gray area will be swept out in the display. This method can be used to select a range for spectral line fitting or collapsing a data cube (in the Collapse/Moments window).

**NOTE:** If the mouse input to the Spectral Profile browser becomes confused hit the **ESC** key several times and it will reset.

### 7.4.4.3 Spectral-Line Fitting

**NOTE:** Interactive spectral line fitting is still under development.

The **Spectral-Line Fitting** tab, shown in Figures 7.26 and 7.27 allows the user to interactively fit a combination of Gaussian and polynomial profiles to the data shown in the Spectral Line Profile tool. The tool includes a number of options, many of which remain under development:
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Figure 7.26: The Spectral Line Fitting tab in the Spectral Profile Tool. The user can fit a combination of a polynomial and multiple Gaussian components, specifying the range to be fit (gray region) manually or with a shift+click+drag. Initial estimates for each component may be entered by hand or specified via an initial estimates GUI. The results are output to a dialog and text file with the fit overplotted (here in blue) on the spectrum (with the possibility to save it to disk).

Figure 7.27: The left panel shows the graphical specification of initial estimates for Gaussian fitting. Slider bars specify the center, FWHM, and peak intensity for the initial estimate. The right panel shows the verbose output of the fitting.

- A drag-down menu at the top of the panel allows the user to pick which data set to fit.
- The spectral range to fit can be specified by either holding shift+left click+dragging out a
region in the main spectral profile window or by typing it manually into the box at the top left of the fitting panel.

- Optionally multiple fits can be carried out once, fitting each spectrum in the region in turn. To enable this, check the “MultiFit” box. (*Under development.*)

- Optionally a polynomial of the specified order may be fit. To do so, check the ”Polynomial” fit check box and then specify the desired order. (*Under Development.*)

- The results may be saved to a text file. This text file should be specified before the fit is carried out. Click ”Save” and then use the dialog to specify the file name. *Note that the fit curve itself becomes a normal spectral profile data set and can be saved to disk using the toolbar (disk icon) after the fit.*

- One or more Gaussians can be fit (*Results are presently most stable for one Gaussian.*). Specify the number of Gaussians and then enter initial estimates for the peak, center, and FWHM in the table below. Any of these values can be fixed for any of the Gaussians being fit. Initial estimates can also be manually specified by clicking Specify Estimates. This brings up an additional GUI window (Figure 7.27), where slides can be used to specify initial estimates for each Gaussian to be fit.

- For plotting purposes, one may wish to oversample the fit (i.e., plot a smooth Gaussian), you can do so by increasing the Fit Samples/Channel to a high number to finely sample the fit when plotting.

**NOTE:** Currently the tool works well for specifying a single Gaussian. Fitting multiple components can become unstable and polynomial and multiple line-of-sight fitting are still under development. This is an area of active development and future releases will offer improved capabilities.

### 7.4.4.4 Line Overlays

CASA ships with a local version of the Splatalogue spectral line database ([www.splatalogue.net](http://www.splatalogue.net)) and this can be used to identify and overplot spectral transitions. This feature, shown in Figure 7.28 allows the user to search Splatalogue over the range of interest.

To overlay spectral lines:

1. Select the **Line Overlays** tab in the Spectral Profiles tab.

2. If you know it, enter the redshift or velocity of your source in the ”Doppler Shift” panel. Otherwise, the lines will be overlaid assuming a redshift of 0.

3. Specify a minimum and maximum frequency range to search, either by typing a range or by holding shift and left click and dragging out a range in the spectrum (you will see a gray box appear). If you don’t specify a range, the tool will search over the frequency range of spectrum.
4. Optionally, you may select an astronomical filter from the list (for example, commonly used extragalactic lines or lines often found in hot cores, see Splatalogue for more information). This is usually a good idea because it pare the potentially very large list of candidate lines to a smaller set of reasonable candidates.

5. Click "Search" and the Spectral Profile will search Splatalogue for a list of Spectral lines that fit that Astronomical Filter in that frequency range for that redshift. A dialog will pop up showing the list of candidate lines.

6. Highlight one or more of these transitions and click "Graph Selected Lines." A set of vertical markers will appear in the main Spectral Profile window at the appropriate (redshifted) frequencies for the line.

We emphasize that this feature remains under active development. Look for improved performance and an expanded feature set in the next release.

**NOTE:** You will want to click "Clear Lines" between searches, especially if you update the redshift.

### 7.4.5 The Brightness Profile Tool

The "line" tool can be used to display 1-dimensional brightness profiles of images. The viewer accepts even more than one line segments such as shown in Fig. 7.29. The "region" dock will then
display a preview of the slice in the “Slice Cut” tab and the full “1-D Slice Tool” can be launched from there. This panel allows one to select the interpolation method to connect the pixels, and a number count for the sampled pixels in between markers. “Automatic” will show all pixels. The x-axis of the display can be either the distance along the slice or the X and Y coordinate projections (e.g. along RA and DEC). All segments are also listed at the bottom with their start and end coordinates, the distance and the position angles of each slice segment. The color tool can be used to give each segment a separate color.

Figure 7.29: 1-dimensional slice of an image. The 1D slicer tool shows the brightness distribution along line segments.

7.4.6 The Collapse/Moments Tool

The CASA Viewer can collapse a data cube into an image, for instance allowing to look at the emission integrated along the $z$ axis or the mean velocity of emission along the line of sight. You can access this functionality via the Collapse/Moments tool (accessed via the Tools menu or the four arrow icons), shown in Figure 7.30.

The tool uses the same format as the Spectral Profile tool and will show the integrated spectrum of whatever region or point is currently selected in the main Display Panel. To create a moment map:

1. Select a range over which to integrate either manually using the left part of the window, by adding an interval and typing in the values into the box or by holding **SHIFT + Left Click** and dragging out the range of interest.

2. Pick the set of algorithms that you will use to collapse the image along the $z$ axis. Clicking toggles each moment method, and the collapse will create a new image for each selected moment. For details on the individual collapse method, see the `immoments` task for more details on each moment.
3. The moment may optionally include or exclude pixels within a certain range (for example, you might include only values with signal-to-noise three or greater when calculating the velocity dispersion). You can enter the values to include or exclude manually in the Thresholding window on the right or you can open a histogram tool to specify this range graphically by clicking Specify Graphically (before this can work, you must click "Include" or "Exclude").

4. The results of the collapse be saved to a file, which consists of a string specifying the specific moment tacked onto a root file name that you can specify using Select Root Output File.

5. When you are satisfied with you chosen options, press Collapse.

NOTE: Even if you don’t save the results of the collapse to a file, you can still save the map later using the Save as . . . entry in the Data pull down menu from the main Viewer Display Panel.

NOTE: This area remains under active development and may still exhibit some stability issues in CASA 4.1.

7.4.7 The Histogram Tool

CASA can calculate and visualize a histogram of pixel values inside a region of interest. To examine this histogram, select Histogram from the Tools drop-down menu or the Histogram icon (looks like a comb). This opens the full histogram tool; more limited versions are accessible from the Region Manager Panel, the graphical color table manipulation tool, and the Collapse/Moments tool.
The Histogram tool, accessed from the Main Toolbar or the Tools drop down menu. Details of the display and included pixels can be manipulated via the menus along the top of the window. The right hand panel allows one to attempt to fit a distribution to the histogram.

The resulting Histogram Tool should look something like Figure 7.31. The menus along the top (or the corresponding mouse clicks) allow one to:

- Zoom to the full range, a selected percentile, or a graphical range.
- Change the display of the histogram to show a log axis, display as either a line plot, an outline, or a filled histogram. Change the number of bins in the histogram, or clear the plot (to start over).
- Configure what data are fed into the histogram. You can use this menu to tell the histogram to track the channel currently selected in the main Viewer Display Panel (click the "Track Channel" box) or to integrate across some range of channels (defaulting to the whole image). You can also switch the 2-D footprint used between the whole Image, the Selected Region, and All Regions.
- Save (via the disk icon) an image of the histogram to a graphical file on disk.

The Histogram Tool also allows you to fit the distribution using either a Gaussian or a Poisson distribution, for example to estimate the noise in the image (a Gaussian will be a good choice to describe the noise in most radio data cubes). You can specify initial estimates or let the program generate initial guesses. The fit is then overplotted on the histogram (colors can be adjusted by
clicking the color wheel icon in the toolbar) and details of the fit are printed to the text window below the fit button.

### 7.4.8 The Two-D Fitting Tool

![The interface to the two dimensional fitting tool](image)

Figure 7.32: The interface to the two dimensional fitting tool (Tools:Fit... or the blue circles icon). The interface allows you to specify and automatically generate (Find Sources) initial estimates, to specify the range of pixel values to be included in the fit, and to specify the output (log file, residual image, and visualization). Click **Fit** to start the fit.

**NOTE:** This functionality is still under very active development. Not all features are functional at this point.

CASA can fit two dimensional Gaussians to an intensity distribution, and the Two-Dimensional Fitting tool in the Viewer exposes this functionality interactively. This tool, accessed by the blue circles icon or the Tools:Fit... menu item, has an interface like that shown in Figure 7.32. The interface exposes several options:

1. You can select whether to fit only the selected region or the whole image plane and specify which channel of the cube you want to operate on. **NOTE:** The two dimensional fitter only operates on a single channel at a time.

2. **Initial Estimates:** The box in the top left corner allows the user to specify initial estimates by feeding in a file. The easiest way to make an appropriate file is to edit an existing one. Even easier, you can use the **Find Sources** button to automatically generate a temporary
file of initial estimates. **NOTE:** This functionality is still under development. When it is working, you click on **Find Sources**

3. **Pixel Range:** You can choose to only include a certain range of pixel intensity values in the fit. For example, you might choose to only fit Gaussians to pixels above a few times the measured noise level. You can use the **Specify Graphically** button to bring up an interactive histogram of the region (a reduced functionality version of the full Histogram Tool).

4. **Output:** You can choose to save the output of the fit as a file to the specified directory and to subtract the fit from the image and to subtract the fit from the original, creating a **Residual Image** that gets stored as a CASA image and automatically loaded into the viewer. This gives a way to tell how well your fit describes the total emission.

5. **Visualization:** You can toggle whether the fit is displayed on the viewer or not and change the color of the marker.

Click **Fit** to start the fit. If the fit does not converge, try improving your initial estimates and fitting again.

### 7.4.9 Interactive Position-Velocity Diagram Creation

Figure 7.33: Interactive creation of position-velocity cuts in the viewer. Use the P/V tool from the Mouse Toolbar to define a cut, then use the pV tool from the Region Manager Panel to adjust the cut (including the width). Click Generate P/V to build the position velocity cut and open it in a new Viewer Display Panel (from which it can be saved to disk).

The route to create position-velocity cuts in the **viewer** is illustrated in Figure 7.33.

1. Select the P/V cut tool from the Mouse Toolbar and use it to draw a line across a data cube along the axis you want to visualize.
2. Open the Region Manager Panel and go to the \( pV \) tab. Highlight the cut you just drew. You should see the end point coordinates listed, along with information on the length and position angle of the cut. You can set the averaging width (in pixels) in a window at the bottom of the tab.

3. When you are satisfied, hit \textbf{Generate P/V}. This will create a new Main Viewer Display Panel showing the position velocity cut. The axes should be Offset and velocity.

The new image can be saved to disk with the \texttt{Data:Save as...} option.

### 7.5 Viewing Measurement Sets

![Load Data - Viewer panel](image)

Figure 7.34: The \textbf{Load Data - Viewer} panel as it appears if you select an MS. The only option available is to load this as a \texttt{Raster Image}. In this example, clicking on the \texttt{Raster Image} button would bring up the displays shown in Figure 7.2.

Visibility data can also be displayed and flagged directly from the viewer. For Measurement Set files the only option for display is 'Raster' (similar to AIPS task \texttt{TVFLG}). An example of MS display is shown in Figure 7.2; loading of an MS is shown in Figure 7.34.
Warning: Only one MS should be registered at a time on a Display Panel. Only one MS can be shown in any case. You do not have to close other images/MSs, but you should at least ‘unregister’ them from the Display Panel used for viewing the MS. If you wish to see other images or MSs at the same time, create multiple Display Panel windows.

7.5.1 Data Display Options Panel for Measurement Sets

The Data Display Options panel provides adjustments for MSs similar to those for images, and also includes flagging options. As with images, this window appears when you choose the Data:Adjust menu or use the wrench icon from the Main Toolbar. It is also shown by default when an MS is loaded. The right panel of Figure 7.2 shows a Data Options window. It has a tab for each open MS, containing a set of categories. The options within each category can be either ‘rolled up’ or expanded by clicking the category label.

For a Measurement Set, the categories are:

- Advanced
- MS and Visibility Selection
- Display Axes
- Flagging Options
- Basic Settings
- Axis Drawing and Labels
- Color Wedge

7.5.1.1 MS Options — Basic Settings

The Basic Settings roll-up is expanded by default. It contains entries similar to those for a raster image (§7.4.1.2). Together with the brightness/contrast and colormap adjustment icons on the Mouse Toolbar of the Display Panel, they are especially important for adjusting the color display of your MS.

The available Basic options are:

- Data minimum/maximum
  This has the same usage as for raster images. Lowering the data maximum will help brighten weaker data values.
- Scaling power cycles
  This has exactly the same usage as for raster images (see §7.4.1.2). Again, lowering this value often helps make weaker data visible. If you want to view several fields with very different amplitudes simultaneously, this is typically one of the best adjustments to make early, together with the Colormap fiddling mouse tool, which is on the middle mouse button by default.
• Colormap

Greyscale or Hot Metal colormaps are generally good choices for MS data.

7.5.1.2 MS Options—MS and Visibility Selections

• Visibility Type
• Visibility Component
• Moving Average Size

This roll-up provides choice boxes for Visibility Type (Observed, Corrected, Model, Residual) and Component (Amplitude, Phase, Real, or Imaginary).

Figure 7.35: The MS for NGC4826 BIMA observations has been loaded into the viewer. We see the first of the spw in the Display Panel, and have opened up MS and Visibility Selections in the Data Display Options panel. The display panel raster is not full of visibilities because spw 0 is continuum and was only observed for the first few scans. This is a case where the different spectral windows have different numbers of channels also.

Changes to Visibility Type or Component (changing from Phase to Amplitude, for example) require the data to be retrieved again from the disk into memory, which can be a lengthy process. When a large MS is first selected for viewing, the user must trigger this retrieval manually by pressing
the **Apply** button (located below all the options), after selecting the data to be viewed (see **Field IDs** and **Spectral Windows**, below).

**Tip:** Changing visibility type between 'Observed' and 'Corrected' can also be used to assure that data and flags are reloaded from disk. You should do this if you’re using another flagging tool such as autoflag simultaneously, so that the viewer sees the other tool’s new edits and doesn’t overwrite them with obsolete flags. The **Apply** button alone won’t reload unless something within the viewer itself requires it; in the future, a button will be provided to reload flags from the disk unconditionally.

You can also choose to view the difference from a running mean or the local RMS deviation of either Phase or Amplitude. There is a slider for choosing the nominal number of time slots in the 'local neighborhood' for these displays.

(Note: **Insufficient Data** is shown in the tracking area during these displays when there is no other unflagged data in the local neighborhood to compare to the point in question. The moving time windows will not extend across changes in either field ID or scan number boundaries, so you may see this message if your scan numbers change with every time stamp. An option will be added later to ignore scan boundaries).

- **Field IDs**
- **Spectral Windows**

You can retrieve and edit a selected portion of the MS data by entering the desired Spectral Window and Field ID numbers into these boxes. **Important:** Especially with large MSs, often the first thing you’ll want to do is to select spectral windows which all have the **same number of channels** and the **same polarization setup**. It also makes sense to edit only a few fields at a time. Doing this will also greatly reduce data retrieval times and memory requirements.

You can separate the ID numbers with spaces or commas; you do not need to enter enclosing brackets. Changes to either entry box will cause the selected MS data to be reloaded from disk.

If you select, say, spectral windows 7, 8, 23, and 24, the animator, slice position sliders, and axis labeling will show these as 0, 1, 2, and 3 (the 'slice positions' or 'pixel coordinates' of the chosen spectral windows). Looking at the position tracking display is the best way to avoid confusion in such cases. It will show something like: **Sp Win 23 (s 2)** when you are viewing spectral window 23 (plane 2 of the selected spectral windows).

Changes to MS selections will not be allowed until you have saved (or discarded) any previous edits you have made (see **Flagging Options -- Save Edits**, below). A warning is printed on the console (not the logger).

Initially, all fields and spectral windows are selected. To revert to this 'unselected' state, choose 'Original' under the wrench icons next to the entry boxes.

See Figure 7.35 for an example showing the use of the **MS and Visibility Selections** controls when viewing an MS.
7.5.1.3 MS Options — Display Axes

This roll-up is very similar to that for images: it allows the user to choose which axes (from Time, Baseline, Polarization, Channel, and Spectral Window) are on the display and the animator. There are also sliders here for choosing positions on the remaining axes. (It’s useful to note that the data is actually stored internally in memory as an array with these five axes).

![Display Axes Roll-up]

Figure 7.36: The MS for NGC4826 from Figure 7.35 now with the Display Axes open in the Data Display Options panel. By default, channels are on the Animation Axis and thus in the tapedeck, while spectral window and polarization are on the Display Axes sliders.

For MSs, changing the choice of axis on one control will automatically swap axes, maintaining different axes on each control. Changing axes or slider/animator positions does not normally require pressing Apply — the new slice is shown immediately. However, the display may be partially or completely grey in areas if the required data is not currently in memory, either because no data has been loaded yet, or because not all the selected data will fit into the allowed memory. Press the Apply button in this case to load the data (see §7.5.1.6 and Max. Visibility Memory at the end of §7.5.1.5).

Within the Display Axes rollup you may also select whether to order the baseline axis by antenna1-antenna2 (the default) or by (unprojected) baseline length.

See Figures 7.36 7.37 showing the use of the Display Axes controls to change the axes on the
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Figure 7.37: The MS for NGC4826, continuing from Figure 7.36. We have now put spectral window on the Animation Axis and used the tapedeck to step to spw 2, where we see the data from the rest of the scans. Now channels is on a Display Axes slider, which has been dragged to show Channel 33.

animation and sliders.

7.5.1.4 MS Options — Flagging Options

These options allow you to edit (flag or unflag) MS data. The Point Tool and Rectangle Region Mouse Tools (§ 7.4.3.2) are used on the display to select the area to edit. When using the Rectangle Region tool, double-click inside the selected rectangle to confirm the edit.

The options below determine how edits will be applied.

- Show Flagged Regions...

You have the option to display flagged regions in the background color (as in TVFLG) or to highlight them with color. In the former case, flagged regions look just like regions of no data. With the (default) color option, flags are shown in shades of blue: darker blue for flags already saved to disk, lighter blue for new flags not yet saved; regions with no data will be shown in black.
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• Flag or Unflag
This setting determines whether selected regions will be flagged or unflagged. This does not affect previous edits; it only determines the effect which later edits will have. Both flagging and unflagging edits can be accumulated and then saved in one pass through the MS.

• Flag/Unflag All...
These flagging extent checkboxes allow you to extend your edit over any of the five data axes. For example, to flag all the data in a given time range, you would check all the axes except Time, and then select the desired time range with the Rectangle Region mouse tool. Such edits will extend along the corresponding axes over the entire selected MS (whether loaded into memory or not) and optionally over unselected portions of the MS as well (Use Entire MS, below). Use care in selecting edit extents to assure that you’re editing all the data you wish to edit.

• Flag/Unflag Entire Antenna?
This control can be used to extend subsequent edits to all baselines which include the desired antenna[s]. For example, if you set this item to ‘Yes’ and then click the point tool on a visibility position with baseline 3-19, the edit would extend over baselines 0-3, 1-3, 2-3, 3-3, 3-4, ..., 3-nAntennas-1. Note that the second antenna of the selection (19) is irrelevant here – you can click anywhere within the ‘Antenna 3 block’, i.e., where the first antenna number is 3, to select all baselines which include antenna 3.

This item controls the edit extent only along the baseline axis. If you wish to flag all the data for a given antenna, you must still check the boxes to flag all Times, Channels, Polarizations and Spectral Windows. There would be no point, however, in activating both this item and the ‘Flag All Baselines’ checkbox. You can flag an antenna in a limited range of times, etc., by using the appropriate checkboxes and selecting a rectangular region of visibilities with the mouse.

Note: You do not need to include the entire ‘antenna block’ in your rectangle (and you may stray into the next antenna if you try). Anywhere within the block will work. To flag higher-numbered antennas, it often helps to zoom in.

• Undo Last Edit

• Undo All Edits
The ‘Undo’ buttons do the expected thing: completely undo the effect of the last edit (or all unsaved edits). Please note, however, that only unsaved edits can be undone here; there is no ability to revert to the flagging state at the start of the session once flags have been saved to disk (unless you have previously saved a ‘flag version’. The flag version tool is not available through the viewer directly).

• Use Entire MS When Saving Edits?
”Yes” means that saving the edits will flag/unflag over the entire MS, including fields (and possibly spectral windows) which are not currently selected for viewing. Specifically, data within time range(s) you swept out with the mouse (even for unselected fields) will be edited.
In addition, if "Flag/Unflag All..." boxes were checked, such edits will extend throughout the MS. Note that only unselected times (fields) can be edited without checking extent boxes for the edits as well. Unselected spectral windows, e.g., will not be edited unless the edit also has "Flag/Unflag All Spectral Windows" checked.

Warning: Beware of checking “All Spectral Windows” unless you have also checked "All Channels” or turned “Entire MS” off; channel edits appropriate to the selected spectral windows may not be appropriate to unselected ones. Set ”Use Entire MS” to “No” if your edits need to apply only to the portion of the MS you have selected for viewing. Edits can often be saved significantly faster this way as well.

Also note that checkboxes apply to individual edits, and must be checked before making the edit with the mouse. “Use Entire MS”, on the other hand, applies to all the edits saved at one time, and must be set as desired before pressing ”Save Edits”.

- Save Edits

MS editing works like a text editor in that you see all of your edits immediately, but nothing is committed to disk until you press “Save Edits”. Feel free to experiment with all the other controls; nothing but ‘Save Edits’ will alter your MS on disk. As mentioned previously, however, there is no way to undo your edits once they are saved, except by manually entering the reverse edits (or restoring a previously-saved 'flag version’).

Also, you must save (or discard) your edits before changing the MS selections. If edits are pending, the selection change will not be allowed, and a warning will appear on the console.

If you close the MS in the viewer, unsaved edits are simply discarded, without prior warning. It’s important, therefore, to remember to save them yourself. You can distinguish unsaved flags (when using the 'Flags In Color' option), because they are in a lighter shade of blue.

The program must make a pass through the MS on disk to save the edits. This can take a little time; progress is shown in the console window.

7.5.1.5  MS Options— Advanced

These settings can help optimize your memory usage, especially for large MSs. A rule of thumb is that they can be increased until response becomes sluggish, when they should be backed down again.

You can run the unix 'top' program and hit 'M' in it (to sort by memory usage) in order to examine the effects of these settings. Look at the amount of RSS (main memory) and SWAP used by the X server and 'casaviewer' processes. If that sounds familiar and easy, then fiddling with these settings is for you. Otherwise, the default settings should provide reasonable performance in most cases.

- Cache size

The value of this option specifies the maximum number of different views of the data to save so that they can be redrawn quickly. If you run an animation or scroll around zoomed data, you will notice that the data displays noticeably faster the second time through because
of this feature. Often, setting this value to the number of animation frames is ideal. However, that on multi-panel displays, each panel counts as one cached image.

Large images naturally take more room than small ones. The memory used for these images will show up in the X server process. If you need more Visibility Memory (below) for a really large MS, it is usually better to forgo caching a large number of views.

- **Max. Visibility Memory**

  This option specifies how many megabytes of memory may be used to store visibility data from the measurement set internally. *Even if you do not adjust this entry, it is useful to look at it to see how many megabytes are required to store your entire (selected) MS in memory.* If the slider setting is above this, the whole selected MS will fit into the memory buffer. Otherwise, some data planes will be ‘grayed out’ (see Apply Button, § 7.5.1.6 below), and the selected data will have to be viewed one buffer at a time, which is somewhat less convenient. In most cases, this means you should **select fewer fields or spectral windows** – see § 7.5.1.2. The 'casaviewer' process contains this buffer memory (it contains the entire viewer, but the memory buffer can take most of the space).

### 7.5.1.6 MS Options — Apply Button

When viewing large MSs the display may be partially or completely grey in areas where the required data is not currently in memory, either because no data has been loaded yet, or because not all the selected data will fit into the allowed memory (see Max. Visibility Memory above). When the cursor is over such an area, the following message shows in the position tracking area:

- press 'Apply' on Adjust panel to load data

Pressing the Apply button (which lies below all the options) will reload the memory buffer so that it includes the slice you are trying to view.

The message **No Data** has a different meaning; in that case, there simply is no data in the selected MS at the indicated position.

For large measurement sets, loading visibility data into memory is the most time-consuming step. Progress feedback is provided in the console window. Again, careful selection of the data to be viewed can greatly speed up retrieval.

### 7.6 Printing from the Viewer

You can select **Data:Print** from the drop down menu or click the **Print** icon to bring up the **Viewer Print Manager**. From this panel, you can **Print** the contents of Display Panel to a hardcopy or **Save** them as an image in a format selected from the drop-down menu at the bottom left of the window. Note that the save feature will overwrite the file in question without prompting.

The Viewer Print Manager allows you to adjust the DPI, orientation, and page format (Output Media) for Postscript or PDF files and to scale the image to a desired pixel size for other images.
Figure 7.38: Printing the display to a hardcopy of a file. From the Viewer Print Manager, located in top right here and accessed by the print icon or from the Data drop down menu, you can use the Save button to save an image or Print directly to a printer. To achieve the best results, it is often helpful to adjust the settings in the Data Display Options and Viewer Canvas Manager, shown at right.

To achieve the best output it is usually advisable to adjust the settings in the Viewer Print Manager, Data Display Options, and Viewer Canvas Manager. For PDF and Postscript output, turning the DPI up all the way yields the best-looking results. For other images, a white background often makes for better looking images than the default black. It is often necessary to increase the Line Width in the Axis Label Properties (in the Data Display Options panel) to ensure that the labels will be visible when printed. Increasing from the default of 1.4 to a value around 2 often works well.

Figure 7.38 shows an example of printing to a file while adjusting the Data Display Options and Viewer Canvas Manager to improve the appearance of the plot.
### 7.7 Image Viewer (imview)

The imview task offers scriptable access to many viewer options. This enables the production of customized plots without invoking the GUI and allows one to open the viewer to a carefully selected state.

**imview** has the following inputs:

<table>
<thead>
<tr>
<th>Input</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>raster</strong></td>
<td>(Optional) Raster filename (string) or complete raster config dictionary. The allowed dictionary keys are file (string), scaling (numeric), range (2 element numeric vector), colormap (string), and colorwedge (bool).</td>
</tr>
<tr>
<td><strong>contour</strong></td>
<td>(Optional) Contour filename (string) or complete contour config dictionary. The allowed dictionary keys are file (string), levels (numeric vector), unit (float), and base (float).</td>
</tr>
<tr>
<td><strong>zoom</strong></td>
<td>(Optional) zoom can specify intermental zoom (integer), zoom region read from a file (string) or dictionary specifying the zoom region. The dictionary can have two forms. It can be either a simple region specified with blc (2 element vector) and trc (2 element vector) [along with an optional coord key (&quot;pixel&quot; or &quot;world&quot;; pixel is the default) or a complete region rectangle e.g. loaded with &quot;rg.fromfiletorecord()&quot;]. The dictionary can also contain a channel (integer) field which indicates which channel should be displayed.</td>
</tr>
<tr>
<td><strong>axes</strong></td>
<td>(Optional) this can either be a three element vector (string) where each element describes what should be found on each of the x, y, and z axes or a dictionary containing fields &quot;x&quot;, &quot;y&quot; and &quot;z&quot; (string).</td>
</tr>
<tr>
<td><strong>out</strong></td>
<td>(Optional) Output filename or complete output config dictionary. If a string is passed, the file extension is used to determine the output type (jpg, pdf, eps, ps, png, ...)</td>
</tr>
</tbody>
</table>
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# xbm, xpm, or ppm). If a dictionary is passed, it can contain the fields, file (string), scale (float), dpi (int), or orient (landscape or portrait). The scale field is used for the bitmap formats (i.e. not ps or pdf) and the dpi parameter is used for scalable formats (pdf or ps).

The **raster** and **contour** parameters specify which images to load and how these images should be displayed. These parameters take python dictionaries as inputs. The fields in these dictionaries specify how the image will be displayed.

An example call to `imview` looks like this:

```python
imview(raster={'file': 'ngc5921.clean.image',
               'range': [-0.01,0.03],
               'colormap': 'Hot Metal 2',
               'scaling': -1},
contour={'file': 'ngc5921.clean.image'},
axes={'x':'Declination'},
zoom={'channel': 7, 'blc': [75,75], 'trc': [175,175], 'coord': 'pixel'},
out='myout.png')
```

The argument to **raster** is enclosed in the curly braces `{ }`. Within these braces are a number of "key":"value" pairs. Each sets an option in the viewer, with the GUI parameter to set defined by the "key" and the value to set it to defined by "value." In the example above, 'file': 'ngc5921.clean.image' sets the file name of the raster image, 'range': [-0.01,0.03] sets the range of pixel values used for the scaling.

**contour** works similar to 'raster' but can accept multiple dictionaries in order to produce multiple contour overlays on a single image. To specify multiple contour overlays, simply pass multiple dictionaries (comma delimited) in to the contour argument:

```python
contour={['file': 'file1.image', 'levels': [1,2,3]},
        {'file': 'file2.image', 'levels': [0.006, 0.008, 0.010] })
```

**zoom** specifies the part of the image to be shown.

**axes** defines what axes are shown. By default, the viewer will show 'x':'Right Ascension', 'y':'Declination' but one may also view position-frequency images.

**out** defines the filename of the output, with the extension setting the file type.

Currently, the following parameters are supported with additional functionality planned for future releases:
raster -- (string) image file to open
   (dict) file (string) => image file to open
   scaling (float) => scaling power cycles
   range (float*2) => data range
   colormap (string) => name of colormap
   colorwedge (bool) => show color wedge?

contour -- (string) file to load as a contour
   (dict) file (string) => file to load
   levels (float*N) => relative levels
   base (numeric) => zero in relative levels
   unit (numeric) => one in the relative levels

zoom -- (int) integral zoom level
   (string) region file to load as the zoom region
   (dict) blc (numeric*2) => bottom left corner
   trc (numeric*2) => top right corner
   coord (string) => pixel or world
   channel (int) => channel to display
   (dict) <region record> => record loaded
      e.g. rg.fromfiletorecord()

axes -- (string*3) dimension to display on the x, y, and z axes
   (dict) x => dimension for x-axes
   y => dimension for y-axes
   z => dimension for z-axes

out -- (string) file with a supported extension
   [jpg, pdf, eps, ps, png, xbm, xpm, ppm]
   (dict) file (string) => filename
   format (string) => valid ext (filename ext overrides)
   scale (numeric) => scale for non-eps, non-ps output
   dpi (numeric) => dpi for eps or ps output
   orient (string) => portrait or landscape

Examples are also found in help imview.

7.8 Measurement Viewer (msview)

The Measurement Viewer msview is mostly a clone of the viewer at this stage. A difference is that msview allows the user to select data before it is loaded into the GUI and displayed. A screenshot is shown in Fig. 7.39 and selection parameters are field, spectral window, time range, uv range, antenna, corr, scan, array, ms selection expression in the usual CASA selection syntax (see Sect. 2.3).
Figure 7.39: Data selection in msview.
Chapter 8

Single Dish Data Processing

The Single Dish part of CASA uses the ATNF Spectral Analysis Package (ASAP), which is imported as the sd tool at the start-up of CASA. ASAP forms the basis for a series of tasks (the “SDtasks”) that encapsulates the functionality within the standard CASA task framework. ASAP was originally developed to support the Australian telescopes such as Mopra, Parkes, and Tidbinbilla. Currently, datasets taken at ALMA, GBT (see the note below for limitation of GBT SDFITS handling), ASTE, NRO 45m and Mopra are supported for data processing within CASA.

For details on ASAP – including the User Guide, Reference Manual, and tutorial – see the ASAP home page at ATNF:


The ASAP tools are prefaced with sd. within CASA, e.g., the ASAP tool scantable becomes sd.scantable. See §8.5 for more information on the tools.

All of the ASAP functionalities are available within the CASA installation. Since we extended ASAP, there are certain functionalities that are available only in the CASA version of ASAP. In the following subsections, we outline how to access ASAP from within CASA and the data flow for standard use cases.

If you run into trouble, be sure to check the list of known issues and features of ASAP and the SDtasks presented in §8.7 first.

8.1 Guidelines for the use of ASAP and SDtasks in CASA

8.1.1 Environment Variables

There are a number of environment variables that the ASAP tools (and thus the SDtasks) use to control the operation. They are located in .asaprc and are described in the ASAP User Guide. Within CASA, they are contained in the Python dictionary sd.rcParams and are accessible through its keys and values. For SDtask users, the most important parameter is verbose, which controls the display of detailed messages from the tools. By default,
produces lots of messages. Also, the `scantable.storage` parameter controls whether scantable operations are done in memory or on disk. The default is

```python
sd.rcParams['scantable.storage'] = 'memory'
```

which is the best choice if there is enough memory than the size of the data to be loaded. On the other hand,

```python
sd.rcParams['scantable.storage'] = 'disk'
```

forces the task to store datasets on disk, which might be necessary when the dataset is large. See §8.5.1 for more details on the ASAP environment variables.

**Important Note:**
User must use `sd.rcParams[scantable.storage] = 'disk'` with care when you call any tool level functions since some functions may overwrite original data even if you set `sd.rcParams['insitu'] = False`, which tells the system not to overwrite the original data (in contrast, setting `sd.rcParams['insitu']` to True forces to overwrite original data). Relevant methods, which may overwrite original data in the above case, are as follows:

- `sd.average_time`
- `sd.merge`
- four operations (+, -, *, /) of `sd.scantable` instance with scalar or array
- `sd.scantable.add`
- `sd.scantable.clip`
- `sd.scantable.flag`
- `sd.scantable.flag_nans`
- `sd.scantable.flag_row`
- `sd.scantable.scale`
- `sd.scantable.recalc_azel`
- any setter functions of `sd.scantable` class (both `setxxx` and `_setxxx` functions)

If you only use the SDtasks, you don’t need to worry about this since the SDtasks are designed to keep the original data unchanged.

### 8.1.2 Assignment

Some ASAP methods and functions require assigning a method to a variable which can then be manipulated. These methods and functions include `sd.scantable` and `sd.selector`, both of which make objects. For example,

```python
s = sd.scantable('foo.PM01.asap', average=False)
```
8.1.3 Dictionaries

Currently, the SDtasks return the Python dictionary for the results of line fitting (in `sdfit`) and region statistics (in `sdstat`). If you invoke these tasks by assigning a variable for the return, you can then access the elements through the keywords, e.g.,

```
CASA <10>: line_stat=sdstat()
Current fluxunit = K
No need to convert fluxunits
Using current frequency frame
Using current doppler convention
```

```
CASA <11>: line_stat
Out[11]:
{'eqw': 70.861755476162784,
 'max': 1.2750182151794434,
 'mean': 0.35996028780937195,
 'median': 0.23074722290039062,
 'min': -0.20840644836425781,
 'rms': 0.53090775012969971,
 'stddev': 0.39102539420127869,
 'sum': 90.350028991699219}
```

One can then use these values in scripts by accessing this dictionary, e.g.,

```
CASA <12>: print "Line max = %5.3f K" % (line_stat['max'])
Line max = 1.275 K
```

8.1.4 Line Formatting

The SDtasks trap leading and trailing whitespace on string parameters (such as `infile`) but ASAP does not, so be careful when setting string parameters. ASAP is case-sensitive, with most parameters being upper-case, such as `ASAP` for the `sd.scantable.save` file format. The SDtasks are generally more forgiving. Also, beware Python’s sensitivity to indentation.

8.1.5 Logging

Before R3.0, all messages from ASAP were written to the standard output (`sys.stdout`) and they disappeared after exiting CASA. After R3.0, the logging system of ASAP is integrated into CASA logging system. Therefore, all outputs from ASAP commands, except for GUI related notifications, are sent to the log file for the current session and they are displayed to the CASA Logger (see §1.5.2).

8.2 Task Interface Changes in CASA 4.2.2

Major interface changes to SDtasks were taken place in CASA 4.2.2 release. The interface of the following tasks are modified: `sdbaseline`, `sdcal`, `sdcal2`, `sdfit`, `sdflag`, `sdgrid`, `sdimaging`,
sdmath, sdplot, sdreduce, sdsave, and sdstat. Additionally, a new task called sdaverage is available. Task sdsMOOTH has been incorporated in the new task and removed.

The tasks with old interfaces are available with name {taskname}old and will be deleted in CASA 4.4 release. Users are advised to update existing scripts.

8.2.1 Overview

1. Changes of data selection parameter names and selection syntax (as in the tasks for interferometer): all tasks
   - scanlist, scans → scan
   - iflist, ifno, ifs → spw
   - pollist, pols → pol
   - beamlist → beam
   - timerange selection is added

2. Changes of parameter names
   - telescopeparm → telescopeparam: sdbaseline, sdcal, sdfit, sdmath, sdplot, sdreduce, sdstat
   - varlist → varnames: sdmath
   - step → width: sdimaging

3. Removal of unnecessary parameters
   - averaging parameters (scanaverage, timeaverage, polaverage, tweight, pweight): sdcal, sdcOADD, sdlIST, sdsave
   - specunit, frame, doppler: sdcal, sdmath, sdcOADD
   - fluxunit, telescopeparm: sdcOADD
   - dochannelmap: sdimaging

4. Addition of parameters
   - averaging parameters (scanaverage, timeaverage, polaverage, tweight, pweight): sdfit, sdstat
     - scanaverage parameter is a sub-parameter of timeaverage (shown only when timeaverage=True)
     - default value of scanaverage is False
   - freq Tol: sdcOADD
   - infiles: sdmath

5. New task
   - sdaverage is created by merging the average function in sdcal and sdsMOOTH
8.2.2 Data selection

The selection syntax for single dish is modified based on the CASA MS data selection syntax.

8.2.2.1 scan, beam

- Only ID number (integer) is supported
- comma-separated values and ranges are supported. spaces on both sides of comma are supported (e.g., ’1, 3, 5’)
- range is specified by ’~’ (e.g., ’3~5’)
- inequality sign is supported (e.g., ’>2’, ’<5’)

8.2.2.2 pol

- ID number selection as in scan and beam is supported
- selection by string (e.g., ’XX’, ’YY’, ’RR’, ’LL’) is not supported

8.2.2.3 field

- ID number selection as in scan and beam is supported
- pattern matching and multiple string selections separated by comma are supported
- regular expression is not supported

8.2.2.4 spw

- spw selection is in the format of multiple ’spwSelection:channelSelection’ separated by comma
- spwSelection supports the following three types:
  - ID number selection as in scan and beam
  - ’*’ or blank string selection (e.g., ’*:200~300’ or ’:200~300’)
  - frequency range selection is supported to select spwID which matches the frequency range (e.g., ’1412~1415MHz’)
- channelSelection
  - ID number selection as in scan and beam is supported
  - multiple ranges can be selected by separating by semicolon (e.g., ’0:50~250;350~462’)
  - selections by channel, frequency, and velocity are supported (e.g., ’110~120GHz’, ’-100~100km/s’)
  - step selection by ’^’ is not supported
8.2.3 Addition of functions

8.2.3.1 sdcoadd

- parameter freqtol is added
- IFs are merged by considering freqtol when merging scantables

8.2.3.2 sdmath

- parameter 'infiles' accepts a list of input file names
- input file names are automatically replaced with the phrases IN0, IN1, ... in expr parameter
- parameter name changed from varlist to varnames

8.3 New Features in CASA 4.3

8.3.1 Proper handling of flagging information

Channel-based (FLAGTRA data column) and row-based (FLAGROW) flag information are properly applied in all relevant SDtasks. Detailed descriptions for each task are as follows. Here ”flagged channels” means channels with a channel-based flag, and ”flagged spectra” means spectra with a row-based flag.

- sdaverage
  [smoothing]
  - the data at flagged channels should be linearly interpolated before FFT
  - smoothing should be made for flagged spectra in rigrid mode
  - smoothing should not be made for flagged spectra in hanning and gaussian mode
  - the output flag values should not be modified
  [averaging]
  - the data with flagged channels or spectra should not be used in averaging procedure
  - the output flag values should be false only in case there is no effective data to be averaged

- sdbaseline
  - the data with flagged channels or spectra should not be used in fitting
  - subtraction of the best fit baseline should be executed for flagged channels, but not for flagged spectra
  - the output flag values should be identical as those in input data
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- **sdcal**
  - for on-source spectra, apply should be executed for flagged channels, but not for flagged spectra
  - for the others (off and tsys), flagged channels/spectra should not be used for calibration procedure
  - the output row-based flag values should be identical as those in input data, while the output channel-based flag should be flagged if the corresponding off-source spectra are flagged

- **sdcal2**
  - for on-source spectra, apply should be executed for flagged channels, but not for flagged spectra
  - for the others (off and tsys), flagged channels/spectra should not be used for calibration procedure
  - for on-source spectra, the output row-based flag values should be identical as those in input data, while the output channel-based flag should be flagged if the corresponding off-source spectra are flagged
  - for off/tsys spectra, the output flag values should follow the way of sdaverage

- **sdcoadd**
  - neither channel/row flags should be modified

- **sdfit**
  - for data averaging, flag values should follow the way of sdaverage
  - for fitting, flagged channels should not be used
  - fitting procedure should not be executed for flagged spectra

- **sdflag**
  - if the spectrum is flagged, channel flags should not be modified

- **sdgrid, sdimaging, sdimprocess**
  - flagged channels/spectra should not be used
  - the output flag values should follow the way of sdaverage

- **sdlist**
  - output should be made for flagged spectra

- **sdmath**
  - calculation procedure should be made for flagged channel, but should be skipped for flagged spectra
the output flag values should be flagged if there is a flagged channel/spectrum in the relevant input data

- `sdscale`
  - scaling calculation should be made for flagged channels but should be skipped for flagged spectra
  - the output flag values should be identical with those in input data

- `sdstat`
  - calculation should not use flagged channels and should be skipped for flagged spectra.
  - the output flag values should be identical with those in input data

- `sdtpimaging`
  - flagged channels/spectra should not be used
  - the output flag values should follow the way of `sdbaseline` (in baselining) and of `sdaverage` (in imaging)

### 8.3.2 Rasterutil

Rasterutil is a module which enables you to select individual raster rows or rasters from raster-scanned dataset. Suppose you have a Scantable 'foo.asap' in which several rasters are scanned. To know how many rasters or raster rows this dataset contains, execute first as follows:

```python
s = sd.scantable('foo.asap', False)
import rasterutil
r = rasterutil.Raster(s)
r.detect()
```

Now you are ready to go ahead. Numbers of raster rows or rasters are shown by typing 'r.nrow' or 'r.nraster', respectively. Once you run `rasterutil.Raster.detect()`, IDs are given in chronological order both for each raster rows and rasters: a raster row can be specified by ID in range from 0 to r.nrow-1, while a raster can be specified by ID from 0 to r.nraster-1.

Here are sample commands to obtain a Scantable that contains specified raster row or raster only or other stuffs to select the specified ones:

```python
srow0 = r.asscantable(rowid=0)  # get Scantable object containing the first raster row
sras1 = r.asscantable(rasterid=1)  # get Scantable object containing the second raster
selrow2 = r.asselector(rowid=2)  # get selector object for selecting the third raster row
taqlras3 = r.astaql(rasterid=3)  # get TaQL query for selecting the 4th raster
```

Other useful commands include `rasterutil.Raster.plot_rows()`, data selected as a raster row or raster can be visualized using asapplotter.
8.4 A Step-by-Step Guide for Reducing and Imaging ALMA Single Dish Data

8.4.1 Introduction and Workflow

Data from the Total Power (TP) array of ALMA is initially in a format called the ASDM. These ASDMs must be converted to ASAP scantables in order to use standard calibration procedures for single dish telescopes;

- Reference (OFF) subtraction
- $T_{\text{sys}}$ calibration
- Baseline subtraction

In the case of continuum observations, the baseline subtraction step can be skipped or be done in the spatial domain. The final imaging part cannot be done in ASAP, and the data must be converted back to Measurement Sets (MSs), and then imaged using a standard CASA task. Below we explain the data reduction in detail, assuming an ASDM format called “foo”, and the dataset contains data from antennas “PM01” and “PM02”.

8.4.2 Create Single Dish Data Format

The ASDMs can be converted to scantables in two ways; directly from the ASDM, or via a MS. We expect that conversion via MS will be used more in the current version of CASA, as the direct conversion can work with only one antenna at a time, while the conversion from a MS can work on multiple antennas simultaneously.

8.4.2.1 Direct Conversion from ASDM

The standard importasdm task in CASA can convert ASDM to scantables by setting singledish=True. It will work with only one antenna at a time, so in case the dataset contains several antennas, the same task must be repeated the same number of times each time specifying the antenna name or ID.

CASA <#>: importasdm(asdm = 'foo', vis = 'foo.PM01.asap',
                        singledish = True, antenna = 'PM01')

CASA <#>: importasdm(asdm = 'foo', vis = 'foo.PM02.asap',
                        singledish = True, antenna = 'PM02')

If the antenna name or ID is not specified, the task will default to the first antenna in the array.
8.4.2.2 Conversion via MS

A conversion to MS first then to scantables may be an efficient choice in the case of multiple antennas.

\[
\text{CASA}<\#:\text{importasdm(asdm = 'foo', vis = 'foo.ms')}\]

will create a MS with all antennas. Then,

\[
\text{CASA}<\#:\text{sdsave(infile = 'foo.ms', splitant = True, outfile = 'foo.asap')}\]

will split the dataset into individual antennas while converting them to scantables. The individual scantables will have names in the format of <outfile>.<antenna name> or <prefix>.<antenna name>.asap if outfile is <prefix>.asap. For the following steps in the reduction, all tasks must be iterated on the scantables of the individual antennas. For simplicity we will just note the CASA input parameters assuming the “PM01” antenna.

8.4.3 Calibration

Data will be calibrated into units of Kelvins, by applying the $T_{\text{sys}}$ calibration and removing the signal from the OFF position. Applying calibration is done with the task \texttt{sdcal2}. \texttt{sdcal2} is an implementation of an interferometry-like calibration scheme, i.e., generate caltables and apply them. \texttt{sdcal2} generates single dish specific caltables for sky (OFF) or $T_{\text{sys}}$ calibration. You can control these behavior using calibration mode (calmode). Available calibration modes are ‘ps’, ‘otf’, ‘otfraster’, and ‘tsys’. In many science observations, calmode is ‘ps’, for Position Switching. If you use calmode = ‘ps’, it will do an (ON-OFF)/OFF calculation by interpolating the nearest (in time) OFFs to the ONs.

calmode = ‘otf’ or ‘otfraster’ are designed for OTF observations without explicit OFF scans, but accept data with OFF scans. If the observing pattern is ‘raster’, you should use the ‘otfraster’ mode to calibrate the data. Otherwise, the ‘otf’ mode should be used.

The generated caltables can be applied to data by using calmode = ‘apply’. You can calibrate data on-the-fly by setting calmode to a composite calmode string separated by comma. For the data already filled $T_{\text{sys}}$, you should use calmode = ‘ps,apply’ (you can replace ‘ps’ with other sky calibration modes), which means doing sky calibration and apply it on-the-fly. In this case, caltable is generated as a temporary plain table and will be deleted at the end.

There are several control parameters for sky and $T_{\text{sys}}$ calibration and application of caltables.

8.4.3.1 Interpolating $T_{\text{sys}}$ into Science SPWs and Apply Calibration

In the current version of ALMA software (as of June 2014), the system temperature ($T_{\text{sys}}$) can only be measured using a spectral setup of 124 channels over a 1.9375 GHz bandwidth (assuming the use of the ACA correlator). The actual science observing is likely to have more channels, in which
case a different spectral window (SPW) is used. In order to calibrate the data, the $T_{\text{sys}}$ from the 124 channel SPW must be interpolated into these science SPWs, both in time and frequency.

First, we must identify which $T_{\text{sys}}$ SPW corresponds to which science SPW. You have to specify a list of spwids for $T_{\text{sys}}$ measurement in tsyspw. Also, you have to set spw mapping between science and $T_{\text{sys}}$ by spwmap. spwmap must be a dictionary with key of spwid for $T_{\text{sys}}$, and value of a list of spwids for science. For example, suppose that $T_{\text{sys}}$ measurements for science windows 17, 19, 21, and 23 are done in spw 9, 11, 13, and 15, respectively. In this case, tsyspw and spwmap should be specified as follows:

```
CASA <#>: sdcal2(infile = 'foo.PM01.asap', calmode = 'ps,tsys,apply',
    tsyspw = '9,11,13,15',
    spwmap = {9:[17],11:[19],13:[21],15:[23]},
    outfile = 'foo.PM01.asap.cal')
```

### 8.4.4 Data Inspection

Look at the calibrated data for obvious or subtle problems. The task `sdplot` will enable looking at time- or scan-integrated spectra. If you need to look at individual data rows, use `sdflag`. Flag as necessary.

```
CASA <#>: sdplot(infile = 'foo.PM01.asap.cal', plottype = 'spectra')
```

or use `plottype = 'pointing'` to look at the data points on the sky. This will be useful also before calibration to check where OFFs and ATMcal scans were done.

### 8.4.5 Flagging

Flagging of data is done using `sdflag`. You can look at individual data rows and choose which to flag using `interactive = T`, or `interactive = F` if you know which scans/spws to flag.

```
CASA <#>: sdflag(infile = 'foo.PM01.asap.cal',
    spw = '17:0~119;3960~4079, 19:0~119;3960~4079,
        21:0~119;3960~4079, 23:0~119;3960~4079',
    overwrite = True)
```

In this case, 120 channels on each side are flagged.

```
CASA <#>: sdflag(infile = 'foo.PM01.asap.cal',
    spw = '17,19,21,23',
    scan = '21~23',
    overwrite = True)
```

In this case, scan IDs 21, 22, and 23 are flagged.
8.4.6 Spectral Baseline Subtraction

Relevant only for the spectral line observation, \texttt{sdbaseline} can identify lines automatically and iteratively (maskmode = 'auto') or you can specify the channel range (maskmode = 'list') to calculate the spectral baselines. The spectral baseline fitting method is controlled by the parameters blfunc and its subparameters. Many fitting methods are available ('poly', 'chebyshev', 'cspline', 'sinusoid', see help of \texttt{sdbaseline} task for more information). Use of zeroth or first order polynomial is good for most cases.

\begin{verbatim}
CASA <#>: sdbaseline(infile = 'foo.PM01.asap.cal',
          maskmode = 'auto',
          thresh = 3.0,
          blfunc = 'poly',
          order = 1,
          outfile = 'foo.PM01.asap.cal.bl')
\end{verbatim}

Look at the baseline-subtracted data with \texttt{sdplot}. Spectral smoothing is done with task \texttt{sdsmooth}.

8.4.7 Batch Reduction: \texttt{sdreduce}

Task \texttt{sdreduce} performs data selection, calibration, and/or spectral baseline fitting for single-dish spectra. This task internally calls the tasks, \texttt{sdcal}, \texttt{sdsmooth}, and \texttt{sdbaseline} and it can be used to run all the three steps in one task execution.

8.4.8 Imaging

8.4.8.1 Convert back to Measurement

Imaging the calibrated single dish dataset is done using a standard CASA task, which takes only MS for input. Therefore the calibrated scantable must first be converted back to a MS.

\begin{verbatim}
CASA <#>: sdsave(infile = 'foo.PM01.asap.cal.bl',
             outfile = 'foo.PM01.asap.cal.bl.ms',
             outform = 'MS2')
\end{verbatim}

8.4.8.2 Concatenating the datasets

Before imaging, the individually calibrated single dish datasets can be combined into one. The individual calibrated MSs have slightly different observing frequencies as the observing frequency is defined in topocentric velocities. The \texttt{freqtol} parameter sets the tolerance for assigning the same SPW ID to datasets with different frequencies. This value should be set to something sensible, as to include all matching SPWs in the individual MSs.
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CASA <#>: concat(vis = ['foo.PM01.asap.cal.bl.ms','foo.PM02.asap.cal.bl.ms'],
             concatvis = 'foo.combined.ms.cal',
             freqtol = '50MHz')

If you do not need to set freqtol, you can skip this step by selecting multiple datasets in sdimaging.

8.4.8.3 Imaging

Now we are ready to image the calibrated TP array dataset. Here we assume that the observation was performed at 115.27 GHz on the \(^{12}\text{CO}(J = 1-0)\) line, on a target with field ID=0 and SPW=9.

CASA <#>: sdimaging(infiles = 'foo.combined.ms.cal',
                      field = '0',
                      spw = '9',
                      restfreq = '115271.204MHz',
                      nchan = 50, start = '115000MHz', width = '10MHz',
                      gridfunction = 'GJINC',
                      imsize = [50,50],
                      cell = ['10arcsec'],
                      phasecenter = 'J2000 12h22m54.9 +15d49m15',
                      outfile = 'foo.image')

This will create an image cube with 50 channels starting from 115000 MHz at 10 MHz frequency resolution. The phasecenter is set to the coordinate of the source (or whatever center coordinate the user wishes to be the center of the map). If the phasecenter is left blank, then the task will roughly calculate the center automatically using the coordinates in the pointing table. The cell parameter specifies the size of 1 pixel, and should be chosen so that it is about 1/4 to 1/3 of the beam size. The gridfunction is the function used to interpolate the observed positions on the sky to a regular image grid. The default is BOX, but the user may choose from several options.

- SF : A prolate spheroidal function. This function can minimize aliasing effects.
- BOX : A pillbox (box-car) function, defaulting to a kernel box size of 1 pixel.
- PB : Uses the primary beam, assuming an Airy disk. If the dataset is from an ALMA 12m antenna, it will use an effective diameter of 10.7m,
- GAUSS : A gaussian function.
- GJINC : A gaussian convolved with a Jinc function. This function can minimize broadening of the effective beam.

Additional subparameters can be specified for SF, GAUSS, and GJINC which controls the size of the functions. The user can choose what velocity frame to create the images in, but in most cases the default (conversion to LSR frame) should be fine.
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8.5 Using The ASAP Toolkit within CASA

ASAP is included with the CASA installation/build. It is loaded upon start-up, and the ASAP functionality is under the Python 'sd' tool. Note: This means that if you are following the ASAP cookbook or documentation, all of the commands should be invoked with a 'sd.' before the native ASAP command.

The ASAP interface is essentially the same as that of the CASA toolkit, that is, there are groups of functionality (aka tools) which have the ability to operate on your data. Type:

```
sd.AsapLogger   sd.asaplinefind   sd.mask_or
sd.__builtins__ sd.asaplog      sd.matplotlib
sd.__class__    sd.asaplog_post_dec sd.merge
sd.__date__     sd.asapmath     sd.new_asaplot
sd.__delattr__  sd.asapplotter sd-opacity
sd.__dict__     sd.average_time sd-opacity_model
sd.__doc__      sd.calfs        sd.os
sd.__file__     sd.calibrate   sd.page
sd.__format__   sd.calns        sd.parameters
sd.__getattribute__ sd.calps      sd.plotter
sd.__init__     sd.coordinate   sd.plotter2
sd.__name__     sd.dosigref    sd.plylab
sd.__new__      sd.dototalpower sd.rc
sd.__package__  sd.edgemarkersd.rcParams
sd.__path__     sd.env         sd.rcParamsDefault
sd.__reduce__   sd.fitter      sd.rcp
sd.__reduce_ex__ sd.flagplotter sd.re
sd.__repr__     sd.get_revision sd.sbseparator
sd.__revision__ sd.gui         sd.scantable
sd.__setattr__  sd.inspect     sd.selector
sd.__sizeof__   sd.interactivemask sd.setup_env
sd.__str__      sd.ipysupport  sd.simplelinefinder
sd.__subclasshook__ sd.is_asap_cli sd.skydip
sd.__version__  sd.is_asap    sd.splitant
sd.is_asap      sd.is_ipython sd.srcype
sd.is_sequence_or_number sd.linecatalog sd.sys
sd.__n_bools    sd.linefinder sd.toggle_verbose
sd.__to_list    sd.list_files sd.unique
sd.almacal      sd.list_rcparameters sd.utils
sd.apexcal      sd.list_scans sd.version
sd.asapfitter    sd.logging    sd.welcome
sd.asapgrid     sd.mask_and   sd.xyplotter
sd.asapgrid2    sd.mask_not
```

...to see the list of tools.

In particular, the following are essential for most reduction sessions:
• **sd.scantable** - the data structure for ASAP and the core methods for manipulating the data; allows importing data, making data selections, basic operations (averaging, baselines, etc.) and setting data characteristics (e.g., frequencies, etc.).

• **sd.selector** - selects a subset of data for subsequent operations

• **sd.fitter** - fit data

• **sd.plotter** - plotting facilities (uses matplotlib)

The **scantable** functions are used most often and can be applied to both the initial scantable and to any spectrum from that scan table. Type

```
sd.scantable.<TAB>
```

(using TAB completion) to see the full list.

### 8.5.1 Environment Variables

The **asaprc** environment variables are stored in the Python dictionary **sd.rcParams** in CASA. This contains a number of parameters that control how ASAP runs, for both tools and tasks. You can see what these are set to by typing at the CASA prompt:

```python
CASA <2>: sd.rcParams
Out[2]:
{'insitu': True,
 'plotter.axesformatting': 'asap',
 'plotter.colours': '',
 'plotter.decimate': False,
 'plotter.ganged': True,
 'plotter.gui': True,
 'plotter.histogram': False,
 'plotter.linestyles': '',
 'plotter.panelling': 's',
 'plotter.papertype': 'A4',
 'plotter.Stacking': 'p',
 'scantable.autoaverage': True,
 'scantable.freqframe': 'LSRK',
 'scantable.history': True,
 'scantable.parallactify': False,
 'scantable.reference': '.*(e|w|_R)$',
 'scantable.save': 'ASAP',
 'scantable.storage': 'memory',
 'scantable.verbosesummary': False,
 'useplotter': True,
 'verbose': False}
```
The use of these parameters is described in detail in the ASAP Users Guide.

These parameters can be changed through the sd.rc function. Use is described in help sd.rc:

CASA <3>: help(sd.rc)
Help on function rc in module asap:

rc(group, **kwargs)
   Set the current rc params. Group is the grouping for the rc, e.g.
   for scantable.save the group is 'scantable', for plotter.stacking, the
   group is 'plotter', and so on. kwargs is a list of attribute
   name/value pairs, e.g.

   rc('scantable', save='SDFITS')

sets the current rc params and is equivalent to

   rcParams['scantable.save'] = 'SDFITS'

Use rcdefaults to restore the default rc params after changes.

Important Note:
User must use sd.rcParams['scantable.storage']='disk' with care when you call any tool level
functions since some functions may overwrite original data even if you set sd.rcParams['insitu']=False,
which tells the system not to overwrite original data (in contrast, setting sd.rcParams['insitu']
to True forces to overwrite original data). Relevant methods, which may overwrite original data in
the above case, are as follows:

- sd.average_time
- sd.merge
- four operations (+, -, *, /) of sd.scantable instance with scalar or array
- sd.scantable.add
- sd.scantable.clip
- sd.scantable.flag
- sd.scantable.flag_nans
- sd.scantable.flag_row
- sd.scantable.scale
- sd.scantable.recalc_azel
- any setter functions of sd.scantable class (both set_xxx and _setxxx functions)
8.5.2 Data Import

Single dish data sets can be loaded into ASAP using the `scantable` module. The ASAP module, `scantable`, imports single dish data sets in a variety of formats (the CASA Measurement Set, NRO data format, RPFITS, and varieties of SDFITS). An example of import command is:

```
CASA <1>: scans = sd.scantable('foo.ms', average=False)
```

Use the `summary` function to examine the data and get basic information:

```
CASA <8>: scans.summary()
```

The output is printed to the logger. For example,

```plaintext
Scan Table Summary

Project: AGBT06A_018_01
Obs Date: 2006/01/19/01:45:58
Observer: Joseph McMullin
Antenna Name:GBTGREENBANK
Data Records: 512 rows
Obs. Type: OffOn:PSWITCHOFF:TPWCAL
Beams: 1
IFs: 8
Polarisations: 2 (circular)
Channels: 8192
Flux Unit: K
Abscissa: Channel
Selection: none

<table>
<thead>
<tr>
<th>Scan Source</th>
<th>Time range</th>
<th>Int[s]</th>
<th>Record SrcType FreqIDs MolIDs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beam Position (J2000)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>21 OrionS 0</td>
<td>2006/01/19/01:45:58.0 - 01:47:58.2</td>
<td>30.03</td>
<td>[PSOFF, PSOFF:CALON] [0, 1, 2, 3] [0]</td>
</tr>
<tr>
<td>22 OrionS 0</td>
<td>2006/01/19/01:48:38.0 - 01:50:38.2</td>
<td>30.03</td>
<td>[PSON, PSON:CALON] [0, 1, 2, 3] [0]</td>
</tr>
<tr>
<td>23 OrionS 0</td>
<td>2006/01/19/01:51:21.0 - 01:53:21.2</td>
<td>30.03</td>
<td>[PSOFF, PSOFF:CALON] [0, 1, 2, 3] [0]</td>
</tr>
<tr>
<td>24 OrionS 0</td>
<td>2006/01/19/01:54:01.0 - 01:56:01.2</td>
<td>30.03</td>
<td>[PSON, PSON:CALON] [0, 1, 2, 3] [0]</td>
</tr>
<tr>
<td>25 OrionS 0</td>
<td>2006/01/19/02:01:47.0 - 02:03:47.2</td>
<td>30.03</td>
<td>[PSOFF, PSOFF:CALON] [4, 5, 6, 7] [1]</td>
</tr>
<tr>
<td>26 OrionS 0</td>
<td>2006/01/19/02:04:27.0 - 02:06:27.2</td>
<td>30.03</td>
<td>[PSON, PSON:CALON] [4, 5, 6, 7] [1]</td>
</tr>
<tr>
<td>27 OrionS 0</td>
<td>2006/01/19/02:07:10.0 - 02:09:10.2</td>
<td>30.03</td>
<td>[PSOFF, PSOFF:CALON] [4, 5, 6, 7] [1]</td>
</tr>
<tr>
<td>28 OrionS 0</td>
<td>2006/01/19/02:09:51.0 - 02:11:51.2</td>
<td>30.03</td>
<td>[PSON, PSON:CALON] [4, 5, 6, 7] [1]</td>
</tr>
</tbody>
</table>
```

--------------------------------------------------------------------------------
CHAPTER 8. SINGLE DISH DATA PROCESSING

FREQUENCIES: 4

<table>
<thead>
<tr>
<th>ID</th>
<th>IFNO(SPW)</th>
<th>#Chans</th>
<th>Frame</th>
<th>Ch0[MHz]</th>
<th>ChanWid[kHz]</th>
<th>Center[MHz]</th>
<th>POLNOs</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>8192</td>
<td>LSRK</td>
<td>45464.3506</td>
<td>6.10423298</td>
<td>45489.3505</td>
<td>[0, 1]</td>
</tr>
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<td>1</td>
<td>1</td>
<td>8192</td>
<td>LSRK</td>
<td>45275.7825</td>
<td>6.10423298</td>
<td>45300.7824</td>
<td>[0, 1]</td>
</tr>
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<td>2</td>
<td>2</td>
<td>8192</td>
<td>LSRK</td>
<td>44049.9264</td>
<td>6.10423298</td>
<td>44074.9263</td>
<td>[0, 1]</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>8192</td>
<td>LSRK</td>
<td>44141.2121</td>
<td>6.10423298</td>
<td>44166.212</td>
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</tr>
<tr>
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<td>12</td>
<td>8192</td>
<td>LSRK</td>
<td>43937.1232</td>
<td>6.10423356</td>
<td>43962.1231</td>
<td>[0, 1]</td>
</tr>
<tr>
<td>5</td>
<td>13</td>
<td>8192</td>
<td>LSRK</td>
<td>42620.4173</td>
<td>6.10423356</td>
<td>42645.4172</td>
<td>[0, 1]</td>
</tr>
<tr>
<td>6</td>
<td>14</td>
<td>8192</td>
<td>LSRK</td>
<td>41569.9768</td>
<td>6.10423356</td>
<td>41594.9767</td>
<td>[0, 1]</td>
</tr>
<tr>
<td>7</td>
<td>15</td>
<td>8192</td>
<td>LSRK</td>
<td>43397.8198</td>
<td>6.10423356</td>
<td>43422.8196</td>
<td>[0, 1]</td>
</tr>
</tbody>
</table>

MOLCEULES:

<table>
<thead>
<tr>
<th>ID</th>
<th>RestFreq</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>[4.5490258e+10]</td>
<td>[]</td>
</tr>
<tr>
<td>1</td>
<td>[4.3963e+10]</td>
<td>[]</td>
</tr>
</tbody>
</table>

8.5.2.1 General descriptions

The followings are some cautions when using import feature.

- It is important to specify the `antenna` parameter when importing a Measurement Set. For example:

  \[
  \text{CASA<1>: scans = sd.scantable( 'foo.ms', average=False, antenna=0 )}
  \]

  The value of the `antenna` parameter can be either id (integer) or name (string). The default value for `antenna` parameter is 0.

- It is important to use the `average=False` parameter setting as the calibration routines supporting GBT data require all of the individual times and phases.

- GBT data may need some pre-processing prior to using ASAP. In particular, the program which converts GBT raw data into CASA Measurement Sets tends to proliferate the number of spectral windows due to shifts in the tracking frequency; this is being worked on by GBT staff.

- ASAP identifies the observing modes of data sets and the source types, e.g., reference and target sources, by identification numbers. The identification numbers are stored at the SRC-TYPE column in scantable. For example, the ids of target sources of the position switched, Nod, and frequency switched switched data are 0, 2, and 3, respectively. The ids of corresponding reference data of position switched and frequency switched switched modes are 1 and 4, respectively.

- Import of Nobeyama Radio Observatory (NRO) data (in both NEWSTAR and NOSTAR formats) is available.
8.5.2.2 Handling ALMA data

- Using `importasdm` task, ASDM data can be imported to ASAP directly. To do that, you should set `singledish=True` and specify id or name of the antenna by the `antenna` parameter (see § 2.2.1). This functionality is still under testing. You can use previous two step process (import ASDM as MS using `importasdm` task first, then import MS as ASAP format) to import ASDM as ASAP format if you have any problem during direct import.

- If the MS data contain data from multiple single dish antennas you need either, to specify the `antenna` parameter when importing data (sd.scantable) or, to split the data by antenna using `sdsave` for further processing in ASAP. This is because ASAP Scottable stores data only from single antenna. The method `sdsave` splits a Measurement Set by antenna ID and save the tables as scantables containing data from each antenna. The individual scantables will have names in the format of `<outfile>.<antenna name>` or `<prefix>.<antenna name>.asap` if outfile is `<prefix>.asap`.

- From 4.1, ASAP does not convert the frequency reference frame to LSRK when importing data. The frequency reference frame of an imported scantable is the same as that of MS or ASDM (usually TOPO).

- In ALMA, the system temperature ($T_{sys}$) is measured in specific spectral windows for calibration. It is necessary to transfer $T_{sys}$ to the spectral windows for target scans. This can be done using `filltsys` module or `sdcal2` task. Usage of `filltsys` is as follows:

  ```python
  import filltsys
  filltsys.fillTsys( filename='mydata.asap', specif=5, tsysif=1, mode='linear' )
  ```

  If the $T_{sys}$ SPW frequency range does not completely cover the science SPW, the user may choose `extrap = True` to extrapolate the $T_{sys}$ to those channels (default = False). Interpolation in the time domain is linear when possible, but in cases where science observations are not bracketed by $T_{sys}$ scans, then the nearest $T_{sys}$ measurement will be used. The filltsys library can only deal with one SPW at a time, so must be looped over the relevant SPWs.

  The parameter `filename` specifies a name of the data to be processed. Data must be in ASAP format. The spectral windows (=IFs) for target scan and calibration scan should be set using `specif` and `tsysif`, respectively. You can identify those information using `sdlist` task. The `mode` is an interpolation mode along frequency axis. Available options are 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic', and any integer specifying an order of the spline interpolation. Note that `filltsys` will overwrite the data specified by `filename`. See § 8.8.1.2 if you want to use `sdcal2`.

8.5.2.3 Import of NRO data

Importing NRO data is available. Here are some notes on import of NRO data.

- Both NEWSTAR and NOSTAR formats are supported.
• Dual-polarization data is supported.

• An individual IFNO is assigned to each of arrays to identify data from an array. Note that arrays are numbered by successive IFNOs. For example, if observation of three arrays, A01, A03, and A05, are stored in a data set, their IFNOs will be 0, 1, and 2, respectively.

• Developments are mainly focused on recent data, especially for new correlator. Although older data (using AOS) can be imported, there may be an inconsistency with original data.

• The parameter, freqref, controls how frequency reference frame is set. The default is 'rest'. The other option is 'vref'. If freqref is 'vref', frequency reference frame takes from VREF field in input NRO data. This parameter is only available in tool level, i.e. sd.scantable.

• There is a problem on imaging multi-beam data.

8.5.3 Scantable Manipulation

Within ASAP, data is stored in a scantable, which holds all of the observational information and provides functionality to manipulate the data and information. The building block of a scantable is an integration which is a single row of a scantable. Each row contains just one spectrum of a beam, IF and polarization.

Once you have a scantable in ASAP, you can select a subset of the data based on scan numbers, or source names; note that each of these selections returns a new 'scantable' with all of the underlying functionality:

```plaintext
CASA <5>: scan27=scans.get_scan(27)  # Get the 27th scan
CASA <6>: scans20to24=scans.get_scan(range(20,25))  # Get scans 20 - 24
CASA <7>: scansOrion=scans.get_scan('Ori*')  # Get all Orion scans
```

To copy a scantable, do:

```plaintext
CASA <15>: ss=scans.copy()
```

8.5.3.1 Data Selection

In addition to the basic data selection above, data can be selected based on IF, beam, polarization, scan number as well as values such as Tsys. To make a selection create a selector object choose among various selection functions, e.g.,

```plaintext
sel = sd.selector()  # initialize a selector object
# sel.<TAB> will list all options
sel.set_ifs(0)  # select only the first IF of the data
scans.set_selection(sel)  # apply the selection to the data
print scans  # shows just the first IF
```
8.5.3.2 State Information

Some properties of a scantable apply to all of the data, such as spectral units, frequency frame, or Doppler type. This information can be set using the `scantable.set_xxxx` methods. These are currently:

```plaintext
CASA <1>: sd.scantable.set_<TAB>
sd.scantable.set_dirframe sd.scantable.set_selection
sd.scantable.set_doppler sd.scantable.set_sourcename
sd.scantable.set_feedtype sd.scantable.set_sourcetype
sd.scantable.set_fluxunit sd.scantable.set_spectrum
sd.scantable.set_freqframe sd.scantable.set_tsys
sd.scantable.set_instrument sd.scantable.set_unit
sd.scantable.set_restfreqs
```

For example, `sd.scantable.set_fluxunit` sets the default units that describe the flux axis:

```plaintext
scans.set_fluxunit('K') # Set the flux unit for data to Kelvin
```

Choices are 'K' or 'Jy'. Note: the `scantable.set_fluxunit` function only changes the name of the current fluxunit. To change fluxunit, use `scantable.convert_flux` as described in § 8.5.4.2 instead (currently it is necessary to do some gymnastics for non-AT telescopes).

Use `sd.scantable.set_unit` to set the units to be used on the spectral axis:

```plaintext
scans.set_unit('GHz')  # Use GHz as the spectral axis for plots
```

The choices for the units are 'km/s', 'channel', or '*Hz' (e.g. 'GHz', 'MHz', 'kHz', 'Hz'). This does the proper conversion using the current frame and Doppler reference as can be seen when the spectrum is plotted.

Set the frame in which the frequency (spectral) axis is defined by `sd.scantable.set_freqframe`:

```plaintext
CASA <2>: help(sd.scantable.set_freqframe)
Help on method set_freqframe in module asap.scantable:

```

set_freqframe(self, frame=None) unbound asap.scantable.scantable method

Set the frame type of the Spectral Axis.
Parameters:
  frame: an optional frame type, default 'LSRK'. Valid frames are:
  'REST', 'TOPO', 'LSRD', 'LSRK', 'BARY',
  'GEO', 'GALACTIC', 'LGROUP', 'CMB'

Examples:
  scan.set_freqframe('BARY')
```

The most useful choices here are `frame = 'LSRK'` and `frame = 'TOPO'` (what ALMA actually observes in). Note that the 'REST' option is not yet available. The Doppler frame is set with `sd.scantable.set_doppler`:
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CASA <3>: help(sd.scantable.set_doppler)
Help on method set_doppler in module asap.scantable:

```
set_doppler(self, doppler='RADIO') unbound asap.scantable.scantable method
    Set the doppler for all following operations on this scantable.
    Parameters:
        doppler: One of 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
```

Finally, there are a number of functions to query the state of the scantable. These can be found in the usual way:

CASA <4>: sd.scantable.get\_<TAB>
sd.scantable.get_abcissa sd.scantable.get_parangle
sd.scantable.get_antennaname sd.scantable.get_restfreqs
sd.scantable.get_azimuth sd.scantable.get_rms
sd.scantable.get_column_names sd.scantable.get_row
sd.scantable.get_coordinate sd.scantable.get_row_selector
sd.scantable.get_direction sd.scantable.get_scan
sd.scantable.get_directionval sd.scantable.get_selection
sd.scantable.get_elevation sd.scantable.get_sourcename
sd.scantable.get_fit sd.scantable.get_spectrum
sd.scantable.get_fluxunit sd.scantable.get_time
sd.scantable.get_inttime sd.scantable.get_tsys
sd.scantable.get_mask sd.scantable.get_tsysspectrum
sd.scantable.get_mask_indices sd.scantable.get_unit
sd.scantable.get_masklist sd.scantable.get_weather

These include functions to get the current values of the states mentioned above, as well as methods to query the number of scans, IFs, and polarizations in the scantable and their designations. See the inline help of the individual functions for more information.

8.5.3.3 Masks

Several functions (fitting, baseline subtraction, statistics, etc.) may be run on a range of channels (or velocity/frequency ranges). You can create masks of this type using the create_mask function:

```
# spave = an averaged spectrum
spave.set_unit('channel')
rmsmask=spave.create_mask([5000,7000])  # create a region over channels 5000-7000
rms=spave.stats(stat='rms',mask=rmsmask)  # get rms of line free region

rmsmask=spave.create_mask([3000,4000],invert=True)  # choose the region
    # *excluding* the specified channels
```

The mask is stored in a simple Python variable (a list) and so can be manipulated using Python facilities.
8.5.3.4 Scantable Management

scantables can be listed via:

CASA <33>: sd.list_scans()
The user created scantables are:
['scans20to24', 's', 'scan27']

As every scantable will consume memory usage, if you will not use it any longer, you can explicitly remove it via:

del <scantable name>

8.5.3.5 Scantable Mathematics

It is possible to do simple mathematics directly on scantables from the CASA command line using the +, −, *, / operators as well as their cousins +=, −=, *=, /=.

CASA <10>: scan2=scan1+2.0 # add 2.0 to data
CASA <11>: scan *= 1.05 # scale spectrum by 1.05

Operands can be a numerical value and one- or two-dimensional Python list. For list operand, its shape should be conform with the shape of spectral data stored in the scantable. Mathematics between two scantables is also available. In that case, scantables must be conform with each other.

NOTE: In scantable mathematics, scantable must be put on the left. For example:

CASA<12>: scan2=scan1+2.0 # this works
CASA<13>: scan2=2.0+scan1 # this causes an error

8.5.3.6 Scantable Save and Export

ASAP can export scantables in a variety of formats, suitable for reading into other packages. The formats are:

- ASAP – This is the internal format used for ASAP. It is the only format that allows the user to restore the data, fits, etc., without losing any information. As mentioned before, the ASAP scantable is a CASA Table (memory-based table). This function just converts it to a disk-based table. You can access it with the CASA browsetable task or any other CASA table tasks.

- SDFITS – The Single Dish FITS format. This format was designed for interchange between packages but few packages can actually read it.
• ASCII – A simple text based format suitable for the user to process using Python or other means.

• Measurement Set (V2: CASA format) – Saves the data in a Measurement Set. All CASA tasks which use an MS should work with this format.

Scantables are exported by the function, `save`:

```
CASA <19>: scans.save(name='foo.output.ms',format='MS2')
```

### 8.5.4 Calibration

For some observatories, the calibration happens transparently as the input data contains the Tsys measurements taken during the observations. The nominal 'Tsys' values may be in Kelvin or Jansky. The user may wish to apply a Tsys correction or apply gain-elevation and opacity corrections.

#### 8.5.4.1 Tsys scaling

If the nominal Tsys measurement at the telescope is wrong due to incorrect calibration, the `scale` function allows it to be corrected.

```
scans.scale(1.05, tsys=True) # by default only the spectra are scaled
# (and not the corresponding Tsys) unless tsys=True
```

#### 8.5.4.2 Flux and Temperature Unit Conversion

The function, `convert_flux`, is available for converting measurements in Kelvin to Jansky (and vice versa). It converts and scales data to the selected units. The user may need to supply the aperture efficiency, telescope diameter, or the Jy/K factor

```
scans.convert_flux(eta=0.48, d=35.) # Unknown telescope
scans.convert_flux(jypk=15) # Unknown telescope (alternative)
scans.convert_flux() # known telescope (mostly AT telescopes)
scans.convert_flux(eta=0.48) # if telescope diameter known
```

#### 8.5.4.3 Gain-Elevation and Atmospheric Optical Depth Corrections

At higher frequencies, it is important to make corrections for atmospheric opacity and gain-elevation effects. **NOTE:** Currently, the MS to scantable conversion does not adequately populate the azimuth and elevation in the `scantable`. As a result, one must calculate these via:
scans.recalc_azel()
Computed azimuth/elevation using
Position: [882590, -4.92487e+06, 3.94373e+06]
Time: 01:48:38 Direction: 05:35:13.5 -05.24.08.2
  => azel: 154.696 43.1847 (deg)
Time: 01:48:38 Direction: 05:35:13.5 -05.24.08.2
  => azel: 154.696 43.1847 (deg)
Time: 01:48:38 Direction: 05:35:13.5 -05.24.08.2
  => azel: 154.696 43.1847 (deg)
Time: 01:48:38 Direction: 05:35:13.5 -05.24.08.2
  => azel: 154.696 43.1847 (deg)
Time: 01:48:38 Direction: 05:35:13.5 -05.24.08.2
  => azel: 154.696 43.1847 (deg)
...  

With the correct Az/El, the effect of atmospheric opacity can be corrected for a given opacity by:

```python
scans.opacity(tau=0.09)  # Opacity from which the correction factor:
# exp(tau*zenith-distance)
```

### 8.5.4.4 Comprehensive calibration function

A new function **calibrate** is introduced for comprehensive calibration. This function calls appropriate calibration scheme by referring the antenna name in the meta-data. The user just specify input scantable and calmode parameter that indicates calibration mode ('ps', 'nod', 'fs', 'otf', or 'otfraster'). For GBT data, **calibrate** function calls one of the functions listed in the next section depending upon the value of **calmode**. For APEX data, **apexcal** function is called, while for ALMA data, that contains string 'ALMA' or 'OSF' in its antenna name, **almacal** function is called. Calibration scheme for these two telescopes are essentially same, and can be expressed as,

\[
T_a = T_{sys} \times (ON - OFF) / OFF,
\]

where OFF scans are interpolated in time if possible.

### 8.5.4.5 Calibration of GBT data

Data from the GBT are uncalibrated and come as sets of integrations representing the different phases of a calibration cycle (e.g., on source, calibration on, on source, calibration off, on reference, calibration on; on reference, calibration off). Currently, there are a number of routines emulating the standard GBT calibration (in GBTIDL):

- **calps** - calibrate position switched data
- **caldfs** - calibrate frequency switched data
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- **calnod** - calibration nod (beam switch) data

All these routines calibrate the spectral data to antenna temperature adopting the GBT calibration method as described in the GBTIDL calibration document available at:


There are two basic steps:

First, determine system temperature using a noise tube calibrator (**sd.dototalpower()**)

For each integration, the system temperature is calculated from CAL noise on/off data as:

\[
T_{sys} = T_{cal} \times \left( \frac{<ref_{calon}>}{<ref_{calon}-ref_{caloff}>} \right) + \frac{T_{cal}}{2}
\]

*ref* which refers to reference data and the spectral data are averaged across the bandpass. Note that the central 80% of the spectra are used for the calculation.

Second, determine antenna temperature (**sd.dosigref()**)

The antenna temperature for each channel is calculated as:

\[
T_a(\nu) = T_{sys} \times \frac{sig(\nu)-ref(\nu)}{ref(\nu)}
\]

where 
\[
sig = \frac{1}{2}(sig_{calon} + sig_{caloff}), ref = \frac{1}{2}(ref_{calon} + ref_{caloff}).
\]

Each calibration routine may be used as:

```python
scans=sd.scantable('inputdata',False) # create a scantable called 'scans'
calibrated_scans = sd.calps(scans,[scanlist]) # calibrate scantable with position-switched # scheme
```

**Note**: For calps and calnod, the **scanlist** must be scan pairs in correct order as these routines only do minimal checking.

### 8.5.5 Averaging

One can average polarizations in a scantable using the **sd.scantable.average_pol** function:

```python
averaged_scan = scans.average_pol(mask,weight)
```

**where**:

**Parameters**:

- **mask**: An optional mask defining the region, where the averaging will be applied. The output will have all specified points masked.
- **weight**: Weighting scheme. 'none' (default), 'var' (1/var(spec weighted), or 'tsys' (1/Tsys**2 weighted).
Example:

```python
spave = stave.average_pol(weight='tsys')
```

One can also average scans over time using `sd.average_time`:

```python
sd.average_time(scantable, mask, scanav, weight, align)
```

where:

**Parameters:**

- **one scan** or comma separated scans
- **compel**: if True, enable averaging of multi-resolution spectra.
- **mask**: an optional mask (only used for 'var' and 'tsys' weighting)
- **scanav**: True averages each scan separately.
  False (default) averages all scans together,
- **weight**: Weighting scheme.
  'none' (mean no weight)
  'var' (1/var(spec) weighted)
  'tsys' (1/Tsys**2 weighted)
  'tint' (integration time weighted)
  'tintsys' (Tint/Tsys**2)
  'median' (median averaging)
- **align**: align the spectra in velocity before averaging. It takes the time of the first spectrum in the first scantable as reference time.

Example:

```python
stave = sd.average_time(scans, weight='tintsys')
```

Note that alignment of the velocity frame should be done before averaging if the time spanned by the scantable is long enough. This is done through the `align=True` option in `sd.average_time`, or explicitly through the `sd.scantable.freq_align` function, e.g.,

CASA <62>: sc = sd.scantable('foo_PM01.asap',False)
CASA <63>: sc.freq_align()
Aligned at reference Epoch 2006/01/19/01:49:23 (UTC) in frame LSRK
CASA <64>: av = sd.average_time(sc)

The time averaging can also be applied to multiple scantables. For example, such data might have been taken on different days. The `sd.average_time` function takes multiple scantables as input. However, if they are taken at significantly different times (different days for example), then `sd.scantable.freq_align` must be used to align the velocity scales to the same time, e.g.,

CASA <65>: sc1 = sd.scantable('foo.scan0.asap',False)
CASA <66>: sc2 = sd.scantable('foo.scan1.asap',False)
CASA <67>: sc1.freq_align()
Aligned at reference Epoch 2006/01/19/01:49:23 (UTC) in frame LSRK
CASA <68>: sc2.freq_align(reftime='2006/01/19/01:49:23')
Aligned at reference Epoch 2006/01/19/01:54:46 (UTC) in frame LSRK
CASA <69>: scav = sd.average_time(sc1,sc2)
### 8.5.6 Spectral Smoothing

Smoothing on data can be done as follows:

```python
scantable.smooth(kernel, # type of smoothing: 'hanning' (default), 'gaussian',
width, # width in pixels (ignored for hanning); FWHM for gaussian.
order, # the order of the polynomial (only valid when kernel='poly')
plot, # plot the original and the smoothed spectra
insitu) # if False (default), do smoothing in-situ; otherwise,
# make new scantable
```

Example:

```python
# spave is an averaged spectrum
spave.smooth('boxcar',5) # do a 5 pixel boxcar smooth on the spectrum
sd.plotter.plot(spave) # should see smoothed spectrum
```

### 8.5.7 Spectral Regridding

Regridding of channels in spectra is available as follows:

```python
scantable.regrid_channel(width, # The channel width (float) of regridded spectra
    # in the current spectral unit.
insitsu) # if False (default), do smoothing in-situ;
# otherwise, make new scantable
```

Example:

```python
# spave is an averaged spectrum
spave.set_unit('channel') # Use channel width
spave.regrid_channel(5) # regrid 5 channels in each spectrum to one channel
sd.plotter.plot(spave) # should see smoothed spectrum
```

### 8.5.8 Baseline Fitting

CASA offers a variety of functions for baseline fitting: polynomial, cubic spline, Chebyshev polynomial and sinusoid are available. The available baseline functions in ASAP are,

- `sd.scantable.poly_baseline` and `sd.scantable.auto_poly_baseline`
- `sd.scantable.cspline_baseline` and `sd.scantable.auto_cspline_baseline`
- `sd.scantable.chebyshev_baseline` and `sd.scantable.auto_chebyshev_baseline`
- `sd.scantable.sinusoid_baseline` and `sd.scantable.auto_sinusoid_baseline`
- `sd.scantable.sub_baseline`
The function, `sd.scantable.poly_baseline`, carries out a polynomial baseline fit, given a mask of channels (if desired):

```python
msk=scans.create_mask([100,400],[600,900])
scans.poly_baseline(msk,order=1)
```

This example fits spectra by the first order polynomial to the selected channels and subtract the polynomial from the full spectrum.

The full set of parameters available in `sd.scantable.poly_baseline` is:

```python
scans.poly_baseline(mask,order,insitu,clipthresh,clipniter,plot,...):
```

Parameters:
- `mask`: an optional mask
- `order`: the order of the polynomial (default is 0)
- `insitu`: if False a new scantable is returned. Otherwise, the scaling is done in-situ
  The default is taken from .asaprc (False)
- `clipthresh`: Clipping threshold. (default is 3.0, unit: sigma)
- `clipniter`: maximum number of iteration of 'clipthresh'-sigma clipping (default is 0)
- `plot`: plot the fit and the residual. In this each individual fit has to be approved, by typing 'y' or 'n'
- `getresidual`: if False, returns best-fit values instead of residual. (default is True)
- `showprogress`: show progress status for large data.
  default is True.
- `minnrow`: minimum number of input spectra to show.
  default is 1000.
- `outlog`: Output the coefficients of the best-fit function to logger (default is False)
- `blfile`: Name of a text file in which the best-fit parameter values to be written
  (default is "": no file/logger output)
- `csvformat`: if True blfile is csv-formatted, default is False.
- `bltable`: name of a baseline table where fitting results
  (coefficients, rms, etc.) are to be written.
  if given, fitting results will NOT be output to scantable (insitu=True) or None will be returned (insitu=False).
  (default is "": no table output)

The parameter `order` defines the order of polynomial fit. The other parameters are common to all baseline functions except for `sd.scantable.sub_baseline`.

Two types of operations are available in baseline functions, i.e., on-the-fly baseline subtraction and generation of a baseline table. When `bltable` is not specified (as in the above example), baseline fits are subtracted from spectra on-the-fly and the results are stored as a scantable.
On the other hand, when `bltable` is specified, the fit parameters are stored in a baseline table, but not actually subtracted from spectra. The baseline table is a CASA Table containing baseline function type, parameters, fitting results (coefficients, rms, etc.) and so on. The function, `sd.scantable.apply_bltable`, is available to apply it to a scantable to invoke baseline subtraction.

e.g.,

```python
msk = scans.create_mask([100,400],[600,900])
scans.poly_baseline(msk,order=1,bltable='foo.blcal')
blls = scans.apply_bltable(inbltable='foo.blcal',insitu=False)
```

The function, `auto_poly_baseline`, is available for automatic selection of line-free channels. It detects the line-free emission in each spectrum and subtracts a polynomial fit to it from the spectrum. Line-free channels are selected using linefinder. Three additional parameters are available to define criteria of line-free channels.

- **edge**: an optional number of channel to drop at the edge of spectrum. If only one value is specified, the same number will be dropped from both sides of the spectrum. Default is to keep all channels. Nested tuples represent individual edge selection for different IFs (a number of spectral channels can be different).
- **threshold**: the threshold used by line finder. It is better to keep it large as only strong lines affect the baseline solution.
- **chan_avg_limit**: the maximum number of consecutive spectral channels to average during the search of weak and broad lines. The default is no averaging (and no search for weak lines). If such lines can affect the fitted baseline (e.g. a high order polynomial is fitted), increase this parameter (usually values up to 8 are reasonable). Most users of this method should find the default value sufficient.

Note, these parameters are in common for all the baseline fitting functions that use linefinder (`auto_*_baseline`).

The functions, `chebyshevBaseline` and `auto_chebyshevBaseline`, are for baselining using the Chebyshev polynomials with or without linefinder, respectively.

The functions, `csplineBaseline` and `auto_csplineBaseline`, are for cubic spline fitting with or without using linefinder, respectively. The unique parameters for these are as follows:

- **npiece**: Number of pieces. (default is 2)

The functions, `sinusoidBaseline` and `auto_sinusoidBaseline`, are for sinusoidal fitting with or without using linefinder, respectively. The unique parameters for these are as follows:
nwave: the maximum wave number of sinusoids within a range of spectral width multiplied by maxwavelength. The default is 3 (i.e., sinusoids with wave number of 0 (constant), 1, 2, and 3 are used for fitting). Also it is possible to explicitly specify all the wave numbers to be used, by giving a list including them (e.g., [0, 1, 2, 15, 16]).

maxwavelength: the longest sinusoidal wavelength. The default is 1.0 (unit: spectral range)

The function, `sd.scantable.sub_baseline`, allows user to specify baseline function type and parameters per spectrum, e.g.,

```python
sub_baseline(blinfo=[{'row':0, 'blfunc':'poly', 'order':5, 'masklist':[[10,350],[352,510]]},
{'row':1, 'blfunc':'cspline', 'npiece':3, 'masklist':[3,16],[19,404],[407,511]}
])
```

In this example, the first spectrum (row=0) is fitted with polynomial of order=5 and the next one (row=1) is fitted with cubic spline consisting of 3 pieces.

The 'goodness' of baseline fitting can be evaluated. A new sd tool function `sd.scantable.calc_aic` is available to calculate several values known as model selection criteria for a given spectrum and a baseline function. It can calculate Akaike Information Criterion (AIC), the corrected Akaike Information Criterion (AICc), Bayesian Information Criterion (BIC) and the Generalised Cross Validation (GCV).

Note the parameter `plot` is ignored for cubic spline and sinusoidal fitting.

### 8.5.9 Line Fitting

Multi-component Gaussian fitting is done by creating a fitting object, specifying fit parameters and finally fitting the data. Fitting can be done on a `scantable` selection or an entire `scantable` using the `auto_fit` function.

```python
# spave is an averaged spectrum
f=sd.fitter() # create fitter object
msk=spave.create_mask([3928,4255]) # create mask region around line
f.set_function(gauss=1) # set a single gaussian component
f.set_scan(spave,msk) # set the scantable and region

f.fit() # fit
f.plot(residual=True) # plot residual
f.get_parameters() # retrieve fit parameters

# 0: peak = 0.786 K, centre = 4091.236 channel, FWHM = 70.586 channel
```
# area = 59.473 K channel
f.store_fit('orions_hc3n_fit.txt')  # store fit
#
# To specify an initial guess:
# set a single gaussian component
# set initial guesses for Gaussian
# for first component (0)
# (peak,center,fwhm)
#
# For multiple components set
# initial guesses for each, e.g.,
# set two gaussian components
# set initial guesses for Gaussian
# for first component (0)
# set initial guesses for Gaussian
# for second component (1)

8.5.10 Plotting

8.5.10.1 ASAP plotter

The ASAP plotter uses the same Python matplotlib library as in CASA (for x-y plots). It is accessed via the:

```
sd.plotter<TAB> # see all functions (omitted here)
sd.plotter.plot(scans) # the workhorse function
sd.plotter.set<TAB>
sd.plotter.set_abcissa sd.plotter.set_layout sd.plotter.set_ordinate
sd.plotter.set_colors sd.plotter.set_legends sd.plotter.set_panelling
sd.plotter.set_colours sd.plotter.set_linestyles sd.plotter.set_range
sd.plotter.set_data sd.plotter.set_margin sd.plotter.set_selection
sd.plotter.set_font sd.plotter.set_mask sd.plotter.set_stacking
sd.plotter.set_histogram sd.plotter.set_mode sd.plotter.set_title
```

Spectra can be plotted at any time when `refresh = True` (default) is selected, and it will attempt to do the correct layout depending on whether it is a set of scans or a single scan. You can switch off verbose plotting by `refresh = False` in tool parameters for faster plotting in scripts:

```
sd.plotter.set_data(scan,refresh=False)  # set scantable to plot
   # this should be done at first.
sd.plotter.set_mode(stacking='time',panelling='if',refresh=False)
sd.plotter.set_range(ystart=-1.0,yend=5.0,refresh=False)
sd.plotter.plot()  # actual plotting
```

The details of the plotter display (matplotlib) are detailed in the earlier section.
8.5.10.2 Line Catalog

ASAP allows loading a custom line catalog in ASCII format. The ASCII text file must have at least 4 columns with Molecule name, frequency in MHz, frequency error and intensity (any units). If the molecule name contains any spaces, they must be wrapped in quotes "". A sample of the ASCII catalog is shown below.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Frequency</th>
<th>Error</th>
<th>Intensity</th>
</tr>
</thead>
<tbody>
<tr>
<td>H2D+</td>
<td>3955.2551</td>
<td>228.8818</td>
<td>-7.1941</td>
</tr>
<tr>
<td>H2D+</td>
<td>12104.7712</td>
<td>177.1558</td>
<td>-6.0769</td>
</tr>
<tr>
<td>H2D+</td>
<td>45809.2731</td>
<td>118.3223</td>
<td>-3.9494</td>
</tr>
<tr>
<td>CH</td>
<td>701.6811</td>
<td>0.0441</td>
<td>-7.1641</td>
</tr>
<tr>
<td>CH</td>
<td>724.7709</td>
<td>0.0456</td>
<td>-7.3912</td>
</tr>
<tr>
<td>CH</td>
<td>3263.7940</td>
<td>0.1000</td>
<td>-6.3501</td>
</tr>
<tr>
<td>CH</td>
<td>3335.4810</td>
<td>0.1000</td>
<td>-6.0304</td>
</tr>
</tbody>
</table>

You can load the ASCII line catalog, for example, if it is called my_custom_linecat.txt, by following command:

```python
mycatlog = sd.linecatlog('my_custom_linecat.txt')
```

Use `sd.plotter.plot_line` to overlay the line catalog on the plot. (Currently over-plotting line catalog works only spectra plotted in frequency.)

```python
scans.set_unit('GHz')
sd.plotter.plot(scans)
sd.plotter.plot_lines(mycatlog)
```

The following are some useful functions to control the line catalog access. See ASAP User Guide for more complete descriptions.

```python
mycatlog.save('my_custom_linecat.tbl') # save to the internal table format
mycatlog.set_frequency_limits(100,115,'GHz') # set a frequency range for line selection
mycatlog.set_name('*OH') # select all alcohols
```

8.5.10.3 Plotter2

Plotter2 is a generic PGLOT-based plotter newly available from this release. Developed as a light-weight plotter to be used in the ALMA SD pipeline, plotter2 can also be used by general CASA users to draw a graph containing multiple viewports in a flexible manner. It can be accessed via:

```python
sd.plotter2<TAB> # see all functions (omitted here)
sd.plotter2.plot # the workhorse function
sd.plotter2.set<TAB>
```
CHAPTER 8. SINGLE DISH DATA PROCESSING

To simply plot a set of data points given as $x=[x_1, x_2, ...]$ and $y=[y_1, y_2, ...]$ on the screen, just type the following lines:

```python
p = sd.plotter2()
p.set_data(x, y)
p.plot()
```

If you want to save the above plot as a file `foo.png`, run `p.save("foo.png")` instead of `p.plot()`.

A bit more complicated plot can be made by using functions of plotter2. Figure 8.1 shows an example. This figure is generated by the following commands:

```python
s = sd.scantable(filename='foo.PM01.asap', average=False)
s_fit = s.cspline_baseline(npiece=5, insitu=False, showprogress=False, getresidual=False)
s_res = s.cspline_baseline(npiece=5, insitu=False, showprogress=False, getresidual=True)
x = range(s.nchan())
p = sd.plotter2() # create a plotter2 object
for i in range(5):
    for j in range(5):
        # get y-value of data -------
        rowid = i*5+j
        y1 = s.get_spectrum(rowid) # raw spectrum
        y2 = s_fit.get_spectrum(rowid) # baseline computed by cubic spline fitting
        y3 = s_res.get_spectrum(rowid) # baseline-subtracted spectrum

        # position of viewports ----
        vp_xmin1 = 0.2*j+0.03
        vp_xmax1 = 0.2*j+0.115
        vp_xmin2 = vp_xmax1
        vp_xmax2 = 0.2*j+0.2
        vp_ymin = 0.2*(4-i)+0.012
        vp ymax = 0.2*(4-i)+0.18

        # setup left panels -------
        # set data and line attributes
```
p.set_vp(vp_xmin1, vp_xmax1, vp_ymin, vp_ymax)
p.set_data(x, y1)  # set first data
p.set_line("blue")  # to be connected with blue line
p.set_data(x, y2)  # set second data
p.set_line("red", 4)  # to be connected with thick red line
(ymin, ymax) = p.get_yrange()  # get y-range of left panel

# channel-masked regions
p.set_xmask(100, 108, "orange", "outline", 1)
p.set_xmask(100, 108, "orange", "hatched", 1, 0.3)
p.set_xmask(353, 401, "orange", "outline", 1)
p.set_xmask(353, 401, "orange", "hatched", 1, 0.3)

# labels and title
p.set_fontsize(0.4)
p.set_xlabel("channel", "normal", 0.45, vp_xmin1, 0.2*(4-i))
p.set_ylabel("Intensity (arbitrary)", "normal", 0.45, \\
    vp_xmin1-0.02, 0.5*(vp_ymin+vp_ymax))
p.set_title("Fit: row = "+str(rowid), "normal", 0.48, \\
    0.5*(vp_xmin1+vp_xmax1), vp_ymax+0.005)

# change background color
if rowid in [4, 13, 14]: p.set_vpbgcolor("lightgray")
# hide some panels
if rowid in [7, 21, 22, 23, 24]: p.hide_vp()

# setup right panels -------
p.set_vp(vp_xmin2, vp_xmax2, vp_ymin, vp_ymax)
p.set_yrange(ymin, ymax)  # make y-range common with left panel

p.set_data(x, y3)
p.set_line("blue")

p.set_fontsize(0.4)
p.set_ynumlocation(""")  # no numbering in y-axis
p.set_title("Reduced: row = "+str(rowid), "normal", 0.48, \\
    0.5*(vp_xmin2+vp_xmax2), vp_ymax+0.005)

# change background color
if rowid in [4, 13, 14]: p.set_vpbgcolor("lightgray")
# hide some panels
if rowid in [7, 21, 22, 23, 24]: p.hide_vp()

p.save("foo.png")

del s, s_fit, s_res, p

For details, see inline help of each plotter2 function.
8.5.11 Setting/Getting Rest Frequencies

The rest frequencies used in the data can be retrieved by `sd.scantable.get_restfreqs()` and set to new values by `sd.scantable.set_restfreqs()`. The CASA version of ASAP now can store multiple rest frequencies for each IF.

```python
scans.get_restfreqs()  # retrieve current rest frequencies
#{0: [45490258000.0]}
```

All of the rest frequencies currently set to the data are listed in python dictionary for each MOLECULE_ID.

Here is an example of setting multiple rest frequencies for spectra of a particular IF:
#Select IFs, then set rest frequencies,
    sel=sd.selector()
    sel.setifs(0)
    scans.set_selection(sel)
    scans.set_restfreqs([45490258000.0, 45590258000.0, 45690258000.0])

NOTE: there is no functionality yet to select a specific rest frequency to apply to a specific line, etc. Currently, the first one in the list of the rest frequencies is used for such calculation.

8.5.12 Single Dish Spectral Analysis Use Case With ASAP Toolkit

Below is a script that illustrates how to reduce single dish data using ASAP within CASA. First a summary of the dataset is given and then the script.

```python
# MeasurementSet Name: /home/rohir3/jocular/SD/OrionS_rawACSmod MS Version 2
# Project: AGBT06A_018_01
# Observation: GBT(1 antennas)
#
#Data records: 256 Total integration time = 1523.13 seconds
# Observed from 01:45:58 to 02:11:21
#
#Fields: 4
# ID Name Right Ascension Declination Epoch
# 0 OrionS 05:15:13.45 -05.24.08.20 J2000
# 1 OrionS 05:35:13.45 -05.24.08.20 J2000
# 2 OrionS 05:15:13.45 -05.24.08.20 J2000
# 3 OrionS 05:35:13.45 -05.24.08.20 J2000
#
#Spectral Windows: (8 unique spectral windows and 1 unique polarization setups)
# SpwID #Chans Frame Ch1(MHz) Resoln(kHz) TotBW(kHz) Ref(MHz) Corrs
# 0 8192 LSRK 45464.3506 6.10423298 50005.8766 45489.3536 RR LL HC3N
# 1 8192 LSRK 45275.7825 6.10423298 50005.8766 45300.7854 RR LL HN15CO
# 2 8192 LSRK 44049.9264 6.10423298 50005.8766 44074.9293 RR LL CH3OH
# 3 8192 LSRK 44141.2121 6.10423298 50005.8766 44166.2151 RR LL HCCC15N
# 12 8192 LSRK 43937.1232 6.10423356 50005.8813 43962.1261 RR LL HNCO
# 13 8192 LSRK 42620.4173 6.10423356 50005.8813 42645.4203 RR LL H15NCO
# 14 8192 LSRK 41569.9768 6.10423356 50005.8813 41594.9797 RR LL HNC18O
# 15 8192 LSRK 43397.8198 6.10423356 50005.8813 43422.8227 RR LL SiO
#
# Scans: 21-24 Setup 1 HC3N et al
# Scans: 25-28 Setup 2 SiO et al

casapath=os.environ['AIPSPATH']

#ASAP script
---
# COMMENTS
import asap as sd #import ASAP package into CASA
```
#Orion-S (SiO line reduction only)

#Notes:
#scan numbers (zero-based) as compared to GBTIDL

#changes made to get to OrionS_rawACSmod
#modifications to label sig/ref positions

```
os.environ['AIPSPATH']=casapath  #set this environment variable back - ASAP changes it
```

```
s=sd.scantable('OrionS_rawACSmod',False)#load the data without averaging
```

```
s.summary()  #summary info
s.set_fluxunit('K')  # make 'K' default unit
scal=sd.calps(s,[20,21,22,23])  # Calibrate HC3N scans
scal.recalc_azel()  # recalculate az/el to
scal.opacity(0.09)  # do opacity correction
sel=sd.selector()  # Prepare a selection
sel.set_ifs(0)  # select HC3N IF
scal.set_selection(sel)  # get this IF
```

Figure 8.2: Multi-panel display of the scantable. Sub-panels are displayed per scan. There are two spectra in each scan indicating two polarization (RR and LL).
Figure 8.3: Two panel plot of the calibrated spectra. The GBT data have a separate scan for the SOURCE and REFERENCE positions so scans 20, 21, 22 and 23 result in these two spectra.

```python
cal = sd.average_time(scal, weight='tintsys')  # average in time
spave = stave.average_pol(weight='tsys')  # average polarizations; Tsys-weighted (1/Tsys**2) average
sd.plotter.plot(spave)  # plot

spave.smooth('boxcar', 5)  # boxcar 5
spave.auto_poly_baseline(order=2)  # baseline fit order=2
sd.plotter.plot(spave)  # plot

spave.set_unit('GHz')
sd.plotter.plot(spave)
sd.plotter.set_histogram(hist=True)  # draw spectrum using histogram
sd.plotter.axhline(color='r', linewidth=2)  # zline
sd.plotter.save('orions_hc3n_reduced.eps')  # save postscript spectrum

rmsmask = spave.create_mask([5000, 7000])  # get rms of line free regions
base_rms = spave.stats(stat='rms', mask=rmsmask)  # rms

linemask = spave.create_mask([3900, 4200])  # LINE
line_max = spave.stats('max', linemask)  # IF[0] = 0.918
line_sum = spave.stats('sum', linemask)  # IF[0] = 64.994
```
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Figure 8.4: Calibrated spectrum with a line at zero (using histograms).

```python
line_median = spave.stats('median', linemask)  # IF[0] = 0.091
line_mean = spave.stats('mean', linemask)     # IF[0] = 0.210

# Fitting
spave.set_unit('channel')  # set units to channel
sd.plotter.plot(spave)    # plot spectrum
f = sd.fitter()            # Fitting
msk = spave.create_mask([3928, 4255])  # create region around line
f.set_function(gauss=1)   # set a single gaussian component
f.set_scan(spave, msk)    # set the data and region for the fitter
f.fit()                   # fit
f.plot(residual=True)     # plot residual

f.get_parameters()        # retrieve fit parameters
# 0: peak = 0.786 K, centre = 4091.236 channel, FWHM = 70.586 channel
# area = 59.473 K channel
f.store_fit('orions_hc3n_fit.txt')  # store fit
# Save the spectrum
spave.save('orions_hc3n_reduced', 'ASCII', True)  # save the spectrum
```
8.6 Single Dish Imaging

Single dish imaging is supported within CASA using standard tasks and tools. The data must be in the Measurement Set format. Once there, you can use the `im` (imager) tool to create images:

**Tool example:**

```python
scans.save('outputms','MS2') # Save your data from ASAP into an MS
im.open('outputms') # open the data set
im.selectvis(nchan=901,start=30,step=1, spwid=0,field=0) # choose a subset of the data
               # (just the key emission channels)
dir='J2000 17:18:29 +59.31.23' # set map center
im.defineimage(nx=150,cellx='1.5arcmin',
                phasenumber=dir,mode='channel',start=30,
                nchan=901,step=1) # define image parameters
               # (note it assumes symmetry if ny,celly
               #  aren't specified)
im.setoptions(ftmachine='sd',cache=1000000000) # choose SD gridding
im.setsdoptions(convsupport=4) # use this many pixels to support the
                                # gridding function used
                                # (default=prolate spheroidal wave function)
im.makeimage(type='singledish',
             image='FLS3a_HI.image') # make the image
```

8.6.1 Single Dish Imaging Use Case With ASAP Toolkit

The data summary and the script are given below.

```bash
# Project: AGBT02A_007_01
# Observation: GBT(1 antennas)

# Telescope Observation Date Observer Project
# GBT [ 4.57539e+09, 4.5754e+09]Lockman AGBT02A_007_01
# GBT [ 4.57574e+09, 4.57575e+09]Lockman AGBT02A_007_02
# GBT [ 4.5831e+09, 4.58313e+09]Lockman AGBT02A_031_12

# Thu Feb 1 23:15:15 2007 NORMAL ms::summary:
# Data records: 76860 Total integration time = 7.74277e+06 seconds
# Observed from 22:05:41 to 12:51:56

# Thu Feb 1 23:15:15 2007 NORMAL ms::summary:
# Fields: 2
# ID Name Right Ascension Declination Epoch
# 0 FLS3a 17:18:00.00 +59.30.00.00 J2000
# 1 FLS3b 17:18:00.00 +59.30.00.00 J2000

# Thu Feb 1 23:15:15 2007 NORMAL ms::summary:
# Spectral Windows: (2 unique spectral windows and 1 unique polarization setups)
```
CHAPTER 8. SINGLE DISH DATA PROCESSING

<table>
<thead>
<tr>
<th># SpwID</th>
<th>#Chans Frame Ch1(MHz)</th>
<th>Resoln(kHz)</th>
<th>TotBW(kHz)</th>
<th>Ref(MHz)</th>
<th>Corrs</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1024 LSRK 1421.89269</td>
<td>2.44140625</td>
<td>2500</td>
<td>1420.64269</td>
<td>XX</td>
</tr>
<tr>
<td>1</td>
<td>1024 LSRK 1419.39269</td>
<td>2.44140625</td>
<td>2500</td>
<td>1418.14269</td>
<td>XX</td>
</tr>
</tbody>
</table>

# FLS3 data calibration
# this is calibration part of FLS3 data
#
casapath=os.environ['AIPSPATH']
import asap as sd
os.environ['AIPSPATH']=casapath

print '---Import---'
s=sd.scantable('FLS3_all_newcal_SP',false) # read in MeasurementSet

print '---Split---'

# splitting the data for each field
s0=s.get_scan('FLS3a*') # split the data for the field of interest
s0.save('FLS3a_HI.asap') # save this scantable to disk (asap format)
del s0 # free up memory from scantable

print '---Calibrate---'
s=sd.scantable('FLS3a_HI.asap') # read in scantable from disk (FLS3a)
s.set_fluxunit('K') # set the brightness units to Kelvin
scanns = s.getscannos() # get a list of scan numbers
sn=list(scanns) # convert it to a list
print "No. scans to be processed:", len(scanns)
res=sd.calfs(s,sn) # calibrate all scans listed using frequency

print '---Save calibrated data---'
res.save('FLS3a_calfs', 'MS2') # Save the dataset as a MeasurementSet

print '---Image data---'
im.open('FLS3a_calfs') # open the data set
im.selectvis(nchan=901,start=30,step=1, spvid=0,field=0) # choose a subset of the data
dir="J2000 17:18:29 +59.31.23' # (just the key emission channels)
im.defineimage(nx=150,cellx='1.5arcmin', phasecenter=dir,mode='channel',start=30, nchan=901,step=1) # define image parameters
im.setoptions(ftmachine='sd',cache=1000000000) # choose SD gridding
im.setsdoptions(convsupport=4) # use this many pixels to support the gridding function used
im.makeimage(type='singledish',image='FLS3a_HI.image') # make the image
Figure 8.5: FLS3a HI emission. The display illustrates the visualization of the data cube (left) and the profile display of the cube at the cursor location (right); the Tools menu of the Viewer Display Panel has a Spectral Profile button which brings up this display. By default, it grabs the left-mouse button. Pressing down the button and moving in the display will show the profile variations.
8.7 Known Issues, Problems, Deficiencies and Features

The Single-Dish calibration and analysis package within CASA is still very much under development. Not surprisingly, there are a number of issues with ASAP and the SDtasks that are known and under repair. Some of these are non-obvious "features" by the way ASAP and sd are implemented, or limitations of the current Python tasking environment. Some are functions that have yet to be implemented. These currently include:

1. \texttt{sd.plotter}
   The method, \texttt{sd.plotter.set\_range()} sets the same range for multiple panels, while we would like it to be able to set the range for each independently, including the default ranges.
   The \texttt{sd.plotter} object remembers things throughout the session and thus can easily get confused. For example, one must reset the range \texttt{sd.plotter.set\_range()} if set manually.
   This behaviour is not always expected, but is a consequence of having \texttt{sd.plotter} be its own object that is fed data and commands.

   Eventually we would like the capability to interactively set things using the plots, like select frequency ranges, identify lines, start fitting.

2. \texttt{sd.selector}
   The selector object only allows one selection of each type. It would be nice to be able to make a union of selections (without resorting to query) for the \texttt{set\_name}. Note that the others like scans and IFs work off lists, which is fine. We should make \texttt{set\_name} work off lists of names.

3. \texttt{sd.scantable}
   The \texttt{scantable.verbosesummary} asaprc parameter (e.g. in \texttt{sd.rcParams}) does nothing.
   GBT data has an undefined parameter fluxunit ("" that should be 'K'), an incorrect freqframe ('LSRK' that is is really 'TOPO'), and reference frequency (set to that of the first IF only).
   The \texttt{sd.scantable.freq\_align} does not yet work correctly.

4. \texttt{sd} general issues
   There should be a \texttt{sdhelp} equivalent of \texttt{toolhelp} and \texttt{tasklist} for the \texttt{sd} tools and tasks.
   The current output of ASAP is verbose and controlled by setting \texttt{sd.rcParams['verbose']=False} (or True). We will make some of the output less cryptic.
   We will strip off leading and trailing whitespace on string parameters.

5. SDtasks general issues
   The SDtasks work with files saved onto disk in one of the scantable supported formats. It might be useful to work with scantables in memory (passing the objects) but this would require changes to the tasking system. Note that this behavior is consistent throughout the casapy tasks.
6. sdcal (and sdreduce)
   `averageall=True` is still experimental since the test was insufficient because of a lack of test data.

7. sdfit
   Only way to handle multi-IFs is to set `fitmode='auto'` (linefinder is applied for each spectra and derives initial guesses). For `fitmode='list'`, there are no way to give initial guesses for each IFs by hand.

8. sdplot
   Only handles the included JPL line catalog. Also, see `sd.plotter` issues above.

9. GBT raw SDFITS data
   The SDtasks and `sd.scantable` are able to handle GBT raw SDFITS data format (version 1.3) data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

**8.8 Single Dish Analysis Tasks**

A set of SDtasks is available for simplifying basic reduction activities. The list currently includes:

- **sdcal** — select, calibrate, and average SD data
- **sdcal2** — generate sky and tsys caltables for SD data, and apply them
- **sdaverage** — average and smooth SD spectra
- **sdbaseline** — fit/remove spectral baselines from SD data
- **sdreduce** — `sdcal`, `sdaverage`, and `sdbaseline` combined to perform standard single dish processing all at once
- **sdcoadd** — merge/co-add multiple SD data
- **sdflag** — channel/row flagging of SD spectra
- **sdflagmanager** — enable list, save, restore, delete and rename flag version files
- **sdfit** — line fitting to SD spectra
- **sdgrid** — convolve map data onto regularly spaced grid
- **sdimaging** — create an image from the total power or spectral data
- **sdlist** — print a summary of a SD dataset
- **sdmath** — do simple arithmetic for SD spectra
• **sdplot** — plotting of SD spectra, including overlay of line catalog data
• **sdsave** — save SD data to a different format
• **sdscale** — scale SD data
• **sdstat** — compute statistics of regions of SD spectra
• **sdtpimaging** — do a simple calibration and create an image from the total power raster scans
• **sdimprocess** — remove the 'scanning noise' from raster scanned image
• **msmoments** — compute moments from spectral data

All of the SDtasks, except those related to imaging (sdtpimaging, sdimprocess, and sdimprocess), work from a file on disk rather than from a scantable in memory as the ASAP toolkit does (see §8.5). Inside the tasks we invoke a call to sd.scantable to read the data. The scantable objects do not persist within CASA after completion of the tasks and are destroyed to free up memory.

Three tasks sdcal, sdaverage, and sdbaseline are the workhorses for the calibration, selection, averaging, baseline fitting, and smoothing. The output datasets for each task are written to a file on disk. Alternatively, one can use the task sdreduce to perform all of the steps in the three tasks described. Its operation is controlled by three main "mode" parameters: calmode (which selects the type of calibration, if any, to be applied), kernel (which selects the smoothing), and blfunc (which selects baseline fitting). There are also parameters controlling the selection such as field, spw, scan, timeaverage, scanaverage, and polaverage. Note that sdreduce can be run with calmode='none' to allow re-selection or writing out of data that is already calibrated. There is a "wiring diagram" of the dataflow and control inputs for sdreduce shown in Figure 8.6.

The SDtasks support the import and export file formats supported by ASAP itself. For import, this includes: ASAP (scantables), MS (CASA Measurement Set), RPFITS, SDFITS (version 1.3) and NRO data format. For export, this includes: ASAP (scantables), MS (CASA Measurement Set), ASCII (text file), SDFITS (a flavor of SD FITS). The sdsave task is available exclusively for exporting with these data selection options. The sdocadd task is available to merge data in separate data files into one. A brief summary of the data in a file is found in the sdlist task help.

Plotting of spectra is handled in the sdplot task. It also offers some selection, averaging, and smoothing options in case you are working from a dataset that has not been split or averaged. Note that there is some rudimentary plotting capability in many of SDtasks, controlled through the plotlevel parameter, to aid in the assessment of the performance of these tasks.

Scaling of the spectra and Tsys is available in the sdscale task. For arithmetic operations of spectra in separate scantables, sdmath has been added.

Calculation of statistics on spectral regions is available in the sdstat task. Results are passed by a Python dictionary return variable. The statistics of spectra can also be calculated via msmoments task. The input of msmoments task must be in CASA Measurement Set format. The task newly creates Measurement Set to store statistics values.
Basic Gaussian line-fitting is handled by the *sdfit* task. It can deal with the simpler cases, and offers some automation as well as interactive selection of fitting region, but more complicated fitting
is best accomplished through the toolkit (sd.fitter).

Basic interactive and non-interactive channel and row flagging are available in the sdflag tasks. The flags in the input file is updated by default, i.e., outfile='' and overwrite=True. Otherwise, a new file is created to store dataset with updated flag information.

Limited total power data analysis functionality is available through the task sdtpimaging. A single dish image data cube can be created using sdimaging, which also handles total power imaging. These tasks directly access the Measurement Set without converting it to scantable format. The sdimprocess is intended to remove the 'scanning noise' from single dish images by either the 'Basket-Weaving' or 'Pressed-out' methods.

The task sdgrid, which convolves map data onto regularly spaced grid, is available. The task can be used as imaging tool although output of this task is not an image but a scantable. Also, it can be regarded as a data averaging tool with various weight.

Although the Measurement Set can store data from multiple antennas even if it consists of only single-dish spectra (auto-correlation data), the scantable cannot distinguish data from multiple antennas. It causes a problem when the user processes the Measurement Set using SDtasks. Therefore, id or name of the antenna that the user want to process must be explicitly specified if the input dataset for SDtasks is Measurement Set. This can be done by antenna parameter. By default (antenna=0), data associate with antenna id 0 is imported. The antenna parameter takes no effect for other input data formats.

8.8.1 SDtask Summaries

The following are the list of parameters and brief descriptions of each of the SDtasks. These descriptions are also contained in help <taskname>.

The interface of the following tasks are modified in CASA 4.2.2: sdbaseline, sdcal, sdcal2, sdfit, sdflag, sdgrid, sdimaging, sdmath, sdplot, sreduce, ssave, and sstat. Additionally, a new task called sdaverage is available. Task sdsmooth has been incorporated in the sdaverage and removed. The tasks with old interfaces are available with name {taskname}old and kept by CASA 4.3 release. Users are advised to update existing scripts.

8.8.1.1 sdcal

Keyword arguments:
infile -- name of input SD dataset
antenna -- select an antenna name or ID
    default: 0
    example: 'PM03'
    NOTE this parameter is effective only for MS input
fluxunit -- units for line flux
    options: 'K','Jy',''
    default: '' (keep current fluxunit in data)
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WARNING: For GBT data, see description below.

>>> fluxunit expandable parameter
telescopeparam -- parameters of telescope for flux conversion
    options: (str) name or (list) list of gain info
    default: '' (none set)
    example: if telescopeparam='', it tries to get the telescope name from the data.
    Full antenna parameters (diameter,ap.eff.) known to ASAP are 'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43', 'CEDUNA','HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
telescopeparam=[104.9,0.43] diameter(m), ap.eff.
telescopeparam=[0.743] gain in Jy/K
telescopeparam='FIX' to change default fluxunit

field -- select data by field IDs and names
    default: '' (use all fields)
    example: field='3C2*' (all names starting with 3C2)
    field='0,4,5~7' (field IDs 0,4,5,6,7)
    field='0,3C273' (field ID 0 or field named 3C273)
    this selection is in addition to the other selections to data

spw -- select data by IF IDs (spectral windows)
    NOTE this task only supports IF ID selction and ignores channel selection.
    default: '' (use all IFs and channels)
    example: spw='3,5,7' (IF IDs 3,5,7; all channels)
    spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
    spw='30~45GHz' (IF IDs with the center frequencies in range 30-45GHz; all channels)
    this selection is in addition to the other selections to data

scan -- select data by scan numbers
    default: '' (use all scans)
    example: scan='21~23' (scan IDs 21,22,23)
    this selection is in addition to the other selections to data

pol -- select data by polarization IDs
    default: '' (use all polarizations)
    example: pol='0,1' (polarization IDs 0,1)
    this selection is in addition to the other selections to data

calmode -- calibration mode
    options: 'ps','nod','otf','otfraster', 'fs','quotient','none'
    default: 'ps'
example: choose mode 'none' if you have already calibrated and want to correct for atmospheric opacity defined by tau.

>>> calmode expandable parameter

calmode expandable parameter

fraction -- edge marker parameter of 'otf' and 'otfraster'. Specify a number of OFF integrations (at each side of the raster rows in 'otfraster' mode) as a fraction of total number of integrations. In 'otfraster' mode, number of integrations to be marked as OFF, n_off, is determined by the following formula,

\[
\text{n}_{\text{off}} = \text{floor}(\text{fraction} \times \text{n}),
\]

where \(n\) is number of integrations per raster row. Note that \(n_{\text{off}}\) from both sides will be marked as OFF so that twice of specified fraction will be marked at most. For example, if you specify fraction='10\%', resultant fraction of OFF integrations will be 20\% at most.

In 'otf' mode, \(n_{\text{off}}\) is given by,

\[
\text{n}_{\text{off}} = \text{floor}(\text{fraction} \times \text{n}),
\]

where \(n\) is number of total integrations. \(n_{\text{off}}\) is used as criterion of iterative marking process. Therefore, resulting total number of OFFs will be larger than \(n_{\text{off}}\). In practice, fraction is a geometrical fraction of edge region. Thus, if integrations are concentrated on edge region (e.g. some of Lissajous patterns), then resulting \(n_{\text{off}}\) may be unexpectedly large.

default: '10\%

options: '20\%' in string style or float value less than 1.0 (e.g. 0.15).

'aio' is available only for 'otfraster'.

noff -- edge marking parameter for 'otfraster'.

It is used to specify a number of OFF scans near edge directly. Value of noff comes before setting by fraction. Note that \(n_{\text{off}}\) from both sides will be marked as OFF so that twice of specified noff will be marked at most.

default: -1 (use fraction)
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options: any positive integer
width -- edge marking parameter for 'otf'.
Pixel width with respect to a median spatial
separation between neighboring two data in time.
Default will be fine in most cases.
default: 0.5
options: float value
elongated -- edge marking parameter for 'otf'.
Set True only if observed area is elongated
in one direction.
options: (bool) True, False
default: False
markonly -- set True if you want to save data just after
edge marking (i.e. uncalibrated data) to see
how OFF scans are defined.
options: (bool) True, False
default: False
plotpointings -- load plotter and plot pointing directions of
ON and OFF scans.
options: (bool) True, False
default: False
tau -- the zenith atmospheric optical depth for correction
default: 0.0 (no correction)
verify -- interactively verify the results of calibration.
When verify = True, for the first six on-source spectra (at
max), spectra before and after the calibration are displayed
in a plot window. At the prompt there are two choices of
action: 'Y' (accept the operation for whole dataset),
'N' (reject the operation and finish task).
Note that when the operation is rejected by 'N',
no operation is done to the spectrum/spectra.
options: (bool) True,False
default: False
outfile -- name of output file
default: '' (<infile>_cal)
outform -- output file format
options: 'ASAP','MS2', 'ASCII','SDFITS'
default: 'ASAP'
NOTE the ASAP format is easiest for further sd
processing; use MS2 for CASA imaging.
If ASCII, then will append some stuff to
the outfile name
overwrite -- overwrite the output file if already exists
options: (bool) True,False
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**Default:** False

**Note:** this parameter is ignored when outform='ASCII'

**Plotlevel -- control for plotting of results**

**Options:** (int) 0, 1, 2, and their negative counterparts

**Default:** 0 (No plotting)

**Example:**
- `plotlevel=1;` plot calibrated spectra
- `plotlevel=2;` additionally list data before and after operation.
- `plotlevel<0` as abs(plotlevel), e.g. `-1` => hardcopy of final plot (will be named `<outfile>_calspec.eps`

**Description:**

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**How to Choose Calmode**

---------------------

For position switching calibration, the user should choose appropriate calibration mode depending on the data. Use case for each mode is as follows:

- **'ps':** position switch (including OTF) with explicit reference (OFF) scans
- **'otf':** non-raster OTF scan without explicit OFFs (e.g. Lissajous, double circle, etc.) intends to calibrate fast scan data
- **'otfraster':** raster OTF scan without explicit OFFs

So, if the data contains explicit reference scans, 'ps' should be used. Otherwise, 'otfraster' and 'otf' are appropriate for raster OTF and non-raster OTF, respectively. In 'otf' and 'otfraster' modes, the task first try to find several integrations near edge as OFF scans, then the data are calibrated using those OFFs. If the observing pattern is raster, you should use the 'otfraster' mode to calibrate data. Otherwise, the 'otf' mode should be used. For detail about edge marking, see inline help of sd.edgemarker module and its methods.

Those modes are designed for OTF observations without explicit OFF scans. However, these modes should work even if explicit reference scans exist. In this case, explicit reference scans will be ignored and scans near edges detected by edge marker will be used as reference.

Except for how to choose OFFs, the procedure to derive calibrated spectra is common for the above three modes. Selected (or preset) OFF integrations are separated by its continuity in time domain, averaged in each segment, then interpolated to timestamps for ON integrations. Effectively, it means that OFF integrations are averaged by each OFF
scans for 'ps' mode, averaged by either ends of each raster row for 'otfraster' mode, averaged by each temporal segments of detected edges for 'otf' mode. The formula for calibrated spectrum is

\[ T_{sys} \times \frac{(ON - OFF)}{OFF}. \]

The 'fs' mode is for frequency switch calibration. Currently, only GBT frequency switch data is supported.

The 'quotient' mode is special mode for "AT" telescopes, namely ANNF MOPRA. It assumes that observing sequence looks like "target, reference, target, reference,..." and it derives calibrated spectrum as

\[ T_{sys} \times \frac{ON}{OFF}, \]

slightly different from position switch modes.

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FLUX UNIT CONVERSION
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The task is able to convert flux unit between K and Jy. To do that, fluxunit and its subparameter telescopeparam must be properly set. The fluxunit should be 'Jy' or 'K' depending on what unit input data is and what unit you want to convert. If given fluxunit is different from the unit of input data, unit conversion is performed. The telescopeparam is used to specify conversion factor. There are three ways to specify telescopeparam: 1) set Jy/K conversion factor, 2) set telescope diameter, D, and aperture efficiency, \( \eta \), separately, and 3) 'FIX' mode (only change the unit without converting spectral data).

If you give telescopeparam as a list, then if the list has a single float it is assumed to be the gain in Jy/K (case 1), if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively (case 2).

See the above parameter description as well as note on 'FIX' mode below for details.

There are two special cases that don't need telescopeparam for unit conversion. Telescope name is obtained from the data.
1) ASAP (sd tool) recognizes the conversion factor (actually D and \( \eta \)) for the "AT" telescopes, namely ATNF MOPRA telescope, until 2004.
2) The task does know D and \( \eta \) for GBT telescope.

If you wish to change the fluxunit, by leaving the sub-parameter telescopeparam unset (telescopeparam=''), it will use internal telescope parameters for flux conversion for the data from AT telescopes and it
will use an approximate aperture efficiency conversion for the GBT data.

Note that sdcal assumes that the fluxunit is set correctly in the data already. If not, then set telescopeparam='FIX' and it will set the default units to fluxunit without conversion. Note also that, if the data in infile is an ms from GBT and the default flux unit is missing, this task automatically fixes the default fluxunit to 'K' before the conversion.

-------
WARNING
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For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

Notes:

- Direction information is not considered when data is averaged. Use sdgrid task if you want to take care of direction information when averaging.

- Calibration of frequency switching data is now fully implemented, including folding.

- Additional calibration algorithms are implemented. These are the 'Chopper-Wheel' calibration and the one adopted in APEX telescope (an advanced version of classical 'Chopper-Wheel' method).

8.8.1.2 sdcal2

Keyword arguments:
infile -- Name of input SD dataset

calmode -- Calibration mode. If you want to generate calibration table or apply existing calibration tables, set calmode to simple string. On the other hand, if you want to calibrate data on-the-fly, you have to set calmode to a composite calmode string separated by comma. So far, sky calibration has three types, 'ps', 'otf', and 'otfraster'. If observation is configured to observe reference position, calmode must be 'ps'. Otherwise, 'otf' or 'otfraster' should be used depending on observing pattern. If observing pattern is raster scan, calmode must be 'otfraster' while 'otf' must be used when observing pattern is non-raster (e.g., Lissajous).

options: 'ps','otf','otfraster','tsys','apply'
default: 'ps'

example: Here is an example for composite calmode.
    'ps,apply' (do sky cal and apply)
    'ps,tsys,apply' (do sky and Tsys cal and apply)

>>> calmode expandable parameter

fraction -- Edge marker parameter of 'otf' and 'otfraster'.
Specify a number of OFF integrations (at each side of the raster rows in 'otfraster' mode) as a fraction of total number of integrations.
In 'otfraster' mode, number of integrations to be marked as OFF, n_off, is determined by the following formula,

\[ n_{off} = \text{floor}(\text{fraction} \times n), \]

where n is number of integrations per raster row. Note that n_off from both sides will be marked as OFF so that twice of specified fraction will be marked at most. For example, if you specify fraction='10%', resultant fraction of OFF integrations will be 20% at most.
In 'otf' mode, n_off is given by,

\[ n_{off} = \text{floor}(\text{fraction} \times n), \]

where n is number of total integrations.

n_off is used as criterion of iterative marking process. Therefore, resulting total number of OFFs will be larger than n_off. In practice, fraction is a geometrical fraction of edge region. Thus, if integrations are concentrated on edge region (e.g. some of Lissajous patterns), then resulting n_off may be unexpectedly large.

default: '10%'
options: '20%' in string style or float value less than 1.0 (e.g. 0.15).
    'auto' is available only for 'otfraster'.

noff -- Edge marking parameter for 'otfraster'.
It is used to specify a number of OFF spectra near edge directly. Value of noff comes before setting by fraction. Note that n_off from both sides will be marked as OFF so that twice of specified noff will be marked at most.
default: -1 (use fraction)
options: any positive integer

width -- Edge marking parameter for 'otf'.
Pixel width with respect to a median spatial separation between neighboring two data in time.
Default will be fine in most cases.
default: 0.5
options: float value

elongated -- Edge marking parameter for 'otf'.
Set True only if observed area is elongated in one direction.
default: False

tsysavg -- Whether Tsys is averaged in spectral axis or not.
default: False
options: (bool) True, False

tsysspw -- list of IF IDs (spectral windows) and their channel ranges of averaging for Tsys calibration.
It does no effect if you don't want to do Tsys calibration. the user is able to specify channel range for averaging (effective if tsysavg is True).
default: '' (auto-detect tsys spws)
example: tsysspw='3,5,7' (IF IDs 3,5,7; all channels)
tsysspw='<2' (IF IDs less than 2; all channels)
tsysspw='1:0~100' (IF ID1; between channels 0 and 100)

applytable -- List of sky/Tsys calibration tables you want to apply.
default: ''

interp -- Interpolation method in time and frequency axis.
Set comma separated method strings if you want to use different interpolation in time and frequency.
options: 'linear', 'cspline', 'nearest', any numeric string indicating an order of polynomial.
default: '' (linear in time and frequency)
example: 'linear,cspline' (linear in time, cubic spline in frequency)
  'linear,3' (linear in time, third order polynomial in frequency)
  'nearest' (nearest in time and frequency)

spwmap -- Dictionary defining transfer of Tsys calibration. Key must be IFNO for Tsys and its value must be a list of IFNOs for science target.
default: {}
example: {1: [5,6], 3: [7,8]}
Tsps in IFNO1 is transferred to IFNO5, 6 
while Tsys in IFNO3 is to IFNO7, 8.

**field** -- select data by field IDs and names 
default: '' (use all fields)
example: field='3C2*' (all names starting with 3C2)
field='0,4,5~7' (field IDs 0,4,5,6,7)
field='0,3C273' (field ID 0 or field named 3C273)
this selection is in addition to the other selections to data

**spw** -- select data by IF IDs (spectral windows) 
NOTE this task only supports IF ID selction and ignores channel 
selection.
default: '' (use all IFs and channels)
example: spw='3,5,7' (IF IDs 3,5,7; all channels)
spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
spw='30~45GHz' (IF IDs with the center frequencies in range 30-45GHz; all channels)
this selection is in addition to the other selections to data

**scan** -- select data by scan numbers 
default: '' (use all scans)
example: scan='21~23' (scan IDs 21,22,23)
this selection is in addition to the other selections to data

**pol** -- select data by polarization IDs 
default: '' (use all polarizations)
example: pol='0,1' (polarization IDs 0,1)
this selection is in addition to the other selections to data

**outfile** -- Name of output file 
NOTE if you omit, behavior of the task depends on calmode. 
If calmode includes 'apply', then omitting outfile indicates 
that infile is overwritten by the calibrated data. In this case, 
you have to set overwrite to True. If calmode doesn't include 
'apply', omitting outfile indicates that the task will use 
default outfile name based on infile and predefined suffix 
('_sky' for sky, '_tsys' for Tsys).
default: '' (<infile>_<suffix> for calibration 
while overwrite infile for apply mode)

**overwrite** -- overwrite the output file if already exists 
options: (bool) True,False
default: False
NOTE this parameter is ignored when outform='ASCII'

**DESCRIPTION:**

Task sdcal2 is an implementation of a calibration scheme like as 
interferometry, i.e., generate caltables and apply them. Available 
calibration modes are 'ps', 'otf', 'otfraster', and 'tsys'. Those
modes generates caltables for sky or Tsys calibration. Those caltables can be applied to the data by using calmode 'apply'.

The above three calibration modes, 'ps', 'otf', and 'otfraster', generate sky calibration tables. The user should choose appropriate calibration mode depending on the data. Use case for each mode is as follows:

'ps': position switch (including OTF) with explicit reference (OFF) spectra
'otf': non-raster OTF scan without explicit OFFs (e.g. Lissajous, double circle, etc.) intends to calibrate fast scan data
'otfraster': raster OTF scan without explicit OFFs

So, if the data contains explicit reference spectra, 'ps' should be used. Otherwise, 'otfraster' and 'otf' are appropriate for raster OTF and non-raster OTF, respectively. In 'otf' and 'otfraster' modes, the task first try to find several integrations near edge as OFF spectra, then the data are calibrated using those OFFs. If the observing pattern is raster, you should use the 'otfraster' mode to calibrate data. Otherwise, the 'otf' mode should be used. For detail about edge marking, see inline help of sd.edgemark module and its methods. Those modes are designed for OTF observations without explicit OFF spectra. However, these modes should work even if explicit reference spectra exist. In this case, these spectra will be ignored and spectra near edges detected by edge marker will be used as reference.

Except for how to choose OFFs, the procedure to derive calibrated spectra is common for the above three modes. Selected (or preset) OFF integrations are separated by its continuity in time domain, averaged in each segment, then interpolated to timestamps for ON integrations. Effectively, it means that OFF integrations are averaged by each OFF spectrum for 'ps' mode, averaged by either ends of each raster row for 'otfraster' mode, averaged by each temporal segments of detected edges for 'otf' mode. The formula for calibrated spectrum is

\[ \text{Tsys} \times \frac{(\text{ON} - \text{OFF})}{\text{OFF}}. \]

You can calibrate data on-the-fly like sdcal task by setting calmode to a composite calmode string separated by comma.
For example, calmode='ps,apply' means doing sky calibration and apply it on-the-fly. In this case, caltable is generated as a temporary plain table and will be deleted at the end. Allowed calibration modes in this task is as follows:

\begin{verbatim}
ps
  generate sky caltable using 'ps' mode
otf
  generate sky caltable using 'otf' mode
otfraster
  generate sky caltable using 'otfraster' mode
tsys
  generate tsys caltable
apply
  apply caltables specified by applytable parameter
ps,apply
  generate temporary sky caltable using 'ps' mode and apply it. also apply caltables specified by applytable
ps,tsys,apply
  generate temporary sky caltable using 'ps' mode as well as temporary tsys caltable, and apply them.
otf,apply
  generate temporary sky caltable using 'otf' mode and apply it. also apply caltables specified by applytable
otf,tsys,apply
  generate temporary sky caltable using 'otf' mode as well as temporary tsys caltable, and apply them.
otfraster,apply
  generate temporary sky caltable using 'otfraster' mode and apply it. also apply caltables specified by applytable
otfraster,tsys,apply
  generate temporary sky caltable using 'otfraster' mode as well as temporary tsys caltable, and apply them.
\end{verbatim}

There are several control parameters for sky/Tsys calibration and application of caltables. See the above parameter description.

In ALMA, Tsys measurement is usually done using different spectral setup from spectral windows for science target. In this case, sdcal2 transfers Tsys values to science spectral windows in the application stage. To do that, the user has to give a list of spectral windows for Tsys measurement as well as mapping between spectral windows for Tsys measurement and science target. These can be specified by parameters 'tsysspw' and 'spwmap', which are defined as subparameters of 'calmode'. For example, suppose that Tsys measurements for science windows 17, 19,
21, and 23 are done in spw 9, 11, 13, and 15, respectively. In this case, tsysspw and spwmap should be specified as follows:

```python
tsysspw = '9,11,13,15'
spwmap = {9:[17],11:[19],13:[21],15:[23]}
```

Below is an example of full specification of task parameters for calmode of 'ps,tsys,apply':

```python
default(sdcal2)
infile = 'foo.asap'
calmode = 'ps,tsys,apply'
spw = ''
tsysspw = '9,11,13,15'
spwmap = {9:[17],11:[19],13:[21],15:[23]}
outfile = 'bar.asap'
sdcal2()
```

Note that, in contrast to applycal task, spwmap must be a dictionary with Tsys spectral window as key and a list of corresponding science spectral window as value. Note also that the parameter 'spw' should not be used to specify a list of spectral windows for Tsys measurement. It is intended to select data to be calibrated so that the list should contain spectral windows for both science target and Tsys measurement. The task will fail if you use 'spw' instead of 'tsysspw'.

For Tsys calibration, the user is able to choose whether Tsys is averaged in spectral axis or not. If tsysavg is False (default), resulting Tsys is spectral value. On the other hand, when tsysavg is True, Tsys is averaged in spectral axis before output. The channel range for averaging is whole channels by default. If channel range is specified by tsysspw string, it is used for averaging. The user can specify channel range with ms selection syntax. For example,

```python
tsysspw = '1:0-100'
```

specifies spw 1 for Tsys calibration and channel range between channel 0 and 100 for averaging. You can specify more than one ranges per spw.

```python
tsysspw = '1:0-100;200-400'
```

In this case, selected ranges are between 0 and 100 plus 200 and 400. Note that even if multiple ranges are selected, the task average whole ranges together and output single averaged value. You can specify multiple spws by separating comma.
tsysspw = ’1:0~100,3:400~500’

Note that specified channel range is ignored if tsysavg is False.

8.8.1.3 sdaverage

Keyword arguments:
infile -- name of input SD dataset
antenna -- select an antenna name or ID
default: 0
example: ‘PM03’
NOTE this parameter is effective only for MS input
field -- select data by field IDs and names
default: ’’ (use all fields)
example: field=’3C2*’ (all names starting with 3C2)
field=’0,4,5~7’ (field IDs 0,4,5,6,7)
field=’0,3C273’ (field ID 0 or field named 3C273)
this selection is in addition to the other selections to data
spw -- select data by IF IDs (spectral windows)
NOTE this task only supports IF ID selection and ignores channel
selection.
default: ’’ (use all IFs and channels)
example: spw=’3,5,7’ (IF IDs 3,5,7; all channels)
spw=’<2’ (IF IDs less than 2, i.e., 0,1; all channels)
spw=’30~45GHz’ (IF IDs with the center frequencies in range 30-45GHz; all channels)
this selection is in addition to the other selections to data
scan -- select data by scan numbers
default: ’’ (use all scans)
example: scan=’21~23’ (scan IDs 21,22,23)
this selection is in addition to the other selections to data
pol -- select data by polarization IDs
default: ’’ (use all polarizations)
example: pol=’0,1’ (polarization IDs 0,1)
this selection is in addition to the other selections to data
timeaverage -- average spectra over time
options: (bool) True, False
default: False

>>>timeaverage expandable parameter
tweight -- weighting for time averaging
options: ’var’ (1/var(spec) weighted)
’tsys’ (1/Tsys**2 weighted)
’tint’ (integration time weighted)
'tintsy' (Tint/Tsys**2)
'median' (median averaging)
default: 'tintsy'

scanaverage -- average spectra within a scan number
when True, spectra are NOT averaged over
different scan numbers.
options: (bool) True, False
default: False

averageall -- average multi-resolution spectra
spectra are averaged by referring
their frequency coverage
default: False

polaverage -- average spectra over polarizations
options: (bool) True, False
default: False

>>>polaverage expandable parameter
pweight -- weighting for polarization averaging
options: 'var' (1/var(spec) weighted)
'tsys' (1/Tsys**2 weighted)
default: 'tsys'

kernel -- type of spectral smoothing kernel
options: '', 'hanning', 'gaussian', 'boxcar', 'regrid'
default: '' (no smoothing)

>>>kernel expandable parameter
kwidth -- width of spectral smoothing kernel
options: (int) in channels
default: 5
example: 5 or 10 seem to be popular for boxcar
ignored for hanning (fixed at 5 chans)
(0 will turn off gaussian or boxcar)
chanwidth -- channel width of regridded spectra
default: '5' (in channels)
example: '500MHz', '0.2km/s'

verify -- interactively verify the results of smoothing for each
spectrum. When verify = True, for each input spectrum,
spectra before and after the operation are displayed
in a plot window. At the prompt there are four choices
of action: 'Y' (accept the operation and continue to
the next input spectrum), 'N' (reject the operation
and continue to the next input spectrum), 'A' (accept
the current operation and continue non-interactively),

...
and 'R' (reject the current operation and exit from operation).
Note that when the operation is rejected by 'N' or 'R', no smoothing is done to the spectrum/spectra.

Options:
- **outfile** -- name of output file
  - Default: '' (<infile>_sm)
- **outform** -- output file format
  - Options: 'ASAP', 'MS2', 'ASCII', 'SDFITS'
  - Default: 'ASAP'
  - NOTE the ASAP format is easiest for further sd processing; use MS2 for CASA imaging.
  - If ASCII, then will append some stuff to the outfile name
- **overwrite** -- overwrite the output file if already exists
  - Options: (bool) True,False
  - Default: False
  - NOTE this parameter is ignored when outform='ASCII'
- **plotlevel** -- control for plotting and summary of smoothing results
  - Options: (int) 0, 1, 2, and their negative counterparts
  - Default: 0 (no plotting)
  - Example: plotlevel=1; plot spectra before and after smoothing
    - plotlevel=2; additionally list data before and after operation.
    - plotlevel<0 as abs(plotlevel), e.g.
    - -1 => hardcopy of final plot (will be named <outfile>_smspec.eps)

AVERAGING OF SPECTRA

Task sdaverage has two modes of averaging spectra, i.e., time and polarization average.

When timeaverage=True, spectra are averaged over time for each IF (spectral window), polarization, and beam, independently. Note that, by default (scanaverage=False), timeaverage=True averages spectra irrespective of scan IDs.

It is possible to average spectra separately for each scan ID by setting a sub-parameter scanaverage=True.

For example, the combination of parameters: scan='0~2', timeaverage=True, and scanaverage=False: averages spectra in scan ID 0 through 2 all together to a spectrum,
scanaverage=True : averages spectra per scan ID and end up with three
spectra from scan 0, 1, and 2.

When polaverage=True, spectra are averaged over polarization for each IF (spectral window) and beam. Note that, so far, time averaging is automatically switched on when polaverage is set to True. This behavior is not desirable and will be discarded in future.

-------
WARNING
-------

For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

8.8.1.4 sdbaseline

Keyword arguments
infile -- name of input SD dataset
antenna -- select an antenna name or ID
  default: 0
  example: 'PM03'
  NOTE this parameter is effective only for MS input
fluxunit -- units for line flux
  options: 'K','Jy',''
  default: '' (keep current fluxunit in data)
  WARNING: For GBT data, see description below.
  >>> fluxunit expandable parameter
  telescopeparam -- parameters of telescope for flux conversion
  options: (str) name or (list) list of gain info
  default: '' (none set)
  example: if telescopeparam='', it tries to get the telescope name from the data.
  Full antenna parameters (diameter,ap.eff.) known to ASAP are
  'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43', 'CEDUNA','HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
  telescopeparam=[104.9,0.43] diameter(m), ap.eff.
  telescopeparam=[0.743] gain in Jy/K
  telescopeparam='FIX' to change default fluxunit
  see description below
field -- select data by field IDs and names
  default: '' (use all fields)
example: field='3C2*' (all names starting with 3C2)
    field='0,4,5~7' (field IDs 0,4,5,6,7)
    field='0,3C273' (field ID 0 or field named 3C273)
this selection is in addition to the other selections to data

spw -- select data by IF IDs (spectral windows)/channels
default: '' (use all IFs and channels)
example: spw='3,5,7' (IF IDs 3,5,7; all channels)
    spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
    spw='30~45GHz' (IF IDs with the center frequencies in range 30-45GHz; all channels)
    spw='0:5~61' (IF ID 0; channels 5 to 61; all channels)
    spw='3:10~20;50~60' (select multiple channel ranges within IF ID 3)
    spw='3:10~20,4:0~30' (select different channel ranges for IF IDs 3 and 4)
    spw='1"4;6:15~48' (for channels 15 through 48 for IF IDs 1,2,3,4 and 6)
this selection is in addition to the other selections to data

>>> spw expandable parameter

restfreq -- the rest frequency
available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz', 'kHz', and 'Hz' are available.
a list can be used to set different rest frequencies for each IF. the length of list input must be number of IFs. dictionary input should be a pair of line name and frequency with keys of 'name' and 'value', respectively. values in the dictionary input follows the same manner as as for single float or string input.
example: 345.796
    '1420MHz'
    [345.8, 347.0, 356.7]
    ['345.8MHz', '347.0MHz', '356.7MHz']
    [{'name': 'CO', 'value': 345}]

frame -- frequency reference frame
options: 'LSRK', 'TOPO', 'LSRD', 'BARY', 'GALACTO', 'LGROUP', 'CMB'
default: '' (keep current frame in data)

doppler -- doppler convention (effective only when spw is in velocity unit)
options: 'RADIO', 'OPTICAL', 'Z', 'BETA', or 'GAMMA'
default: '' (keep current doppler setting in data)

timerange -- select data by time range
default: '' (use all)
example: timerange = 'YYYY/MM/DD/hh:mm:ss~YYYY/MM/DD/hh:mm:ss'
Note: YYYY/MM/DD can be dropped as needed:
timerange='09:14:00~09:54:00' # this time range
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```
timerange='09:44:00' # data within one integration of time
timerange='>10:24:00' # data after this time
timerange='09:44:00+00:13:00' # data 13 minutes after time
this selection is in addition to the other selections to data
```

```
scan -- select data by scan numbers
  default: '' (use all scans)
ex: scan='21~23' (scan IDs 21,22,23)
this selection is in addition to the other selections to data
```

```
pol -- select data by polarization IDs
  default: '' (use all polarizations)
ex: pol='0,1' (polarization IDs 0,1)
this selection is in addition to the other selections to data
```

```
tau -- the zenith atmospheric optical depth for correction
  default: 0.0 (no correction)
```

```
maskmode -- mode of setting additional channel masks
```

```
options: 'auto', 'list', or 'interact'
  default: 'auto'
ex: maskmode='auto' runs linefinder to detect line regions
to be excluded from fitting. this mode requires three
expandable parameters: thresh, avg_limit, and edge.
USE WITH CARE! May need to tweak the expandable parameters.
maskmode='list' uses the given masklist only: no additional
masks applied.
maskmode='interact' allows users to manually modify the
mask regions by dragging mouse on the spectrum plotter GUI.
use LEFT or RIGHT button to add or delete regions,
respectively.
```

```
>>> maskmode expandable parameters
thresh -- S/N threshold for linefinder. a single channel S/N ratio
  above which the channel is considered to be a detection.
  default: 5
avg_limit -- channel averaging for broad lines. a number of
  consecutive channels not greater than this parameter
  can be averaged to search for broad lines.
  default: 4
edge -- channels to drop at beginning and end of spectrum
  default: 0
ex: edge=[1000] drops 1000 channels at beginning AND end.
  edge=[1000,500] drops 1000 from beginning and 500
  from end.
```

Note: For bad baselines threshold should be increased,
and avg_limit decreased (or even switched off completely by
setting this parameter to 1) to avoid detecting baseline
undulations instead of real lines.
blfunc -- baseline model function
options: 'poly', 'chebyshev', 'cspline', or 'sinusoid'
default: 'poly'
example: blfunc='poly' uses a single polynomial line of
any order which should be given as an expandable
parameter 'order' to fit baseline.
blfunc='chebyshev' uses Chebyshev polynomials.
blfunc='cspline' uses a cubic spline function, a piecewise
cubic polynomial having C2-continuity (i.e., the second
derivative is continuous at the joining points).
blfunc='sinusoid' uses a combination of sinusoidal curves.

>>> blfunc expandable parameters

order -- order of baseline model function
options: (int) (<0 turns off baseline fitting)
default: 5
example: typically in range 2-9 (higher values
seem to be needed for GBT)

npiece -- number of the element polynomials of cubic spline curve
options: (int) (<0 turns off baseline fitting)
default: 2

applyfft -- automatically set wave numbers of sinusoidal functions
for fitting by applying some method like FFT.
options: (bool) True, False
default: True

fftmethod -- method to be used when applyfft=True. Now only
'fft' is available and it is the default.

ffththresh -- threshold to select wave numbers to be used for
sinusoidal fitting. both (float) and (str) accepted.
given a float value, the unit is set to sigma.
for string values, allowed formats include:
'xsigma' or 'x' (= x-sigma level. e.g., '3sigma'), or
'topx' (= the x strongest ones, e.g. 'top5').
default is 3.0 (unit: sigma).

addwn -- additional wave number(s) of sinusoids to be used
for fitting.
(list) and (int) are accepted to specify every
wave numbers. also (str) can be used in case
you need to specify wave numbers in a certain range.
default: [0] (i.e., constant is subtracted at least)
example: 0
[0,1,2]
'a-b' (= a, a+1, a+2, ..., b-1, b),
'<a' (= 0,1,...,a-2,a-1),
'>=a' (= a, a+1, ... up to the maximum wave
number corresponding to the Nyquist
frequency for the case of FFT).

rejwn -- wave number(s) of sinusoid NOT to be used for fitting.
  can be set just as addwn but has higher priority:
  wave numbers which are specified both in addwn
  and rejwn will NOT be used.
  default: []

cliptresh -- clipping threshold for iterative fitting
  default: 3

clipniter -- maximum iteration number for iterative fitting
  default: 0 (no iteration, i.e., no clipping)

verify -- interactively verify the results of operation for each spectrum.
  When verify = True, for each input spectrum, spectra
  before and after the operation are displayed in a plot
  window. At the prompt there are four choices of action:
  'Y' (accept the operation and continue to the next input
  spectrum), 'N' (reject the operation and continue to the
  next input spectrum), 'A' (accept the current operation
  and continue non-interactively), and 'R' (reject the
  current operation and exit from operation).
  Note that when the operation is rejected by 'N' or 'R',
  no operation is done to the spectrum/spectra.
  options: (bool) True,False
  default: False

NOTE: Currently available only when blfunc='poly'

verbose -- output fitting results to logger. if False, the fitting results
  including coefficients, residual rms, etc., are not output to
  the CASA logger, while the processing speed gets faster.
  options: (bool) True, False
  default: True

bloutput -- output fitting results to a text file. if False, the fitting
  results including coefficients, residual rms, etc., are not
  output to a text file (<outfile>_blparam.txt), while
  the processing speed gets faster.
  options: (bool) True, False
  default: True

blformat -- format of the logger output and text file specified with bloutput
  options: '', 'csv'
  default: '' (same as in the past, easy to read but huge)

showprogress -- show progress status for large data
  options: (bool) True, False
  default: True

>>> showprogress expandable parameter

minnrow -- minimum number of input spectra to show progress status
  default: 1000

outfile -- name of output file
default: '' (<infile>_bs)

outform -- output file format
options: 'ASAP', 'MS2', 'ASCII', 'SDFITS'
default: 'ASAP'
NOTE the ASAP format is easiest for further sd processing; use MS2 for CASA imaging.
If ASCII, then will append some stuff to the outfile name

overwrite -- overwrite the output file if already exists
options: (bool) True, False
default: False
NOTE this parameter is ignored when outform='ASCII'

plotlevel -- control for plotting of results.
options: 0, 1, 2, or <0
default: 0 (no plotting)
example: 0 (no plotting)
1 (some)
2 (more)
-1 (hardcopy) as abs(plotlevel), e.g.
-1 => hardcopy of final plot (will be named <outfile>_bspec.eps)

-----------
DESCRIPTION
-----------

Task sdbaseline performs baseline fitting/removal for single-dish spectra. The fit parameters, terms and rms of baseline are saved to an ascii file, '<outfile>_blparam.txt' if bloutput is True.

BASELINE MODEL FUNCTION
------------------------
The list of available model functions are shown above (see Keyword arguments section). In general 'cspline' or 'chebyshev' are recommended since they are more stable than others. 'poly' will work for lower order but will be unstable for higher order fitting. 'sinusoid' is kind of special mode that will be useful for the data that clearly shows standing wave in the spectral baseline.

SIGMA CLIPPING (ITERATIVE FITTING)
-----------------------------------
In general least square fitting is strongly affected by an extreme data so that the resulting fit makes worse. Sigma clipping is an iterative
baseline fitting with data clipping based on a certain threshold. Threshold is set as a certain factor times rms of the resulting (baseline subtracted) spectra. If sigma clipping is on, baseline fit/removal is performed several times. After each baseline subtraction, the data whose absolute value is above threshold are detected and those data are excluded from the next round of fitting. By using sigma clipping, extreme data are excluded from the fit so that resulting fit is more robust.

The user is able to control a multiplication factor using parameter clipthresh for clipping threshold based on rms. Actual threshold for sigma clipping will be (clipthresh) x (rms of spectra). Also, the user can specify number of maximum iteration to the parameter clipniter.

In general, sigma clipping will lower the performance since it increases number of fits per spectra. However, it is strongly recommended to turn on sigma clipping unless you are sure that the data is free from any kind of extreme values that may affect the fit.

-------------
FLUX UNIT CONVERSION
-------------
The task is able to convert flux unit between K and Jy. To do that, fluxunit and its subparameter telescopeparam must be properly set. The fluxunit should be 'Jy' or 'K' depending on what unit input data is and what unit you want to convert. If given fluxunit is different from the unit of input data, unit conversion is performed.

The telescopeparam is used to specify conversion factor. There are three ways to specify telescopeparam: 1) set Jy/K conversion factor, 2) set telescope diameter, D, and aperture efficiency, eta, separately, and 3) 'FIX' mode (only change the unit without converting spectral data).

If you give telescopeparam as a list, then if the list has a single float it is assumed to be the gain in Jy/K (case 1), if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively (case 2).

See the above parameter description as well as note on 'FIX' mode below for details.

There are two special cases that don’t need telescopeparam for unit conversion. Telescope name is obtained from the data.
1) ASAP (sd tool) recognizes the conversion factor (actually D and eta) for the "AT" telescopes, namely ATNF MOPRA telescope, until 2004.
2) The task does know D and eta for GBT telescope.

If you wish to change the fluxunit, by leaving the sub-parameter telescopeparam unset (telescopeparam='',) it will use internal telescope
parameters for flux conversion for the data from AT telescopes and it will use an approximate aperture efficiency conversion for the GBT data.

Note that sdbaseline assumes that the fluxunit is set correctly in the data already. If not, then set telescopeparam='FIX' and it will set the default units to fluxunit without conversion. Note also that, if the data in infile is an ms from GBT and the default flux unit is missing, this task automatically fixes the default fluxunit to 'K' before the conversion.

-------
WARNING
-------

For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

8.8.1.5 sdreduce

Keyword arguments:
infile -- name of input SD dataset
 default: 0
 example: 'PM03'
NOTE this parameter is effective only for MS input
antenna -- select an antenna name or ID
 default: 0
 example: 'PM03'
fluxunit -- units for line flux
 options: 'K','Jy',''
 default: '' (keep current fluxunit in data)
 WARNING: For GBT data, see description below.
>>> fluxunit expandable parameter
telescopeparam -- parameters of telescope for flux conversion
 options: (str) name or (list) list of gain info
 default: '' (none set)
 example: if telescopeparam='', it tries to get the telescope name from the data.
 Full antenna parameters (diameter,ap.eff.) known to ASAP are
 'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43',
 'CEDUNA','HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
telescopeparam=[104.9,0.43] diameter(m), ap.eff.
telescopeparam=[0.743] gain in Jy/K
telescopeparam='FIX' to change default fluxunit
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see description below

field -- select data by field IDs and names
default: '' (use all fields)
example: field='3C2*' (all names starting with 3C2)
field='0,4,5~7' (field IDs 0,4,5,6,7)
field='0,3C273' (field ID 0 or field named 3C273)
this selection is in addition to the other selections to data

spw -- select data by IF IDs (spectral windows)/channels
NOTE channel range selections are interpreted as mask regions to
INCLUDE in BASELINE fit, and ignored in the other operations.
when maskmode is 'auto' or 'interact', the channel mask
will be applied first before fitting as base mask
default: '' (use all IFs and channels)
example: spw='3,5,7' (IF IDs 3,5,7; all channels)
spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
spw='30~45GHz' (IF IDs with the center frequencies in range 30-45GHz; all channels)
spw='0:5~61' (IF ID 0; channels 5 to 61)
spw='3:10~20;50~60' (select multiple channel ranges within IF ID 3)
spw='3:10~20;4:0~30' (select different channel ranges for IF IDs 3 and 4)
spw='1~4;6:15~48' (for channels 15 through 48 for IF IDs 1,2,3,4 and 6)
this selection is in addition to the other selections to data

>>> spw expandable parameters
restfreq -- the rest frequency
available type includes float, int, string, list of float,
list of int, list of string, and list of dictionary. the
default unit of restfreq in case of float, int, or string
without unit is Hz. string input can be a value only
(treated as Hz) or a value followed by unit for which 'GHz',
'MHz','kHz',and 'Hz' are available.
a list can be used to set different rest frequencies for
each IF. the length of list input must be number of IFs.
dictionary input should be a pair of line name and
frequency with keys of 'name' and 'value', respectively.
values in the dictionary input follows the same manner as
as for single float or string input.
example: 345.796
'1420MHz'
[345.8, 347.0, 356.7]
['345.8MHz', '347.0MHz', '356.7MHz']
[['name':'CO','value':345]]

frame -- frequency reference frame
options: 'LSRK', 'TOPO', 'LSRD', 'BARY', 'GALACTO', 'LGROUP', 'CMB'
default: '' (keep current frame in data)
doppler -- doppler convention (effective only when spw is in
velocity unit)
options: 'RADIO', 'OPTICAL', 'Z', 'BETA', or 'GAMMA'
default: '' (keep current doppler setting in data)

timerange -- select data by time range
default: '' (use all)
example: timerange = 'YYYY/MM/DD/hh:mm:ss~YYYY/MM/DD/hh:mm:ss'
Note: YYYY/MM/DD can be dropped as needed:
timerange='09:14:00~09:54:00' # this time range
timerange='09:44:00' # data within one integration of time
timerange='>10:24:00' # data after this time
timerange='09:44:00+00:13:00' #data 13 minutes after time
this selection is in addition to the other selections to data

scan -- select data by scan numbers
default: '' (use all scans)
example: scan='21~23' (scan IDs 21, 22, 23)
this selection is in addition to the other selections to data

pol -- select data by polarization IDs
default: '' (use all polarizations)
example: pol='0,1' (polarization IDs 0,1)
this selection is in addition to the other selections to data

calmode -- calibration mode
options: 'ps','nod','otf','otfraster',
'fs','quotient','none'
default: 'none'
example: choose mode 'none' if you have already calibrated
and want to correct for atmospheric opacity defined
by tau, subtract baseline or average/smooth spectra.

>>> calmode expandable parameter

fraction -- edge marker parameter of 'otf' and 'otfraster'.
Specify a number of OFF integrations (at each
side of the raster rows in 'otfraster' mode)
as a fraction of total number of integrations.
In 'otfraster' mode, number of integrations
to be marked as OFF, n_off, is determined by
the following formula,

    n_off = floor(fraction * n),

where n is number of integrations per raster
row. Note that n_off from both sides will be
marked as OFF so that twice of specified
fraction will be marked at most. For example,
if you specify fraction='10%', resultant
fraction of OFF integrations will be 20% at most.
In 'otf' mode, n_off is given by,

\[ n_{\text{off}} = \text{floor}(\text{fraction} \times n), \]

where \( n \) is number of total integrations. 
\( n_{\text{off}} \) is used as criterion of iterative marking process. Therefore, resulting total number of OFFs will be larger than \( n_{\text{off}} \). In practice, fraction is a geometrical fraction of edge region. Thus, if integrations are concentrated on edge region (e.g. some of Lissajous patterns), then resulting \( n_{\text{off}} \) may be unexpectedly large.

default: '10%'
options: '20%' in string style or float value less than 1.0 (e.g. 0.15).
        'auto' is available only for 'otfraster'.

\( n_{\text{off}} \) -- edge marking parameter for 'otfraster'.
It is used to specify a number of OFF scans near edge directly. Value of \( n_{\text{off}} \) comes before setting by fraction. Note that \( n_{\text{off}} \) from both sides will be marked as OFF so that twice of specified \( n_{\text{off}} \) will be marked at most.
default: -1 (use fraction)
options: any positive integer

\( \text{width} \) -- edge marking parameter for 'otf'.
Pixel width with respect to a median spatial separation between neighboring two data in time. Default will be fine in most cases.
default: 0.5
options: float value

\( \text{elongated} \) -- edge marking parameter for 'otf'.
Set True only if observed area is elongated in one direction.
options: (bool) True, False
default: False

\( \text{markonly} \) -- set True if you want to save data just after edge marking (i.e. uncalibrated data) to see how OFF scans are defined.
options: (bool) True, False
default: False

\( \text{plotpointings} \) -- load plotter and plot pointing directions of ON and OFF scans.
options: (bool) True, False
default: False

tau -- the zenith atmospheric optical depth for correction
default: 0.0 (no correction)

average -- averaging on spectral data
options: (bool) True, False
default: False

>>>average expandable parameter

timeaverage -- average spectra over time
options: (bool) True, False
default: False
example: if True, this happens after calibration
tweight -- weighting for time averaging (effective only when
timeaverage=True)
options: 'var' (1/var(spec) weighted)
'tsys' (1/Tsys**2 weighted)
'tint' (integration time weighted)
'tintsys' (Tint/Tsys**2)
'median' (median averaging)
default: 'tintsys'

scanaverage -- average spectra within a scan number (effective
only when timeaverage=True)
when True, spectra are NOT averaged over
different scan numbers.
options: (bool) True, False
default: False

averageall -- average multi-resolution spectra (effective only
when timeaverage=True)
spectra are averaged by referring their frequency
coverage
default: False

polaverage -- average spectra over polarizations
options: (bool) True, False
default: False

pweight -- weighting for polarization averaging (effective only
when polaverage=True)
options: 'var' (1/var(spec) weighted)
'tsys' (1/Tsys**2 weighted)
default: 'tsys'

kernel -- type of spectral smoothing kernel
options: 'none', 'hanning', 'gaussian', 'boxcar', 'regrid', 'none'
default: 'none' (no smoothing)
>>> kernel expandable parameter
    kwidth -- width of spectral smoothing kernel
    options: (int) in channels
    default: 5
    example: 5 or 10 seem to be popular for boxcar
              ignored for hanning (fixed at 5 chans)
              (0 will turn off gaussian or boxcar)
    chanwidth -- channel width of regridded spectra
    default: '5' (in channels)
    example: '500MHz', '0.2km/s'

maskmode -- mode of setting additional channel masks
    options: 'auto', 'list', or 'interact'
    default: 'auto'
    example: maskmode='auto' runs linefinder to detect line regions
             to be excluded from fitting. this mode requires three
             expandable parameters: thresh, avg_limit, and edge.
             USE WITH CARE! May need to tweak the expandable parameters.
             maskmode='list' uses the given masklist only: no additional
             masks applied.
             maskmode='interact' allows users to manually modify the
             mask regions by dragging mouse on the spectrum plotter GUI.
             use LEFT or RIGHT button to add or delete regions,
             respectively.

>>> maskmode expandable parameters
    thresh -- S/N threshold for linefinder. a single channel S/N ratio
              above which the channel is considered to be a detection.
    default: 5
    avg_limit -- channel averaging for broad lines. a number of
                 consecutive channels not greater than this parameter
                 can be averaged to search for broad lines.
    default: 4
    edge -- channels to drop at beginning and end of spectrum
    default: 0
    example: edge=[1000] drops 1000 channels at beginning AND end.
             edge=[1000,500] drops 1000 from beginning and 500
             from end.
    Note: For bad baselines threshold should be increased,
    and avg_limit decreased (or even switched off completely by
    setting this parameter to 1) to avoid detecting baseline
    undulations instead of real lines.

blfunc -- baseline model function
options: 'poly', 'chebyshev', 'cspline', or 'sinusoid'
default: 'none' (no baseline fit)
example: blfunc='poly' uses a single polynomial line of any order which should be given as an expandable parameter 'order' to fit baseline.
blfunc='chebyshev' uses Chebyshev polynomials.
blfunc='cspline' uses a cubic spline function, a piecewise cubic polynomial having C2-continuity (i.e., the second derivative is continuous at the joining points).
blfunc='sinusoid' uses a combination of sinusoidal curves.

>>> blfunc expandable parameters
order -- order of baseline model function
options: (int) (<0 turns off baseline fitting)
default: 5
example: typically in range 2-9 (higher values seem to be needed for GBT)
npiece -- number of the element polynomials of cubic spline curve
options: (int) (<0 turns off baseline fitting)
default: 2
applyfft -- automatically set wave numbers of sinusoidal functions for fitting by applying some method like FFT.
options: (bool) True, False
default: True
fftmethod -- method to be used when applyfft=True. Now only 'fft' is available and it is the default.
fftthresh -- threshold to select wave numbers to be used for sinusoidal fitting. both (float) and (str) accepted.
given a float value, the unit is set to sigma.
for string values, allowed formats include: 'xsigma' or 'x' (= x-sigma level. e.g., '3sigma'), or 'topx' (= the x strongest ones, e.g. 'top5').
default is 3.0 (unit: sigma).
addwn -- additional wave number(s) of sinusoids to be used for fitting.
(list) and (int) are accepted to specify every wave numbers. also (str) can be used in case you need to specify wave numbers in a certain range.
default: [0] (i.e., constant is subtracted at least)
example: 0
    [0,1,2]
    'a-b' (= a, a+1, a+2, ..., b-1, b),
    '<a' (= 0,1,...,a-2,a-1),
    '>=a' (= a, a+1, ... up to the maximum wave number corresponding to the Nyquists frequency for the case of FFT).
rejwn -- wave number(s) of sinusoid NOT to be used for fitting. can be set just as addwn but has higher priority: wave numbers which are specified both in addwn and rejwn will NOT be used. default: []

clipthresh -- clipping threshold for iterative fitting default: 3

clipniter -- maximum iteration number for iterative fitting default: 0 (no iteration, i.e., no clipping)

verifycal -- interactively verify the results of calibration See description of verify parameter in the task, sdcal, for details. options: (bool) True, False default: False

verifysm -- interactively verify the results of smoothing for each spectrum. See description of verify parameter in the task, sdaverage, for details. options: (bool) True, False default: False

Note: verification is not yet available for kernel='regrid'

verifybl -- interactively verify the results of baseline fitting for each spectrum. See description of verify parameter in the task, sdbaseline, for details. options: (bool) True, False default: False

NOTE: Currently available only when blfunc='poly'

verbosebl -- output fitting results to logger. If False, the fitting results including coefficients, residual rms, etc., are not output to the CASA logger, while the processing speed gets faster. options: (bool) True, False default: True

bloutput -- output fitting results to a text file. if False, the fitting results including coefficients, residual rms, etc., are not output to a text file (<outfile>_blparam.txt), while the processing speed gets faster. options: (bool) True, False default: True

blformat -- format of the logger output and text file specified with bloutput options: '', 'csv' default: '' (same as in the past, easy to read but huge)

showprogress -- show progress status for large data options: (bool) True, False
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default: True

>>> showprogress expandable parameter

minrow -- minimum number of input spectra to show progress status
    default: 1000

outfile -- name of output file
    default: '' (<infile>_cal)

outform -- output file format
    options: 'ASAP','MS2', 'ASCII','SDFITS'
    default: 'ASAP'

    NOTE the ASAP format is easiest for further sd
    processing; use MS2 for CASA imaging.
    If ASCII, then will append some stuff to
    the outfile name

overwrite -- overwrite the output file if already exists
    options: (bool) True,False
    default: False

    NOTE this parameter is ignored when outform='ASCII'

plotlevel -- control for plotting and summary of results
    options: (int) 0, 1, 2, and their negative counterparts
    default: 0 (no plotting)

    example: plotlevel=1; show plot of spectra (see description of
    the parameter in sdcal, sdaverage, and sdbaseline)
    plotlevel=2; additionally list data before and after operation.
    plotlevel<0 as abs(plotlevel), e.g.
    -1 => hardcopy of final plot at each step, i.e.,
    <outfile>_calspec.eps, <outfile>_smspec.eps, and/or
    <outfile>_bsspec.eps

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WARNING
-------

For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the
data filler is available. However, the functionality is not well
tested yet, so that there may be unknown bugs.

8.8.1.6 sdcoadd

------------------

Keyword arguments
------------------

infiles -- list of names of input SD dataset
    antenna -- select an antenna name or ID
    default: 0
example: 'PM03'

NOTE this parameter is effective only for MS input

freqtol -- Frequency shift tolerance for considering data to be in the same spwid. The number of channels must also be the same.
default: `'` == 0 Hz (combine spwid only when frequencies are the same)
extype: freqtol='10MHz' will not combine spwid unless they are within 10 MHz.

Note: This option is useful to combine spectral windows with very slight frequency differences caused by Doppler tracking, for example.

outfile -- name of output file
default: `'` (<infile>_coadd)

outform -- format of output file
options: 'ASCII','SDFITS','MS','ASAP'
default: 'ASAP'
extype: the ASAP format is easiest for further sd processing; use MS for CASA imaging.

If ASCII, then will append some stuff to the outfile name

overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: False

NOTE this parameter is ignored when outform='ASCII'

------
WARNING
------

For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

8.8.1.7 sdflag

Keyword arguments:
infile -- name of input SD dataset
antenna -- select an antenna name or ID
default: 0
eexample: 'PM03'

NOTE this parameter is effective only for MS input

mode -- type of flag operation
options: (str) 'manual', 'clip', 'interactive', 'rowid'
default: 'manual'

>>> common data selection parameters for all modes except mode='rowid'
field -- select data by field IDs and names
default: '' (use all fields)
example: field='3C2*' (all names starting with 3C2)
        field='0,4,5~7' (field IDs 0,4,5,6,7)
        field='0,3C273' (field ID 0 or field named 3C273)
this selection is in addition to the other selections to data

spw -- select data by IF IDs (spectral windows)/channels
NOTE channel range selection is valid only in mode='manual'
default: '' (use all IFs and channels)
example: spw='3,5,7' (IF IDs 3,5,7; all channels)
        spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
        spw='115GHz' (IF IDs with the center frequencies in range 30-45GHz; all)
        spw='0:5~61' (IF ID 0; channels 5 to 61)
        spw='3:10^20;50^60' (select multiple channel ranges within IF ID 3)
        spw='3:10^20,4:0^30' (select different channel ranges for IF IDs 3 and 4)
        spw='1;4;6:15^48' (for channels 15 through 48 for IF IDs 1,2,3,4 and 6)
this selection is in addition to the other selections to data

timerange -- select data by time range
default: '' (use all)
example: timerange = 'YYYY/MM/DD/hh:mm:ss~YYYY/MM/DD/hh:mm:ss'
Note: YYYY/MM/DD can be dropped as needed:
timerange='09:14:00~09:54:00' # this time range
        timerange='09:44:00' # data within one integration of time
        timerange='>10:24:00' # data after this time
        timerange='09:44:00+00:13:00' # data 13 minutes after time
this selection is in addition to the other selections to data

scan -- select data by scan numbers
default: '' (use all scans)
example: scan='21~23' (scan IDs 21,22,23)
this selection is in addition to the other selections to data

pol -- select data by polarization IDs
default: '' (use all polarizations)
example: pol='0,1' (polarization IDs 0,1)
this selection is in addition to the other selections to data

beam -- select data by beam IDs
default: '' (use all beams)
example: beam='0,1' (beam IDs 0,1)
this selection is in addition to the other selections to data

>>> common data parameters for all modes except mode='interactive'
unflag -- flag or unflag
default: False (flag selected data)
options: (bool) True, False

>>> mode='manual' expandable parameters
restfreq -- the rest frequency (effective only when spw selection is in velocity unit.)
available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz', 'kHz', and 'Hz' are available.
a list can be used to set different rest frequencies for each IF. the length of list input must be number of IFs. dictionary input should be a pair of line name and frequency with keys of 'name' and 'value', respectively. values in the dictionary input follows the same manner as
as for single float or string input.
example: 345.796
'1420MHz'
[345.8, 347.0, 356.7]
['345.8MHz', '347.0MHz', '356.7MHz']
[{"name":"CO","value":345}]

frame -- frequency reference frame (effective only when spw selection is in velocity or frequency unit.)
options: 'LSRK', 'TOPO', 'LSRD', 'BARY', 'GALACTO', 'LGROUP', 'CMB'
default: '' (keep current frame in data)

doppler -- doppler convention (effective only when spw is in velocity unit)
options: 'RADIO', 'OPTICAL', 'Z', 'BETA', or 'GAMMA'
default: '' (keep current doppler setting in data)

>>> mode='clip' expandable parameters

clipminmax -- range of data that will NOT be flagged
default: [] (means no clip operation)
exmaple: [0.0, 1.5]

clipoutside -- clip OUTSIDE the range?
options: (bool) True, False
default: True
example: clipoutside=False means flag data WITHIN the range.

>>> mode='interactive' expandable parameters

showflagged -- show flagged data on plots
options: (bool) True, False
default: False

>>> mode='rowid' expandable parameters

row -- select data by row IDs to apply flag/unflag in the input scannable
Note, this parameter is effective only when one or more row IDs are given explicitly
default: '' (means no selection)
exmaple: '200-300,400-500' (rows 200 to 300 and 400 to 500)
outfile -- name of output file
    default: ''
    Note: by default (outfile=''), actual output file name is set as follows:
    (1) if overwrite=True (default), infile (input) will be overwritten.
        WARNING: If the formats of input and ouput files are different,
        this causes complete loss of input file.
    (2) if overwrite=False, outfile will be <infile>_f.

outform -- output file format
    options: 'ASAP','MS2', 'ASCII','SDFITS'
    default: 'ASAP'
    NOTE the ASAP format is easiest for further sd processing; use MS2 for CASA imaging.
    If ASCII, then will append some stuff to the outfile name
    WARNING: Be sure outform is same as the input file format when you overwrite the input file by overwrite=True and outfile='' (default).

overwrite -- overwrite the output file if already exists
    options: (bool) True,False
    default: True
    WARNING: input file is overwritten if overwrite=True and outfile='' (default).
    This causes the complete loss of input file if the formats of input and ouput files are different.

plotlevel -- control for plotting of results
    options: (int) 0, 1, 2, and their negative counterparts
    default: 0 (no plotting)
    example: plotlevel=1; plot spectra and flagged channels before and after current operation. asked if you accept the flag for each spw. also, the first spectrum after the operation is plotted.
    plotlevel=2; additionally list scantable before and after operation.
    plotlevel<0 as abs(plotlevel), e.g.
    -1 => hardcopy of final plot (will be named <outfile>_flag.eps)

Interactive flag operations on the Flag plotter

When sdflag is executed with interactive=True, interactive flag operation is available on a plotter, Flag Plotter, as shown in Figure 8.7. Flag Plotter uses the matplotlib plotting library to display its plots. You can find information on matplotlib at http://matplotlib.sourceforge.net. Note the plotter is loaded after non-interactive flag operation, if any of maskflag, flagrow, or clip is specified.

The Flag Plotter has two rows of buttons at the bottom to control its operation – in particular, to determine flagging and unflagging behaviors. When no button in the toolbar is depressed, the Flag Plotter is in spectral value mode. Click on a spectrum to select it and drag the mouse to
Figure 8.7: The Flag plotter. The **bottom set of buttons** are the standard `matplotlib` toolbar. See the caption of Figure 3.5 for detailed descriptions. The **upper set of buttons in the lower left** are: 1) **region**. Press this to begin marking regions (rather than zooming or panning). 2) **panel**. Press this to begin marking panels to select the whole spectrum. 3,4,5,6) **clear**, **flag**, **unflag**, **statistics**. Click on these to clear, flag, unflag, or calculate statistics of the data within the marked regions and spectra. 7) **notation**. Press this to begin editing notes on the plotter. 8,9) +, −. Click to move to the next or previous page in a series of iterated plots. The page counter on their left shows the current page number. Finally, the **Quit** is on the bottom right.

print the spectral value at the channel position of mouse. The value is printed to the bottom right corner of plotter window.

The buttons on the lower row are the standard `matplotlib` navigation buttons. See §3.3.2.1 about details of their capabilities.

In a row above it, there are a set of the other buttons (left to right):

- **region** — If depressed lets you mark channel regions in the panels. This is done by left-clicking the mouse twice at start and end channels of a region to mark. The marked regions are indicated with gray boxes. Clicking the button again will un-depress it and go back to the default spectral value mode. You can Mark multiple regions before doing something.

- **panel** — If depressed lets you mark spectra in the panels. This is done by left-clicking the mouse on panels you want to mark the whole spectrum. The marked panels are colored in
gray. Clicking the button again will un-depress it and go back to the default spectral value mode. You can Mark multiple spectra before doing something.

- **clear** — Click this to forget marked regions and spectra.

- **flag** — Click this to Flag the points in marked regions and spectra.

- **unflag** — Click this to Unflag any flagged point that would be in marked regions and spectra (even if invisible).

- **statistics** — Click this to print out statics of marked regions and spectra to the logger.

- **notation** — If depressed lets you edit texts on the plotter. Clicking the button again will un-depress it and go back to the default spectral value mode. See §8.8.1.14 for details.

- **+ and −** — Step to the next or previous plot in an iteration. The page counter on their left shows the current page number.

- **Quit** — Click this to close Flag Plotter.

To operate flagging and unflagging interactively, press **region** button (which will appear to depress), then mark channel regions by left-clicking the mouse at start and end channels of the region (each selection will add an additional region), and/or press **panel** button (which will appear to depress), then mark spectra by left-clicking on their panels (each selection will add an additional spectrum). You can get rid of all your regions and spectra by clicking **clear** button. Once regions and spectra are marked, click on one of **flag**, **unflag**, and **statistics** button to take the action.

### 8.8.1.8 sdflagmanager

**Keyword arguments:**

- **infile** -- name of input SD dataset
  
  default: ''
  
  example: infile='ngc5921.asap'

- **mode** -- Flag version operation
  
  default: 'list'; to list existing flagtables
  
  'save' to copy flag columns of infile to a flag file
  
  'restore' to place the specified flag file into infile
  
  'delete' to delete the specified flag file
  
  'rename' to rename the specified flag file

```bash
>>> mode expandable parameters
```

- **versionname** -- Flag version name
  
  default: none; example: versionname='original_data'
  
  No imbedded blanks in the versionname

- **comment** -- Short description of a versionname, when mode is 'save' or 'rename'
  
  default: ''
  
  example: comment='Clip above 1.85'
comment = versionname
oldname -- When mode='rename', the flag file to rename
merge -- merge operation
options: 'or','and', but not recommended for now.

8.8.1.9  sdfit

-----------------------------
Keyword arguments
-----------------------------
infile -- name of input SD dataset
antenna -- select an antenna name or ID
   default: 0
   example: 'PM03'
   NOTE this parameter is effective only for MS input
fluxunit -- units for line flux
   options: 'K','Jy',''
   default: '' (keep current fluxunit in data)
   WARNING: For GBT data, see description below.
   >>> fluxunit expandable parameter
   telescopeparam -- parameters of telescope for flux conversion
   options: (str) name or (list) list of gain info
   default: '' (none set)
   example: if telescopeparam='', it tries to get the telescope
   name from the data.
   Full antenna parameters (diameter,ap.eff.) known
   to ASAP are
   'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43',
   'CEDUNA','HOBART'. For GBT, it fixes default fluxunit
   to 'K' first then convert to a new fluxunit.
   telescopeparam=[104.9,0.43] diameter(m), ap.eff.
   telescopeparam=[0.743] gain in Jy/K
   telescopeparam='FIX' to change default fluxunit
   see description below
field -- select data by field IDs and names
   default: '' (use all fields)
   example: field='3C2*' (all names starting with 3C2)
   field='0,4,5~7' (field IDs 0,4,5,6,7)
   field='0,3C273' (field ID 0 or field named 3C273)
   this selection is in addition to the other selections to data
spw -- select data by IF IDs (spectral windows)/channels
   default: '' (use all IFs and channels)
   example: spw='3,5,7' (IF IDs 3,5,7; all channels)
   spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
spw='30~45GHz' (IF IDs with the center frequencies in range 30-45GHz; all channels)
spw='0:5~61' (IF ID 0; channels 5 to 61; all channels)
spw='3:10~20;50~60' (select multiple channel ranges within IF ID 3)
spw='3:10~20,4:0~30' (select different channel ranges for IF IDs 3 and 4)
spw='1~4;6:15~48' (for channels 15 through 48 for IF IDs 1,2,3,4 and 6)
this selection is in addition to the other selections to data

>>> spw expandable parameter
restfreq -- the rest frequency
available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz', 'kHz', and 'Hz' are available.
a list can be used to set different rest frequencies for each IF. the length of list input must be number of IFs. dictionary input should be a pair of line name and frequency with keys of 'name' and 'value', respectively. values in the dictionary input follows the same manner as as for single float or string input.
example: 345.796
   1420MHz
[345.8, 347.0, 356.7]
['345.8MHz', '347.0MHz', '356.7MHz']
[{'name': 'CO', 'value': 345}]
frame -- frequency reference frame
options: 'LSRK', 'TOPO', 'LSRD', 'BARY', 'GALACTO', 'LGROUP', 'CMB'
default: '' (keep current frame in data)
doppler -- doppler convention (effective only when spw is in velocity unit)
options: 'RADIO', 'OPTICAL', 'Z', 'BETA', or 'GAMMA'
default: '' (keep current doppler setting in data)
scan -- select data by scan numbers
default: '' (use all scans)
example: scan='21~23' (scan IDs 21,22,23)
this selection is in addition to the other selections to data
pol -- select data by polarization IDs
default: '' (use all polarizations)
example: pol='0,1' (polarization IDs 0,1)
this selection is in addition to the other selections to data
fitfunc -- function for fitting
options: 'gauss' (Gaussian), 'lorentz' (Lorentzian)
default: 'gauss'
fitmode -- mode for fitting
options: 'auto', 'list', or 'interact'
default: 'auto'
example: 'auto' will use the linefinder to fit for lines
using the following parameters
'list' will use maskline to define regions to
fit for lines with nfit in each
'interact' allows adding and deleting mask
regions by drawing rectangles on the plot
with mouse. Draw a rectangle with LEFT-mouse
to ADD the region to the mask and with RIGHT-mouse
to DELETE the region.

>>> fitmode expandable parameters
thresh -- S/N threshold for linefinder. a single channel S/N ratio
above which the channel is considered to be a detection.
default: 5
min_nchan -- minimum number of consecutive channels required to
pass threshold
default: 3
avg_limit -- channel averaging for broad lines. a number of
consecutive channels not greater than this parameter
can be averaged to search for broad lines.
default: 4
box_size -- running mean box size specified as a fraction
of the total spectrum length
default: 0.2
edge -- channels to drop at beginning and end of spectrum
default: 0
example: edge=[1000] drops 1000 channels at beginning AND end.
edge=[1000,500] drops 1000 from beginning and 500
from end

Note: For bad baselines threshold should be increased,
and avg_limit decreased (or even switched off completely by
setting this parameter to 1) to avoid detecting baseline
undulations instead of real lines.

nfit -- list of number of gaussian/lorentzian lines to fit in in maskline
region (ignored when fitmode='auto')
default: 0 (no fitting)
example: nfit=[1] for single line in single region,
nfit=[2] for two lines in single region,
nfit=[1,1] for single lines in each of two regions, etc.

outfile -- name of output file
default: no output fit file
example: 'mysd.fit'
overwrite -- overwrite the output file if already exists
    options: (bool) True, False
    default: False
plotlevel -- control for plotting of results
    options: 0, 1, 2
    default: 0 (no plotting)
    example: plotlevel=0 no plotting
              plotlevel=1 plots fit
              plotlevel=2 plots fit and residual
              no hardcopy available for fitter
WARNING: be careful plotting OTF data with lots of fields

------
Returns
------
a Python dictionary of line statistics
    keys: 'peak', 'cent', 'fwhm', 'nfit'
    example: each value is a list of lists with one list of
             2 entries [fitvalue, error] per component.
             e.g. xstat['peak']=[[234.9, 4.8],[234.2, 5.3]]
             for 2 components.

--------
DESCRIPTION
--------

Task sdfit is a basic line-fitter for single-dish spectra.
It assumes that the spectra have been calibrated in sdcal
or sdreduce.

Furthermore, it assumes that any selection of scans, IFs,
polarizations, and time and channel averaging/smoothing has
also already been done (in other sd tasks) as there are no controls
for these. Note that you can use sdsave to do selection, writing
out a new scantable.

Note that multiple scans, IFs, and polarizations can in principle
be handled, but we recommend that you use scan, field, spw, and pol
to give a single selection for each fit.

Currently, you can choose Gaussian or Lorentzian profile as a
fitting model.

For complicated spectra, sdfit does not do a good job of
"auto-guessing" the starting model for the fit. We recommend
you use sd.fitter in the toolkit which has more options, such as fixing components in the fit and supplying starting guesses by hand.

-------
FITMODE
-------
As described in the parameter description section, sdfit implements three fitting mode, 'auto', 'list', and 'interact'. Only difference between these modes are a method to set initial guess for line fitting. In 'auto' mode, initial guess is automatically set by executing line finder. On the other hand, 'list' and 'interact' allow to set initial guess manually. In these modes, only controllable parameter for the guess is range of the line region and number of lines per region. In 'list' mode, the user must give line region via spw parameter by using ms selection syntax while number of lines per region can be specified via nfit parameter. For example,

\[
\text{spw} = '17:1500\sim2500' \\
\text{nfit} = [1]
\]

will set line region between channels 1500 and 2500 for spw 17, and indicate that there is only one line in this region. Specifying single region with multiple line is also possible but is not recommended. In 'interact' mode, spectral data to be fitted will be displayed with pre-defined line region specified by spw parameter. The user is able to customize line region interactively.

-------------------
FLUX UNIT CONVERSION
-------------------
The task is able to convert flux unit between K and Jy. To do that, fluxunit and its subparameter telescopeparam must be properly set. The fluxunit should be 'Jy' or 'K' depending on what unit input data is and what unit you want to convert. If given fluxunit is different from the unit of input data, unit conversion is performed. The telescopeparam is used to specify conversion factor. There are three ways to specify telescopeparam: 1) set Jy/K conversion factor, 2) set telescope diameter, D, and aperture efficiency, \(\eta\), separately, and 3) 'FIX' mode (only change the unit without converting spectral data). If you give telescopeparam as a list, then if the list has a single float it is assumed to be the gain in Jy/K (case 1), if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively (case 2). See the above parameter description as well as note on 'FIX' mode below.
There are two special cases that don't need telescopeparam for unit conversion. Telescope name is obtained from the data.

1) ASAP (sd tool) recognizes the conversion factor (actually D and \( \eta \)) for the "AT" telescopes, namely ATNF MOPRA telescope, until 2004.

2) The task does know D and \( \eta \) for GBT telescope.

If you wish to change the fluxunit, by leaving the sub-parameter `telescopeparam` unset (\texttt{telescopeparam=''}), it will use internal telescope parameters for flux conversion for the data from AT telescopes and it will use an approximate aperture efficiency conversion for the GBT data.

Note that sdcal assumes that the fluxunit is set correctly in the data already. If not, then set `telescopeparam='FIX'` and it will set the default units to fluxunit without conversion.

Note also that, if the data in infile is an ms from GBT and the default flux unit is missing, this task automatically fixes the default fluxunit to 'K' before the conversion.

-------------------------------------
AVERAGING OF SPECTRA
-------------------------------------

Task sdfit has two averaging modes, i.e., time and polarization average.

When \texttt{timeaverage=True}, spectra are averaged over time for each IF (spectral window), polarization, and beam, independently. Note that, by default (\texttt{scanaverage=False}), \texttt{timeaverage=True} averages spectra irrespective of scan IDs.

It is possible to average spectra separately for each scan ID by setting a sub-parameter \texttt{scanaverage=True}.

For example, the combination of parameters: \texttt{scan='0~2', timeaverage=True, and scanaverage=False}: averages spectra in scan ID 0 through 2 all together to a spectrum,
\texttt{scanaverage=True} : averages spectra per scan ID and end up with three spectra from scan 0, 1, and 2.

When \texttt{polaverage=True}, spectra are averaged over polarization for each IF (spectral window) and beam. Note that, so far, time averaging is automatically switched on when \texttt{polaverage} is set to True. This behavior is not desirable and will be discarded in future.

-------
WARNING
-------
For the GBT raw SDFITS format data as input: SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

8.8.1.10 sdgrid

Keyword arguments:

infiles -- a list of names of input SD datasets. In case input is a single dataset, its name can be given as a string.
   example: 'testimage.asap' ['testimage1.asap','testimage2.asap']

antenna -- select an antenna name or ID
   default: -1
   example: 'PM03'
   NOTE this parameter is effective only for MS input

spw -- select data by IF IDs (spectral windows)
   NOTE this task only supports IF ID selection and ignores channel selection.
   default: '-1' (only process IFNO in the first row)
   example: spw='3,5,7' (IF IDs 3,5,7; all channels)
            spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
            spw='30~45GHz' (IF IDs with the center frequencies in range 30-45GHz; all channels)
   this selection is in addition to the other selections to data

scan -- select data by scan numbers
   default: '' (use all scans)
   example: scan='21~23' (scan IDs 21,22,23)
   this selection is in addition to the other selections to data

pol -- select data by polarization IDs
   default: '' (use all polarizations)
   example: pol='0,1' (polarization IDs 0,1)
   this selection is in addition to the other selections to data

gridfunction -- gridding function
   options: 'BOX' (Box-car), 'SF' (Spheroidal), 'GAUSS' (Gaussian), 'PB' (Primary-beam)
            'GJINC' (Gaussian*Jinc)
   default: 'BOX'
   example: 'SF'

>>> gridfunction expandable parameter:
convsupport -- convolution support for 'SF'
   default: -1 (use default for each gridfunction)
   example: 3

truncate -- truncation radius of convolution kernel.
   effective only for 'GAUSS' and 'GJINC'.
default: '-1' (use default for each gridfunction)
example: 3, '20arcsec', '3pixel'
gwidth -- HWHM for gaussian. Effective only for 'GAUSS' and 'GJINC'.
default: '-1' (use default for each gridfunction)
example: 3, '20arcsec', '3pixel'
jwidth -- Width of jinc function. Effective only for 'GJINC'.
default: '-1' (use default for each gridfunction)
example: 3, '20arcsec', '3pixel'
weight -- weight type (both lower-case and upper-case are acceptable)
options: 'UNIFORM', 'TSYS' (1/Tsys**2 weighted)
'TINT' (integration time weighted)
'TINTSYS' (Tint/Tsys**2)
default: 'UNIFORM'
clipminmax -- do min/max clipping if True
default: False
outfile -- name of output file
default: '' (outfile will be set to infile[0]'+'.grid')
example: 'mydata.asap.grid'
overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: False
NOTE this parameter is ignored when outform='ASCII'

npix -- x and y image size in pixels, symmetric for single value
default: -1 (automatically calculated from cell size and the data)
example: npix=200 (equivalent to [200,200])
cell -- x and y cell size. default unit arcsec
default: '' (automatically calculated from npix if it is set, otherwise '1.0arcmin')
example: cell=['0.2arcmin', 0.2arcmin']
    cell='0.2arcmin' (equivalent to example above)
    cell=12.0 (interpreted as '12.0arcsec'='0.2arcmin')
center -- grid center
default: '' (automatically calculated from the data)
example: 'J2000 13h44m00 -17d02m00'
    ['05:34:48.2', '-05.22.17.7'] (in J2000 frame)
    [1.46, -0.09] (interpreted as radian in J2000 frame)
plot -- Plot result or not
default: False (not plot)
example: if True, result will be plotted
DESCRIPTION:

The sdgrid task performs spatial gridding according to the user specification of spatial grid, convolution function, etc.

For grid configuration, the task supplements necessary information by referring input data if any of gridding parameter ('npix', 'cell', or 'center') is not specified by the user. If 'center' is default value (empty string), central position of the grid will be set to the center of observed area, i.e. $x=0.5*(xmax+xmin)$, $y=0.5*(ymax+ymin)$. If either 'cell' or 'npix' is set, unspecified one will be calculated from the others. In that case, total extent of the grid will be set to cover all observed position. If neither 'cell' nor 'npix' is set, cell size will be set to 1.0 arcmin and number of pixel will be calculated based on that cell size.

Currently, only J2000 frame is supported.

The parameter gridfunction sets gridding function for imaging. Currently, the task supports 'BOX' (Box-car), 'SF' (Prolate Spheroidal Wave Function), 'GAUSS' (Gaussian), 'GJINC' (Gaussian* Jinc), where $Jinc(x) = \frac{J_1(\pi x/c)}{\pi x/c}$ with a first order Bessel function $J_1$, and 'PB' (Primary Beam, not implemented yet). For 'PB', correct antenna informations should be included in input file.

There are four subparameters for gridfunction: convsupport, truncate, gwidth, and jwidth. The convsupport is an integer specifying cut-off radius for 'SF' in units of pixel. By default (convsupport=-1), the cut-off radius is set to 3 pixels. The truncate is a cut-off radius for 'GAUSS' or 'GJINC'. It accepts integer, float, and string values of numeric plus unit. Allowed units are angular units such as 'deg', 'arcmin', 'arcsec', and 'pixel'. Default unit is 'pixel' so that string without unit or numerical values (integer or float) will be interpreted as radius in pixel. Default value for truncate, which is used when negative radius is set, is $3*\text{HWHM}$ for 'GAUSS' and radius at first null for 'GJINC'. The gwidth is the HWHM of gaussian for 'GAUSS' and 'GJINC'. Default value is $\sqrt{\ln(2)}$ pixel for 'GAUSS' and $2.52*\sqrt{\ln(2)}$ pixel for 'GJINC'. The jwidth specifies width of the jinc function (parameter 'c' in the definition above). Default is 1.55 pixel. Both gwidth and jwidth allow integer, float, or string of numeric plus unit. Default values for gwidth and jwidth are taken from Mangum et al. (2007). Formula for 'GAUSS' and 'GJINC' are taken from Table 1 in the paper, and are written as below using gwidth and jwidth:
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GAUSS: \( \exp(-(|r|/gwidth)^2) \)

GJINC: \( J_1(\pi*|r|/jwidth)/(\pi*|r|/jwidth) * \exp(-(|r|/gwidth)^2) \)

Boolean parameter 'plot' controls whether gridded result is plotted or not. If True, color map of gridded data will be shown. Pixel center and observed position are overlayed as blue dot and red dot, respectively. Currently, channel averaged value will be plotted.


8.8.1.11 sdimaging

Keyword arguments:

infiles -- a list of names of input SD Measurementsets
  example: 'm100.PM01.ms'
  ['m100.PM01.ms','m100.PM03.ms']; multiple MSes

outfile -- name of output image
  default: ''
  example: 'mySDimage.im'

overwrite -- overwrite the output file if already exists
  options: (bool) True,False
  default: False (do NOT overwrite)
  example: if True, existing file will be overwritten

field -- select data by field IDs and names
  If field string is a non-negative integer, it is assumed to be a field index otherwise, it is assumed to be a field name
  default: '' (use all fields)
  example: field='3C2*' (all names starting with 3C2)
  field='0,4,5~7' (field IDs 0,4,5,6,7)
  field='0,3C273' (field ID 3 or field named 3C273)
  For multiple MS input, a list of field strings can be used:
  field = ['0~2','0~4'] (field ids 0-2 for the first MS and 0-4 for the second)
  field = '0~2' (field ids 0-2 for all input MSes)
  this selection is in addition to the other selections to data

spw -- select data by spectral window IDs/channels
  NOTE: channels de-selected here will contain all zeros if selected by the parameter mode subparameters.
  default: '' (use all IFs and channels)
  example: spw='3,5,7' (IF IDs 3,5,7; all channels)
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\[ \text{spw='<2'} \text{ (IF IDs less than 2, i.e., 0,1; all channels)} \]
\[ \text{spw='30~45GHz'} \text{ (IF IDs with the center frequencies in range 30-45GHz; all channels)} \]
\[ \text{spw='0:5~61'} \text{ (IF ID 0; channels 5 to 61; all channels)} \]
\[ \text{spw='3:10~20;50~60'} \text{ (select multiple channel ranges within IF ID 3)} \]
\[ \text{spw='3:10~20,4:0~30'} \text{ (select different channel ranges for IF IDs 3 and 4)} \]
\[ \text{spw='1^4;6:15~48'} \text{ (for channels 15 through 48 for IF IDs 1,2,3,4 and 6)} \]

For multiple MS input, a list of spw strings can be used:
\[ \text{spw=['0','0~3']} \text{ (spw ids 0 for the first MS and 0-3 for the second)} \]
\[ \text{spw='0~3'} \text{ (spw ids 0-3 for all input MSes)} \]

this selection is in addition to the other selections to data

**antenna** -- select data by antenna names or IDs

If antenna string is a non-negative integer, it is assumed to be an antenna index, otherwise, it is considered an antenna name.

default: '' (all baselines, i.e. all antenna in case of auto data)

example: antenna='PM03'

For multiple MS input, a list of antenna strings can be used:

antenna=['5','6'] (antenna id5 for the first MS and 6 for the second)

antenna='5' (antenna index 5 for all input MSes)

this selection is in addition to the other selections to data

**scan** -- select data by scan numbers

default: '' (use all scans)

example: scan='21~23' (scan IDs 21,22,23)

For multiple MS input, a list of scan strings can be used:

scan=['0~100','10~200'] (scan ids 0-100 for the first MS and 10-200 for the second)

scan='0~100 (scan ids 0-100 for all input MSes)

this selection is in addition to the other selections to data

**mode** -- spectral gridding type

options: 'channel', 'velocity', 'frequency'

default: 'channel'

>>> mode expandable parameters

**nchan** -- Total number of channels in the output image.

default: -1; Automatically selects enough channels to cover data selected by 'spw' consistent with 'start' and 'width'.

It is often easiest to leave nchan at the default value.

eample: nchan=100

**start** -- First channel, velocity, or frequency.

For mode='channel'; This selects the channel index number from the MS (0 based) that you want to correspond to the first channel of the output cube. The output cube will be in frequency space with the first channel having the frequency of the MS channel selected by start. start=0 refers to the first channel in the first selected spw, even if that channel is de-selected in the spw parameter.
Channels de-selected by the spw parameter will be filled with zeros if included by the start parameter. For example, spw=3~8:3~100 and start=2 will produce a cube that starts on the third channel (recall 0 based) of spw index 3, and the first channel will be blank.

default: ' ' (the first input channel of first input spw)

easy example: start=100 (mode='channel')

start='22.3GHz' (mode='frequency')

start='5.0km/s' (mode='velocity')

width -- Output channel width

For mode='channel', default=1; width>1 indicates channel averaging

example: width=4.

For mode='velocity' or 'frequency', default=''; width of first input channel, or more precisely, the difference in frequencies between the first two selected channels.

-- For example if channels 1 and 3 are selected with spw, then the default width will be the difference between their frequencies, and not the width of channel 1.

-- Similarly, if the selected data has uneven channel-spacing, the default width will be picked from the first two selected channels. In this case, please specify the desired width.

When specifying the width, one must give units

examples: width='1.0km/s', or width='24.2kHz'.

Setting width>0 gives channels of increasing frequency for mode='frequency', and increasing velocity for mode='velocity'.

veltype -- Velocity reference frame of output image

Options: 'radio','optical','true','relativistic'

default: 'radio'

outframe -- velocity reference frame of output image

Options: '','LSRK','LSRD','BARY','GEO','TOPO','GALACTO','LGROUP','CMB'

default: ' '; same as input data or 'LSRK' for multiple-MS inputs

example: frame='bary' for Barycentric frame

gridfunction -- gridding function for imaging

options: 'BOX' (Box-car), 'SF' (Spheroidal),

'PB' (Primary-beam), 'GAUSS' (Gaussian),

'GJINC' (Gaussian*Jinc)

default: 'BOX'

example: 'SF'

>>> gridfunction expandable parameter:

consupport -- convolution support for 'SF'

default: -1 (use default for each gridfunction)

example: 3

truncate -- truncation radius of convolution kernel.

effective only for 'GAUSS' and 'GJINC'.
default: '-1' (use default for each gridfunction)
example: 3, '20arcsec', '3pixel'
gwidth -- HWHM for gaussian. Effective only for
'GAUSS' and 'GJINC'.
default: '-1' (use default for each gridfunction)
example: 3, '20arcsec', '3pixel'
jwidth -- Width of jinc function. Effective only for 'GJINC'.
default: '-1' (use default for each gridfunction)
example: 3, '20arcsec', '3pixel'
imsize -- x and y image size in pixels, symmetric for single value
default: [] (=cover all pointings in MS)
example: imsize=200 (equivalent to [200,200])
cell -- x and y cell size. default unit arcmin
default: '' (= 1/3 of FWHM of primary beam)
example: cell=['0.2arcmin, 0.2arcmin']
cell='0.2arcmin' (equivalent to example above)
phasetcenter -- image phase center: direction measure or field ID
default: '' (= the center of pointing directions in POINTING table of infiles)
example: 6 (field id), 'J2000 13h44m00 -17d02m00',
'AZEL -123d48m29 15d41m41'
ephemsrncname -- ephemeris source name for moving source
default: ''
if the source name in the data matches one of the known
solar objects by the system, this task automatically set
the source name.
example: 'mars'
pointingcolumn -- pointing data column to use
option: 'direction', 'target', 'pointing_offset', 'source_offset', encoder'
default: 'direction'
restfreq -- specify rest frequency to use for output image
default: '' (refer input data)
example: 1.0e11, '100GHz'
stokes -- stokes parameters or polarization types to image
default: '' (= Stokes I)
example: 'XX'
minweight -- Minimum weight ratio to the median of weight used in
weight correction and weight based masking
default: 0.1
example: minweight = 0.

Gridding Kernel
The parameter gridfunction sets gridding function (convolution kernel) for imaging. Currently, the task supports 'BOX' (Box-car), 'SF' (Prolate Spheroidal Wave Function), 'GAUSS' (Gaussian), 'GJINC' (Gaussian*Jinc), where Jinc(x) = J_1(pi*x/c)/(pi*x/c) with a first order Bessel function J_1, and 'PB' (Primary Beam). For 'PB', correct antenna informations should be included in input file.

There are four subparameters for gridfunction: convsupport, truncate, gwidth, and jwidth. The convsupport is an integer specifying cut-off radius for 'SF' in units of pixel. By default (convsupport=-1), the cut-off radius is set to 3 pixels. The truncate is a cut-off radius for 'GAUSS' or 'GJINC'. It accepts integer, float, and string values of numeric plus unit. Allowed units are angular units such as 'deg', 'arcmin', 'arcsec', and 'pixel'. Default unit is 'pixel' so that string without unit or numerical values (integer or float) will be interpreted as radius in pixel. Default value for truncate, which is used when negative radius is set, is 3*HWHM for 'GAUSS' and radius at first null for 'GJINC'. The gwidth is the HWHM of gaussian for 'GAUSS' and 'GJINC'. Default value is sqrt(log(2)) pixel for 'GAUSS' and 2.52*sqrt(log(2)) pixel for 'GJINC'. The jwidth specifies width of the jinc function (parameter 'c' in the definition above). Default is 1.55 pixel. Both gwidth and jwidth allows integer, float, or string of numeric plus unit. Default values for gwidth and jwidth are taken from Mangum et al. (2007). Formula for 'GAUSS' and 'GJINC' are taken from Table 1 in the paper, and are written as below using gwidth and jwidth:

\[
\text{GAUSS:}\ \exp[-\log(2)*(|r|/\text{gwidth})^2]
\]

\[
\text{GJINC:}\ J_1(\pi|r|/\text{jwidth})/\pi|r|/\text{jwidth}) \times \exp[-\log(2)*(|r|/\text{gwidth})^2]
\]


Mask in Output Image

The parameter minweight defines a threshold of weight values to mask. The pixels in outfile whose weight is smaller than minweight*median(weight) are masked out. The task also creates a weight image with the name outfile.weight.
8.8.1.12 sdlist

--------------
Keyword arguments
--------------
infile -- name of input SD dataset
    default: none - must input file name
    example: 'mysd.asap'
    See sdcal for allowed formats.
antenna -- select an antenna name or id (only effective for MS input)
    default: 0
    example: 'PM03'
    NOTE this parameter is effective only for MS input
outfile -- name of output file for summary list
    default: '' (no output file)
    example: 'mysd_summary.txt'
overwrite -- overwrite the output file if already exists
    options: (bool) True,False
    default: False

----------
DESCRIPTION
----------
Task sdlist lists the scan summary of the dataset after importing
as a scantable into ASAP. It will optionally output this summary
as file.

-------
WARNING
-------
For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the
data filler is available. However, the functionality is not well
tested yet, so that there may be unknown bugs.

8.8.1.13 sdmath

Keyword arguments:
infiles -- a list of names of input SD datasets
    The file names will automatically replace the phrases
    IN0, IN1, ... in expr parameter.
expr -- mathematical expression using scantables
varnames -- a python dictionary of variables in expr and their values.
    Keys must be coincide with variables used in expr.
Values are substituted in each value in expr.

antenna -- select an antenna name or ID
default: 0
example: 'PM03'
NOTE this parameter is effective only for MS input

fluxunit -- units for line flux
options: 'K','Jy',''
default: '' (keep current fluxunit in data)
WARNING: For GBT data, see description below.

>>> fluxunit expandable parameter


telescopeparam -- parameters of telescope for flux conversion
options: (str) name or (list) list of gain info
default: '' (none set)
example: if telescopeparam='', it tries to get the telescope
name from the data.
Full antenna parameters (diameter,ap.eff.) known
to ASAP are
'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43',
'CEDUNA','HOBART'. For GBT, it fixes default fluxunit
to 'K' first then convert to a new fluxunit.
telescopeparam=[104.9,0.43] diameter(m), ap.eff.
telescopeparam=[0.743] gain in Jy/K
telescopeparam='FIX' to change default fluxunit
see description below

field -- select data by field IDs and names
default: '' (use all fields)
example: field='3C2*' (all names starting with 3C2)
field='0,4,5~7' (field IDs 0,4,5,6,7)
field='0,3C273' (field ID 0 or field named 3C273)
this selection is in addition to the other selections to data

spw -- select data by IF IDs (spectral windows)
NOTE this task only supports IF ID selection and ignores channel
selection.
default: '' (use all IFs and channels)
example: spw='3,5,7' (IF IDs 3,5,7; all channels)
spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
spw='30~45GHz' (IF IDs with the center frequencies in range 30-45GHz; all channels)
this selection is in addition to the other selections to data

scan -- select data by scan numbers
default: '' (use all scans)
example: scan='21~23' (scan IDs 21,22,23)
this selection is in addition to the other selections to data

pol -- select data by polarization IDs
default: '' (use all polarizations)
example: pol='0,1' (polarization IDs 0,1)
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this selection is in addition to the other selections to data

outfile -- name of output file
default: '' (must be specified)

outform -- output file format
options: 'ASAP','MS2', 'ASCII','SDFITS'
default: 'ASAP'

NOTE the ASAP format is easiest for further sd
processing; use MS2 for CASA imaging.
If ASCII, then will append some stuff to
the outfile name

overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: False

NOTE this parameter is ignored when outform='ASCII'

DESCRIPTION:

Task sdmath execute a simple arithmetic (i.e., subtraction, addition,
multiplication, and division) expression for single dish spectra.
The spectral data file can be any of the formats supported by
ASAP (scantable, MS, rpfits, and SDFITS). In the expression,
these file names should be put inside of single or double quotes.

You can use variables in the expression. If you want to use, you
must define varnames dictionary. Name of variables should be simple,
e.g. V0, V1, etc., to avoid unexpected error. Keys of varnames must
be name of variables that you used in the expression, and their
values will be substituted for variables in the expression. Allowed
type for the value is numerical values, one- or two-dimensional lists
(Python list or numpy.ndarray), and filename strings that indicate
spectral data or ASCII text, which is space-separated list of
numerical values consisting of adequate number of rows and columns.
In case you give a list of file names in infiles, they are
automatically referred to as IN0, IN1, etc. in expr and you can not
use IN0, IN1, etc. as variable names in varnames.

The fluxunit can be set, otherwise, the current settings of the first
spectral data in the expression are used.
Other selections (e.g. scan No, . IF, Pol) also apply to all
the spectral data in the expression, so if any of the data does
not contains selection, the task will produce no output.

WARNING for the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the
data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

Example:

```bash
# do on-off/off calculation
expr='("orion_on_data.asap"-"orion_off_data.asap")/"orion_off_data.asap"
outfile='orion_cal.asap'
sdmath()

# do on-off/off calculation (using infiles)
infiles = ["orion_on_data.asap", "orion_off_data.asap"]
expr='(IN0-IN1)/IN1'
outfile='orion_cal.asap'
sdmath()

# do on-off/off calculation using varnames
varnames={} (this can be skipped if you executed inp(sdmath) or default(sdmath).)
varnames['V0']="orion_on_data.asap"
varnames['V1']="orion_off_data.asap"
varnames['V2']=1.0
expr='V0/V1-V2'
outfile='orion_cal.asap'
sdmath()

# do on-off/off calculation using varnames (in pythonic way)
sdmath(varnames={'V0':'orion_on_data.asap','V1':'orion_off_data.asap','V2':1.0}, expr='V0/V1-V2', outfile='orion_cal.asap')
```

# interpretation of ASCII file value for varnames
If the contents of input ASCII file is shown as,

```
0.5 0.3 0.2
1.0 0.2 0.9
```

it is interpreted as a list `[[0.5, 0.3, 0.2],[1.0, 0.2, 0.9]]`.

### 8.8.1.14 sdplot

Keyword arguments:

- **infilen** -- name of input SD dataset
- **antenna** -- select an antenna name or ID
  - default: 0
  - example: 'PM03'
NOTE this parameter is effective only for MS input

fluxunit -- units for line flux
options: 'K', 'Jy', ''
default: '' (keep current fluxunit in data)
WARNING: For GBT data, see description below.

>>> fluxunit expandable parameter

telescopeparam -- parameters of telescope for flux conversion
options: (str) name or (list) list of gain info
default: '' (none set)
example: if telescopeparam='', it tries to get the telescope name from the data.
Full antenna parameters (diameter,ap.eff.) known to ASAP are
'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43', 'CEDUNA', 'HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
telescopeparam=[104.9,0.43] diameter(m), ap.eff.
telescopeparam=[0.743] gain in Jy/K
telescopeparam='FIX' to change default fluxunit see description below

specunit -- units for spectral axis
options: (str) 'channel', 'km/s', 'GHz', 'MHz', 'kHz', 'Hz'
default: '' (=current)
example: this will be the units for masklist

>>> specunit expandable parameter

restfreq -- the rest frequency
available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz', 'kHz', and 'Hz' are available.
a list can be used to set different rest frequencies for each IF. the length of list input must be number of IFs.
dictionary input should be a pair of line name and frequency with keys of 'name' and 'value', respectively. values in the dictionary input follows the same manner as for single float or string input.
example: 345.796
   '1420MHz'
[345.8, 347.0, 356.7]
['345.8MHz', '347.0MHz', '356.7MHz']
[{'name':'CO','value':345}]

frame -- frequency reference frame
options: 'LSRK', 'TOPO', 'LSRD', 'BARY', 'GALACTO', 'LGROUP', 'CMB'
CHAPTER 8. SINGLE DISH DATA PROCESSING

**default:** ’’ (keep current frame in data)

**doppler** -- doppler convention (effective only when spw is in velocity unit)

**options:** ’RADIO’, ’OPTICAL’, ’Z’, ’BETA’, or ’GAMMA’

**default:** ’’ (keep current doppler setting in data)

**field** -- select data by field IDs and names

**default:** ’’ (use all fields)

**example:** field=’3C2*’ (all names starting with 3C2)

field=’0,4,5~7’ (field IDs 0,4,5,6,7)

field=’0,3C273’ (field ID 0 or field named 3C273)

this selection is in addition to the other selections to data

**spw** -- select data by IF IDs (spectral windows)

**NOTE** this task only supports IF ID selection and ignores channel selection.

**default:** ’’ (use all IFs and channels)

**example:** spw=’3,5,7’ (IF IDs 3,5,7; all channels)

spw=’<2’ (IF IDs less than 2, i.e., 0,1; all channels)

spw=’30~45GHz’ (IF IDs with the center frequencies in range 30-45GHz; all channels)

this selection is in addition to the other selections to data

**scan** -- select data by scan numbers

**default:** ’’ (use all scans)

**example:** scan=’21~23’ (scan IDs 21,22,23)

this selection is in addition to the other selections to data

**pol** -- select data by polarization IDs

**default:** ’’ (use all polarizations)

**example:** pol=’0,1’ (polarization IDs 0,1)

this selection is in addition to the other selections to data

**beam** -- select data by beam IDs

**default:** ’’ (use all beams)

**example:** beam=’0,1’ (beam IDs 0,1)

this selection is in addition to the other selections to data

**timeaverage** -- average spectra over time

**options:** (bool) True, False

**default:** False

>>>timeaverage expandable parameter

**tweight** -- weighting for time averaging

**options:** ’var’ (1/var(spec) weighted)

’tsys’ (1/Tsys**2 weighted)

’tint’ (integration time weighted)

’tintsys’ (Tint/Tsys**2)

’median’ (median averaging)

**default:** ’tintsys’

**scanaverage** -- average spectra within a scan number

when True, spectra are NOT averaged over different scan numbers.
CHAPTER 8. SINGLE DISH DATA PROCESSING

options: (bool) True, False
default: False
polaverage -- average spectra over polarizations
  options: (bool) True, False
default: False
>>>polaverage expandable parameter
  pweight -- weighting for polarization average
    options: 'var' (1/var(spec) weighted)
             'tsys' (1/Tsys**2 weighted)
    default: 'tsys'
kernl -- type of spectral smoothing
    options: 'hanning', 'gaussian', 'boxcar', 'none'
default: 'none'
>>>kernl expandable parameter
  kwidth -- width of spectral smoothing kernel
    options: (int) in channels
    default: 5
    example: 5 or 10 seem to be popular for boxcar
             (0 will turn off gaussian or boxcar)
    ignored for hanning (fixed at 5 chans)
plottype -- type of plot
    options: 'spectra', 'totalpower', 'pointing', 'azel', 'grid'
default: 'spectra'
>>> plottype expandable parameters
  stack -- code for stacking on single plot for spectral plotting
    options: 'p', 'b', 'i', 't', 's', 'r' or
             'pol', 'beam', 'if', 'time', 'scan', 'row'
    default: 'p'
    example: maximum of 16 stacked spectra
    stack by pol, beam, if, time, scan
    Note stack selection is ignored when panel='r'.
panel -- code for splitting into multiple panels for spectral plotting
    options: 'p', 'b', 'i', 't', 's', 'r' or
             'pol', 'beam', 'if', 'time', 'scan', 'row'
    default: 'i'
    example: maximum of 16 panels
    panel by pol, beam, if, time, scan
    Note panel selection is ignored when stack='r'.
frange -- range for flux axis of plot for spectral plotting
    options: (list) [min, max]
    default: [] (full range)
    example: flrange=[-0.1,2.0] if 'K'
    assumes current fluxunit
sprange -- range for spectral axis of plot
    options: (list) [min, max]
CHAPTER 8. SINGLE DISH DATA PROCESSING

default: [] (full range)
example: sprange=[42.1,42.5] if 'GHz'
asumes current specunit

linecat -- control for line catalog plotting for spectral plotting
options: (str) 'all', 'none' or by molecule
default: 'none' (no lines plotted)
example: linecat='SiO' for SiO lines
         linecat='*OH' for alcohols
uses sprange to limit catalog
WARNING: specunit must be in frequency (*Hz)
to plot from the line catalog!
         and must be 'GHz' or 'MHz' to use
sprange to limit catalog

linedop -- doppler offset for line catalog plotting (spectral plotting)
options: (float) doppler velocity (km/s)
default: 0.0
example: linedop=-30.0

center -- the central direction of gridding
default: '' (map center)
example: 'J2000 19h30m00 -40d00m00'
Note currently only supports 'J2000' as direction frame

cell -- x and y cell size of gridding
default: [] (map extent divided by # of subplots in x and y)
exmple: cell=['1.0arcmin', '1.0arcmin']
        cell='1.0arcmin' (equivalent to the example above)
Note default number of subplots is 1 x 1 in plottype='grid'.

subplot -- number of subplots (row and column) on a page
NOTICE plotter will slow down when a large number is specified
default: -1 (auto. for plottype='spectra', 1x1 for plottype='grid')
example: 23 (2 rows by 3 columns)
colormap -- the colours to be used for plot lines.
default: None
example: colormap="green red black cyan magenta" (html standard)
         colormap="g r k c m" (abbreviation)
         colormap="#008000 #00FFFF #FF0090" (RGB tuple)
The plotter will cycle through these colours when lines are overlaid (stacking mode).
linestyles -- the linestyles to be used for plot lines.
default: None
example: linestyles="line dashed dotted dashdot dashdotdot dashdashdot".
The plotter will cycle through these linestyles when lines are overlaid (stacking mode).
WARNING: Linestyles can be specified only one color has been set.

linewidth -- width of plotted lines.
default: 1
example: linewidth=1 (integer)
    linewidth=0.75 (double)

histogram -- plot histogram
    options: (bool) True, False
    default: False

scanpattern -- plot additional lines on the plot to indicate scan patterns
    when plottype='pointing'
    options: (bool) True, False
    default: False

header -- print header information on the plot
    options: (bool) True, False
    default: True

The header information is printed only on the logger when
    plottype = 'azel' and 'pointing'.

>>> header expandable parameter

headsize -- header font size
    options: (int)
    default: 9

plotstyle -- customise plot settings
    options: (bool) True, False
    default: False

>>> plotstyle expandable parameter

margin -- a list of subplot margins in figure coordinate (0-1),
    i.e., fraction of the figure width or height.
    The order of elements should be:
    [left, bottom, right, top, horizontal space btw panels,
    vertical space btw panels]
    example: margin = [0.125, 0.1, 0.9, 0.9, 0.2, 0.2]

legendloc -- legend location on the axes (0-10)
    options: (integer) 0 -10
    see help of "sd.plotter.set_legend" for
    the detail of location. Note that 0 ('best')
    is very slow.
    default: 1 ('upper right')

outfile -- file name for hardcopy output
    options: (str) filename.eps,.ps,.png
    default: '' (no hardcopy)
    example: 'specplot.eps','specplot.png'
    Note this autodetects the format from the suffix (.eps,.ps,.png).

overwrite -- overwrite the output file if already exists
    options: (bool) True, False
    default: False
Task sdplot displays single-dish spectra, total power, or pointing direction of input data. It assumes that the spectra have been calibrated. It does allow selection of scans, IFs, polarizations, and some time and channel averaging/smoothing options also, but does not write out this data.

This task adds an additional toolbar to Matplotlib plotter. See the cookbook for details of its capability.

*** Data selection ***
This task allows data selection via field name, scan, IF, polarization and beam IDs. Selection of field allows pattern matching using asterisk, e.g., 'FLS3a*'. Selection of scans, IFs, polarizations, and beams, is possible by a CASA type selection syntax using a string of comma separated numbers with operators, i.e., '~', '>', and '<'. For example, the following selection
scan = "<3,7~9,15"
is to select scan IDs 0, 1, 2, 7, 8, 9, and 15.

AVERAGING OF SPECTRA

Task sdplot has two averaging modes, i.e., time and polarization average.

When timeaverage=True, spectra are averaged over time for each IF (spectral window), polarization, and beam, independently. Note that, by default (scanaverage=False), timeaverage=True averages spectra irrespective of scan IDs. It is possible to average spectra separately for each scan ID by setting a sub-parameter scanaverage=True. For example, the combination of parameters: scan='0~2', timeaverage=True, and scanaverage=False: averages spectra in scan ID 0 through 2 all together to a spectrum, scanaverage=True : averages spectra per scan ID and end up with three spectra from scan 0, 1, and 2.

When polaverage=True, spectra are averaged over polarization for each IF (spectral window) and beam. Note that, so far, time averaging is automatically switched on when polaverage is set to True. This behavior is not desirable and will be discarded in future.
*** available plottypes ***

* plottype = 'spectra' plots single dish spectra. Multiple scans, IFs, polarizations, and beams can be handles through stacking and panelling.
  
  This task uses the JPL line catalog as supplied by ASAP. If you wish to use a different catalog, or have it plot the line IDs from top or bottom (rather than alternating), then you will need to explore the sd toolkit also.

* plottype = 'grid' plots spectra based on their pointing direction. The spectra are gridded by direction before plotting. Multiple IFs and polarizations are not handled in this mode. Only the first IF and polarization is gridded and plotted if data includes multiple IDs after selections are applied. Hence, over plotting is not available.

Currently most of the parameters are ignored in the following modes.

* plottype='totalpower' is used to plot the total power data. and only plot option is amplitude versus data row number.
* plottype='azel' plots azimuth and elevation tracks of the source.
* plottype='pointing' plots antenna pointings.

*** control of plot lines in 'spectra' and 'grid' plottype ***

Note that colormap and linestyles cannot be controlled at a time. The 'linestyles' is ignored if both of them are specified. Some plot options, like changing titles, legends, fonts, and the like are not supported in this task. You should use sd.plotter from the ASAP toolkit directly for this.

ASAP recognizes the data of the "AT" telescopes, but currently does not know about the GBT or any other telescope. This task does know about GBT. Telescope name is obtained from the data. If you wish to change the fluxunit (see below), and telescopeparam='', for the AT telescopes it will use internal telescope parameters for flux conversion. For GBT, it will use an approximate aperture efficiency conversion. If you give telescopeparam a list, then if the list has a single float it is assumed to be the gain in Jy/K, if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively.

WARNING: be careful plotting otf data with lots of fields!

WARNING for the GBT raw SDFITS format data as input: SIdtasks are able to handle GBT raw SDFITS format data since the
data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

GUI Plot Control on ASAP Plotter

The principal ways to plot single dish spectra are using the sdplot task and sd.plotter toolkit. These task and toolkit load ASAP Plotter which uses the matplotlib plotting library to display plots. You can find information on matplotlib at [http://matplotlib.sourceforge.net/](http://matplotlib.sourceforge.net/).

Figure 8.8: The toolbars on ASAP plotter. The **bottom set of buttons** are the standard matplotlib toolbar. See the caption of Figure 3.5 for detailed descriptions. The **upper set of buttons** are: 1) **notation**. Press this to begin editing notes on the plotter. 2) **statistics**. Press this to begin printing statistics to the logger. 3,4) +, −. Click to move to the next or previous page in a series of iterated plots. The page counter on their left shows the current page number. Finally, the **Quit** is on the bottom right.

The ASAP Plotter has two rows of buttons at the bottom to control interactive operations as shown in Figure 8.8. When none of the button is depressed, the ASAP Plotter is in spectral value mode. Click on a spectrum to select it and drag the mouse to print the spectral value at the channel position of mouse. The value is printed to the bottom right corner of plotter window.

The buttons on the lower row are the standard matplotlib navigation buttons. See § 3.3.2.1 about details of their capabilities.

In a row above it, there are a set of the other buttons (left to right):

- **notation** — If depressed lets you edit texts on the plotter. See below for details of text edition. Clicking the button again will un-depress it and go back to the default spectral value mode.

- **statistics** — If depressed lets you print statics of a selected regions of scantable to the logger. See below for details of region selection. Clicking the button again will un-depress it and go back to the default spectral value mode.

- + and − — Step to the next or previous plot in an iteration. The page counter on their left shows the current page number.

- **Quit** — Click this to close ASAP Plotter.

**Editing texts on the plotter**
When the notation button is depressed, it lets you edit texts on the plotter. Left-click at a position on the plotter to print a new text, and the Notation window is loaded (Figure 8.9). Type the arbitrary text in the text box, select an anchor, and press the print button to print it at the position you clicked. There are three choices of anchors: figure, panel, and data. The figure or panel locates the text at a fixed position in the figure or subplot, respectively. Its relative position to the figure or subplot boundaries doesn’t change when you resize the plotter. On the other hand, the text is fixed on a position in the data coordinate of subplot, when data is selected as the anchor. The text moves along with plotted spectra as you pan the subplot.

You can modify or delete texts you added on the plotter. To do it, right-click on a text to show a menu with Modify and Delete. When Modify is selected, the Notation window is loaded to modify the selected text. Click on Delete and confirm the operation in a pop-up dialog to delete the text. Clicking the notation button again will un-depress it and go back to the default spectral value mode.

**Printing statistics of scantable**

When statistics button is depressed, it lets you print statistics of a selected channel region of the scantable plotted. The statistics values are printed to the logger. You can select a channel region by left- or right-clicking and dragging the mouse to draw a rectangle. Draw it with left-mouse to print statistics within the region, while do with right-mouse to print statistics excluding the region. Clicking the statistics button again will un-depress it and go back to the default spectral value mode.

**8.8.1.15 sdsave**

-----------------

**Keyword arguments**

-----------------

inflie -- name of input SD dataset
splitant -- split output file by antenna. this parameter is only effective for MS input.

options: (bool) True, False
CHAPTER 8. SINGLE DISH DATA PROCESSING

>>> splitant expandable parameter
antenna -- select an antenna name or ID. this parameter is effective only for MS input.
default: 0
example: antenna=0 specifies antenna by id
antenna='PM03' specifies antenna by name

getpt -- fill DIRECTION column properly (True), or reuse POINTING table in original MS (False). this parameter is only effective for MS input.
options: (bool) True, False
default: True

field -- select data by field IDs and names
default: '' (use all fields)
example: field='3C2*' (all names starting with 3C2)
field='0,4,5~7' (field IDs 0,4,5,6,7)
field='0,3C273' (field ID 0 or field named 3C273)
this selection is in addition to the other selections to data

spw -- select data by IF IDs (spectral windows)/channels
default: '' (use all IFs and channels)
example: spw='3,5,7' (IF IDs 3,5,7; all channels)
spw='<2' (IF IDs less than 2, i.e., 0,1; all channels)
spw='30~45GHz' (IF IDs with the center frequencies in range 30-45GHz; all channels)
spw='0:5~61' (IF ID 0; channels 5 to 61; all channels)
spw='3:10~20;50~60' (select multiple channel ranges within IF ID 3)
spw='3:10~20,4:0~30' (select different channel ranges for IF IDs 3 and 4)
spw='1~4;6:15~48' (for channels 15 through 48 for IF IDs 1,2,3,4 and 6)
this selection is in addition to the other selections to data

timerange -- select data by time range
default: '' (use all)
example: timerange = 'YYYY/MM/DD/hh:mm:ss~YYYY/MM/DD/hh:mm:ss'
Note: YYYY/MM/DD can be dropped as needed:
timerange='09:14:00~09:54:00' # this time range
timerange='09:44:00' # data within one integration of time
timerange='>10:24:00' # data after this time
timerange='09:44:00+00:13:00' # data 13 minutes after time
this selection is in addition to the other selections to data

scan -- select data by scan numbers
default: '' (use all)
example: scan='21~23' (scan IDs 21,22,23)
this selection is in addition to the other selections to data

pol -- select data by polarization IDs
default: '' (use all polarizations)
example: pol='0,1' (polarization IDs 0,1)
this selection is in addition to the other selections to data
beam -- select data by beam IDs
   default: '' (use all beams)
   example: beam='0,1' (beam IDs 0,1)
   this selection is in addition to the other selections to data
restfreq -- the rest frequency
   available type includes float, int, string, list of float,
   list of int, list of string, and list of dictionary. the
default unit of restfreq in case of float, int, or string
   without unit is Hz. string input can be a value only
   (treated as Hz) or a value followed by unit for which 'GHz',
   'MHz','kHz',and 'Hz' are available.
a list can be used to set different rest frequencies for
each IF. the length of list input must be number of IFs.
dictionary input should be a pair of line name and
frequency with keys of 'name' and 'value', respectively.
   values in the dictionary input follows the same manner as
   as for single float or string input.
   example: 345.796
   '1420MHz'
   [345.8, 347.0, 356.7]
   ['345.8MHz', '347.0MHz', '356.7MHz']
   [{'name': 'CO', 'value': 345}]
outfile -- name of output file
   default: '' (<infile>_saved)
   NOTE actual output file name(s) will be modified if splitant
   is True as antenna names are to be included. If outfile has a
   suffix '.asap' or '.ASAP', antenna name will be inserted before
   the suffix like 'out.antName.asap', otherwise, antenna name
   will be simply appended to outfile like 'out.sdfits.antName'.
outform -- output file format
   options: 'ASAP', 'MS2', 'ASCII', 'SDFITS'
   default: 'ASAP'
   NOTE the ASAP format is easiest for further sd
   processing; use MS2 for CASA imaging.
   If ASCII, then will append some stuff to
   the outfile name
fillweight -- fill WEIGHT and SIGMA column for output MS
   default: True
   options: True, False

-------
DESCRIPTION
-------

Task sdsave writes the single dish data to a disk file in
specified format (ASAP, MS2, SDFITS, ASCII). It is possible to
save the subset of the data by selecting field names, spw ids, time ranges, scan numbers, and polarization ids. The ASAP (scantable) format is recommended for further analysis using Sd tool or tasks except imaging. For further imaging using imager or sdtpimaging, save the data to the Measurement Set (MS2).

Note that setting getpt=False needs a lot of attention. If you set getpt=False, the task retrieves pointing direction from MS’s FIELD table, which might not be correct for single dish observation, instead to check MS’s POINTING table, which is the default behavior of the task (getpt=True). To compensate this, absolute path to MS’s POINTING table is stored, and it will be used for POINTING table when the data is converted back to MS format. In general, getpt=False is faster especially for large data. However, MS created from Scantable cannot have correct POINTING table if original MS’s POINTING table doesn’t exist. Such situation will happen when original MS is removed or renamed, or imported Scantable is moved to other computer alone.

--------
WARNING
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For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

8.8.1.16  sdscale

--------------------
Keyword arguments
--------------------

infile -- name of input SD dataset
antenna -- select an antenna name or ID
    default: 0
    example: 'PM03'
    NOTE this parameter is effective only for MS input
factor -- scaling factor. float, one- or two-dimensional float list, or filename storing scaling factor are acceptable
    default: 1.0 (no scaling)
    example: see description below
scaletsys -- scaling of associated Tsys
    options: (bool) True, False
CHAPTER 8. SINGLE DISH DATA PROCESSING

default: True
outfile -- name of output file
default: outfile=''' (<infile>_scaled<factor>)
example: 'scaled.asap'
overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: False
NOTE this parameter is ignored when outform='ASCII'

---------
DESCRIPTION
---------
Task sdscale performs scaling of single-dish spectra.
Associated Tsys is also scaled if scaletsys is True.
Tsys informations are written in the casalogger and they are automatically stored in 'casapy.log'.
The infile can be any of ASAP, MS, SDFITS, or RPFITS format.
If outfile name is given or outfile=''(default), the scaled data is written to a new file with the same format as the input data
(Note: in case of the RPFITS format input data, it will be written to SDFITS format).

The scaling factor, factor, accepts both scalar type and list type value. The list must be one or two dimensional. If factor is one dimensional, its length must coincide with a number of spectral channel. If factor is two dimensional, its shape must be (n, 1) or (n, m), where n is a number of spectrum, while m is a number of channel for each spectrum. In addition, the factor can be an ASCII filename that stores a space-separated list of scaling factor consisting of adequate number of rows and columns. For example, if the contents of input ASCII file is shown as,

\[
\begin{align*}
0.5 & \ 0.3 & \ 0.2 \\
1.0 & \ 0.2 & \ 0.9 \\
\end{align*}
\]

it is interpreted as a list \[[0.5, 0.3, 0.2], [1.0, 0.2, 0.9]\].

-------
WARNING
-------
For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.
8.8.1.17  sdstat

------------------------
How to use return values
------------------------

xstat = sdstat();
print "rms = ",xstat[‘rms’]

these can be used for testing in scripts or for regression

‘max_abscissa’ and ‘min_abscissa’ refer to the channel of max and min intensity.
‘totint’ is the integrated intensity (sum*channel).
‘eqw’ is equivalent width (totint/mag) where mag is either max or min depending on which has greater magnitude.
Note that ‘max_abscissa’, ‘min_abscissa’, ‘totint’ and ‘eqw’ are quantities (python dictionaries with keys, ‘unit’ and ‘value’).

------------------------
AVERAGING OF SPECTRA
------------------------

Task sdstat has two averaging modes, i.e., time and polarization average.

When timeaverage=True, spectra are averaged over time for each IF (spectral window), polarization, and beam, independently. Note that, by default (scanaverage=False), timeaverage=True averages spectra irrespective of scan IDs.
It is possible to average spectra separately for each scan ID by setting a sub-parameter scanaverage=True.
For example, the combination of parameters: scan=’0~2’, timeaverage=True, and scanaverage=False: averages spectra in scan ID 0 through 2 all together to a spectrum,
scanaverage=True : averages spectra per scan ID and end up with three spectra from scan 0, 1, and 2.

When polaverage=True, spectra are averaged over polarization for each IF (spectral window) and beam. Note that, so far, time averaging is automatically switched on when polaverage is set to True. This behavior is not desirable and will be discarded in future.

------------------------
FLUX UNIT CONVERSION
------------------------

The task is able to convert flux unit between K and Jy. To do that, fluxunit and its subparameter telescopeparam must be properly set.
The fluxunit should be 'Jy' or 'K' depending on what unit input data is and what unit you want to convert. If given fluxunit is different from the unit of input data, unit conversion is performed. The telescopeparam is used to specify conversion factor. There are three ways to specify telescopeparam: 1) set Jy/K conversion factor, 2) set telescope diameter, D, and aperture efficiency, eta, separately, and 3) 'FIX' mode (only change the unit without converting spectral data). If you give telescopeparam as a list, then if the list has a single float it is assumed to be the gain in Jy/K (case 1), if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively (case 2).

See the above parameter description as well as note on 'FIX' mode below for details.

There are two special cases that don't need telescopeparam for unit conversion. Telescope name is obtained from the data.
1) ASAP (sd tool) recognizes the conversion factor (actually D and eta) for the "AT" telescopes, namely ATNF MOPRA telescope, until 2004.
2) The task does know D and eta for GBT telescope.

If you wish to change the fluxunit, by leaving the sub-parameter telescopeparam unset (telescopeparam='') it will use internal telescope parameters for flux conversion for the data from AT telescopes and it will use an approximate aperture efficiency conversion for the GBT data.

Note that xxx assumes that the fluxunit is set correctly in the data already. If not, then set telescopeparam='FIX' and it will set the default units to fluxunit without conversion.

Note also that, if the data in infile is an ms from GBT and the default flux unit is missing, this task automatically fixes the default fluxunit to 'K' before the conversion.

------------------
Keyword arguments
------------------

infile -- name of input SD dataset
    default: none - must input file name
    example: 'mysd.asap'
    See sdcal for allowed formats.

antenna -- select an antenna name or ID
    default: 0
    example: 'PM03'
    NOTE this parameter is effective only for MS input

fluxunit -- units for line flux
    options: 'K','Jy','

default: '' (keep current fluxunit in data)
WARNING: For GBT data, see description below.

>>> fluxunit expandable parameter

**telescopeparam** -- parameters of telescope for flux conversion

*options: (str) name or (list) list of gain info*

default: '' (none set)

*example: if telescopeparam='', it tries to get the telescope name from the data.*

Full antenna parameters (diameter,ap.eff.) known to ASAP are

`'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43', 'CEDUNA','HOBART'`. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.

`telescopeparam=[104.9,0.43]` diameter(m), ap.eff.
`telescopeparam=[0.743]` gain in Jy/K
`telescopeparam='FIX'` to change default fluxunit

see description below

**field** -- select data by field IDs and names

default: '' (use all fields)

*example: field='3C2*' (all names starting with 3C2)*

`field='0,4,5~7'` (field IDs 0,4,5,6,7)
`field='0,3C273'` (field ID 0 or field named 3C273)

this selection is in addition to the other selections to data

**spw** -- select data by IF IDs (spectral windows)/channels

default: '' (use all IFs and channels)

*example: spw='3,5,7' (IF IDs 3,5,7; all channels)*

`spw='<2'` (IF IDs less than 2, i.e., 0,1; all channels)
`spw='30~45GHz'` (IF IDs with the center frequencies in range 30-45GHz; all channels)
`spw='0:5~61'` (IF ID 0; channels 5 to 61)
`spw='3:10~20;50~60'` (select multiple channel ranges within IF ID 3)
`spw='3:10~20,4:0~30'` (select different channel ranges for IF IDs 3 and 4)
`spw='1~4;6:15~48'` (for channels 15 through 48 for IF IDs 1,2,3,4 and 6)

this selection is in addition to the other selections to data

>>> spw expandable parameter

**restfreq** -- the rest frequency

available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz','kHz', and 'Hz' are available.

a list can be used to set different rest frequencies for each IF. the length of list input must be number of IFs. dictionary input should be a pair of line name and
frequency with keys of 'name' and 'value', respectively. Values in the dictionary input follows the same manner as for single float or string input.

e.g.,

\[
\begin{align*}
\text{345.796} & = 1420MHz \\
[345.8, 347.0, 356.7] & = ['345.8MHz', '347.0MHz', '356.7MHz'] \\
[\{'name': 'CO', 'value': 345\}] & = \text{frame -- frequency reference frame} \\
\text{options: 'LSRK', 'TOPO', 'LSRD', 'BARY', 'GALACTO', 'LGROUP', 'CMB'} & = \text{doppler -- doppler convention (effective only when spw is in velocity unit)} \\
\text{options: 'RADIO', 'OPTICAL', 'Z', 'BETA', or 'GAMMA'} & = \text{timerange -- select data by time range} \\
\text{default: '' (use all data)} & = \text{scan -- select data by scan numbers} \\
\text{default: '' (use all scans)} & = \text{pol -- select data by polarization IDs} \\
\text{default: '' (use all polarizations)} & = \text{beam -- select data by beam IDs} \\
\text{default: '' (use all beams)} & = \text{timeaverage -- average spectra over time} \\
\text{options: (bool) True, False} & = \text{tweight -- weighting for time averaging} \\
\text{options: 'var' (1/var(spec) weighted)} & = \text{tintsys'} (Tint/Tsys**2)
\end{align*}
\]
'median'  (median averaging)
default: 'tintsy'

scanaverage -- average spectra within a scan number
when True, spectra are NOT averaged over
different scan numbers.
options: (bool) True, False
default: False

polaverage -- average spectra over polarizations
options: (bool) True, False
default: False

>>> polaverage expandable parameter

pweight -- weighting for polarization averaging
options: 'var' (1/var(spec) weighted)
         'tsys' (1/Tsys**2 weighted)
default: 'tsys'

interactive -- determines interactive masking
options: (bool) True, False
default: False
example: interactive=True allows adding and deleting mask
         regions by drawing rectangles on the plot with mouse.
         Draw a rectangle with LEFT-mouse to ADD the region to
         the mask and with RIGHT-mouse to DELETE the region.

outfile -- name of output file (ASCII) to save statistics
default: '' (no output statistics file)
example: 'stat.txt'

format -- format string to print statistic values
default: '3.3f'

overwrite -- overwrite the statistics file if already exists
options: (bool) True, False
default: False

-------------------------------------------------------------------
Returns: a Python dictionary of line statistics
keys: 'rms', 'stddev', 'max', 'min', 'max_abscissa',
      'min_abscissa', 'sum', 'median', 'mean', 'totint', 'eqw'

------
WARNING
------

For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the
data filler is available. However, the functionality is not well
tested yet, so that there may be unknown bugs.
8.8.1.18 sdtpimaging

Keyword arguments:

infile -- name of an input SD Measurementset
    example: 'm100.tp.ms'

calmode -- SD calibration mode (currently only baseline subtraction)
    options: 'baseline','none'
    default: 'none'
    example: choose mode 'none' if you have
             already calibrated and want to do
             plotting nd/or imaging

>>> calmode='baseline' expandable parameters

masklist -- numbers of integrations from each edge of each scan
    to be included for baseline fitting
    default: [] (no edge. should define positive number)
    example: [30,30] or [30]
             used first 30 rows and last 30 rows of each scan
             for the baseline

blpoly -- polynomial order of the baseline fit
    default: (int) 1
    example: any number >=0

backup -- set True to create backup of input data
    options: (bool) True, False
    default: True

flaglist -- list of scan numbers to flag (ranges can be accepted)
    default: [] (use all scans)
    example: [[0,3],80]
             flag the scan range [0,3] = [0,1,2,3] and scan 80

antenna -- select data based on antenna names or IDs
    default: '' (use all antennas)
    example: antenna='0,1' (antenna ID 0 and 1)
             antenna='DV01'

    WARNING: currently baseline subtraction properly
             only one of the antennas.

spw -- spectral window ID for imaging (should have only one channel)
    default: 0
    example: spw=11 (SPW ID 11)

stokes -- stokes parameters or polarization types to image
    default: '' (Stokes I)
    example: stokes='XX' (image plane of linear polarization, XX)
             stokes='XXYY' (image cube with XX and YY image in each plane)
             stokes='I' (Stokes I image = total intensity)

createimage -- do imaging?
    options: (bool) True, False
    default: False
>>> createimage=True expandable parameters
outfile -- name of output image
   default: ''
   example: 'mySDimage.im'
imsize -- x and y image size in pixels, symmetric for single value
   default: [256,256]
   example: imsize=200 (equivalent to [200,200])
cell -- x and y cell size. default unit arcmin
   default: '1.0arcmin'
   example: cell=['0.2arcmin, 0.2arcmin']
   cell='0.2arcmin' (equivalent to example above)
phasename -- image phase center: direction measure or field ID
   default: 0
   example: 'J2000 13h44m00 -17d02m00', 'AZEL -123d48m29 15d41m41'
ephemsrcname -- ephemeris source name of moving source to use to
   correct movements of source direction during observation.
   default: ''
   if the source name in the data matches one of the
   known solar objects by the system, this task
   automatically set the source name.
   example: 'mars'
pointingcolumn -- pointing data column to use
   options: 'direction', 'target', 'pointing_offset',
   'source_offset', 'encoder'
   default: 'direction'
griddifying -- gridding function for imaging
   options: 'BOX' (Box-car), 'SF' (Spheroidal),
   'PB' (Primary-beam), 'GAUSS' (Gaussian),
   'GJINC' (Gaussian*Jinc)
   default: 'BOX'
   example: 'SF'
plotlevel -- control for plotting of results
   options: (int) 0=none, 1=some, 2=more, <0=hardcopy
   default: 0 (no plotting)
   example: plotlevel<0 as abs(plotlevel), e.g.
   -1: hardcopy plot
      (will be named <infile>_scans.eps)
   1: plot raw data, calibrated data
      (for calmode='baseline')
      plot raw or if exist calibrated data
      (for calmode='none')
   2: plot raw data, progressively display baseline
      fitting for each scan, and final calibrated data
      (for calmode='baseline')
CHAPTER 8. SINGLE DISH DATA PROCESSING

--------------
Gridding Kernel
--------------

The parameter gridfunction sets gridding function (convolution kernel) for imaging. Currently, the task supports 'BOX' (Box-car), 'SF' (Prolate Spheroidal Wave Function), 'GAUSS' (Gaussian), 'GJINC' (Gaussian*Jinc), where $J_1(x) = J_1(\pi x / c) / (\pi x / c)$ with a first order Bessel function $J_1$, and 'PB' (Primary Beam). For 'PB', correct antenna informations should be included in input file. Sub-parameters for convolution functions cannot be specified in this task. To customize your convolution function, please do imaging using sdimaging task or imager tool.

8.8.1.19 sdimprocess

--------------
Keyword arguments
--------------
infiles -- name or list of names of input SD (FITS or CASA) image(s)
mode -- image processing mode
  options: 'basket' (FFT-based Basket-Weaving),
  'press' (Pressed-out method)
  default: 'basket'
>>>mode expandable parameter
direction -- scan direction of each input image in unit of degree
  default: []
  example: direction=[0.0, 90.0] means that the first image has scan direction along longitude axis while the second image is along latitude axis.
masklist -- mask width for Basket-Weaving on percentage
  default: 1.0 (1.0% of map size)
numpoly -- order of polynomial fit in Pressed-out method
  default: 2
beamsize -- beam size for Pressed-out method
  default: 0.0
  example: beamsize=10.0 is interpreted as '10arcsec'. beamsize='1arcmin' specifies beam size as quantity.
smoothsize -- smoothing beam size in Pressed-out method.
  if numeric value is given, it is interpreted in unit of beam size specified by the parameter beamsize
  default: 2.0
example: smoothsize=2.0 means that smoothing beam size is
2.0 * beamsize.
smoothsize='1arcmin' sets smoothsize directly.
tmax -- maximum threshold value for processing
  default: 0.0 (no threshold in maximum)
  example: 10.0 (mask data larger value than 10.0)
tmin -- minimum threshold value for processing
  default: 0.0 (no threshold in minimum)
  example: -10.0 (mask data smaller value than -10.0)
outfile -- name of output file. output file is in CASA image format.
  default: '' (use default name 'sdimprocess.out.im')
  example: 'output.im'
overwrite -- overwrite the output file if already exists
  options: (bool) True, False
  default: False

--------
DESCRIPTION
--------
Task sdimprocess is used to remove a scanning noise that appears
as a striped noise pattern along the scan direction in a raster
scan data.

By default, the scanning noise is removed by using the FFT-based
'Basket-Weaving' method (Emerson &amp; Grave 1988) that requires
multiple images that observed exactly the same area with different
scanning direction. If only one image is available, the 'Pressed-out'
method (Sofue &amp; Reich 1979) can be used to remove the scanning
effect.

For 'Basket-Weaving', scanning directions must have at least two
different values. Normally, the scanning direction should be
specified for each input image. Otherwise, specified scanning
directions will be used iteratively. The masklist is a width of
masking region in the Fourier plane. It is specified as a fraction
(percentage) of the image size.

For 'Pressed-out', the scanning direction must be unique. There are
two ways to specify a size of smoothing beam used for process. One
is to specify smoothing size directly. To do this, smoothsize should
be specified as string that consists of a numerical value and an unit
(e.g. '10.0arcsec'). A value of beamsize will be ignored in this case.
Another way to specify smoothing size is to set an observed beam size
and indicate smoothing size as a scale factor of the observed beam
size. In this case, the beamsize is interpreted as the observed beam size.
size, and the smoothsize is the scale factor. If the beamsize is provided as float value, its unit is assumed to 'arcsec'. It is also possible to set the beamsize as string consisting of the numerical value and the unit. The smoothsize must be float value.

The infiles only allows an image data (CASA or FITS), and not does not work with MS or Scantable. The direction is an angle with respect to the horizontal direction, and its unit is degree. Any value may be interpreted properly, but the value ranging from 0.0 to 180.0 will be secure. The tmax and the tmin is used to specify a threshold that defines a range of spectral values used for processing. The data point that has the value larger than tmax or smaller than tmin will be excluded from the processing. The default (0.0) is no threshold. The outfile specifies an output CASA image name. If the outfile is empty, the default name ('sdimprocess.out.im') will be used.

8.8.1.20  msmoments

Keyword arguments:
infile -- Name of input MS data
default: none; example: infile="OrionS_rawACSmad"
moments -- List of moments you would like to compute
default: 0 (integrated spectrum); example: moments=[0,1]
see list above
antenna -- antenna name or id that the user wants to compute moments
default: '' (all antennae)
field -- field name or id that the user wants to compute moments
default: '' (all fields)
spw -- spectral window id that the user wants to compute moments
default: '' (all spectral windows)

includemask -- List of masks to include
default: [-1] (include all channels); example=[2,100]
excludemask -- List of masks to exclude
default: [-1] (don't exclude channels); example=[100,200]
outfile -- Output MS file name (or root for multiple moments)
default: 'source.moment' (input+auto-determined suffix); example: outfile='source_moment'
overwrite -- Overwrite existing output files
default: false

Task msmoments computes moments from spectral data stored in MS. The task is defined in analogy with immoments task, so that you can calculate any moments that is available for immoments task. Currently, the task only accepts MS with FLOAT_DATA column.
The spectral moment distributions at each row in input MS are determined. Input MS must have FLOAT_DATA column, i.e. autocorrelation data. See the cookbook and User Reference Manual for mathematical details.

The main control of the calculation is given by parameter `moments`:

- `moments=-1` - mean value of the spectrum
- `moments=0` - integrated value of the spectrum
- `moments=1` - intensity weighted coordinate; traditionally used to get 'velocity fields'
- `moments=2` - intensity weighted dispersion of the coordinate; traditionally used to get "velocity dispersion"
- `moments=3` - median of I
- `moments=4` - median coordinate
- `moments=5` - standard deviation about the mean of the spectrum
- `moments=6` - root mean square of the spectrum
- `moments=7` - absolute mean deviation of the spectrum
- `moments=8` - maximum value of the spectrum
- `moments=9` - coordinate of the maximum value of the spectrum
- `moments=10` - minimum value of the spectrum
- `moments=11` - coordinate of the minimum value of the spectrum

Note that includemask and excludemask cannot set simultaneously.

Example for finding the 1-moment, intensity-weighted coordinate, often used for finding velocity fields.

```plaintext
msmoments( infile="mydata", moment=1, outfile="velocityfields" )
```

### 8.8.1.21 Tasks with old interfaces

The following tasks with old interfaces are available with name `{taskname}old` and kept by CASA 4.3 release. Users are advised to update existing scripts.

- `sdcalold`
CHAPTER 8. SINGLE DISH DATA PROCESSING

Keyword arguments:

infile -- name of input SD dataset
antenna -- antenna name or id (only effective for MS input).
fluxunit -- units for line flux
    options: 'K','Jy',''
    default: '' (keep current fluxunit)
WARNING: For GBT data, see description below.

>>> fluxunit expandable parameter


telescopeparm -- the telescope characteristics
    options: (str) name or (list) list of gain info
    default: '' (none set)
    example: if telescopeparm='', it tries to get the telescope
     name from the data.
Full antenna parameters (diameter,ap.eff.) known
    to ASAP are
    'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43',
    'CEDUNA','HOBART'. For GBT, it fixes default fluxunit
to 'K' first then convert to a new fluxunit.
telescopeparm=[104.9,0.43] diameter(m), ap.eff.
telescopeparm=[0.743] gain in Jy/K
    telescopeparm='FIX' to change default fluxunit
    see description below

specunit -- units for spectral axis
    options: (str) 'channel','km/s','GHz','MHz','kHz','Hz'
    default: '' (=current)
    example: this will be the units for masklist

frame -- frequency frame for spectral axis
    options: (str) 'LSRK','REST','TOPO','LSRD','BARY',
    'GEO','GALACTO','LGROUP','CMB'
    default: currently set frame in scantable
WARNING: frame='REST' not yet implemented

doppler -- doppler mode
    options: (str) 'RADIO','OPTICAL','Z','BETA','GAMMA'
    default: currently set doppler in scantable

calmode -- calibration mode
    options: 'ps','nod','otf','otfraster',
    'fs','fsotf','quotient','none'
    default: 'none'
    example: choose mode 'none' if you have
    already calibrated and want to
    try averaging
WARNING: 'fsotf' is not implemented yet

>>> calmode expandable parameter
fraction -- Edge marker parameter of 'otf' and 'otfraster'. Specify a number of OFF integrations (at each side of the raster rows in 'otfraster' mode) as a fraction of total number of integrations. In 'otfraster' mode, number of integrations to be marked as OFF, n_off, is determined by the following formula,

\[ n_{\text{off}} = \text{floor}(\text{fraction} \times n), \]

where \( n \) is number of integrations per raster row. Note that n_off from both sides will be marked as OFF so that twice of specified fraction will be marked at most. For example, if you specify fraction='10%', resultant fraction of OFF integrations will be 20% at most.

In 'otf' mode, n_off is given by,

\[ n_{\text{off}} = \text{floor}(\text{fraction} \times n), \]

where \( n \) is number of total integrations. n_off is used as criterion of iterative marking process. Therefore, resulting total number of OFFs will be larger than n_off. In practice, fraction is a geometrical fraction of edge region. Thus, if integrations are concentrated on edge region (e.g. some of Lissajous patterns), then resulting n_off may be unexpectedly large.

default: '10%

options: '20%' in string style or float value less than 1.0 (e.g. 0.15).

'auto' is available only for 'otfraster'.

noff -- Edge marking parameter for 'otfraster'. It is used to specify a number of OFF spectra near edge directly. Value of noff comes before setting by fraction. Note that n_off from both sides will be marked as OFF so that twice of specified noff will be marked at most.

default: -1 (use fraction)

options: any positive integer

width -- Edge marking parameter for 'otf'. Pixel width with respect to a median spatial separation between neighboring two data in time.
CHAPTER 8. SINGLE DISH DATA PROCESSING

Default will be fine in most cases.
default: 0.5
options: float value
elongated -- Edge marking parameter for 'otf'.
    Set True only if observed area is elongated
    in one direction.
default: False
markonly -- Set True if you want to save data just after
    edge marking (i.e. uncalibrated data) to see
    how OFF spectra are defined.
default: False
scanlist -- list of scan numbers to process
default: [] (use all scans)
example: [21,22,23,24]
    this selection is in addition to field, iflist, and pollist
field -- selection string for selecting scans by name
default: '' (no name selection)
example: 'FLS3a*
    this selection is in addition to scanlist, iflist, and pollist
iflist -- list of IF id numbers to select
default: [] (use all IFs)
example: [15]
    this selection is in addition to scanlist, field, and pollist
pollist -- list of polarization id numbers to select
default: [] (use all polarizations)
example: [1]
    this selection is in addition to scanlist, field, and iflist
channelrange -- channel range selection
default: [] (use all channel)
example: [0,5000]
    Note that specified values are recognized as 'channel'
    regardless of the value of specunit
scanaverage -- average integrations within scans
options: (bool) True,False
default: False
timeaverage -- average times for multiple scan cycles
options: (bool) True,False
default: False
    example: if True, this happens after calibration

>>>timeaverage expandable parameter
tweight -- weighting for time average
options: 'none'
    'var'  (1/var(spec) weighted)
    'tsys'  (1/Tsys**2 weighted)
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'tint' (integration time weighted)
'tintsy' (Tint/Tsys**2)
'median' (median averaging)

default: 'none'

averageall -- average multi-resolution spectra
spectra are averaged by referring
to their frequency coverage

default: False

polaverage -- average polarizations
options: (bool) True,False
default: False

>>>polaverage expandable parameter
pweight -- weighting for polarization average
options: 'none'
'var' (1/var(spec) weighted)
'tsys' (1/Tsys**2 weighted)

default: 'none'

tau -- atmospheric optical depth

default: 0.0 (no correction)
verify -- interactively verify the results of calibration. Only
effective if calmode = 'ps' (but not for ALMA data),
'otf', and 'nod'.
When verify = True, spectra before and after calibration
are displayed in a plot for six spectra in scantable.
At the prompt there are two choices of action:
'Y' (accept the calibration) and 'N' (reject the calibration).
Note that when calibration is rejected by 'N', no
calibration is done to the whole scantable.

options: (bool) True,False

default: False

outfile -- Name of output file

default: '' (<infile>_cal)

outform -- format of output file
options: 'ASCII','SDFITS','MS','ASAP'
default: 'ASAP'

example: the ASAP format is easiest for further sd
processing; use MS for CASA imaging.
If ASCII, then will append some stuff to
the outfile name

overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: False
WARNING: if outform='ASCII', this parameter is ignored
plotlevel -- control for plotting of results
    options: (int) 0=none, 1=some, 2=more, <0=hardcopy
    default: 0 (no plotting)
    example: plotlevel<0 as abs(plotlevel), e.g.
        -1 => hardcopy of final plot (will be named
        <outfile>_calspec.eps)
WARNING: be careful plotting in fsotf mode!

DESCRIPTION:

-------
OVERVIEW
-------
Task sdcalold performs data selection, calibration for single-dish
spectra. By setting calmode='none', one can run sdcalold on already
calibrated data, for further selection, averaging and atmospheric
optical depth correction. To save the output spectra in a certain
range of channels, you set the range in channelrange.

If you give multiple IFs in iflist, then your scantable will have
multiple IFs by default. Averaging of multi-resolution (multi-IFs)
spectra can be achieved by setting a sub-parameter in timeaverage,
averageall, to True. It handles multi-IFs by selecting overlaps in
frequency coverage and assigning new IFs in the output spectra.

------------------------
FLUX UNIT CONVERSION
------------------------
The task is able to convert flux unit between K and Jy. To do that,
fluxunit and its subparameter telescopeparm must be properly set.
The fluxunit should be 'Jy' or 'K' depending on what unit input
data is and what unit you want to convert. If given fluxunit is
different from the unit of input data, unit conversion is performed.
The telescopeparm is used to specify conversion factor. There are
three ways to specify telescopeparm: 1) set Jy/K conversion factor,
2) set telescope diameter, D, and aperture efficiency, eta,
separately, and 3) 'FIX' mode (only change the unit without
converting spectral data). If you give telescopeparm as a list,
then if the list has a single float it is assumed to be the gain
in Jy/K (case 1), if two or more elements they are assumed to be
telescope diameter (m) and aperture efficiency respectively
(case 2).
See the above parameter description as well as note on 'FIX' mode below for details.

There are two special cases that don’t need telescopeparm for unit conversion. Telescope name is obtained from the data.
1) ASAP (sd tool) recognizes the conversion factor (actually D and \( \eta \)) for the "AT" telescopes, namely ATNF MOPRA telescope, until 2004.
2) The task does know D and \( \eta \) for GBT telescope.
If you wish to change the fluxunit, by leaving the sub-parameter telescopeparm unset (telescopeparm=’’), it will use internal telescope parameters for flux conversion for the data from AT telescopes and it will use an approximate aperture efficiency conversion for the GBT data.

Note that sdcalold assumes that the fluxunit is set correctly in the data already. If not, then set telescopeparm=’FIX’ and it will set the default units to fluxunit without conversion. Note also that, if the data in infile is an ms from GBT and the default flux unit is missing, this task automatically fixes the default fluxunit to ’K’ before the conversion.

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HOW TO CHOOSE CALMODE
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For position switching calibration, the user should choose appropriate calibration mode depending on the data. Use case for each mode is as follows:

'ps': position switch (including OTF) with explicit reference (OFF) spectra
'otf': non-raster OTF scan without explicit OFFs (e.g. Lissajous, double circle, etc.) intends to calibrate fast scan data
'otfraster': raster OTF scan without explicit OFFs

So, if the data contains explicit reference spectra, ’ps’ should be used. Otherwise, ’otfraster’ and ’otf’ are appropriate for raster OTF and non-raster OTF, respectively. In ’otf’ and ’otfraster’ modes, the task first try to find several integrations near edge as OFF integrations, then the data are calibrated using those OFFs. If the observing pattern is raster, you should use the ’otfraster’ mode to calibrate data. Otherwise, the ’otf’ mode should be used. For detail about edge marking, see inline help of sd.edgemark module and its methods.
Those modes are designed for OTF observations without explicit OFF integrations. However, these modes should work even explicit reference spectra exist. In this case, those spectra will be ignored and integrations near edges detected by edge marker will be used as reference.

Except for how to choose OFFs, the procedure to derive calibrated spectra is common for the above three modes. Selected (or preset) OFF integrations are separated by its continuity in time domain, averaged in each segment, then interpolated to timestamps for ON integrations. Effectively, it means that OFF integrations are averaged by each OFF spectrum for 'ps' mode, averaged by either ends of each raster row for 'otfraster' mode, averaged by each temporal segments of detected edges for 'otf' mode. The formula for calibrated spectrum is

$$T_{sys} \times (ON - OFF) / OFF.$$  

The 'fs' mode is for frequency switch calibration. Currently, only GBT frequency switch data is supported.

The 'quotient' mode is special mode for "AT" telescopes, namely ANNF MOPRA. It assumes that observing sequence looks like "target, reference, target, reference,..." and it derives calibrated spectrum as

$$T_{sys} \times ON / OFF,$$

slightly different from position switch modes.

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WARNING
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For the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

* sdcal2old

Keyword arguments:
infile -- name of input SD dataset
antenna -- antenna name or id (only effective for MS input).
specunit -- units for spectral axis
options: (str) 'channel', 'km/s', 'GHz', 'MHz', 'kHz', 'Hz', ''
default: '' (=current)
example: this will be the units for maskflag

>>> specunit expandable parameters

restfreq -- rest frequency
available type includes float, int, string, list of float,
list of int, list of string, and list of dictionary. the
default unit of restfreq in case of float, int, or string
without unit is Hz. string input can be a value only
(treated as Hz) or a value followed by unit for which 'GHz',
'MHz', 'kHz', and 'Hz' are available.
a list can be used to set different rest frequencies for
each IF. the length of list input must be nIF. dictionary
input should be a pair of molecule name and frequency with
keys of 'name' and 'value', respectively. values in the
dictionary input follows the same manner as for single
float or string input.
example: 345.796
'1420MHz'
[345.8, 347.0, 356.7]
['345.8MHz', '347.0MHz', '356.7MHz']
[{'name':'CO','value':345}]

frame -- frequency frame for spectral axis
options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY',
'GEO', 'GALACTO', 'LGROUP', 'CMB'
default: currently set frame in scantable
WARNING: frame='REST' not yet implemented
doppler -- doppler mode
options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
default: currently set doppler in scantable
mode -- type of flag operation
options: (str) 'manual', 'clip', 'interactive', 'rowid'
default: 'manual'

>>> common data selection parameters for all modes except mode='rowid'

scans -- list of scan numbers to process
default: [] (use all scans)
example: [21, 22, 23, 24]
this selection is in addition to field, timerange,
ifs and pols
field -- selection string for selecting scans by name
default: '' (no name selection)
example: 'FLS3a*'
this selection is in addition to scans, timerange,
ifs, and pols
CHAPTER 8. SINGLE DISH DATA PROCESSING

timerange -- Select data based on time range:
default = '' (all); example,
timerange = 'YYYY/MM/DD/hh:mm:ss~YYYY/MM/DD/hh:mm:ss'
Note: YYYY/MM/DD can be dropped as needed:
timerange='09:14:0~09:54:0' # this time range
timerange='09:44:00' # data within one integration of time
timerange='>10:24:00' # data after this time
timerange='09:44:00+00:13:00' # data 13 minutes after time

ifs -- list of IF id numbers to select
default: [] (use all IFs)
example: [15]
this selection is in addition to scans, field,
timerange, ifs, and pols

pols -- list of polarization id numbers to select
default: [] (use all polarizations)
example: [1]
this selection is in addition to scans, field,
timerange, and ifs

>>> mode='manual' expandable parameters
maskflag -- list of mask regions to apply flag/unflag
Note, this parameter is ignored if one or more rows are
given in flagrow, or clip=True.
default: [] (entire spectrum)
example: [[1000,3000],[5000,7000]]

>>> mode='clip' expandable parameters
clipminmax -- range of data that will NOT be flagged
default: [] (means no clip operation)
example: [0.0,1.5]
clipoutside -- clip OUTSIDE the range ?
options: (bool)True,False
default: True
example: clipoutside=False means flag data WITHIN the range.

>>> mode='interactive' expandable parameters
showflagged -- show flagged data on plots
default: False

>>> mode='rowid' expandable parameters
rows -- a list of row IDs to apply flag/unflag in the input scannable
Note, this parameter is effective only when one or more row
IDs are given explicitly
default: [] (means no selection)
example: [0, 2, 3]
unflag -- flag or unflag
default: False (flag selected data)
options: (bool) True, False

outfile -- Name of output file
default: ''
Note: by default (outfile=''), actual output file name is set as follows:
(1) if overwrite=True (default), infile (input) will be overwritten.
WARNING: If the formats of input and output files are different,
this causes complete loss of input file.
(2) if overwrite=False, outfile will be <infile>_f.

outform -- Format of output file
options: 'ASCII','SDFITS','MS','ASAP'
default: 'ASAP'
example: the ASAP format is easiest for further sd
processing; use MS for CASA imaging.
WARNING: Be sure outform is same as the input file format when you
overwrite the input file by overwrite=True and outfile='' (default)

overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: True
WARNING: input file is overwritten if overwrite=True and outfile='' (default)
This causes the complete loss of input file if the formats of
input and output files are different.

plotlevel -- Control for plotting of results
options: (int) 0=none, 1=some, 2=more, <0=hardcopy
default: 0 (no plotting)
example: plotlevel<0 as abs(plotlevel), e.g.
-1 => hardcopy of final plot (will be named
<outfile>_flag.eps)
WARNING: be careful plotting in fsotf mode!

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DESCRIPTION:

Task sdflag2old performs either interactive or non-interactive channel/row
based flagging on spectra.
Currently, there are three ways of non-interactive flagging available:
(1) channel or row based flagging by selecting spectra by field,
lists of scan numbers, IF numbers, and polarization idices in mode='manual',
(2) channel based flagging by specifying a range of spectral values in
mode='clip', and
(3) row based flagging by specifying a list of row numbers in
mode='rowid'. Note this is an EXPERT mode.

In mode='manual', the channel based flagging are invoked when regions
in channel, frequency, or velocity are specified to maskflag parameter.
Otherwise, the whole channels are flagged for the selected spectra. Note the mode='rowid' is an EXPERT mode since it might not be straightforward for general users to select data by row IDs in scantable.

Interactive flagging is available when mode='interactive'. The available ways of interactive flagging include:
(1) row based flagging by selecting 'panel' and (2) channel based flagging by selecting 'region's of channels on Flag plotter. See the cookbook for details of how to select channel regions and spectra on the plotter.

NOTE the task sdflag2old only modifies flag information, FLAGROW and FLAGTRA, in the input scantable, and does not filter rows in the dataset unlike sdflagold task.

If plotlevel>=1, the task asks you if you really apply the flags before it is actually written to the data with a plot indicating flagged regions. Please note that this task is still experimental.

WARNING for overwrite option:
Be sure 'outform' is the same as data format of input file when you overwrite it. The default value of the option 'overwrite' is True in this task, thereby the current dataset (infile) is overwritten unless a different file name is set to outfile. There is a known issue in overwriting infile. If 'outform' differs to the data format of infile, the data is overwritten with the new data format (specified by 'outform') and the data in the original format will be lost.

WARNING for the GBT raw SDFITS format data as input: SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

- sdsmoothold

Keyword arguments:
  infile -- name of input SD dataset
  antenna -- antenna name or id (only effective for MS input).
  scanaverage -- average integrations within scans
    options: (bool) True,False
default: False
    example: if True, this happens in read-in
    For GBT, set False!
  scanlist -- list of scan numbers to process
default: [] (use all scans)
example: [21,22,23,24]
this selection is in addition to field, iflist, and pollist

field -- selection string for selecting scans by name
default: '' (no name selection)
example: 'FLS3a*'
this selection is in addition to scanlist, iflist, pollist

iflist -- list of IF id numbers to select
default: [] (use all IFs)
example: [15]
this selection is in addition to scanlist, field, and pollist

pollist -- list of polarization id numbers to select
default: [] (use all polarizations)
example: [1]
this selection is in addition to scanlist, field, and iflist

kernel -- type of spectral smoothing
options: 'hanning','gaussian','boxcar','regrid'
default: 'hanning'

>>>kernel expandable parameter
kwidth -- width of spectral smoothing kernel
options: (int) in channels
default: 5
example: 5 or 10 seem to be popular for boxcar
        ignored for hanning (fixed at 5 chans)
        (0 will turn off gaussian or boxcar)
chanwidth -- channel width of regridded spectra
default: '5' (in channels)
exmple: '500MHz', '0.2km/s'

verify -- interactively verify the results of smoothing for each
spectrum.
When verify = True, for each input spectrum, spectra
before and after the smoothing are displayed in a plot
window. At the prompt there are four choices of action:
'Y' (accept the smoothing and continue to the next input
spectrum), 'N' (reject the smoothing and continue to the
next input spectrum), 'A' (accept the current smoothing
and continue non-interactively), and 'R' (reject the
current smoothing and exit from smoothing).
Note that when the smoothing is rejected by 'N' or 'R',
no smoothing is done to the spectrum/spectra.
options: (bool) True,False
default: False
Note: verification is not yet available for kernel='regrid'
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outfile -- Name of output ASAP format (scantable) file
    default: '' (<infile>_sm)

outform -- format of output file
    options: 'ASCII', 'SDFITS', 'MS', 'ASAP'
    default: 'ASAP'
    example: the ASAP format is easiest for further sd
    processing; use MS for CASA imaging.
    If ASCII, then will append some stuff to
    the outfile name

overwrite -- overwrite the output file if already exists
    options: (bool) True, False
    default: False
    WARNING: if outform='ASCII', this parameter is ignored

plotlevel -- control for plotting of results
    options: (int) 0=none, 1=some, 2=more, <0=hardcopy
    default: 0 (no plotting)
    example: plotlevel<0 as abs(plotlevel), e.g.
        -1 => hardcopy of final plot (will be named
        <outfile>_smspec.eps)

DESCRIPTION:

Task sdsmoothold performs smoothing of the single-dish spectra.
Set plotlevel >= 1 to plot spectrum before and after smoothing.

WARNING for the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the
data filler is available. However, the functionality is not well
tested yet, so that there may be unknown bugs.

• sdbaselineold

Keyword arguments:
infile -- name of input SD dataset
antenna -- antenna name or id (only effective for MS input).
fluxunit -- units for line flux
    options: 'K', 'Jy', ''
    default: '' (keep current fluxunit)
    WARNING: For GBT data, see description below.
    >>> fluxunit expandable parameter
telescopeparm -- the telescope characteristics
    options: (str) name or (list) list of gain info
    default: '' (none set)
    example: if telescopeparm='', it tries to get the telescope
name from the data.
Full antenna parameters (diameter, ap.eff.) known to ASAP are 'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43', 'CEDUNA', 'HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
telescopeparm=[104.9, 0.43] diameter(m), ap.eff.
telescopeparm=[0.743] gain in Jy/K
telescopeparm='FIX' to change default fluxunit
see description below

specunit -- units for spectral axis
options: (str) 'channel', 'km/s', 'GHz', 'MHz', 'kHz', 'Hz', ''
default: '' (=current)
example: this will be the units for masklist

>>> specunit expandable parameters

restfreq -- rest frequency
available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz', 'kHz', and 'Hz' are available.
a list can be used to set different rest frequencies for each IF. the length of list input must be nIF. dictionary input should be a pair of molecule name and frequency with keys of 'name' and 'value', respectively. values in the dictionary input follows the same manner as for single float or string input.
example: 345.796
'1420MHz'
[345.8, 347.0, 356.7]
['345.8MHz', '347.0MHz', '356.7MHz']
[{'name':'CO', 'value':345}]

frame -- frequency frame for spectral axis
options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY', 'GEO', 'GALACTO', 'LGROUP', 'CMB'
default: currently set frame in scantable
WARNING: frame='REST' not yet implemented

doppler -- doppler mode
options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
default: currently set doppler in scantable

scanlist -- list of scan numbers to process
default: [] (use all scans)
example: [21, 22, 23, 24]
this selection is in addition to field, iflist, and pollist
field -- selection string for selecting scans by name
    default: '' (no name selection)
    example: 'FLS3a*'
    this selection is in addition to scanlist, iflist, and pollist
iflist -- list of IF id numbers to select
    default: [] (use all IFs)
    example: [15]
    this selection is in addition to scanlist, field, and pollist
pollist -- list of polarization id numbers to select
    default: [] (use all polarizations)
    example: [1]
    this selection is in addition to scanlist, field, and iflist
tau -- atmospheric optical depth
    default: 0.0 (no correction)
masklist -- list or string of mask regions to INCLUDE in BASELINE fit
    a string masklist allows per IF selection of channels.
    default: [] (entire spectrum)
    example: [[1000,3000],[5000,7000]]
        '0:1000~3000;5000~7000, 1:200~350;450~600'
        when maskmode is 'auto' or 'interact', this mask
        will be applied first before fitting as base mask
maskmode -- mode of setting additional channel masks
    options: (str) 'auto','list','interact'
    default: 'auto'
    example: maskmode='auto' runs linefinder to detect line regions
        to be excluded from fitting. this mode requires three
        expandable parameters: thresh, avg_limit, and edge.
        USE WITH CARE! May need to tweak the expandable parameters.
        maskmode='list' uses the given masklist only: no additional
        masks applied.
        maskmode='interact' allows users to manually modify the
        mask regions by dragging mouse on the spectrum plotter GUI.
        use LEFT or RIGHT button to add or delete regions, respectively.

>>> maskmode expandable parameters
thresh -- S/N threshold for linefinder
    default: 5
    example: a single channel S/N ratio above which the channel is
        considered to be a detection
avg_limit -- channel averaging for broad lines
    default: 4
    example: a number of consecutive channels not greater than
        this parameter can be averaged to search for broad lines
edge -- channels to drop at beginning and end of spectrum
default: 0
example: [1000] drops 1000 channels at beginning AND end
[1000,500] drops 1000 from beginning and 500 from end
Note: For bad baselines threshold should be increased,
and avg_limit decreased (or even switched off completely by
setting this parameter to 1) to avoid detecting baseline
undulations instead of real lines.

blfunc -- baseline model function
options: (str) 'poly','chebyshev','cspline','sinusoid'
default: 'poly'
example: blfunc='poly' uses a single polynomial line of
any order which should be given as an expandable
parameter 'order' to fit baseline.
blfunc='chebyshev' uses Chebyshev polynomials.
blfunc='cspline' uses a cubic spline function, a piecewise
cubic polynomial having C2-continuity (i.e., the second
derivative is continuous at the joining points).
blfunc='sinusoid' uses a combination of sinusoidal curves.

>>> blfunc expandable parameters
order -- order of baseline polynomial
options: (int) (<0 turns off baseline fitting)
default: 5
example: typically in range 2-9 (higher values
seem to be needed for GBT)

npiece -- number of the element polynomials of cubic spline curve
options: (int) (<0 turns off baseline fitting)
default: 2

applyfft -- automatically set wave numbers of sinusoidal functions
for fitting by applying some method like FFT.
options: (bool) True, False
default: True

fftmethod -- method to be used when applyfft=True. Now only
'fft' is available and it is the default.

fftthresh -- threshold to select wave numbers to be used for
sinusoidal fitting. both (float) and (str) accepted.
given a float value, the unit is set to sigma.
for string values, allowed formats include:
'xsigma' or 'x' (= x-sigma level. e.g., '3sigma'), or
'topx' (= the x strongest ones, e.g. 'top5').
default is 3.0 (unit: sigma).

addwn -- additional wave number(s) of sinusoids to be used
for fitting.
(list) and (int) are accepted to specify every
wave numbers. also (str) can be used in case
you need to specify wave numbers in a certain range,
e.g., 'a-b' (= a, a+1, a+2, ..., b-1, b),
    '<a' (= 0,1,...,a-2,a-1),
    '>=a' (= a, a+1, ... up to the maximum wave number corresponding to the Nyquist frequency for the case of FFT).
    default: [0] (i.e., constant is subtracted at least)
rejwn -- wave number(s) of sinusoid NOT to be used for fitting. can be set just as addwn but has higher priority:
    wave numbers which are specified both in addwn and rejwn will NOT be used.
    default: []
clipthresh -- clipping threshold for iterative fitting
    default: 3
clipniter -- maximum iteration number
    default: 0 (no iteration, i.e., no clipping)
verify -- interactively verify the results of baseline fitting for each spectrum.
    When verify = True, for each input spectrum the baseline fit function and spectra before and after the fit are displayed in a plot window. At the prompt there are four choices of action: 'Y' (accept the fit and continue to the next input spectrum), 'N' (reject the fit and continue to the next input spectrum), 'A' (accept the current fit and continue non-interactively), and 'R' (reject the current fit and exit from baseline fitting).
    Note that when the baseline fit is rejected by 'N' or 'R', no baseline fit is applied to the spectrum/spectra.
    options: (bool) True,False
    default: False
NOTE: Currently available only when blfunc='poly' verbose -- output fitting results to logger
    default: True
    example: If False, the fitting results including coefficients, residual rms, etc., are not output to the CASA logger, while the processing speed gets faster
bloutput -- output fitting results to a text file
    default: True
    example: If False, the fitting results including coefficients, residual rms, etc., are not output to a text file (<outfile>_blparam.txt), while the processing speed gets faster
blformat -- format of the logger output and text file specified with bloutput
    options: '', 'csv'
    default: '' (same as in the past, easy to read but huge)
showprogress -- show progress status for large data
    default: True
minnrow -- minimum number of input spectra to show progress status
default: 1000

outfile -- Name of output file
default: '' (<infile>_bs)

outform -- format of output file
options: 'ASCII','SDFITS','MS','ASAP'
default: 'ASAP'
example: the ASAP format is easiest for further sd processing; use MS for CASA imaging.
If ASCII, then will append some stuff to the outfile name

overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: False
WARNING: if outform='ASCII', this parameter is ignored

plotlevel -- control for plotting of results
options: (int) 0=none, 1=some, 2=more, <0=hardcopy
default: 0 (no plotting)
example: plotlevel<0 as abs(plotlevel), e.g.
-1 => hardcopy of final plot (will be named <outfile>_bspec.eps)
WARNING: be careful plotting in fsotf mode!

DESCRIPTION:

Task sdbaselineold performs baseline fitting/removal for single-dish spectra.
The fit parameters, terms and rms of base-line are saved to an ascii file, '<outfile>_blparam.txt'.

ASAP recognizes the data of the "AT" telescopes, but currently does not know about the GBT or any other telescope. This task does know about GBT. Telescope name is obtained from the data. If you wish to change the fluxunit (see below), and telescopeparm='', for the AT telescopes it will use internal telescope parameters for flux conversion. For GBT, it will use an approximate aperture efficiency conversion. If you give telescopeparm a list, then if the list has a single float it is assumed to be the gain in Jy/K, if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively.

Note that sdbaselineold assumes that the fluxunit is set correctly in the data already. If not, then set telescopeparm='FIX' and it
will set the default units to fluxunit without conversion.
NOTE: If the data in infile is an ms from GBT and the default flux unit is missing and this task automatically fixes the default fluxunit to 'K' before the conversion.

WARNING for the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

- sdreduceold

Keyword arguments:
infile -- name of input SD dataset
antenna -- antenna name or id (only effective for MS input).
fluxunit -- units for line flux
  options: 'K','Jy',''
  default: '' (keep current fluxunit)
  WARNING: For GBT data, see description below.
  >>> fluxunit expandable parameter
telescopeparm -- the telescope characteristics
  options: (str) name or (list) list of gain info
  default: '' (none set)
  example: if telescopeparm='', it tries to get the telescope name from the data.
  Full antenna parameters (diameter,ap.eff.) known to ASAP are
  'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43',
  'CEDUNA','HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
  telescopeparm=[104.9,0.43] diameter(m), ap.eff.
  telescopeparm=[0.743] gain in Jy/K
  telescopeparm='FIX' to change default fluxunit
  see description below

specunit -- units for spectral axis
  options: (str) 'channel','km/s','GHz','MHz','kHz','Hz',''
  default: '' (=current)
  example: this will be the units for masklist
  >>> specunit expandable parameters
restfreq -- rest frequency
  available type includes float, int, string, list of float,
  list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only
(treated as Hz) or a value followed by unit for which 'GHz', 'MHz', 'kHz', and 'Hz' are available.
a list can be used to set different rest frequencies for each IF. the length of list input must be nIF. dictionary input should be a pair of molecule name and frequency with keys of 'name' and 'value', respectively. values in the dictionary input follows the same manner as for single float or string input.
example: 345.796
'1420MHz'
[345.8, 347.0, 356.7]
['345.8MHz', '347.0MHz', '356.7MHz']
[{'name': 'CO', 'value': 345}]
frame -- frequency frame for spectral axis
options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY', 'GEO', 'GALACTO', 'LGROUP', 'CMB'
default: currently set frame in scantable
WARNING: frame='REST' not yet implemented
doppler -- doppler mode
options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
default: currently set doppler in scantable
calmode -- calibration mode
options: 'ps', 'nod', 'fs', 'fsotf', 'quotient', 'none'
default: 'none'
example: choose mode 'none' if you have already calibrated and want to try baselines or averaging

>>> calmode expandable parameter
fraction -- Edge marking parameter for 'otf' and 'otfraster'.
specify a number of OFF scans as a fraction of total number of data points.
default: '10%'
options: '20%' in string style or float value less than 1.0 (e.g. 0.15).
'auto' is available only for 'otfraster'.
noff -- Edge marking parameter for 'otfraster'.
It is used to specify a number of OFF scans near edge directly. Value of noff comes before setting by fraction.
default: -1 (use fraction)
options: any positive integer
width -- Edge marking parameter for 'otf'.
Pixel width with respect to a median spatial separation between neighboring two data in time.
Default will be fine in most cases.
default: 0.5
options: float value

elongated -- Edge marking parameter for 'otf'.
Set True only if observed area is elongated
in one direction.
default: False

markonly -- Set True if you want to save data just after
edge marking (i.e. uncalibrated data) to see
how OFF scans are defined.
default: False

scanlist -- list of scan numbers to process
default: [] (use all scans)
example: [21,22,23,24]
this selection is in addition to field, iflist, and pollist

field -- selection string for selecting scans by name
default: '' (no name selection)
example: 'FLS3a*'  
this selection is in addition to scanlist, iflist, and pollist

iflist -- list of IF id numbers to select
default: [] (use all IFs)
example: [15]
this selection is in addition to scanlist, field, and pollist

pollist -- list of polarization id numbers to select
default: [] (use all polarizations)
example: [1]
this selection is in addition to scanlist, field, and iflist

channelrange -- channel range selection
default: [] (use all channel)
example: [0,5000]
Note that specified values are recognized as 'channel'
regardless of the value of specunit

average -- averaging on spectral data
options: (bool) True,False
default: False

>>>average expandable parameter

scanaverage -- average integrations within scans
options: (bool) True,False
default: False
example: if True, this happens in read-in
For GBT, set False!

timeaverage -- average times for multiple scan cycles
options: (bool) True,False
default: False
example: if True, this happens after calibration
tweight -- weighting for time average
options: 'none'
    'var'  (1/var(spec) weighted)
    'tsys' (1/Tsys**2 weighted)
    'tint' (integration time weighted)
    'tintsys' (Tint/Tsys**2)
    'median' (median averaging)
default: 'none'

averageall -- average multi-resolution spectra
spectra are averaged by referring their frequency coverage
default: False

polaverage -- average polarizations
options: (bool) True,False
default: False

pweight -- weighting for polarization average
options: 'none'
    'var'  (1/var(spec) weighted)
    'tsys' (1/Tsys**2 weighted)

tau -- atmospheric optical depth
default: 0.0 (no correction)

kernel -- type of spectral smoothing
options: 'none','hanning','gaussian','boxcar','regrid'
default: 'none' (no smoothing)

>>>kernel expandable parameter
kwidth -- width of spectral smoothing kernel
options: (int) in channels
default: 5
example: 5 or 10 seem to be popular for boxcar ignored for hanning (fixed at 5 chans)
(0 will turn off gaussian or boxcar)

chanwidth -- channel width of regridded spectra
default: '5' (in channels)
example: '500MHz', '0.2km/s'

masklist -- list or string of mask regions to INCLUDE in BASELINE fit
a string masklist allows per IF selection of channels.
default: [] (entire spectrum)
example: [[1000,3000],[5000,70000]]
    '0:1000~3000;5000~7000, 1:200~350;450~600'
when maskmode is 'auto' or 'interact', this mask will be applied first before fitting as base mask

maskmode -- mode of setting additional channel masks for baselining
options: (str) 'auto','list','interact'
default: 'auto'

example: maskmode='auto' runs linefinder to detect line regions to be excluded from fitting. This mode requires three expandable parameters: thresh, avg_limit, and edge.
USE WITH CARE! May need to tweak the expandable parameters.

maskmode='list' uses the given masklist only: no additional masks applied.

maskmode='interact' allows users to manually modify the mask regions by dragging mouse on the spectrum plotter GUI. use LEFT or RIGHT button to add or delete regions, respectively.

>>> maskmode expandable parameters

thresh -- S/N threshold for linefinder
default: 5
ex ample: a single channel S/N ratio above which the channel is considered to be a detection

avg_limit -- channel averaging for broad lines
default: 4
example: a number of consecutive channels not greater than this parameter can be averaged to search for broad lines

edge -- channels to drop at beginning and end of spectrum
default: 0
example: [1000] drops 1000 channels at beginning AND end
[1000,500] drops 1000 from beginning and 500 from end

Note: For bad baselines threshold should be increased, and avg_limit decreased (or even switched off completely by setting this parameter to 1) to avoid detecting baseline undulations instead of real lines.

blfunc -- baseline model function
options: (str) 'none','poly','chebyshev','cspline','sinusoid'
default: 'none' (no baselining)
ex ample: blfunc='poly' uses a single polynomial line of any order which should be given as an expandable parameter 'order' to fit baseline.
blfunc='chebyshev' uses Chebyshev polynomials.
blfunc='cspline' uses a cubic spline function, a piecewise cubic polynomial having C2-continuity (i.e., the second derivative is continuous at the joining points).
blfunc='sinusoid' uses a combination of sinusoidal curves.

>>> blfunc expandable parameters

order -- order of baseline polynomial
options: (int) (<0 turns off baseline fitting)
default: 5  
extample: typically in range 2-9 (higher values  
seem to be needed for GBT)

npiece -- number of the element polynomials of cubic spline curve  
options: (int) (<0 turns off baseline fitting)  
default: 2

applyfft -- automatically set wave numbers of sinusoidal functions  
for fitting by applying some method like FFT.  
options: (bool) True, False  
default: True

fftmethod -- method to be used when applyfft=True. Now only  
'fft' is available and it is the default.

fftthresh -- threshold to select wave numbers to be used for  
sinusoidal fitting. both (float) and (str) accepted.  
given a float value, the unit is set to sigma.  
for string values, allowed formats include:  
'xsigma' or 'x' (= x-sigma level. e.g., '3sigma'), or  
'topx' (= the x strongest ones, e.g. 'top5').  
default is 3.0 (unit: sigma).

addwn -- additional wave number(s) of sinusoids to be used  
for fitting.  
(list) and (int) are accepted to specify every  
wave numbers. also (str) can be used in case  
you need to specify wave numbers in a certain range,  
e.g., 'a-b' (= a, a+1, a+2, ..., b-1, b),  
'<a' (= 0,1,...,a-2,a-1),  
'>=a' (= a, a+1, ... up to the maximum wave  
number corresponding to the Nyquist  
frequency for the case of FFT).

default: []

rejwn -- wave number(s) of sinusoid NOT to be used for fitting.  
(can be set just as addwn but has higher priority:  
wave numbers which are specified both in addwn  
and rejwn will NOT be used.  
default: []

cliplthresh -- clipping threshold for iterative fitting  
default: 3

clipniter -- maximum iteration number  
default: 0 (no iteration, i.e., no clipping)

verificcal -- interactively verify the results of calibration  
See description of verify parameter in the task, sdcalold,  
for details.  
options: (bool) True,False  
default: False
verifysm -- interactively verify the results of smoothing for each spectrum.
See description of verify parameter in the task, sdsmoothold, for details.
options: (bool) True,False
default: False
Note: verification is not yet available for kernel='regrid'
verifybl -- interactively verify the results of baseline fitting for each spectrum.
See description of verify parameter in the task, sdbaselineold, for details.
options: (bool) True,False
default: False
NOTE: Currently available only when blfunc='poly'
verbosebl -- output fitting results to logger
default: True
example: If False, the fitting results including coefficients, residual rms, etc., are not output to the CASA logger, while the processing speed gets faster
bloutput -- output fitting results to a text file
default: True
example: If False, the fitting results including coefficients, residual rms, etc., are not output to a text file (<outfile>_blparam.txt), while the processing speed gets faster
blformat -- format of the logger output and text file specified with bloutput
options: '', 'csv'
default: '' (same as in the past, easy to read but huge)
showprogress -- show progress status for large data
default: True
minnrow -- minimum number of input spectra to show progress status
default: 1000
outfile -- Name of output file
default: '' (<infile>_cal)
outform -- format of output file
options: 'ASCII','SDFITS','MS','ASAP'
default: 'ASAP'
example: the ASAP format is easiest for further sd processing; use MS for CASA imaging.
If ASCII, then will append some stuff to the outfile name
overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: False
WARNING: if outform='ASCII', this parameter is ignored
plotlevel -- control for plotting of results
    options: (int) 0=none, 1=some, 2=more, <0=hardcopy
    default: 0 (no plotting)
    example: plotlevel<0 as abs(plotlevel), e.g.
             -1 => hardcopy of final plot (will be named
                <outfile>_calspec.eps)
    WARNING: be careful plotting in fsotf mode!

DESCRIPTION:

Task sdreduceold performs data selection, calibration, and/or spectral baseline fitting for single-dish spectra. This task internally calls the tasks, sdcalold, sdsmoothold, and sdbaselineold and it can be used to run all the three steps in one task execution.
By setting calmode='none', one can run sdreduceold on already calibrated data, for further selection, averaging and atmospheric optical depth correction. To save the output spectra in a certain range of channels, you set the range in channelrange.

If you give multiple IFs in iflist, then your scantable will have multiple IFs by default. Averaging of multi-resolution (multi-IFs) spectra can be achieved by setting a sub-parameter of average, averageall, to True. It handles multi-IFs by selecting overlaps in frequency coverages and assigning new IFs in the output spectra.

ASAP recognizes the data of the "AT" telescopes, but currently does not know about the GBT or any other telescope. This task does know about GBT. Telescope name is obtained from the data.
If you wish to change the fluxunit (see below), by leaving the sub-parameter telescopeparm unset (telescopeparm=''), it will use internal telescope parameters for flux conversion for the data from AT telescopes and it will use an approximate aperture efficiency conversion for the GBT data.
If you give telescopeparm a list, then if the list has a single float it is assumed to be the gain in Jy/K, if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively.

Note that sdreduceold assumes that the fluxunit is set correctly in the data already. If not, then set telescopeparm='FIX' and it will set the default units to fluxunit without conversion.
NOTE: If the data in infile is an ms from GBT and the default flux unit is missing and this task automatically fixes the default fluxunit to 'K' before the conversion.
WARNING for the GBT raw SDFITS format data as input: SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

- `sdflagold`

**Keyword arguments:**
- `infile` -- name of input SD dataset
- `antenna` -- antenna name or id (only effective for MS input).
- `specunit` -- units for spectral axis
  - Options: (str) 'channel', 'km/s', 'GHz', 'MHz', 'kHz', 'Hz', ''
  - Default: '' (=current)
  - Example: this will be the units for maskflag

```python
>>> specunit expandable parameters
restfreq -- rest frequency
```

- `restfreq` -- rest frequency
  - Available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. The default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz', 'kHz', and 'Hz' are available.
  - A list can be used to set different rest frequencies for each IF. The length of list input must be nIF. Dictionary input should be a pair of molecule name and frequency with keys of 'name' and 'value', respectively. Values in the dictionary input follows the same manner as for single float or string input.
  - Example: 345.796
    - 1420MHz
  - [345.8, 347.0, 356.7]
  - ['345.8MHz', '347.0MHz', '356.7MHz']
  - [{'name':'CO','value':345}]

- `frame` -- frequency frame for spectral axis
  - Options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY', 'GEO', 'GALACTO', 'LGROUP', 'CMB'
  - Default: currently set frame in scantable
  - WARNING: frame='REST' not yet implemented

- `doppler` -- doppler mode
  - Options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
  - Default: currently set doppler in scantable

- `scanlist` -- list of scan numbers to be considered for flagging and output
  - Default: [] (use all scans)
  - Example: [21,22,23,24]
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this selection is in addition to field and iflist

field -- field name for selecting scans to be considered for flagging and output
default: ' ' (no name selection)
example: 'FLS3a*

iflist -- list of IF id numbers to be considered for flagging and output
default: [] (use all IFs)
example: [15]

pollist -- list of polarization id numbers to be considered for flagging and output
default: [] (use all polarizations)
example: [1]

maskflag -- list of mask regions to apply flag/unflag
    Note, this parameter is ignored if one or more rows are
given in flagrow, or clip=True.
default: [] (entire spectrum)
example: [[1000,3000],[5000,7000]]

flagrow -- list of row numbers to apply flag/unflag
    Note, this parameter is effective only when one or more row
    numbers are given explicitly and also clip=False
default: [] (no row selection)
example: [0, 2, 3]

clip -- flag data that are outside a specified range
    options: (bool)True,False
default: False
>
>>> clip expandable parameters

clipminmax -- range of data that will NOT be flagged
default: [] means do not use clip option
example: [0.0,1.5]

clipoutside -- clip OUTSIDE the range ?
    options: (bool)True,False
default: True
example: clipoutside=False means flag data WITHIN the range.

flagmode -- flag mode
default: 'flag'
    options: 'flag','unflag'

interactive -- determines interactive flagging
    options: (bool) True,False
default: False
>
>>> interactive expandable parameters

showflagged -- show flagged data on plots
default: False

outfile -- Name of output file
default: ' '
Note: by default (outfile=''), actual output file name is set as follows:
(1) if overwrite=True (default), infile (input) will be overwritten.
WARNING: If the formats of input and output files are different, this causes complete loss of input file.
(2) if overwrite=False, outfile will be <infile>_f.

outform -- format of output file
options: 'ASCII','SDFITS','MS','ASAP'
default: 'ASAP'
example: the ASAP format is easiest for further sd processing; use MS for CASA imaging.
WARNING: Be sure outform is same as the input file format when you overwrite the input file by overwrite=True and outfile=''. (default)

overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: True
WARNING: input file is overwritten if overwrite=True and outfile=''. (default)
This causes the complete loss of input file if the formats of input and output files are different.

plotlevel -- control for plotting of results
options: (int) 0=none, 1=some, 2=more, <0=hardcopy
default: 0 (no plotting)
example: plotlevel<0 as abs(plotlevel), e.g.
-1 => hardcopy of final plot (will be named <outfile>_flag.eps)
WARNING: be careful plotting in fsotf mode!

-----------------------------------------------------------------------------------

DESCRIPTION:

Task sdflagold performs both interactive and non-interactive channel/row based flagging on spectra.
Currently, the available ways of non-interactive flagging include:
(1) channel based flagging by specifying a range of spectral values with clip=True, (2) row based flagging by specifying a list of row numbers to the flagrow parameter, and (3) channel based flagging by specifying regions in channel to the maskflag parameter.
These three ways of flagging can not be executed simultaneously. If more than one parameter above are specified, the task looks for them in the above order and operates the first specified way of flagging operation.

Interactive flagging is available when interactive=True. The available ways of interactive flagging include:
(1) row based flagging by selecting 'panel' and (2) channel
based flagging by selecting 'region's of channels on Flag plotter. Note that the Flag plotter is loaded after carrying out the non-interactive flag operation if any specified. See the cookbook for details of how to select channel regions and spectra on the plotter.

If plotlevel>=1, the task asks you if you really apply the flags before it is actually written to the data with a plot indicating flagged regions. Please note that this task is still experimental.

WARNING for overwrite option:
Be sure 'outform' is the same as data format of input file when you overwrite it. Since CASA 3.1, the default value of the option 'overwrite' has been changed to True, thereby the current dataset (infile) is overwritten unless a different file name is set to outfile. There is a known issue in overwriting infile. If 'outform' differs to the data format of infile, the data is overwritten with the new data format (specified by 'outform') and the data in the original format will be lost.

WARNING for the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

• sdflag2old

Keyword arguments:
infile -- name of input SD dataset
antenna -- antenna name or id (only effective for MS input).
specunit -- units for spectral axis
options: (str) 'channel','km/s','GHz','MHz','kHz','Hz',''
default: '' (=current)
example: this will be the units for maskflag
>>> specunit expandable parameters
restfreq -- rest frequency
available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz','kHz', and 'Hz' are available.
a list can be used to set different rest frequencies for each IF. the length of list input must be nIF. dictionary input should be a pair of molecule name and frequency with keys of 'name' and 'value', respectively. values in the
dictionary input follows the same manner as for single float or string input.
example: 345.796
    '1420MHz'
    [345.8, 347.0, 356.7]
    ['345.8MHz', '347.0MHz', '356.7MHz']
    [{'name': 'CO', 'value': 345}]
frame -- frequency frame for spectral axis
    options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY', 'GEO', 'GALACTO', 'LGROUP', 'CMB'
    default: currently set frame in scantable
WARNING: frame='REST' not yet implemented
doppler -- doppler mode
    options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
    default: currently set doppler in scantable
mode -- type of flag operation
    options: (str) 'manual', 'clip', 'interactive', 'rowid'
    default: 'manual'

>>> common data selection parameters for all modes except mode='rowid'
scans -- list of scan numbers to process
    default: [] (use all scans)
    example: [21, 22, 23, 24]
    this selection is in addition to field, timerange, ifs and pols
field -- selection string for selecting scans by name
    default: '' (no name selection)
    example: 'FLS3a*'
    this selection is in addition to scans, timerange, ifs, and pols
timerange -- Select data based on time range:
    default = '' (all); example,
timerange = 'YYYY/MM/DD/hh:mm:ss~YYYY/MM/DD/hh:mm:ss'
    Note: YYYY/MM/DD can be dropped as needed:
timerange='09:14:0~09:54:0' # this time range
timerange='09:44:00' # data within one integration of time
timerange='>10:24:00' # data after this time
timerange='09:44:00+00:13:00' # data 13 minutes after time
ifs -- list of IF id numbers to select
    default: [] (use all IFs)
    example: [15]
    this selection is in addition to scans, field, timerange, ifs, and pols
pols -- list of polarization id numbers to select
    default: [] (use all polarizations)
example: [1]
this selection is in addition to scans, field, timerange, and ifs

>>> mode='manual' expandable parameters
maskflag -- list of mask regions to apply flag/unflag
Note, this parameter is ignored if one or more rows are given in flagrow, or clip=True.
default: [] (entire spectrum)
example: [[1000,3000],[5000,7000]]

>>> mode='clip' expandable parameters
clipminmax -- range of data that will NOT be flagged
default: [] (means no clip operation)
example: [0.0,1.5]
clipoutside -- clip OUTSIDE the range?
options: (bool)True,False
default: True
example: clipoutside=False means flag data WITHIN the range.

>>> mode='interactive' expandable parameters
showflagged -- show flagged data on plots
default: False

>>> mode='rowid' expandable parameters
rows -- a list of row IDs to apply flag/unflag in the input scannable
Note, this parameter is effective only when one or more row IDs are given explicitly
default: [] (means no selection)
example: [0, 2, 3]

unflag -- flag or unflag
default: False (flag selected data)
options: (bool) True, False

outfile -- Name of output file
default: ''
Note: by default (outfile=''), actual output file name is set as follows:
(1) if overwrite=True (default), infile (input) will be overwritten.
WARNING: If the formats of input and ouput files are different,
this causes complete loss of input file.
(2) if overwrite=False, outfile will be <infile>_f.

outform -- format of output file
options: 'ASCII','SDFITS','MS','ASAP'
default: 'ASAP'
example: the ASAP format is easiest for further sd processing; use MS for CASA imaging.
WARNING: Be sure outform is same as the input file format when you overwrite the input file by overwrite=True and outfile=''' (default)
overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: True
WARNING: input file is overwritten if overwrite=True and outfile=’’ (default).
This causes the complete loss of input file if the formats of input and output files are different.

plotlevel -- control for plotting of results
options: (int) 0=none, 1=some, 2=more, <0=hardcopy
default: 0 (no plotting)
example: plotlevel<0 as abs(plotlevel), e.g.
-1 => hardcopy of final plot (will be named <outfile>_flag.eps)
WARNING: be careful plotting in fsotf mode!

---------------------------------------------

DESCRIPTION:

Task sdflag2old performs either interactive or non-interactive channel/row based flagging on spectra.
Currently, there are three ways of non-interactive flagging available:
(1) channel or row based flagging by selecting spectra by field, lists of scan numbers, IF numbers, and polarization indices in mode='manual',
(2) channel based flagging by specifying a range of spectral values in mode='clip', and
(3) row based flagging by specifying a list of row numbers in mode='rowid'. Note this is an EXPERT mode.

In mode='manual', the channel based flagging are invoked when regions in channel, frequency, or velocity are specified to maskflag parameter. Otherwise, the whole channels are flagged for the selected spectra. Note the mode='rowid' is an EXPERT mode since it might not be straight forward for general users to select data by row IDs in scantable.

Interactive flagging is available when mode='interactive'. The available ways of interactive flagging include:
(1) row based flagging by selecting 'panel' and (2) channel based flagging by selecting 'region's of channels on Flag plotter.
See the cookbook for details of how to select channel regions and spectra on the plotter.

NOTE the task sdflag2old only modifies flag information, FLAGROW and FLAGTRA, in the input scantable, and does not filter rows in the dataset unlike sdflagold task.
If plotlevel>=1, the task asks you if you really apply the flags before it is actually written to the data with a plot indicating flagged regions.
Please note that this task is still experimental.

WARNING for overwrite option:
Be sure 'outform' is the same as data format of input file when you overwrite it. The default value of the option 'overwrite' is True in this task, thereby the current dataset (infile) is overwritten unless a different file name is set to outfile.
There is a known issue in overwriting infile. If 'outform' differs to the data format of infile, the data is overwritten with the new data format (specified by 'outform') and the data in the original format will be lost.

WARNING for the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

- sdfitold

Keyword arguments:
infile -- name of input SD dataset
    default: none - must input file name
    example: 'mysd.asap'
        See sdcalold for allowed formats.
antenna -- antenna name or id (only effective for MS input).
fluxunit -- units for line flux
    options: (str) 'K','Jy',''
    default: '' (keep current fluxunit)
        WARNING: For GBT data, see description below.
    >>> fluxunit expandable parameter
    telescopeparm -- the telescope characteristics
        options: (str) name or (list) list of gain info
        default: '' (none set)
        example: if telescopeparm='', it tries to get the telescope name from the data.
        Full antenna parameters (diameter,ap.eff.) known to ASAP are
        'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43', 'CEDUNA','HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
        telescopeparm=[104.9,0.43] diameter(m), ap.eff.
        telescopeparm=[0.743] gain in Jy/K
        telescopeparm='FIX' to change default fluxunit
see description below

**specunit** -- units for spectral axis
- options: (str) 'channel', 'km/s', 'GHz', 'MHz', 'kHz', 'Hz', ''
- default: '' (=current)
- example: this will be the units for maskline

>>> **specunit** expandable parameters

restfreq -- rest frequency
- available type includes float, int, string, list of float, list of int, list of string, and list of dictionary. the default unit of restfreq in case of float, int, or string without unit is Hz. string input can be a value only (treated as Hz) or a value followed by unit for which 'GHz', 'MHz', 'kHz', and 'Hz' are available.
- a list can be used to set different rest frequencies for each IF. the length of list input must be nIF. dictionary input should be a pair of molecule name and frequency with keys of 'name' and 'value', respectively. values in the dictionary input follows the same manner as for single float or string input.
- example: 345.796
  - '1420MHz'
  - [345.8, 347.0, 356.7]
  - ['345.8MHz', '347.0MHz', '356.7MHz']
  - [{'name': 'CO', 'value': 345}]

frame -- frequency frame for spectral axis
- options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY', 'GEO', 'GALACTO', 'LGROUP', 'CMB'
- default: currently set frame in scantable
- WARNING: frame='REST' not yet implemented

**doppler** -- doppler mode
- options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
- default: currently set doppler in scantable

**scanlist** -- list of scan numbers to process
- default: [] (use all scans)
- example: [21, 22, 23, 24]

**field** -- selection string for selecting scans by name
- default: '' (no name selection)
- example: 'FLS3a*'
- this selection is in addition to scanlist and iflist

**iflist** -- list of IF id numbers to select
- default: [] (use all IFs)
- example: [15]

**pollist** -- list of polarization id numbers to select
- default: [] (use all polarizations)
example: [1]

fitfunc -- function for fitting
options: (str) 'gauss','lorentz'
default: 'gauss'

fitmode -- mode for fitting
options: (str) 'list','auto','interact'
default: 'auto'

example: 'list' will use maskline to define regions to
fit for lines with nfit in each
'auto' will use the linefinder to fit for lines
using the following parameters
'interact' allows adding and deleting mask
regions by drawing rectangles on the plot
with mouse. Draw a rectangle with LEFT-mouse
to ADD the region to the mask and with RIGHT-mouse
to DELETE the region.

>>> fitmode expandable parameters
thresh -- S/N threshold for linefinder
default: 5
example: a single channel S/N ratio above which the channel is
considered to be a detection

min_nchan -- minimum number of consecutive channels for linefinder
default: 3
example: minimum number of consecutive channels required to pass threshold

avg_limit -- channel averaging for broad lines
default: 4
example: a number of consecutive channels not greater than
this parameter can be averaged to search for broad lines

box_size -- running mean box size
default: 0.2
example: a running mean box size specified as a fraction
of the total spectrum length

derge -- channels to drop at beginning and end of spectrum
default: 0
example: [1000] drops 1000 channels at beginning AND end
[1000,500] drops 1000 from beginning and 500 from end

Note: For bad baselines threshold should be increased,
and avg_limit decreased (or even switched off completely by
setting this parameter to 1) to avoid detecting baseline
undulations instead of real lines.

maskline -- list of mask regions to INCLUDE in LINE fitting
default: all
example: maskline=[[3900,4300]] for a single region, or
    maskline=[[3900,4300],[5000,5400]] for two, etc.
invertmask -- invert mask (EXCLUDE masklist instead)
    options: (bool) True, False
    default: False
    example: invertmask=True, then will make one region that is
    the exclusion of the maskline regions
nfit -- list of number of gaussian/lorentzian lines to fit in in maskline region (in
    default: 0 (no fitting)
    example: nfit=[1] for single line in single region,
    nfit=[2] for two lines in single region,
    nfit=[1,1] for single lines in each of two regions, etc.
outfile -- name of output file for fit results
    default: no output fit file
    example: 'mysd.fit'
overwrite -- overwrite the outfile if already exists
    options: (bool) True, False
    default: False
plotlevel -- control for plotting of results
    options: (int) 0=none, 1=some, 2=more
    default: 0 (no plotting)
    example: plotlevel=1 plots fit
    plotlevel=2 plots fit and residual
    no hardcopy available for fitter
    WARNING: be careful plotting OTF data with lots of fields

Returns a Python dictionary of line statistics
    keys: 'peak','cent','fwhm','nfit'
    example: each value is a list of lists with one list of
    2 entries [fitvalue,error] per component.
    e.g. xstat['peak']=[[234.9, 4.8],[234.2, 5.3]]
    for 2 components.

DESCRIPTION:

Task sdfitold is a basic line-fitter for single-dish spectra.
It assumes that the spectra have been calibrated in sdcalold
or sdreduceold.

Furthermore, it assumes that any selection of scans, IFs,
polarizations, and time and channel averaging/smoothing has
also already been done (in other sd tasks) as there are no controls
for these. Note that you can use sdsaveold to do selection, writing
out a new scantable.

Note that multiple scans, IFs, and polarizations can in principle be handled, but we recommend that you use scanlist, field, iflist, and pollist to give a single selection for each fit.

Currently, you can choose Gaussian or Lorentzian profile as a fitting model.

For complicated spectra, sdfitold does not do a good job of "auto-guessing" the starting model for the fit. We recommend you use sd.fitter in the toolkit which has more options, such as fixing components in the fit and supplying starting guesses by hand.

WARNING: sdfitold will currently return the fit for the first row in the scantable.

ASAP recognizes the data of the "AT" telescopes, but currently does not know about the GBT or any other telescope. This task does know about GBT. Telescope name is obtained from the data. If you wish to change the fluxunit (see below), and telescopeparm='', for the AT telescopes it will use internal telescope parameters for flux conversion. For GBT, it will use an approximate aperture efficiency conversion. If you give telescopeparm a list, then if the list has a single float it is assumed to be the gain in Jy/K, if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively.

WARNING for the GBT raw SDFITS format data as input: SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

• sdgridold

Keyword arguments:

infiles -- name of input SD dataset. can be list.
example: 'testimage.asap'
          ['testimage1.asap','testimage2.asap']
antenna -- select data based on antenna name(s) or id(s)
default: -1
example: 0, 'DV01'
scanlist -- list of scan numbers to process
default: [] (use all scans)
example: [21, 22, 23, 24]

\textbf{ifno} -- IFNO to be gridded
\begin{itemize}
  \item default: -1 (only process IFNO in the first row)
  \item example: 1
\end{itemize}

\textbf{pollist} -- POLNO to be gridded
\begin{itemize}
  \item default: [] (all polarizations)
  \item example: 1, [0, 1]
\end{itemize}

\textbf{gridfunction} -- gridding function
\begin{itemize}
  \item options: 'BOX' (Box-car), 'SF' (Spheroidal), 'GAUSS' (Gaussian), 'PB' (Primary-beam), 'GJINC' (Gaussian*Jinc)
  \item default: 'BOX'
  \item example: 'SF'
\end{itemize}

\textbf{>>> gridfunction expandable parameter:}
\begin{itemize}
  \item \textbf{convsupport} -- convolution support for 'SF'
    \begin{itemize}
      \item default: -1 (use default for each gridfunction)
      \item example: 3
    \end{itemize}
  \item \textbf{truncate} -- truncation radius of convolution kernel. Effective only for 'GAUSS' and 'GJINC'.
    \begin{itemize}
      \item default: '-1' (use default for each gridfunction)
      \item example: 3, '20arcsec', '3pixel'
    \end{itemize}
  \item \textbf{gwidth} -- HWHM for gaussian. Effective only for 'GAUSS' and 'GJINC'.
    \begin{itemize}
      \item default: '-1' (use default for each gridfunction)
      \item example: 3, '20arcsec', '3pixel'
    \end{itemize}
  \item \textbf{jwidth} -- Width of jinc function. Effective only for 'GJINC'.
    \begin{itemize}
      \item default: '-1' (use default for each gridfunction)
      \item example: 3, '20arcsec', '3pixel'
    \end{itemize}
\end{itemize}

\textbf{weight} -- weight type (both lower-case and upper-case are acceptable)
\begin{itemize}
  \item options: 'UNIFORM', 'TSYS' (1/Tsys**2 weighted), 'TINT' (integration time weighted), 'TINTSYS' (Tint/Tsys**2)
  \item default: 'UNIFORM'
\end{itemize}

\textbf{clipminmax} -- do min/max clipping if True
\begin{itemize}
  \item default: False
\end{itemize}

\textbf{outfile} -- output data name
\begin{itemize}
  \item default: '' (outfile will be set to infile[0]+'.grid')
  \item example: 'mydata.asap.grid'
\end{itemize}

\textbf{overwrite} -- overwrite option for outfile
\begin{itemize}
  \item default: False (not overwrite)
  \item options: True, False
  \item example: if True, existing file will be overwritten
npix -- x and y image size in pixels, symmetric for single value
default: -1 (automatically calculated from cell size and the data)
example: npix=200 (equivalent to [200, 200])
cell -- x and y cell size. default unit arcsec
default: '' (automatically calculated from npix if it is set, otherwise '1.0arcmin')
example: cell= ['0.2arcmin', '0.2arcmin']
cell='0.2arcmin' (equivalent to example above)
cell=12.0 (interpreted as '12.0arcsec'='0.2arcmin')
center -- grid center
default: '' (automatically calculated from the data)
example: 'J2000 13h44m00 -17d02m00'
        ['05:34:48.2', '-05.22.17.7'] (in J2000 frame)
        [1.46, -0.09] (interpreted as radian in J2000 frame)
plot -- Plot result or not
default: False (not plot)
example: if True, result will be plotted

DESCRIPTION:

The sdgridold task performs spatial gridding according to the user specification of spatial grid, convolution function, etc.

For grid configuration, the task supplements necessary information by referring input data if any of gridding parameter ('npix', 'cell', or 'center') is not specified by the user. If 'center' is default value (empty string), central position of the grid will be set to the center of observed area, i.e. x=0.5*(xmax+xmin), y=0.5*(ymax+ymin). If either 'cell' or 'npix' is set, unspecified one will be calculated from the others. In that case, total extent of the grid will be set to cover all observed position. If neither 'cell' nor 'npix' is set, cell size will be set to 1.0 arcmin and number of pixel will be calculated based on that cell size.

Currently, only J2000 frame is supported.

The parameter gridfunction sets gridding function for imaging. Currently, the task supports 'BOX' (Box-car), 'SF' (Prolate Spheroidal Wave Function), 'GAUSS' (Gaussian), 'GJINC' (Gaussian* Jinc), where Jinc(x) = J_1(pi*x/c)/(pi*x/c) with a first order Bessel function J_1, and 'PB' (Primary Beam, not implemented yet). For 'PB', correct antenna informations should be included in input file.
There are four subparameters for gridfunction: convsupport, truncate, gwidth, and jwidth. The convsupport is an integer specifying cut-off radius for 'SF' in units of pixel. By default (convsupport=-1), the cut-off radius is set to 3 pixels. The truncate is a cut-off radius for 'GAUSS' or 'GJINC'. It accepts integer, float, and string values of numeric plus unit. Allowed units are angular units such as deg, arcmin, arcsec, and pixel. Default unit is 'pixel' so that string without unit or numerical values (integer or float) will be interpreted as radius in pixel. Default value for truncate, which is used when negative radius is set, is 3*HWHM for 'GAUSS' and radius at first null for 'GJINC'. The gwidth is the HWHM of gaussian for 'GAUSS' and 'GJINC'. Default value is sqrt(log(2)) pixel for 'GAUSS' and 2.52*sqrt(log(2)) pixel for 'GJINC'. The jwidth specifies width of the jinc function (parameter 'c' in the definition above). Default is 1.55 pixel. Both gwidth and jwidth allow integer, float, or string of numeric plus unit. Default values for gwidth and jwidth are taken from Mangum et al. (2007). Formula for 'GAUSS' and 'GJINC' are taken from Table 1 in the paper, and are written as below using gwidth and jwidth:

GAUSS: \[ \text{exp}\left[-\left(|r|/\text{gwidth}\right)^2\right] \]

GJINC: \[ \frac{J_1(\pi*|r|/\text{jwidth})}{(\pi*|r|/\text{jwidth})} * \text{exp}\left[-\left(|r|/\text{gwidth}\right)^2\right] \]

Boolean parameter 'plot' controls whether gridded result is plotted or not. If True, color map of gridded data will be shown. Pixel center and observed position are overlayed as blue dot and red dot, respectively. Currently, channel averaged value will be plotted.


Keyword arguments:

- **infiles** -- a list of names of input SD (MS) datasets
  
  example: 'm100.PM01.ms'
  
  
  example: ['m100.PM01.ms','m100.PM03.ms']; multiple MSes

- **specunit** -- units for spectral axis
  
  options: (str) 'channel','km/s','GHz','MHz','kHz','Hz'
  
  default: 'channel'

  example: this will be the units for nchan, start, and step

- **restfreq** -- rest frequency
  
  default: '' (refer input data)

  example: 1.0e11, '100GHz'
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scanlist -- list of scan numbers to process
  default: [] (use all scans)
  example: [21,22,23,24]
  this selection is in addition to field and spw

field -- field id or selection string for selecting scans by name
  default: -1 (all fields)
  example: 'FLS3a', 0
  this selection is in addition to scanlist and spw

spw -- a list of spectral window id
  default: 0
  example: 1
  [1, 3, 5]
  this selection is in addition to scanlist and field

antenna -- select data based on antenna name(s) or id(s)
  default: -1 (all baselines, i.e. all antenna in case of auto data)
  example: 0, 'DV01'

stokes -- select data based on stokes or polarization type
  default: '' (= Stokes I)
  example: 'XX'

gridfunction -- gridding function for imaging
  options: 'BOX' (Box-car), 'SF' (Spheroidal),
    'PB' (Primary-beam), 'GAUSS' (Gaussian),
    'GJINC' (Gaussian*Jinc)
  default: 'BOX'
  example: 'SF'

>>> gridfunction expandable parameter:
    convsupport -- convolution support for 'SF'
      default: -1 (use default for each gridfunction)
      example: 3

    truncate -- truncation radius of convolution kernel.
      effective only for 'GAUSS' and 'GJINC'.
      default: '-1' (use default for each gridfunction)
      example: 3, '20arcsec', '3pixel'

    gwidth -- HWHM for gaussian. Effective only for
      'GAUSS' and 'GJINC'.
      default: '-1' (use default for each gridfunction)
      example: 3, '20arcsec', '3pixel'

    jwidth -- Width of jinc function. Effective only for
      'GJINC'.
      default: '-1' (use default for each gridfunction)
      example: 3, '20arcsec', '3pixel'

minweight -- Minimum weight ratio to the median of weight used in
  weight correction and weight based masking
  default: 0.1
  example: minweight = 0.
CHAPTER 8. SINGLE DISH DATA PROCESSING

outfile -- output image name
    default: none
    example: 'mySDimage.im'

overwrite -- overwrite option for outfile
    default: False (not overwrite)
    options: True, False
    example: if True, existing file will be overwritten

imsizer -- x and y image size in pixels, symmetric for single value
    default: [] (=cover all pointings in MS)
    example: imsize=200 (equivalent to [200,200])

cell -- x and y cell size. default unit arcmin
    default: '' (= 1/3 of FWHM of primary beam)
    example: cell=['0.2arcmin', 0.2arcmin']
    cell='0.2arcmin' (equivalent to example above)

dochannelmap -- channel map image or total power image
    default: False (total power)
    options: True (channel map), False
    >>> dochannelmap=True expandable parameters

    nchan -- number of spectral channel for created image
        default: 1
        example: 100

    start -- reference value of start channel (in units of specunit)
        default: 0 (0th channel if specunit='channel')
        example: 100

    step -- width of each spectral channel for created image
        default: 1 (width of 1 channel if specunit='channel')
        example: 100

outframe -- Velocity reference frame of output image
    Options: ',','LSRK','LSRD','BARY','GEO','TOPO','GALACTO','LGROUP','CMB'
    default: ''; same as input data or 'LSRK' for multiple-MS inputs
    example: frame='bary' for Barycentric frame

phasedcenter -- image phase center: direction measure or field id
    default: '' (= the center of pointing directions in
        POINTING table of infiles)
    example: 6 (field id), 'J2000 13h44m00 -17d02m00',
        'AZEL -123d48m29 15d41m41'

ephemsrcname -- ephemeris source name for moving source
    default: ''
    if the source name in the data matches one of the known
    solar objects by the system, this task automatically set
    the source name.
    example: 'moon'

pointingcolumn -- pointing data column to use
    option: 'direction', 'target', 'pointing_offset', 'source_offset', 'encoder
CHAPTER 8. SINGLE DISH DATA PROCESSING

default: 'direction'

DESCRIPTION:

Task sdimagingold create image from input single-dish data. The input can be either total power and spectral data. Currently, this task directly accesses the Measurement Set data only because of the data access efficiency. So it differs from other single-dish tasks that mostly operate on the ASAP scantable data format.

Units of spectral axis can be specified via a parameter specunit. Allowed values for specunit are 'channel', 'GHz', 'MHz', 'kHz', 'Hz', and 'km/s'. This parameter is also used as the units of the parameter start and step that specify reference value of start channel and width of each spectral channel for channel map, respectively. The parameter nchan specifies number of channels for created image. If you set nchan as -1, the task selects existing all channels and combine those data into one channel to create continuum image.

You can specify field id or name directly. By default, field is set to -1 that means the task selects all fields in the data.

Selection of the antennas can be made by setting antennaid(s) or antenna name(s) in string (e.g. '0', 'DV01', etc.) or integer (e.g. 0). Default value, -1, means that the task selects data from all baseline, i.e., data from all antenna when data only contains auto-correlation.

The parameter gridfunction sets gridding function for imaging. Currently, the task supports 'BOX' (Box-car), 'SF' (Prolate Spheroidal Wave Function), 'GAUSS' (Gaussian), 'GJINC' (Gaussian*Jinc), where Jinc(x) = J_1(pi*x/c)/(pi*x/c) with a first order Bessel function J_1, and 'PB' (Primary Beam). For 'PB', correct antenna informations should be included in input file.

There are four subparameters for gridfunction: convsupport, truncate, gwidth, and jwidth. The convsupport is an integer specifying cut-off radius for 'SF' in units of pixel. By default (convsupport=-1), the cut-off radius is set to 3 pixels. The truncate is a cut-off radius for 'GAUSS' or 'GJINC'. It accepts integer, float, and string values of numeric plus unit. Allowed units are angular units such as 'deg', 'arcmin', 'arcsec', and 'pixel'. Default unit is 'pixel' so that string without unit or numerical values (integer or float) will be interpreted as radius in pixel. Default value
for truncate, which is used when negative radius is set, is $3\times$HWHM for 'GAUSS' and radius at first null for 'GJINC'. The gwidth is the HWHM of gaussian for 'GAUSS' and 'GJINC'. Default value is \sqrt{\log(2)} pixel for 'GAUSS' and 2.52*\sqrt{\log(2)} pixel for 'GJINC'. The jwidth specifies width of the jinc function (parameter 'c' in the definition above). Default is 1.55 pixel. Both gwidth and jwidth allows integer, float, or string of numeric plus unit. Default values for gwidth and jwidth are taken from Mangum et al. (2007). Formula for 'GAUSS' and 'GJINC' are taken from Table 1 in the paper, and are written as below using gwidth and jwidth:

\[
\text{GAUSS: } \exp\left[-\log(2) \times \left(\frac{|r|}{\text{gwidth}}\right)^2\right]
\]

\[
\text{GJINC: } J_1(\pi \times \frac{|r|}{\text{jwidth}}) \div \frac{\pi \times |r|}{\text{jwidth}} \times \exp\left[-\log(2) \times \left(\frac{|r|}{\text{gwidth}}\right)^2\right]
\]


The parameter minweight defines a threshold of weight values to mask. The pixels in outfile whose weight is smaller than minweight*median(weight) are masked out. The task also creates a weight image with the name outfile.weight.

• sdmathold

Keyword arguments:

expr -- Mathematical expression using scantables
varlist -- Dictionary of variables in expr and their values.
Keys must be coincide with variables used in expr.
Values are substituted in each value in expr.
antenna -- antenna name or id (only effective for MS input).
fluxunit -- units for line flux
options: 'K','Jy',''
default: '' (keep current fluxunit)
WARNING: For GBT data, see description below.

>>> fluxunit expandable parameter
telescopeparm -- the telescope characteristics
options: (str) name or (list) list of gain info
default: '' (none set)
example: if telescopeparm='', it tries to get the telescope name from the data.
Full antenna parameters (diameter,ap.eff.) known to ASAP are
'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43',
'CEDUNA', 'HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
telescopeparm=[104.9, 0.43] diameter(m), ap.eff.
telescopeparm=[0.743] gain in Jy/K

`telescopeparm='FIX'` to change default fluxunit
see description below

```
specunit -- units for spectral axis
    options: (str) 'channel', 'km/s', 'GHz', 'MHz', 'kHz', 'Hz'
    default: '' (=current)
    example: this will be the units for masklist

frame -- frequency frame for spectral axis
    options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY', 'GEO', 'GALACTO', 'LGROUP', 'CMB'
    default: currently set frame in scantable
WARNING: frame='REST' not yet implemented

doppler -- doppler mode
    options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
    default: currently set doppler in scantable

scanlist -- list of scan numbers to process
    default: [] (use all scans)
    example: [21, 22, 23, 24]
    this selection is in addition to field, iflist, and pollist

field -- selection string for selecting scans by name
    default: '' (no name selection)
    example: 'FLS3a*'
    this selection is in addition to scanlist, iflist, and pollist

iflist -- list of IF id numbers to select
    default: [] (use all IFs)
    example: [15]
    this selection is in addition to scanlist, field, and pollist

pollist -- list of polarization id numbers to select
    default: [] (use all polarizations)
    example: [1]
    this selection is in addition to scanlist, field, and iflist

outfile -- Name of output file
    default: '' (must be specified)

outform -- format of output file
    options: 'ASCII', 'SDFITS', 'MS', 'ASAP'
    default: 'ASAP'
    example: the ASAP format is easiest for further sd processing; use MS for CASA imaging.
    If ASCII, then will append some stuff to the outfile name

overwrite -- overwrite the output file if already exists
```
Task sdmathold execute a mathematical expression for single dish spectra. The spectral data file can be any of the formats supported by ASAP (scantable, MS, rpfits, and SDFITS). In the expression, these file names should be put inside of single or double quotes.

You can use variables in the expression. If you want to use, you must define varlist dictionary. Name of variables should be simple, e.g. V0, V1, etc., to avoid unexpected error. Keys of varlist must be name of variables that you used in the expression, and their values will be substituted for variables in the expression. Allowed type for the value is numerical values, one- or two-dimensional lists (Python list or numpy.ndarray), and filename strings that indicate spectral data or ASCII text, which is space-separated list of numerical values consisting of adequate number of rows and columns.

The fluxunit, specunit, and frame can be set, otherwise, the current settings of the first spectral data in the expression are used. Other selections (e.g. scan No, . IF, Pol) also apply to all the spectral data in the expression, so if any of the data does not contains selection, the task will produce no output.

WARNING for the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

Example:
# do on-off/off calculation
expr='("orion_on_data.asap"-"orion_off_data.asap")/"orion_off_data.asap"
outfile='orion_cal.asap'
sdmathold()

# do on-off/off calculation using varlist
expr='V0/V1-V2'
varlist['V0']='orion_on_data.asap'
varlist['V1']='orion_off_data.asap'
varlist['V2']=1.0
outfile='orion_cal.asap'
sdmathold()

# interpretation of ASCII file value for varlist
If the contents of input ASCII file is shown as,

0.5 0.3 0.2
1.0 0.2 0.9

it is interpreted as a list [[0.5, 0.3, 0.2],[1.0, 0.2, 0.9]].

• sdplotold

Keyword arguments:

infile -- name of input SD dataset
antenna -- antenna name or id (only effective for MS input).
fluxunit -- units for line flux
    options: 'K','Jy',''
    default: '' (keep current fluxunit)
    WARNING: For GBT data, see description below.

>>> fluxunit expandable parameter

   telescopeparm -- the telescope characteristics
    options: (str) name or (list) list of gain info
    default: '' (none set)
    example: if telescopeparm='', it tries to get the telescope
             name from the data.
    Full antenna parameters (diameter,ap.eff.) known
    to ASAP are
        'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43',
        'CEDUNA','HOBART'. For GBT, it fixes default fluxunit
        to 'K' first then convert to a new fluxunit.
    telescopeparm=[104.9,0.43] diameter(m), ap.eff.
    telescopeparm=[0.743] gain in Jy/K
    telescopeparm='FIX' to change default fluxunit
    see description below

specunit -- units for spectral axis
    options: (str) 'channel','km/s','GHz','MHz','kHz','Hz'
    default: '' (=current)
    example: this will be the units for masklist

>>> specunit expandable parameter

restfreq -- rest frequency
    default: '' (use current setting)
    example: 4.6e10 (float value in Hz),
             '46GHz' (string with unit),
             ['345.8GHz', 347.0e9, 356.7e9] (for each IF)
frame -- frequency frame for spectral axis
   options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY',
            'GEO', 'GALACTO', 'LGROUP', 'CMB'
   default: currently set frame in scantable
   WARNING: frame='REST' not yet implemented

doppler -- doppler mode
   options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
   default: currently set doppler in scantable

scanlist -- list or string of scan numbers to process
   default: [] (use all scans)
   example: [21, 22, 23, 24], or "21~24"
   this selection is in addition to field, iflist, pollist,
   and beamlist

field -- selection string for selecting scans by name
   default: '' (no name selection)
   example: 'FLS3a*' 
   this selection is in addition to scanlist, iflist, pollist,
   and beamlist

iflist -- list or string of IF id numbers to select
   default: [] (use all IFs)
   example: [15]
   this selection is in addition to scanlist, field, pollist,
   and beamlist

pollist -- list or string of polarization id numbers to select
   default: [] (use all polarizations)
   example: [1]
   this selection is in addition to scanlist, field, iflist,
   and beamlist

beamlist -- list or string of beam id numbers to select
   default: [] (use all beams)
   example: [1]
   this selection is in addition to scanlist, field, iflist,
   and pollist

scanaverage -- average integs within scans
   options: (bool) True, False
   default: False

timeaverage -- average times for multiple scan cycles
   options: (bool) True, False
   default: False
   example: if True, this happens after calibration
   >>>timeaverage expandable parameter

tweight -- weighting for time average
   options: 'var' (1/var(spec) weighted)
            'tsys' (1/Tsys**2 weighted)
'tint' (integration time weighted)
'tintsys' (Tint/Tsys**2)
'median' (median averaging)

default: 'tintsys'

polaverage -- average polarizations
options: (bool) True,False
default: False

>>polaverage expandable parameter

pweight -- weighting for polarization average
options: 'var' (1/var(spec) weighted)
'tsys' (1/Tsys**2 weighted)

default: 'tsys'

kernel -- type of spectral smoothing
options: 'hanning','gaussian','boxcar', 'none'
default: 'none'

>>kernel expandable parameter

kwidth -- width of spectral smoothing kernel
options: (int) in channels
default: 5

example: 5 or 10 seem to be popular for boxcar
ignored for hanning (fixed at 5 chans)
(0 will turn off gaussian or boxcar)

plottype -- type of plot
options: 'spectra','totalpower','pointing','azel','grid'
default: 'spectra'

>>plottype expandable parameters

stack -- code for stacking on single plot for spectral plotting
options: 'p','b','i','t','s','r' or
'pol', 'beam', 'if', 'time', 'scan', 'row'
default: 'p'

example: maximum of 16 stacked spectra
stack by pol, beam, if, time, scan

Note stack selection is ignored when panel='r'.

panel -- code for splitting into multiple panels for spectral plotting
options: 'p','b','i','t','s','r' or
'pol', 'beam', 'if', 'time', 'scan', 'row'
default: 'i'

example: maximum of 16 panels
panel by pol, beam, if, time, scan

Note panel selection is ignored when stack='r'.

flrange -- range for flux axis of plot for spectral plotting
options: (list) [min,max]
default: [] (full range)

example: flrange=[-0.1,2.0] if 'K'

assumes current fluxunit
sprange -- range for spectral axis of plot
options: (list) [min,max]
default: [] (full range)
example: sprange=[42.1,42.5] if 'GHz'
  assumes current specunit

linecat -- control for line catalog plotting for spectral plotting
options: (str) 'all','none' or by molecule
default: 'none' (no lines plotted)
example: linecat='SiO' for SiO lines
      linecat='*OH' for alcohols
      uses sprange to limit catalog
WARNING: specunit must be in frequency (*Hz)
  to plot from the line catalog!
  and must be 'GHz' or 'MHz' to use
  sprange to limit catalog

tlinedop -- doppler offset for line catalog plotting (spectral plotting)
options: (float) doppler velocity (km/s)
default: 0.0
example: linedop=-30.0

tcenter -- the central direction of gridding
default: '' (map center)
example: 'J2000 19h30m00 -40d00m00'
  Note currently only supports 'J2000' as direction frame

tcell -- x and y cell size of gridding
default: [] (map extent devided by # of subplots in x and y)
example: cell=['1.0arcmin','1.0arcmin']
      cell='1.0arcmin' (equivalent to the example above)
  Note default number of subplots is 1 x 1 in plottype='grid'.

tsubplot -- number of subplots (row and column) on a page
NOTICE plotter will slow down when a large number is specified
default: -1 (auto. for plottype='spectra', 1x1 for plottype='grid')
example: 23 (2 rows by 3 columns)

tcolormap -- the colours to be used for plot lines.
default: None
example: colormap="green red black cyan magenta" (html standard)
      colormap="g r k c m" (abbreviation)
      colormap="#008000 #00FFFF #FF0090" (RGB tuple)
The plotter will cycle through these colours when lines are overlaid (stacking mode).

linestyles -- the linestyles to be used for plot lines.
default: None
example: linestyles="line dashed dotted dashdot dashdotdot dashdashdot".The plotter will cycle through these linestyles when lines are overlaid (stacking mode).
WARNING: Linestyles can be specified only one color has been set.
linewidth -- width of plotted lines.
  default: 1
  example: linewidth=1 (integer)
          linewidth=0.75 (double)

histogram -- plot histogram
  options: (bool) True, False
  default: False

scanpattern -- plot additional lines on the plot to indicate scan patterns
  when plottype='pointing'
  options: (bool) True, False
  default: False

header -- print header information on the plot
  options: (bool) True, False
  default: True
  The header information is printed only on the logger when
  plottype = 'azel' and 'pointing'.

headsize -- header font size
  options: (int)
  default: 9

plotstyle -- customise plot settings
  options: (bool) True, False
  default: False

margin -- a list of subplot margins in figure coordinate (0-1),
  i.e., fraction of the figure width or height.
  The order of elements should be:
  [left, bottom, right, top, horizontal space btw panels,
   vertical space btw panels]
  example: margin = [0.125, 0.1, 0.9, 0.9, 0.2, 0.2]

legendloc -- legend location on the axes (0-10)
  options: (integer) 0 -10
  see help of "sd.plotter.set_legend" for
  the detail of location. Note that 0 ('best')
  is very slow.
  default: 1 ('upper right')

outfile -- file name for hardcopy output
  options: (str) filename.eps,.ps,.png
  default: '' (no hardcopy)
  example: 'specplot.eps','specplot.png'
  Note this autodetects the format from the suffix (.eps,.ps,.png).

overwrite -- overwrite the output file if already exists
  options: (bool) True,False
  default: False
DESCRIPTION:

Task sdplotold displays single-dish spectra, total power, or pointing direction of input data. It assumes that the spectra have been calibrated. It does allow selection of scans, IFs, polarizations, and some time and channel averaging/smoothing options also, but does not write out this data.

This task adds an additional toolbar to Matplotlib plotter. See the cookbook for details of its capability.

*** Data selection ***
This task allows data selection via field name, scan, IF, polarization and beam IDs. Selection of field allows pattern matching using asterisk, e.g., 'FLS3a*'. Selection of scans, IFs, polarizations, and beams, is possible either by a list of IDs or by a CASA type selection syntax using a string of comma separated numbers with operators, i.e., '~', '>', '>=' ', '<', and '<='. For example, the following two selections are equivalent:
scanlist = [0, 1, 2, 7, 8, 9, 15]
scanlist = "<3,7~9,15"

*** control of plot lines in 'spectra' and 'grid' plottype ***
Note that colormap and linestyles cannot be controlled at a time. The 'linestyles' is ignored if both of them are specified. Some plot options, like changing titles, legends, fonts, and the like are not supported in this task. You should use sd.plotter from the ASAP toolkit directly for this.

*** available plottypes ***
* plottype = 'spectra' plots single dish spectra. Multiple scans, IFs, polarizations, and beams can be handles through stacking and panelling. This task uses the JPL line catalog as supplied by ASAP. If you wish to use a different catalog, or have it plot the line IDs from top or bottom (rather than alternating), then you will need to explore the sd toolkit also.
* plottype = 'grid' plots spectra based on their pointing direction. The spectra are gridded by direction before plotting. Multiple IFs and polarizations are not handled in this mode. Only the first IF and polarizaion is gridded and plotted if data includes multiple IDs after selections are applied. Hence, over
plotting is not available

Currently most of the parameters are ignored in the following modes.

* plottype='totalpower' is used to plot the total power data.
  and only plot option is amplitude versus data row number.
* plottype='azel' plots azimuth and elevation tracks of the source.
* plottype='pointing' plots antenna poinitings.

ASAP recognizes the data of the "AT" telescopes, but currently
does not know about the GBT or any other telescope. This task
does know about GBT. Telescope name is obtained from the data.
If you wish to change the fluxunit (see below), and telescopeparm=’’,
for the AT telescopes it will use internal telescope parameters for
flux conversion. For GBT, it will use an approximate aperture
efficiency conversion. If you give telescopeparm a list,
then if the list has a single float it is assumed to
be the gain in Jy/K, if two or more elements they are assumed
to be telescope diameter (m) and aperture efficiency
respectively.

WARNING: be careful plotting otf data with lots of fields!

WARNING for the GBT raw SDFITS format data as input:
SDtasks are able to handle GBT raw SDFITS format data since the
data filler is available. However, the functionality is not well
tested yet, so that there may be unknown bugs.

• sdsaveold

Keyword arguments:
infile -- name of input SD dataset
splitant -- split output file by antenna (True,False)
  (only effective for MS input).
    default: False
    >>>splitant expandable parameter
    antenna -- antenna name or id (only effective for MS input).
      default: 0
getpt -- fill DIRECTION column properly, or reuse POINTING table
  in original MS (only effective for MS input)
    default: True
    options: True (fill DIRECTION column properly)
           False (reuse POINTING table in original MS)
rowlist -- list of row numbers to process
    default: [] (use all rows)
example: [0,2,4,6]
For expert users only!
this selection is applied first, and then followed by
the selection with scans, fields, ifs, and polarizations.

scanlist -- list of to process
default: [] (use all scans)
example: [21,22,23,24]
this selection is in addition to field, iflist, and pollist

field -- selection string for selecting scans by name
default: '' (no name selection)
example: 'FLS3a*' 
this selection is in addition to scanlist, iflist, and pollist

iflist -- list of IF id numbers to select
default: [] (use all IFs)
example: [15] 
this selection is in addition to scanlist, field, and pollist

pollist -- list of polarization id numbers to select
default: [] (use all polarizations)
example: [1] 
this selection is in addition to scanlist, field, and iflist

scanaverage -- average integrations within scans
options: (bool) True,False
default: False
example: if True, average integrations before it is saved

timeaverage -- average times for multiple scan cycles
options: (bool) True,False
default: False
>>>timeaverage expandable parameter

tweight -- weighting for time average
options: 'none'
    'var'  (1/var(spec) weighted)
    'tsys'  (1/Tsys**2 weighted)
    'tint'  (integration time weighted)
    'tintsys'  (Tint/Tsys**2)
    'median'  ( median averaging)
default: 'none'

polaverage -- average polarizations
options: (bool) True,False
default: False
>>>polaverage expandable parameter

pweight -- weighting for polarization average
options: 'none'
    'var'  (1/var(spec) weighted)
    'tsys'  (1/Tsys**2 weighted)
restfreq -- rest frequencies of output data
Available types are a number, string, a list of numbers or strings (see examples below), and list of dictionaries.
The default unit of restfreq is Hz, if not specified.
A list can be used to set different rest frequencies to each IF. the length of list input must be nIF. Dictionary input should be a pair of molecule name and frequency with keys of 'name' and 'value', respectively. The 'value's in the dictionary input follows the same manner as for single float or string input.
default: '' (use current setting)
example: 4.6e10 (float value in Hz),
'46GHz' (string with unit),
['345.8GHz', '347.0GHz', 356.7e9] (for each IF)
[{'name':'CO','value':345e9}] (a value with name)

outfile -- name of output dataset
default: ''
NOTE actual output file name(s) will be modified if splitant is True as antenna names are to be included. If outfile has a suffix '.asap' or '.ASAP', antenna name will be inserted before the suffix like 'out.antName.asap', otherwise, antenna name will be simply appended to outfile like 'out.sdfits.antName'.

outform -- output data format
default: 'ASAP'
Options: 'ASAP', 'MS2', 'SDFITS', 'ASCII'

overwrite -- overwrite the output file if already exists
options: (bool) True,False
default: False
WARNING: if outform='ASCII', this parameter is ignored

DESCRIPTION:
Task sdsaveold writes the single dish data to a disk file in specified format (ASAP, MS2, SDFITS, ASCII). It is possible to save the subset of the data by selecting row numbers, scan numbers, IF ids, polarization ids and field names. The ASAP (scantable) format is recommended for further analysis using Sd tool. For further imaging using imager, save the data to the Measurement Set (MS2).

Note that setting getpt=False needs a lot of attention.
If you set getpt=False, the task retrieves pointing direction from MS's FIELD table, which might not be correct for single dish observation, instead to check MS's POINTING table, which is the default behavior of the task (getpt=True). To compensate this,
absolute path to MS’s POINTING table is stored, and it will be used for POINTING table when the data is converted back to MS format. In general, getpt=False is faster especially for large data. However, MS created from Scantable cannot have correct POINTING table if original MS’s POINTING table doesn’t exist. Such situation will happen when original MS is removed or renamed, or imported Scantable is moved to other computer alone.

WARNING for the GBT raw SDFITS format data as input: SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

- `sdstatold`

Keyword arguments:

infile -- name of input SD dataset
  default: none - must input file name
  example: 'mysd.asap'
  See `sdcalold` for allowed formats.

antenna -- antenna name or id (only effective for MS input).

fluxunit -- units for line flux
  options: (str) 'K','Jy',''
  default: '' (keep current fluxunit)
  WARNING: For GBT data, see description below.

>>> fluxunit expandable parameter

telescopeparm -- the telescope characteristics
  options: (str) name or (list) list of gain info
  default: '' (none set)
  example: if telescopeparm='', it tries to get the telescope name from the data.
  Full antenna parameters (diameter,ap.eff.) known to ASAP are
  'ATPKSMB', 'ATPKSHOH', 'ATMOPRA', 'DSS-43', 'CEDUNA','HOBART'. For GBT, it fixes default fluxunit to 'K' first then convert to a new fluxunit.
  telescopeparm=[104.9,0.43] diameter(m), ap.eff. telescopeparm=[0.743] gain in Jy/K
  telescopeparm='FIX' to change default fluxunit
  see description below

specunit -- units for spectral axis
  options: (str) 'channel','km/s','GHz','MHz','kHz','Hz',''
  default: '' (=current)

>>> specunit expandable parameter
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restfreq -- rest frequency
  default: '' (use current setting)
  example: 4.6e10 (float value in Hz),
            '46GHz' (string with unit),
            ['345.8GHz', 347.0e9, 356.7e9] (for each IF)
            ['name':'CO','value':345e9] (a value with name)

frame -- frequency frame for spectral axis
  options: (str) 'LSRK', 'REST', 'TOPO', 'LSRD', 'BARY',
           'GEO', 'GALACTO', 'LGROUP', 'CMB'
  default: currently set frame in scantable
  WARNING: frame='REST' not yet implemented

doppler -- doppler mode
  options: (str) 'RADIO', 'OPTICAL', 'Z', 'BETA', 'GAMMA'
  default: currently set doppler in scantable

scanlist -- list of scan numbers to process
  default: [] (use all scans)
  example: [21, 22, 23, 24]
  this selection is in addition to field, iflist, and pollist

field -- selection string for selecting scans by name
  default: '' (no name selection)
  example: 'FLS3a*'  
  this selection is in addition to scanlist, iflist, and pollist

iflist -- list of IF id numbers to select
  default: [] (use all IFs)
  example: [15]
  this selection is in addition to field, scanlist, and pollist

pollist -- list of polarization id numbers to select
  default: [] (use all pols)
  example: [1]
  this selection is in addition to field, scanlist, and iflist

masklist -- list of mask regions to INCLUDE in stats
  default: [] (whole spectrum)
  example: [4000, 4500] for one region
           [[1000, 3000], [5000, 7000]]
           these must be pairs of [lo, hi] boundaries

invertmask -- invert mask (EXCLUDE masklist instead)
  options: (bool) True, False
  default: false

interactive -- determines interactive masking
  options: (bool) True, False
  default: False
  example: interactive=True allows adding and deleting mask
           regions by drawing rectangles on the plot with mouse.
           Draw a rectangle with LEFT-mouse to ADD the region to
           the mask and with RIGHT-mouse to DELETE the region.
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outfile -- name of output file for line statistics
    default: '' (no output statistics file)
    example: 'stat.txt'

format -- format string to print statistic values
    default: '3.3f'

overwrite -- overwrite the statistics file if already exists
    options: (bool) True,False
    default: False

Returns: a Python dictionary of line statistics
    keys: 'rms','stddev','max','min','max_abscissa',
          'min_abscissa','sum','median','mean','totint','eqw'
    example: xstat=sdstatold(); print "rms = ",xstat['rms']
    these can be used for testing in scripts or
    for regression

    'max_abscissa' and 'min_abscissa' refer to the abscissa
    (channel/frequency/velocity) of max and min intensity.
    'totint' is the integrated intensity (sum*dx)
    where dx is the abscissa interval in 'specunit'.
    'eqw' is equivalent width (totint/mag) where mag
    is either max or min depending on which has
    greater magnitude.
    Note that 'max_abscissa', 'min_abscissa', 'totint'
    and 'eqw' are quantities (python dictionaries with
    keys, 'unit' and 'value').

DESCRIPTION:

Task sdstatold computes basic statistics (rms,mean,median,sum)
for single-dish spectra. It assumes that the spectra have
been calibrated. Furthermore, it assumes that any
time and channel averaging/smoothing has also already been done as
there are no controls for these.

Note that multiple scans and IFs can in principle be handled, but
we recommend that you use scanlist, field, iflist, and pollist to give
a single selection for each run.

WARNING: If you do have multiple scantable rows, then the returning
values will be lists.

ASAP recognizes the data of the "AT" telescopes, but currently
does not know about the GBT or any other telescope. This task does know about GBT. Telescope name is obtained from the data. If you wish to change the fluxunit (see below), and telescopeparm='', for the AT telescopes it will use internal telescope parameters for flux conversion. For GBT, it will use an approximate aperture efficiency conversion. If you give telescopeparm a list, then if the list has a single float it is assumed to be the gain in Jy/K, if two or more elements they are assumed to be telescope diameter (m) and aperture efficiency respectively.

WARNING for the GBT raw SDFITS format data as input: SDtasks are able to handle GBT raw SDFITS format data since the data filler is available. However, the functionality is not well tested yet, so that there may be unknown bugs.

8.8.2 Single Dish Analysis Use Cases With SDTasks

8.8.2.1 GBT Position Switched Data Analysis

As an example, the following illustrates the use of the SDtasks for the Orion data set, which contains the HCCCN line in one of its IFs. This walk-through contains comments about setting parameter values and some options during processing.

```
# ORION-S SDtasks Use Case
# Position-Switched data
# Version TT 2008-10-14 (updated)
# Version STM 2007-03-04
#
# This is a detailed walk-through for using the SDtasks on a test dataset.
#
import time
import os

# This is the environment variable pointing to the head of the CASA tree that you are running
casapath=os.environ['AIPSPATH']
```
# This bit removes old versions of the output files
os.system('rm -rf sdusecase_orions* ')
#
# This is the path to the OrionS GBT ms in the data repository
 datapath= casapath+ '/data/regression/ATST5/OrionS/OrionS_rawACSmod'
#
# The following will remove old versions of the data and
# copy the data from the repository to your
# current directory. Comment this out if you already have it
# and don’t want to recopy
os.system('rm -rf OrionS_rawACSmod')
copystring='cp -r '+datapath+' .'
os.system(copystring)

# Now is the time to set some of the more useful
# ASAP environment parameters (the ones that the
# ASAP User Manual claims are in the .asaprc file).
# These are in the Python dictionary sd.rcParams
# You can see what’s in it by typing:
#sd.rcParams
# One of them is the ‘verbose’ parameter which tells
# ASAP whether to spew lots of verbiage during processing
# or to keep quiet. The default is
#sd.rcParams['verbose']=True
# You can make ASAP run quietly (with only task output) with
#sd.rcParams['verbose']=False

# Another key one is to tell ASAP to save memory by
# going off the disk instead. The default is
#sd.rcParams['scantable.storage']='memory'
# but if you are on a machine with small memory, do
#sd.rcParams['scantable.storage']='disk'

# You can reset back to defaults with
#sd.rcdefaults

# ORION-S HC3N
# Position-Switched data
#
startTime=time.time()
startProc=time.clock()
### List data

# List data

### List the contents of the dataset

# First reset parameter defaults (safe)

```python
default('sdlist')
```

# You can see its inputs with
# inp('sdlist')
# or just
# inp
# now that the defaults('sdlist') set the
# taskname='sdlist'
#
# Set the name of the GBT ms file

```python
infile = 'OrionS_rawACSmod'
```

# Set an output file in case we want to
# refer back to it

```python
outfile = 'sdusecase_orions_summary.txt'
```

sdlist()

# You could also just type
# go

# You should see something like:

```
# Scan Table Summary
# Project: AGBT06A_018_01
# Obs Date: 2006/01/19/01:45:58
# Observer: Joseph McMullin
# Antenna Name: GBT@GREENBANK
# Data Records: 512 rows
# Obs. Type: OffOn:PSWITCHOFF:TPWCAL
# Beams: 1
# IFs: 8
# Polarisations: 2 (circular)
# Channels: 8192
# Flux Unit: K
# Abscissa: Channel
# Selection: none
#
# Scan Source Time range Int[s] Record SrcType FreqIDs MolIDs
# Beam Position (J2000)
# --------------------------------------------------------------------------------
# 21 OrionS 2006/01/19/01:45:58.0 - 01:47:58.2 30.03 64 [PSOFF, PSOFF:CALON] [0, 1, 2, 3] [0]
# 0 J2000 05:15:13.5 -05.24.08.6
# 22 OrionS 2006/01/19/01:48:38.0 - 01:50:38.2 30.03 64 [PSON, PSON:CALON] [0, 1, 2, 3] [0]
```
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# 0 J2000 05:35:13.4 -05.24.07.8
# 23 OrionS 2006/01/19/01:51:21.0 - 01:53:21.2 30.03 64 [PSOFF, PSOFF:CALON] [0, 1, 2, 3] [0]
# 0 J2000 05:15:13.6 -05.24.08.5
# 24 OrionS 2006/01/19/01:56:01.2 - 01:58:01.2 30.03 64 [PSON, PSON:CALON] [0, 1, 2, 3] [0]
# 0 J2000 05:35:13.4 -05.24.08.1
# 25 OrionS 2006/01/19/02:03:47.0 - 02:05:47.2 30.03 64 [PSOFF, PSOFF:CALON] [4, 5, 6, 7] [1]
# 0 J2000 05:15:13.5 -05.24.08.1
# 26 OrionS 2006/01/19/02:06:27.2 - 02:08:27.2 30.03 64 [PSON, PSON:CALON] [4, 5, 6, 7] [1]
# 0 J2000 05:35:13.3 -05.24.08.1
# 27 OrionS 2006/01/19/02:09:10.2 - 02:11:10.2 30.03 64 [PSOFF, PSOFF:CALON] [4, 5, 6, 7] [1]
# 0 J2000 05:15:13.5 -05.24.08.4
# 28 OrionS 2006/01/19/02:11:51.2 - 02:13:51.2 30.03 64 [PSON, PSON:CALON] [4, 5, 6, 7] [1]
# 0 J2000 05:35:13.3 -05.24.08.1

# FREQUENCIES: 4
# ID IFNO(SPW) #Chans Frame Ch0[MHz] ChanWid[kHz] Center[MHz] POLNOs
# 0 0 8192 LSRK 45464.3506 6.10423298 45489.3505 [0, 1]
# 1 1 8192 LSRK 45275.7825 6.10423298 45300.7824 [0, 1]
# 2 2 8192 LSRK 44049.9264 6.10423298 44074.9263 [0, 1]
# 3 3 8192 LSRK 44141.2121 6.10423298 44166.2120 [0, 1]
# 4 12 8192 LSRK 43937.1232 6.10423298 43962.1231 [0, 1]
# 5 13 8192 LSRK 42620.4173 6.10423298 42645.4172 [0, 1]
# 6 14 8192 LSRK 41569.9768 6.10423298 41594.9767 [0, 1]
# 7 15 8192 LSRK 43397.8198 6.10423298 43422.8196 [0, 1]
# 8 16 8192 LSRK 42322.3504 6.10423298 42347.3503 [0, 1]

# MOLECULES:
# ID RestFreq Name
# 0 [4.5490258e+10] []
# 1 [4.3963e+10] []

# The HC3N and CH3OH lines are in IFs 0 and 2 respectively
# of scans 20,21,22,23. We will pull these out in our
# calibration.

####################################
# Calibrate data
####################################

# We will use the sdreduce task to calibrate the data.
# Set the defaults
default('sdreduce')

# You can see the inputs with
# inp

# Set our infile (which would have been set from our run of
# sdlist if we were not cautious and reset defaults).
infile = 'OrionS_rawACSmod'
fluxunit = 'K'

# This is position-switched data so we tell sdreduce this
calmode = 'ps'

# For GBT data, it is safest to not have scantable pre-average
# integrations within scans.
average = True
scanaverage = False

# We do want sdreduce to average up scans and polarization after
# calibration however. The averaging of scans are weighted by
# integration time and Tsys, and the averaging of polarization
# by Tsys.
timeaverage = True
tweight = 'tintsys'
polaverage = True
pweight = 'tsys'
# Do an atmospheric optical depth (attenuation) correction
# Input the zenith optical depth at 43 GHz
tau = 0.09

# Select our scans and IFs (for HC3N)
scan = '21~24'
spw = '0'

# We do not require selection by field name (they are all
# the same except for on and off)
field = ''

# We will do some spectral smoothing
# For this demo we will use boxcar smoothing rather than
# the default
# kernel='hanning'
# We will set the width of the kernel to 5 channels
kernel = 'boxcar'
kwidth = 5

# We wish to fit out a baseline from the spectrum
# The GBT has particularly nasty baselines :(
# We will let ASAP use auto_poly_baseline mode
# but tell it to drop the 1000 edge channels from
# the beginning and end of the spectrum.
# A 2nd-order polynomial will suffice for this test.
# You might try higher orders for fun.
mask = 'auto'
blfunc = 'poly'
order = 2
edge = [100]

# We will not give it regions as an input mask
# though you could, with something like
maskmode = 'list'
spw = '0:1000~3000;5000~7000'

# By default, we will not get plots in sdreduce (but
# can make them using sdplot).
plotlevel = 0
# But if you wish to see a final spectrum, set
# plotlevel = 1
# or even
# plotlevel = 2
# to see intermediate plots and baselining output.

# Now we give the name for the output file
outfile = 'sdusecase_orions_hc3n.asap'

# We will write it out in ASAP scantable format
outform = 'asap'

# You can look at the inputs with
# inp

# Before running, lets save the inputs in case we want
# to come back and re-run the calibration.
saveinputs('sdreduce','sdreduce.orions.save')
# These can be recovered by
# execfile 'sdreduce.orions.save'

# We are ready to calibrate
sdreduce()

# Note that after the task ran, it produced a file
# sdreduce.last which contains the inputs from the last
# run of the task (all tasks do this). You can recover
# this (any time before sdreduce is run again) with
# execfile 'sdreduce.last'

# List data
# List the contents of the calibrated dataset
# Set the input to the just created file
infile = outfile
outfile = ''
sdlist()
# You should see:

# When you copy this file (and adjust for line lengths) you can
# # read it like this:  
# # (The #'s indicate comments)
#
# # Scan Table Summary
# # --------------------------------------------------------------------------------
# # Project: AGBT06A_018_01
# # Obs Date: 2006/01/19/01:45:58
# # Observer: Joseph McMullin
# # Antenna Name: GBT@GREENBANK
# # Data Records: 1 rows
# # Obs. Type: OffOn:PSWITCHOFF:TPWCAL
# # Beams: 1
# # IFs: 8
# # Polarisations: 1 (stokes)
# # Channels: 8192
# # Flux Unit: K
# # Abscissa: Channel
# # Selection: none
#
# # Scan Source Time range Int[s] Record SrcType FreqIDs MolIDs 
# # Beam Position (J2000)
# # --------------------------------------------------------------------------------
# # 0 OrionS 2006/01/19/01:52:04.6 - 02:00:05.1 480.48 1 [PSON] [0] [0]
# # 0 J2000 05:35:13.4 -05.24.07.8
# # --------------------------------------------------------------------------------
# # FREQUENCIES: 1
# # ID IFNO(SPW) #Chans Frame Ch0[MHz] ChanWid[kHz] Center[MHz] POLNOs
# # 0 0 8192 LSRK 45464.3506 6.10423298 45489.3505 [0]
# # --------------------------------------------------------------------------------
# # MOLECULES:
# # ID RestFreq Name
# # 0 [4.5490258e+10] []
# # 1 [4.3963e+10] []
# # --------------------------------------------------------------------------------

# Note that our scans are now collapsed (timeaverage=True) but
# we still have our IF 0

#+++++++++++++++++++++++++++++++++++++++++++++++++++++++
# Plot data
#+++++++++++++++++++++++++++++++++++++++++++++++++++++++
default('sdplot')

# The file we produced after calibration
# (if we hadn’t reset defaults it would have
# been set - note that sdplot,sdfit,sdstat use
# infile as the input file, which is the output
# file of sreduce).
infile = 'sdusecase_orions_hc3n.asap'
# Let's just go ahead and plot it up as-is
sdplot()

# Looks ok. Plot with x-axis in GHz
specunit = 'GHz'
sdplot()

# Note that the rest frequency in the scantable
# is set correctly to the HCCCN line at 45.490 GHz.
# So you can plot the spectrum in km/s
specunit = 'km/s'
sdplot()

# Zoom in
sprange = [-100,50]
sdplot()

# Let's plot up the lines to be sure
# We have to go back to GHz for this
# (known deficiency in ASAP)
specunit = 'GHz'
sprange = [45.48,45.51]
linecat = 'all'
sdplot()

# Too many lines! Focus on the HC3N ones
linecat = 'HCCCN'
sdplot()

# Finally, we can convert from K to Jy
# using the aperture efficiencies we have
# coded into the sdtasks
# For GBT data, do not set telescopeparm
fluxunit = 'Jy'
telescopeparm = ''
sdplot()

# Let's save this plot
outfile = 'sdusecase_orions_hc3n.eps'
sdplot()

# Off-line Statistics
# Now do some region statistics
# First the line-free region
# Set parameters
default('sdstat')
infile = 'sdusecase_orions_hc3n.asap'

# Keep the default spectrum and flux units
# K and channel
fluxunit = ''
specunit = ''

# Pick out a line-free region
# You can bring up a default sdplot again
# to check this
spw = '0:5000~7000'

# You can check with
# inp

# sdstat returns some results in
# the Python dictionary. You can assign
# this to a variable
off_stat=sdstat()

# and look at it
off_stat
# which should give
# {'eqw': 38.563105620704945,
# 'max': 0.15543246269226074,
# 'mean': -0.0030361821409314871,
# 'median': -0.0032975673675537109,
# 'min': -0.15754437446594238,
# 'rms': 0.047580458223819733,
# 'stddev': 0.047495327889919281,
# 'sum': -6.0754003524780273}

#You see it has some keywords for the various
#stats. We want the standard deviation about
#the mean, or 'stddev'
print "The off-line std. deviation = ",off_stat['stddev']
# which should give
# The off-line std. deviation = 0.0474953278899

# or better formatted (using Python I/O formatting)
print "The off-line std. deviation = %5.3f K" %}
(off_stat['stddev'])
# which should give
# The off-line std. deviation = 0.047 K

#-------------------------------
# On-line Statistics
#-------------------------------
# Now do the line region
# Continue setting or resetting parameters
spw = '0:3900~4200'

line_stat = sdstat()

# look at these
line_stat
# which gives
# {'eqw': 73.335154614280981,
# 'max': 0.92909121513366699,
# 'mean': 0.22636228798819946,
# 'median': 0.10317134857177734,
# 'min': -0.13283586502075195,
# 'rms': 0.35585442185401917,
# 'stddev': 0.27503398060798645,
# 'sum': 68.135047912597656}

# of particular interest are the max value
print "The on-line maximum = %5.3f K" % (line_stat['max'])
# which gives
# The on-line maximum = 0.929 K

# and the estimated equivalent width (in channels)
# which is the sum/max
print "The estimated equivalent width = %5.1f channels" %\n     (line_stat['eqw'])
# which gives
# The estimated equivalent width = 73.3 channels

#-------------------------------
# Line Fitting
#-------------------------------
# Now we are ready to do some line fitting
# Default the parameters
default('sdfit')

# Set our input file
infile = 'sdusecase_orions_hc3n.asap'

# Stick to defaults
# fluxunit = 'K', specunit = 'channel'
fluxunit = ''

# We will try auto-fitting first
fitmode = 'auto'
# A single Gaussian
nfit = [1]
# Leave the auto-parameters to their defaults for
# now, except ignore the edge channels
edge = [1000]

# Let’s see a plot while doing this
plotlevel = 1

# Save the fit output in a file
outfile = 'sdusecase_orions_hc3n.fit'

# Go ahead and do the fit
fit_stat = sdfit()

# If you had verbose mode on, you probably saw something
# like:
#
# 0: peak = 0.811 K , centre = 4091.041 channel, FWHM = 72.900 channel
#  area = 62.918 K channel
#

# The fit is output in the dictionary

fit_stat
#
# '{cent': [[4091.04052734375, 0.72398632764816284]],
# 'fwhm': [[72.899894714355469, 1.7048574686050415]],
# 'nfit': 1,
# 'peak': [[0.81080442667007446, 0.016420882195234299]]}
#
# So you can write them out or test them:
print "The line-fit parameters were:"
print "  maximum = %6.3f +/- %6.3f K" %
  (fit_stat['peak'][0][0],fit_stat['peak'][0][1])
print "  center = %6.1f +/- %6.1f channels" %
  (fit_stat['cent'][0][0],fit_stat['cent'][0][1])
print "FWHM = %6.2f +/- %6.2f channels" %
(fit_stat['fwhm'][0][0],fit_stat['fwhm'][0][1])
#
# Which gives:
# The line-fit parameters were:
# maximum = 0.811 +/- 0.016 K
# center = 4091.0 +/- 0.7 channels
# FWHM = 72.90 +/- 1.70 channels
#
# We can do the fit in km/s also
specunit = 'km/s'
%
# For some reason we need to help it along with a mask
%
maskline = [-50,0]

outfile = 'sdusecase_orions_hc3n_kms.fit'
fit_stat_kms = sdfit()
# Should give (if in verbose mode)
# 0: peak = 0.811 K , centre = -27.134 km/s, FWHM = 2.933 km/s
# area = 2.531 K km/s
#
#
# with
fit_stat_kms
# giving
# {'cent': [[-27.133651733398438, 0.016480101272463799]],
# 'fwhm': [[2.93294358253479, 0.038807671517133713]],
# 'nfit': 1,
# 'peak': [[0.81080895662307739, 0.0092909494414925575]]}

print "The line-fit parameters were:"
print " maximum = %6.3f +/- %6.3f K" %
(fit_stat_kms['peak'][0][0],fit_stat_kms['peak'][0][1])
print " center = %6.2f +/- %6.2f km/s" %
(fit_stat_kms['cent'][0][0],fit_stat_kms['cent'][0][1])
print " FWHM = %6.4f +/- %6.4f km/s" %
(fit_stat_kms['fwhm'][0][0],fit_stat_kms['fwhm'][0][1])
#
# The line-fit parameters were:
# maximum = 0.811 +/- 0.009 K
# center = -27.13 +/- 0.02 km/s
# FWHM = 2.9329 +/- 0.0388 km/s

########################################################################
8.8.2.2 Imaging of Total Power Raster Scans

This example illustrates the use of \texttt{sdtpimaging} for the total power raster scans of the Moon taken at ATF.

Figure 8.10: Total power data display using \texttt{sdtpimaging}, with \texttt{calmode='baseline'}. The top panel shows uncalibrated data versus row numbers. The middle panel shows baseline fitting of each scan (only shown here the last scan). The bottom panel shows the calibrated (baseline subtracted) data.
# The data used here (uid__X1e1_X3197_X1.ms) is the total power
# raster scans of the Moon taken at ATF (with both antennas).
# It is in MS format which was converted from the ASDM format.

# Do data plotting only
default(sdtpimaging)
inp()
plotlevel = 2
# select antenna 1 (Vertex antenna)
antenna = '1'
infile = 'uid__X1e1_X3197_X1.ms'
sdtpimaging()

# Now, rerun sdtpimaging to do actual data reduction (applying
# baseline subtraction from each scan, and then do imaging).
#
# Do baseline subtraction
calmode='baseline'
masklist = [30] # use 30 data points from each end of scan for fitting

# Do imaging
createimage = True
outfile = 'moon.im'
imagesize = [200,200]
cell = [0.2] # in arcmin
phasesource = 'AZEL 187d54m22s 41d03m0s'
ephemsrcname = 'moon' # specify ephemeris source name (can be omitted)
plotlevel = 1
#plotlevel = 2 to see progress of each fitting
sdtpimaging()
Chapter 9

Simulation

New in 4.2:

• Single dish imaging in CASA is in a state of vigorous development. At the time of the 4.2 release, single dish imaging is accomplished in simalma and simobserve by calling the sdimaging task with spheroidal gridding, and particular choices of cell size and convolution support. These parameters may change as ALMA best practice is refined. CAVEAT: There is currently a 15% uncertainty in the beam area and resulting absolute flux scale for simulated single dish images.

• The simalma task has been significantly improved in 4.2 - in particular, the dryrun parameter allows users to test their input parameters and skymodel, and see a report of what simalma will do without waiting. The report includes information on the multiple requested components (12m interferometric array, 7m interferometric array, and total power) and whether any input parameters are likely to cause issues in simulation. The task’s parameters have been changed to more easily support multiple configurations and components in a single run. CAVEAT: The tools do not split up long observations at this time, so a 12h or longer observation will include time at low elevation, which may be unrealistic. The user may need to manually split up their simulation, and use multiple calls to simobserve with shorter totaltime.

• simobserve adds noise to an observation by default - in previous CASA versions the default was noiseless.

• simanalyze now defaults to producing a dirty image niter=0.

The tasks available for simulating observations are:

• simobserve — simulate an interferometer or total power observation (§ 9.1)
• simanalyze — image and analyze simulated data sets (§ 9.1)
• simalma — simulate an ALMA observation including multiple configurations of the 12-m interferometric array, the 7-m ACA, and total power measurements. Generate a combined image from the simulated data sets (§ 9.2)
The capability of simulating observations and data sets from the JVLA and ALMA are an important use-case for CASA. This not only allows one to get an idea of the capabilities of these instruments for doing science, but also provides benchmarks for the performance and utility of the software to process “realistic” data sets (with atmospheric and instrumental effects). Simulations can also be used to tune parameters of the data reduction and therefore help to optimize the process. CASA can calculate visibilities (create a measurement set) for any interferometric array, and calculate and apply calibration tables representing some of the most important corrupting effects. simobserve can also simulate total power observations, which can be combined with interferometric data in simanalyze (i.e. one would run simobserve twice, simanalyze once). The task simalma is a task to simulate an ALMA observation, including ALMA 12-m, ACA 7-m and total power arrays, and generate a combined image. simalma also attempts to provide useful feedback on those different observation components, to help the user better understand the observing considerations.

CASA’s simulation capabilities continue to be improved with each CASA release. For the most current information, please refer to [http://www.casaguides.nrao.edu](http://www.casaguides.nrao.edu), and click on “Simulating Observations in CASA”. Following general CASA practice, the greatest flexibility and richest functionality is at the Toolkit level. The most commonly used procedures for interferometric and single dish simulation are encapsulated in the simobserve task.

### 9.1 Simulating ALMA observations with simobserve and simanalyze

The simobserve inputs are (submenus expand slightly differently for thermalnoise=manual and single dish observing):

```plaintext
project = 'sim'  # root prefix for output file names
skymodel = ''  # model image to observe
inbright = ''  # scale surface brightness of brightest pixel e.g. "1.2Jy/pixel"
indirection = ''  # set new direction e.g. "J2000 19h00m00 -40d00m00"
inwidth = ''  # set new channel width e.g. "10MHz" (required even for 2D model)
incenter = ''  # set new frequency of center channel e.g. "89GHz"
complist = ''  # componentlist to observe
compwidth = '8GHz'  # bandwidth of components
setpointings = True  # integration (sampling) time
integration = '10s'  # integration (sampling) time
```

Inside the Toolkit:

The simulator methods are in the sm tool. Many of the other tools are also helpful when constructing and analyzing simulations.
This task takes an input model image or list of components, plus a list of antennas (locations and sizes), and simulates a particular observation (specified by mosaic setup and observing cycles and times). The output is a measurement set suitable for further analysis in CASA.

The simanalyze inputs are:

- `project` = 'sim'  
  # root prefix for output file names
- `image` = True  
  # (re)image $project.*.ms to $project.image
- `vis` = 'default'  
  # Measurement Set(s) to image
- `modelimage` = ''  
  # lower resolution prior
  # image to use in clean e.g. existing total
  # power image
This task analyzes one or more measurement sets - interferometric and/or single dish. The output is a synthesized image created from those visibilities, a difference image between the synthesized image and your sky model convolved with the output synthesized beam, and a fidelity image. (see ALMA memo 398 for description of fidelity, which is approximately the output image divided by the difference between input and output)
9.2 Simulating ALMA observations with simalma

The task **simalma** simulates an ALMA observation by ALMA 12-m, ACA-7m and total power arrays. It takes an input model image or a list of components, plus configurations of ALMA antennas (locations and sizes), and simulates a particular ALMA observation (specified by mosaic setup and observing cycles and times). The outputs are measurement sets. The task optionally generates synthesized images from the measurement sets as **simanalyze** does.

Technically speaking, **simalma** internally calls **simobserve** and **simanalyze** as many times as necessary to simulate and analyze an ALMA observation. Some of the simulation (**simobserve**) and imaging (**simanalyze**) parameters are automatically set to values typical of ALMA observations in **simalma** (see §9.2.1 for more details). Thus, it has a simpler task interface compared to **simobserve** plus **simanalyze** at the cost of limited flexibility. If you want to have more control on simulation setup, it is available by manually running **simobserve** and **simanalyze** multiple times or by using **sm** tools.

The **simalma** inputs are:

```
project = '' # root prefix for output file names
project = 'sim' # root prefix for output file names
dryrun = False # dryrun=True will only
  # produce the informative report, not run
  # simobserve/analyze
skymodel = '' # model image to observe
inbright = '' # scale surface brightness of
  # brightest pixel e.g. "1.2Jy/pixel"
indirection = '' # set new direction
  # e.g. "J2000 19h00m00 -40d00m00"
inell = '' # set new cell/pixel size e.g. "0.1arcsec"
incenter = '' # set new frequency of center
  # channel e.g. "89GHz" (required even
  # for 2D model)
inwidth = '' # set new channel width
  # e.g. "1GHz" (required even for 2D model)
complist = '' # componentlist to observe
compwidth = '8GHz' # bandwidth of components
setpointings = True
integration = '10s' # integration (sampling) time
direction = '' # "J2000 19h00m00 -40d00m00"
  # or "" to center on model
mapsize = ['', ''] # angular size of map or "" to cover model
graphics = 'both' # display graphics at each
  # stage to [screen|file|both|none]
verbose = False
overwrite = False # overwrite files starting with $project
```
The task `simalma` is designed as a task that is invoked only once for a simulation setup. It always sets up skymodel and pointings. That means that `simalma` is not supposed to be run multiple times for a project, unlike `simobserve` and `simanalyze`. The task `simalma` may ignore or overwrite the old results when it is run more than once with the same project name.

There are options in `simalma` to simulate observation of ACA 7-m and total power arrays, to apply thermal noise, and/or to generate images from simulated measurement sets. One inputs a vector of configurations, and a corresponding vector of `totaltime` to observe each component. Thermal noise is added to visibilities when `pwv > 0`. The ATM atmospheric model is constructed from the characteristics of the ALMA site and a user defined Precipitable Water Vapour (`pwv`) value. Set `pwv = 0` to omit the thermal noise. Finally, when `image = True`, synthesized images are generated from the simulated measurement sets.

### 9.2.1 Implementation details

As mentioned in the previous section, `simalma` automatically sets some of the simulation and imaging parameters to values typical of ALMA observations. The implementations of antenna configurations, pointings, integration time, and imaging in CASA 4.2.0 are described in this section.

**Antenna Configuration:**

The configurations of the ALMA 12-m and 7-m arrays are defined by the `antennalist` parameter, which can be a vector. Each element of the vector can be either the name of an antenna configuration file or a desired resolution, e.g., `alma;cycle1;5arcsec`. Some examples:

- `antennalist = ['alma.cycle2.5.cfg','aca.cycle2.i.cfg']; totaltime = ['20min','2h']`: Will observe the 12-m array in configuration C32-5 for 20 minutes and the ACA 7-m array for 2 hours.

- `antennalist = ['alma;cycle2;0.5arcsec','aca.i.cfg']; totaltime = ['20min','2h']`: Will observe the 12-m array in whatever cycle 2 configuration yields a zenith synthesized beam as close as possible to 0.5 arcsec (at the center frequency of your skymodel) for 20 minutes and the ACA 7-m array for 2 hours.

- `antennalist = ['alma.cycle1.2.cfg','aca.cycle2.i.cfg']; totaltime = '20min'`: Will observe the 12-m array in cycle 1 configuration for 20 minutes and the ACA 7-m array for the default of 2×(12-m time) = 1h20min. This parameter setting will also generate a warning that the user is combining configurations from different ALMA Cycles (but the simulation will run despite that).

Total power can either be included along with interferometric configurations e.g. `antennalist = ['alma.cycle1.2.cfg','aca.cycle2.i.cfg','alma.tp.cfg']`, or by using the `tpnant` and `tptime` parameters. The latter is preferred since it allows greater control (in particular the number of total power antennas to use – if more than one is used, multiple total power observations will be generated and combined in imaging).

**Field Setup:**
There are two ways to setup pointings, i.e., Rectangle Setup and Multi-Pointing.

In the Rectangle Setups, pointings are automatically calculated from the pointing centre (direction) and the map size. A rectangular map region is covered by a hexagonal grid (maptype = 'alma') with Nyquist sampling, i.e., 0.48PB spacing (where PB ≡ 1.2λ/D), in both ALMA 12-m and ACA 7-m array simulations. A slightly larger area is mapped in ACA total power simulations for later combination with interferometer visibilities. The map area is extended by 1 PB in each direction and covered by a lattice grid with 0.225PB spacing.

In Multi-Pointing, a list of pointings is defined in the direction parameter or read from a file (when setpointings = False). The ALMA 12-m and ACA 7-m arrays observe the specified directions. The ACA total power simulations map either (1) square regions of 2 PB extent centred at each of the pointings, or (2) a rectangle region that covers all the pointings. Either (1) or (2), whichever can be done with the smaller number of points, is selected. The pointing spacing in total power simulations is, again, 0.225PB in lattice grids.

It is advisable that for Total Power Simulations, the field is chosen sufficiently large, maybe padding at least 1-2 primary beams on each side.

**Integration time:**

The total observation time of each component or configuration is defined by the totaltime parameter as noted above. A scalar will trigger use of the Cycle 2 default time multipliers, 1:0.5:2:4 for the first 12-m configuration, any additional 12-m configurations, any 7-m configuration, and any total power observation.

In general, the integration time (dump interval) of simulations is defined by the integration parameter with an exception. Since the ACA total power array always observes larger areas compared to the ALMA 12-m and ACA 7-m arrays, it is possible that the ACA total power array cannot cover all pointings in the given observation time. In such a case, the integration time in the total power simulation is scaled so that the all pointings are observed at least once in its observation time, i.e., integration_TP = tptime / (the number of total power pointings).

**Imaging and combination of ALMA with ACA:**

The CLEAN algorithm is used in simalma to generate images from visibilities. The visibilities are weighted to UV-plane using Briggs weighting.

When ACA observations are simulated, visibilities of ACA 7-m are weighted by the relative sensitivities to ALMA 12-m visibilities, and both data sets are concatenated before imaging. The relative weight of ACA 7-m visibilities is defined in proportion to the difference of beam area, i.e., \((7/12)^2 = 0.34\). This is because simalma uses a bandwidth and an integration time common to both ALMA 12-m and ACA 7-m simulations.

The interferometer and total power images are combined using feather task when total power observations are included. The total power image is scaled by the interferometer primary beam coverage before combination. The final image product is the combined image corrected for the interferometer primary beam coverage. The output image of the feather task is divided by the interferometer primary beam coverage in the final step.
Chapter 10

Parallel Processing in CASA

Since CASA 4.0.0, a parallelized execution of a full data analysis from data import to imaging is possible. This functionality continues to be under development and should still be regarded as experimental but users are encouraged to try this feature if they have access to computers with a solid state disk (SSD) or RAID arrays with a Lustre file system or other fast file systems. On normal SATA disks, there is no benefit from parallelization as the file system cannot feed more than one CASA instance.

The following deal with the handling of visibility data. For imaging with pclean, see §5.10.

10.1 The CASA parallelization scheme

In order to run one analysis on multiple processors, one can parallelize the work by dividing the data into several parts (“partitioning”) and then run a CASA instance on each part or have non-trivially parallelized algorithms which make use of several processors within a single CASA instance. Non-trivial parallelization is presently only implemented in certain sections of the imaging code of CASA based on OpenMP.

All other parallelization is achieved by partitioning the MeasurementSet (MS) of interest using the task partition. The resulting partitioned MS is called a “multi-MS” or “MMS”. Logically, an MMS has the same structure as an MS but internally it is a group of several MSs which are virtually concatenated. Virtual concatenation of several MSs or MMSs into an MMS can also be achieved via the new (in CASA 4.0) task virtualconcat.

Due to the virtual concatenation, the main table of an MMS appears like the union of the main tables of all the member MSs such that when the MMS is accessed like a normal MS, processing can proceed sequentially as usual. Each member MS or “subMS” of an MMS, however, is at the same time a valid MS on its own and can be processed as such. This is what happens when the MMS is accessed by a parallelized task. The partitioning of the MMS is recognized and work is started in parallel on the separate subMSs.

The internal structure of an MMS can be inspected using the new task listpartition.
10.2 Multi-MS-compatible tasks in CASA 4.1

The following tasks in CASA 4.1 have been tested to work with MMSs as input:

- applycal
- bandpass
- clean
- concat (produces an output MS, not an MMS)
- fixplanets
- flagdata
- flagmanager
- fluxscale
- gaincal
- gencal
- listobs
- listpartition
- listvis
- listhistory
- partition (repartitioning of an MMS is also possible)
- pclean
- plotms
- setjy
- split (produces MS by default, an MMS if parameter keepmms=True)
- uvcontsub (produces an MMS)
- virtualconcat (produces an MMS)
- vishead
- wvrgcal

Of these, the following tasks will work in a parallelized way on MMSs to speed up processing:
• applycal
• flagdata
• partition (repartitioning of an MMS is also possible)
• pclean
• setjy (when parameter usescratch=True)
• split (when parameter keepmms=True)
• uvcontsub

You can find an example of a parallelized analysis in the regression script

    alma-m100-analysis-hpc-regression.py

in a subdirectory of your CASA distribution.

10.3 Parallelization control

10.3.1 Requirements

The following requirements are necessary for all the nodes to be included in the cluster:

• Password-less ssh access from the controller (user) machine into all the hosts to be included in the cluster
  
  NOTE: This is not necessary when using only localhost, i.e. if the cluster is deployed only on the machine where casapy is running.

• All the input files must be located in a shared file-system, accessible from all the nodes comprising the cluster, and mounted in the same path of the file-system

• Mirrored CASA installation w.r.t. the CASA installation in the controller (user) machine, so that the following environmental variables are pointing to valid installations: PATH, LD_LIBRARY_PATH, IPYTHONDIR, CASAPATH, CASAARCH, PYTHONHOME, __CASAPY_PYTHONDIR, PGPLOT_DEV, PGPLOT_DIR, PGPLOT_FONT.
10.3.2 Configuration and Start-Up

The “cluster”, i.e. the collection of CASA instances which will run the jobs from parallelized tasks, is set up automatically when it is used the first time, typically when running `partition`. The setup of this default cluster is derived from the properties of the host on which CASA is running. Presently the settings are such that up to 90% of the processors and the available RAM is used. There has to be at least 512 MB per CASA engine. But even if there are not enough resources available, at least one engine is deployed. In that case, the cluster is bypassed and jobs are simply run sequentially.

If the user wants to override these settings, this is possible by creating a “cluster configuration file” with one line per host to be used in the following format:

```
<hostname>, <number of engines>, <work directory>, <RAM usage>, <RAM per engine>
```

The comma-separated parameters have the following meaning:

1. **hostname**: Hostname of the target node where the cluster is deployed
   
   **NOTE**: The hostname has to be provided w/o quotes

2. **number of engines**: Supports in turns 3 different formats
   
   - If provided as an integer >1: It is interpreted as the actual user-specified maximum number of engines
   - If provided as an integer =0: It will deploy as maximum engines as possible according to the idle CPU capacity available at the target node
   - If provided as a float between 0 and 1: It is interpreted as the percentage of idle CPU capacity that the cluster can use in total at the target node

3. **work directory**: Area in which the cluster will put intermediate files such as log files, configuration files, and monitoring files
   
   **NOTE1**: This area has to be accessible from the controller (user) machine, and mounted in the same path of the filesystem
   
   **NOTE2**: The path name has to be provided w/o quotes

4. **RAM usage**: (optional) can be given in three different formats
   
   - If provided as an integer >0 or not at all (default): will deploy as many engines as possible using up to 90% of the free RAM available at target node
   - If provided as an integer >1: interpreted as the actual user-specified maximum amount of RAM (MB) to be used in total at the target node
   - If provided as a float between 0 and 1: interpreted as the percentage of free RAM that the cluster can use in total at the target node
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5. **RAM per engine**: (optional) integer, the required memory per engine in MB (default is 512MB)

It is also possible to add comments, by using the `#` character at the beginning of the line.

Example:

```
# CASA cluster configuration file
orion, 10, /home/jdoe/test/myclusterhome1
m42, 4, /home/jdoe/test/myclusterhome2, 0.6, 1024
antares, 0.6, /home/jdoe/test/myclusterhome3, 0, 2048
```

Will set up a cluster comprised of three nodes, deploying the engines per node as follows:

- At host “orion” up to 10 engines will be deployed with working directory `/home/jdoe/test/myclusterhome1` and using as much free RAM available as possible (up to 90% by default), taking into account that each engine can use up to 512 MB (the default and minimum)

- At host “m42”: It will deploy up to 4 engines, with working directory `/home/jdoe/test/myclusterhome2`, and using at the most 60% of the free RAM available, taking into account that each engine can use up to 1024 MB.

- At host “antares”: It will deploy as many engines as possible, with working directory `/home/jdoe/test/myclusterhome3`, using up to 60% of the idle CPU capacity / cores, and as much free RAM available as possible (up to 90% by default), taking into account that each engine can use up to 2048 MB.

Using such a configuration file named, say, “cluster-config.txt”, the cluster can then be created after CASA startup using the commands

```
from simple_cluster import *
s = simple_cluster()
s.init_cluster('cluster-config.txt', 'mycluster')
```

It will be used by all subsequent calls to parallelized tasks.

To stop an existing cluster without exiting casa type

```
s.stop_cluster()
```

Otherwise, any cluster will be terminated anyway when CASA exits.
10.3.3 Monitoring

The CASA cluster framework comes with a monitoring service that produces a monitoring.log file, in the same directory where the controller instance (user CASA terminal) is started.

This file is updated whilst the cluster is actively carrying out tasks, and shows information per engine, and total per host included in the cluster.

Therefore, it is possible to monitor the cluster by screening this file in a separated terminal, e.g. using the operating system command `watch`:

```
watch cat monitoring.log
```

will result in the screen output

```
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
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<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>hpc01</td>
<td>0</td>
<td>running</td>
<td>75</td>
<td>2</td>
<td>26</td>
<td>334</td>
<td>2</td>
<td>11</td>
<td></td>
<td>0 flagdata X54.0005.ms</td>
<td></td>
</tr>
<tr>
<td>hpc01</td>
<td>1</td>
<td>running</td>
<td>61</td>
<td>2</td>
<td>9</td>
<td>197</td>
<td>0</td>
<td>21</td>
<td></td>
<td>0 flagdata X54.0007.ms</td>
<td></td>
</tr>
<tr>
<td>hpc01</td>
<td>2</td>
<td>running</td>
<td>74</td>
<td>2</td>
<td>28</td>
<td>386</td>
<td>3</td>
<td>13</td>
<td></td>
<td>0 flagdata X54.0003.ms</td>
<td></td>
</tr>
<tr>
<td>hpc01</td>
<td>3</td>
<td>scheduled</td>
<td>24</td>
<td>1</td>
<td>0</td>
<td>101</td>
<td>0</td>
<td>0</td>
<td></td>
<td>0 flagdata X54.0012.ms</td>
<td></td>
</tr>
<tr>
<td>hpc01</td>
<td>4</td>
<td>running</td>
<td>82</td>
<td>2</td>
<td>28</td>
<td>386</td>
<td>3</td>
<td>13</td>
<td></td>
<td>0 flagdata X54.0001.ms</td>
<td></td>
</tr>
<tr>
<td>hpc01</td>
<td>5</td>
<td>scheduled</td>
<td>21</td>
<td>1</td>
<td>0</td>
<td>13</td>
<td>0</td>
<td>0</td>
<td></td>
<td>0 flagdata X54.0011.ms</td>
<td></td>
</tr>
<tr>
<td>------</td>
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<td>-------------</td>
<td>-----------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>hpc01</td>
<td>Total</td>
<td></td>
<td>337</td>
<td>10</td>
<td>1318</td>
<td>8</td>
<td>61</td>
<td>0</td>
<td></td>
<td>-----------------</td>
<td>-----------------</td>
</tr>
</tbody>
</table>
```

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Appendix A

Obtaining, Installing, and Customizing CASA

A.1 Installation On Linux

To install CASA for Linux, we have packaged up a binary distribution of CASA which is available as a downloadable tar file. We believe this binary distribution works with most Linux distributions. While the binary distribution is the only supported public distribution, most CASA developers use RPMs for many third-party packages installed with yum to do development on RedHat Enterprise Linux. Installing the developer RPMs requires root access and we only provide developer support for organizations which have a cooperative agreement to participate in the development of CASA. We are currently working on the development of a distribution for developers similar to our standard binary distribution, but it is not yet ready for testing.

A.1.1 Installation

You do not have to have root or sudo permission, you can easily install CASA, delete it, move it, and it works for many versions of Linux. The one caveat is that CASA on Linux currently will not run if the Security-Enhanced Linux option of the linux operating system is set to enforcing. For the non-root install to work, SELinux must be set to disabled or permissive (in /etc/selinux/config) or you must run (as root):

```
setsebool -P allow_execheap=1
```

Otherwise, you will encounter errors like:

```
casapy: error while loading shared libraries:
  /opt/casa/casapy-20.0.5653-001/lib/liblapack.so.3.1.1:
  cannot restore segment prot after reloc: Permission denied
```
The non-root installation is thought to work on a wide variety of linux platforms, see Sect. 1.2 for the latest supported OSs.

A.1.1.1 Using more than one Linux version of CASA

Up to CASA 3.3.0, CASA .rpm files prohibited installing more than one CASA release at a time. Starting with CASA 3.4.0, CASA .rpm files allow previously installed CASA releases to remain installed.

To start a specific CASA version, type

```bash
  casapy --release <VERSION>
```

or

```bash
  casapy -r <VERSION>
```

where `\<VERSION\>` is a placeholder for the CASA version to be invoked, e.g. 3.3.0.

The programs asdm2MS, casabrowser, casalogger, casaplotms, casapy, casapyinfo, and casaviewer all take the two new command line options: -r and --release. These options allow users to select a CASA program to run from the installed CASA releases.

A.1.2 Unsupported platforms

The non-root install may work on other platforms not listed, please let us know if you find that this binary distribution of CASA works on other linux platforms. Also note, that the plotting tasks like plotxy and plotcal are the ones that typically give problems for new platforms, so a check of these after attempting an unsupported platform installation is advisable.

A.1.3 Download & Unpack

You can download the distribution tar file from

[http://casa.nrao.edu/casa_obtaining.shtml](http://casa.nrao.edu/casa_obtaining.shtml)

This directory will contain two tar files one will be the 32-bit version of CASA and the other will be the 64-bit version of CASA. The file name of the 64-bit version ends with -64b.tar.gz. After downloading the appropriate tar file, untar it with

```bash
  tar -zxf casapy-*.tar.gz
```

This will extract a directory with the same basename as the tar file. Change to that directory and add it to your path with, for example,
APPENDIX A. APPENDIX: OBTAINING, INSTALLING, AND CUSTOMIZING CASA

PATH=‘pwd’:PATH.

After that, you should be able to start CASA by running

casapy

A.2 Installation on Mac OS

CASA for Macintosh is distributed as self-contained Macintosh application. For installation purposes, this means that you can install CASA by simply dragging the application to your hard disk. It should be as easy as copying a file.

1. Download the CASA disk image for your OS version from our download site

   http://casa.nrao.edu/casa_obtaining.shtml

2. Open the disk image file (if your browser does not do so automatically).

3. Drag the CASA application to the Applications folder of your hard disk.

4. Eject the CASA disk image.

5. Double-click the CASA application to run it for the first time. This ensures everything is properly updated if you had installed a previous version.

6. Optional: Create symbolic links to the CASA version and its executables (Administrator privileges are required), which will allow you to run casapy, casaviewer, casaplotms, etc. from any terminal command line. To do so, run !create-symlinks from a CASA prompt. (In CASA 4.2.2 and earlier you will be prompted to update the symlinks on double clicking the Application)

You may need to unload the dbus before the copy will work

launchctl remove org.freedesktop.dbus-session
launchctl remove org.freedesktop.dbus-system

Versions after 12115 are 64bit only and will not work on older mac intel machines The first time you launch the CASA application, it will prompt you to set up an alias to the casapy command. You will be taken through the process of creating several casapy symbolic links, it is advisable to do so as this will allow you to run casapy from a terminal window by typing casapy. Additionally, the viewer (casaviewer), table browser (casabrowser), plotms (casaplotms), and buildmytasks will also be available via the command line. Creating the symbolic links will require that you have administrator privileges.
A.2.0.1 Using more than one Mac version of CASA

By dragging the CASA.app into the Applications folder, any previous version of CASA will be replaced. If one would like to keep older versions, one can simply rename them, e.g., to CASA-3.3.0.app. Double clicking any of the CASA*.app applications will prompt to update the symlinks to that specific CASA version. So any startup of casapy, casaviewer, casaplotms will point to that version. If one decides to switch to a different version, just double click the respective CASA*.app and follow the instructions to update the symlinks.

A.3 Startup

in a terminal type

casapy

and the world of CASA will open its doors for you.

There are a number of options to casapy (see casapy --help): Options are:

```
--rcdir directory
--logfile logfilename specify the name of the log file if other than casapy-DATE.log
--maclogger will use the Mac Console program for the logger
--log2term output the logger text in the terminal
--nologger run without launching the logger
--nologfile does not create a logfile
--nogui will not open the logger GUI
--colors=[NoColor|Linux|LightBG] selects color theme for prompt task inputs
--noipython does not launch ipython (useful when combined with the -c option)
--release <VERSION> launches CASA version <VERSION> when installed as Linux rpm
-r <VERSION> alias for --release
-c filename-or-expression execute a CASA python script from the command line
--help print this text and exit
```

E.g. you can execute a CASA script script.py directly with the command

casapy -c script.py

You can also launch the plotms and viewer GUIs separately without starting CASA itself. To do so, type:

casaplotms
to launch `plotms` and

`casaviewer`

for the viewer.

### A.4 Startup Customization

There are two initialization files that are loaded upon startup. The first is loaded very early in the startup of `casapy`:

```
~/.casa/prelude.py
```

This allows for limited customization of the `casapy` environment, e.g. setting the path to an alternate logger. The second startup file should be used for most purposes:

```
~/.casa/init.py
```

This file is loaded just before the `casapy` prompt is display. This is the place where users can load their own python modules and casa tasks. For example `~/.casa/init.py` might contain:

```
import os
sys.path.insert(1, os.environ['HOME'] + os.sep + "python")
import analysisUtils as aU
```

and `analysisUtils.py` might contain:

```
import numpy as np
from mpfit import mpfit
from pylab import *
from numpy.fft import fft
from scipy import polyfit
import taskinit as ti
from importasdm import importasdm
```

Many options can also be set in the file

```
~/.casarc
```

E.g.
# Set these so that bug(), ask(), etc. know who you are
#
userinfo.name: Sheila User
userinfo.email: suser@nrao.edu
userinfo.org: NRAO

#NOTE: Fill this value in as appropriate - the units are MB
#It is important that you not set this value larger than your actual
#physical memory
#system.resources.memory: 2000
#help.popup.type: mb3long

catalog
  catalog.gui.auto: T
  catalog.confirm: T
  catalog.view.PostScript: ghostview
  catalog.edit.ascii: xterm -e vi

#logger
  #logger.file: ./aips++.log
  #logger.height: 12
  logger.default: screen

#progress meter GUI pop-ups - disable
  progress.show: F

#toolmanager - disable
  toolmanager.gui.auto: F

#Use current working directory for cache/scratch files
user.aipsdir: .
user.cache: .
user.directories.work: .
user.initfiles: almainit.g

#viewer
  display.axislabels: on
  display.colormaps.defaultcolormap: Hot Metal 1

#development
  #ms.async: ddd ./ms %s
A.5 Updating the data repository

Each CASA release for Linux comes with an up to date data repository (containing information such as observatory coordinates, calibrator models, leap second tables, etc.). However, the files that make up the data repository are updated regularly. Therefore, if you install (or have installed) a release that is a few weeks to a month old, it makes sense to update the data repository because it is very easy.

To do so, issue the following command from the CASA prompt:

```
CASA <2>: !update-data
```
Appendix B

Python and CASA

CASA uses Python, IPython and matplotlib within the package. IPython is an enhanced, interactive shell to Python which provides many features for efficient command line interaction, while matplotlib is a Python 2-D plotting library for publication quality figures in different hardcopy formats.

From www.python.org: "Python is an interpreted, interactive, object-oriented programming language". Python is used as the underlying command line interface/scripting language to CASA. Thus, CASA inherits the features and the annoyances of Python. For example, since Python is inherently 0-based in its indexing of arrays, vectors, etc, CASA is also 0-based; any Index inputs (e.g., start (for start channel), fieldIndex, antennaID, etc) will start with 0. Another example is that indenting of lines means something to Python, of which users will have to be aware.

Currently, CASA uses python 2.6 (2.5 for the Mac OS 10.5 version). Some key links to python are:

- [http://python.org](http://python.org) – Main Python page
- [http://docs.python.org/release/2.6.5/tutorial/index.html](http://docs.python.org/release/2.6.5/tutorial/index.html) – Python Reference
- [http://docs.python.org/release/2.6.5/reference/index.html](http://docs.python.org/release/2.6.5/reference/index.html) – Python Tutorial
- [http://ipython.scipy.org](http://ipython.scipy.org) – IPython page

Each of the features of these components behave in the standard way within CASA. In the following sections, we outline the key elements for analysis interactions; see the Python references and the IPython page for the full suite of functionality.

B.1 Python Packages

The following python packages are included in CASA: ipython, nose, pyfits, pytz, dbus, numpy, scientific python, twisted, zope.interface, foolscap, matplotlib, scipy. For their version numbers, please
B.2 Automatic parentheses

Automatic parenthesis is enabled for calling functions with argument lists; this feature is intended to allow less typing for common situations. IPython will display the interpretation of the line, beneath the one typed, as indicated by the '-------->'. Default behavior in CASA is to have automatic parenthesis enabled.

B.3 Indentation

Python pays attention to indentation of lines in scripts or when you enter them interactively. It uses indentation to determine the level of nesting in loops. Be careful when cutting and pasting, if you get the wrong indentation, then unpredictable things can happen (usually it just gives an error).

A blank line can be used to return the indentation to a previous level. For example, expanded parameters in tasks cause indentation in subsequent lines in the interface. For example, the following snippet of inputs from clean can be cut and pasted without error due to the blank line after the indented parameters:

```python
mode = 'channel'  # Type of selection
nchan = -1        # Number of channels to select
start = 0         # Start channel
step = 1          # Increment between channels/velocity
width = 1         # Channel width
alg = 'clark'     # Algorithm to use
```

If the blank line were not there, an error would result if you pasted this at the casapy prompt.

B.4 Lists and Ranges

Sometimes, you need to give a task a list of indices. For example, some tasks and tools expect a comma-separated Python list, e.g.

```python
scanlist = [241, 242, 243, 244, 245, 246]
```

You can use the Python `range` function to generate a list of consecutive numbers, e.g.

```python
scanlist = range(241,247)
```

giving the same list as above, e.g.
CASA <1>: scanlist=range(241,247)
CASA <2>: print scanlist
[241, 242, 243, 244, 245, 246]

Note that `range` starts from the first limit and goes to one below the second limit (Python is 0-based, and `range` is designed to work in loop functions). If only a single limit is given, the first limit is treated as 0, and the one given is used as the second, e.g.

CASA <3>: iflist=range(4)
CASA <4>: print iflist
[0, 1, 2, 3]

You can also combine multiple ranges by summing lists

CASA <5>: scanlist=range(241,247) + range(251,255)
CASA <6>: print scanlist
[241, 242, 243, 244, 245, 246, 251, 252, 253, 254]

### B.5 Dictionaries

Python dictionaries are data structures that contain `key:value` pairs, sort of like a hash array. These are useful to store mini-databases of things. In CASA, the parameter values are kept in a dictionary behind the scenes.

To initialize a dictionary, say we call it `mydict`, for use:

CASA <7>: mydict = {}

To add members:

CASA <8>: mydict['source'] = '0137+331'
CASA <9>: mydict['flux'] = 5.4

To see its contents:

CASA <10>: mydict
Out[10]: {'flux': 5.4000000000000004, 'source': '0137+331'}
CASA <11>: print mydict
{'source': '0137+331', 'flux': 5.4000000000000004}

To access a specific entry:

CASA <12>: print mydict['flux']
5.4
B.5.1 Saving and Reading Dictionaries

To save a simple dictionary to a file:

```python
CASA <13>: dictfile = open('mydictfile.py','w')
CASA <14>: print >>dictfile,"mydict = ",mydict
CASA <15>: dictfile.close()
CASA <16>: !cat mydictfile.py
IPython system call: cat mydictfile.py
mydict = { 'source': '0137+331', 'flux': 5.4000000000000004}

CASA <17>: mydict = {}
CASA <18>: run mydictfile.py
CASA <19>: mydict
Out[19]: {'flux': 5.4000000000000004, 'source': '0137+331'}
```

More complex dictionaries, like those produced by `imstat` that contain NumPy arrays, require a different approach to save. The `pickle` module lets you save general data structures from Python. For example:

```python
CASA <20>: import pickle
CASA <21>: xstat
Out[21]:
{ 'blc': array([0, 0, 0, 0]),
  'blcf': '15:24:08.404, +04.31.59.281, I, 1.41281e+09Hz',
  'flux': array([ 4.0795296]),
  'max': array([ 0.05235516]),
  'maxpos': array([134, 134, 0, 38]),
  'maxposf': '15:21:53.976, +05.05.29.998, I, 1.41374e+09Hz',
  'mean': array([ 1.60097857e-05]),
  'medabsdevmed': array([ 0.0127436]),
  'median': array([ -1.17422514e-05]),
  'min': array([-0.0104834]),
  'minpos': array([160, 1, 0, 30]),
  'minposf': '15:21:27.899, +04.32.14.923, I, 1.41354e+09Hz',
  'npts': array([ 3014656.]),
  'quartile': array([ 0.002584881]),
  'rms': array([ 0.0020222]),
  'sigma': array([ 0.0020222]),
  'sum': array([ 48.26399646]),
  'sumsq': array([ 12.32857318]),
  'trc': array([255, 255, 0, 45]),
  'trcf': '15:19:52.390, +05.35.44.246, I, 1.41391e+09Hz'}
```

```python
CASA <22>: mydict
Out[22]: {'flux': 5.4000000000000004, 'source': '0137+331'}
```

```python
CASA <23>: pickfile = 'myxstat.pickle'
CASA <24>: f = open(pickfile,'w')
CASA <25>: p = pickle.Pickler(f)
```
CASA <26>: p.dump(xstat)
CASA <27>: p.dump(mydict)
CASA <28>: f.close()

The dictionaries are now saved in pickle file myxstat.pickle in the current directory.

To retrieve:

CASA <29>: xstat2 = {}
CASA <30>: mydict2 = {}
CASA <31>: f = open(pickfile)
CASA <32>: u = pickle.Unpickler(f)
CASA <33>: xstat2 = u.load()
CASA <34>: mydict2 = u.load()
CASA <35>: f.close()
CASA <36>: xstat2
Out[36]:
{'blc': array([0, 0, 0, 0]),
 'blcf': '15:24:08.404, +04.31.59.181, I, 1.41281e+09Hz',
 'flux': array([4.0795296]),
 'max': array([0.05235516]),
 'maxpos': array([134, 134, 0, 38]),
 'maxposf': '15:21:53.976, +05.05.29.998, I, 1.41374e+09Hz',
 'mean': array([1.60097857e-05]),
 'medabsdevmed': array([0.00127436]),
 'median': array([-1.17422514e-05]),
 'min': array([-0.0104834]),
 'minpos': array([160, 1, 0, 30]),
 'minposf': '15:21:27.899, +04.32.14.923, I, 1.41354e+09Hz',
 'npts': array([3014656.]),
 'quartile': array([0.00254881]),
 'rms': array([0.00202226]),
 'sigma': array([0.00202222]),
 'sum': array([48.26399646]),
 'sumsq': array([12.32857318]),
 'trc': array([255, 255, 0, 45]),
 'trcf': '15:19:52.390, +05.35.44.246, I, 1.41391e+09Hz'}

CASA <37>: mydict2
Out[37]: {'flux': 5.4000000000000004, 'source': '0137+331'}

Thus, you can make scripts that save information and use it later, like for regressions.

Note that these examples use Python file-handling and IO, as well as importing modules such as pickle. See your friendly Python reference for more on this kind of stuff. It’s fairly obvious how it works.
B.6 Control Flow: Conditionals, Loops, and Exceptions

There are a number of ways to control the flow of execution in Python, including conditionals (if), loops (for and while), and exceptions (try). We will discuss the first two below.

B.6.1 Conditionals

The standard if block handles conditional execution or branches in Python:

```python
if <expression>:
    <statements>
elif <expression>:
    <statements>
elif <expression>:
    <statements>
...
else:
    <statements>
```

Insert a `pass` statement if you want no action to be taken for a particular clause. The `<expression>` should reduce down to True or False.

For example,

```python
if ( importmode == 'vla' ):
    # Import the data from VLA Export to MS
    default('importvla')
    print "Use importvla to read VLA Export and make an MS"
    archivefiles = datafile
    vis = msfile
    bandname = exportband
    autocorr = False
    antnamescheme = 'new'
    project = exportproject
    importvla()

elif ( importmode == 'fits' ):
    # Import the data from VLA Export to MS
    default('importuvfits')
    print "Use importuvfits to read UVFITS and make an MS"
    fitsfile = datafile
    vis = msfile
    importuvfits()
else:
```

```
# Copy from msfile
print "Copying "+datafile+" to "+msfile
os.system('cp -r '+datafile+' '+msfile)
vis = msfile

chooses branches based on the value of the **importmode** Python variable (set previously in script).

## B.6.2 Loops

The **for** loop

```python
for iter in seq:
    <statements>
```

iterates over elements of a sequence **seq**, assigning each in turn to **iter**. The sequence is usually a list of values.

For example,

```python
splitms = 'polcal_20080224.cband_all.split.ms'
srclist = ['0137+331','2136+006','2202+422','2253+161','0319+415','0359+509']
spwlist = ['0','1']
for src in srclist:
    for spwid in spwlist:
        imname = splitms + '.' + src + '.' + spwid + '.clean'
        clean(vis=splitms,field=src,spw=spwid,imagename=imname,
               stokes='IQUV',psfmode='hogbom',imagermode='csclean',
               imsize=[288,288],cell=[0.4,0.4],niter=1000,
               threshold=1.3,mask=[134,134,154,154])
```

# Done with spw

# Done with sources

As usual, blocks are closed by blank lines of the previous indentation level.

You can use the **range** ([B.4]) Python function to generate a numerical loop:

```python
vis = 'polcal_20080224.cband_all.ms'
for i in range(0,6):
    fld = str(i)
    plotxy(vis,field=fld,xaxis='uvdist',yaxis='amp')
```

# Done with fields [0, 1, 2, 3, 4, 5]
There is also a while loop construct

```python
while <expression>:
    <statements>
```

which executes the statement block while the <expression> is True. The while loop can also take an else block.

For example,

```python
# Do an explicit set of clean iterations down to a limit
prevrms = 1.e10
while rms > 0.001 :
    clean(vis=splitms,field=src,spw=spwid,imagename=imname,
         stokes='IQUV',psfmode='hogbom',imagermode='csclean',
         imsize=[288,288],cell=[0.4,0.4],niter=200,
         threshold=1.3,mask=[134,134,154,154])

    offstat=imstat(imname+'.residual',box='224,224,284,284')
    rms=offstat['sigma'][0]
    if rms > prevrms:
        break # the rms has increased, stop

    prevrms = rms

# Clean until the off-source rms residual, reaches 0.001 Jy
```

Note that you can exit a loop using the break statement, as we have here when the rms increases.

### B.7 System shell access

For scripts, the os.system methods are the preferred way to access system shell commands (see §[B.7.1](#)).

In interactive mode, any input line beginning with a ’!’ character is passed verbatim (minus the ’!’) to the underlying operating system. Several common commands (ls, pwd, less) may be executed with or without the ’!’ . Note that the cd command must be executed without the ’!’ , and the cp command must use ’!’ as there is a conflict with the cp tool in casapy.

For example:

```
CASA [1]: pwd
/export/home/corsair-vmI/jmcnulli/data
CASA [2]: ls n*
ngc5921.ms ngc5921.py
CASA [3]: !cp -r ../*.py .
```
B.7.1 Using the os.system methods

To use this, you need the os package. This should be loaded by default by casapy, but if not you can use

    import os

in your script.

For example, in our scripts we use this to clean up any existing output files

    # The prefix to use for all output files
    prefix='ngc5921.usecase'

    # Clean up old files
    os.system('rm -rf '+prefix+'*')

Note that the os package has many useful methods. You can see these by using tab-completion:

```
CASA <2>: os.<tab>
```

```
os.EX_CANTCREAT os._Environ os.fdatasync os.remove
os.EX_CONFIG os.__all__ os.fdopen os.removedirs
os.EX_DATAERR os.__builtins__ os.fork os.rename
os.EX_IOERR os.__class__ os.forkpty os.renames
os.EX_NOHOST os.__delattr__ os.fpathconf os.rmdir
os.EX_NOINPUT os.__dict__ os.fstat os.sep
os.EX_NOPERM os.__doc__ os.fstatvfs os.setegid
os.EX_NOUSER os.__file__ os.fsync os.seteuid
os.EX_OK os.__getattribute__ os.ftruncate os.setgid
os.EX_OSERR os.__hash__ os.getcwd os.setgroups
os.EX_TEMPFAIL os.__init__ os.getcwd os.setpgid
os.EX_USAGE os.__new__ os.getegid os.setpgrp
os.F_OK os._copy_reg os.getgid os.spawnl
os.NGROUPS_MAX os._exists os.geteuid os.spawnlpe
os.O_APPEND os._exit os.getlogin os.spawnlp
os.O_CREAT os._get_exports_list os.getpid os.spawnve
os.O_DIRECT os._make_stat_result os.getppid os.spawnv
os.O_DIRSYNC os._make_statvfs_result os.getpgrp os.spawnvpe
os.O_NOCTTY os.abort os.getpwuid os.stat
os.O_NOLOCK os._chown os.getstat(os.stat_result)
```
APPENDIX B. APPENDIX: PYTHON AND CASA

B.7.2 Directory Navigation

In addition, filesystem navigation is aided through the use of bookmarks to simplify access to frequently-used directories:

```
CASA [4]: cd /home/ballista/jmcmulli/other_data
CASA [4]: pwd
/home/ballista/jmcmulli/other_data
CASA [5]: bookmark other_data
CASA [6]: cd /export/home/corsair-vml/jmcmulli/data
CASA [7]: pwd
/export/home/corsair-vml/jmcmulli/data
CASA [8]: cd -b other_data
(bookmark:data) -> /home/ballista/jmcmulli/other_data
```

For python scripts, there is a special command to change a directory.

```
os.system('cd ~/directory')
```

will NOT work but the following will:
os.chdir('~/directory')

### B.7.3 Shell Command and Capture

See also §B.9 for the use of the command history.

1. `sx shell_command, !!shell_command` - this captures the output to a list

```casa
CASA [1]: sx pwd # stores output of 'pwd' in a list
       Out[1]: ['/home/basho3/jmcmulli/pretest']

CASA [2]: !!pwd # !! is a shortcut for 'sx'
       Out[2]: ['/home/basho3/jmcmulli/pretest']

CASA [3]: sx ls v* # stores output of 'pwd' in a list
       Out[3]: ['vla_calplot.jpg',
                'vla_calplot.png',
                'vla_mssplot_calcs.jpg',
                'vla_mssplot_calcs.png',
                'vla_plotcal_bpass.jpg',
                'vla_plotcal_bpass.png',
                'vla_plotcal_fcal.jpg',
                'vla_plotcal_fcal.png',
                'vla_plotvis.jpg',
                'vla_plotvis.png']

CASA [4]: x=_ # remember '_' is a shortcut for the output from the last command

CASA [5]: x
       Out[5]: ['vla_calplot.jpg',
                'vla_calplot.png',
                'vla_mssplot_calcs.jpg',
                'vla_mssplot_calcs.png',
                'vla_plotcal_bpass.jpg',
                'vla_plotcal_bpass.png',
                'vla_plotcal_fcal.jpg',
                'vla_plotcal_fcal.png',
                'vla_plotvis.jpg',
                'vla_plotvis.png']

CASA [6]: y=Out[2] # or just refer to the enumerated output

CASA [7]: y
       Out[7]: ['/home/basho3/jmcmulli/pretest']
```

2. `sc - captures the output to a variable; options are '-l' and '-v'

```casa
CASA [1]: sc x=pwd # capture output from 'pwd' to the variable 'x'
```
B.8 Logging

There are two components to logging within CASA. Logging of all command line inputs is done via IPython.

Upon startup, CASA will log all commands to a file called ipython.log. This file can be changed via the use of the ~/.casa/ipython/ipythonrc file. This log file can be edited and re-executed as appropriate using the execfile feature (§B.12).

Logging can be turned on and off using the logon, logoff commands.

The second component is the output from applications which is directed to the file ./casapy.log. See §1.5.2 for more on the casalogger.

B.9 History and Searching

Numbered input/output history is provided natively within IPython. Command history is also maintained on-line.

CASA [2]: x
Out[2]: '/home/basho3/jmcmulli/pretest'

CASA [3]: sc -l x=pwd # capture the output from 'pwd' to the variable 'x' but
  # split newlines into a list (similar to sx command)
  
CASA [4]: x
Out[4]: ['/home/basho3/jmcmulli/pretest']

CASA [5]: sc -v x=pwd # capture output from 'pwd' to a variable 'x' and
  # show what you get (verbose mode)
  
x ==
  '/home/basho3/jmcmulli/pretest'

CASA [6]: x
Out[6]: '/home/basho3/jmcmulli/pretest'
Command history can be accessed via the 'hist' command. The history is reset at the beginning of every CASA session, that is, typing 'hist' when you first start CASA will not provide any commands from the previous session. However, all of the commands are still available at the command line and can be accessed through the up or down arrow keys, and through searching.

The history can be saved as a script or used as a macro for further use:

CASA [24]: save script.py 13:16
File 'script.py' exists. Overwrite (y/[N])? y
The following commands were written to file 'script.py':
  x=1

```plaintext
y=3*x
z=x**2+y**2
CASA [25]: !more script.py
x=1
y=3*x
z=x**2+y**2
```

Note that the history commands will be saved up to, but not including the last value (i.e., history commands 13-16 saves commands 13, 14, and 15).

There are two mechanisms for searching command history:

1. Previous/Next: use Ctrl-p (previous,up) and Ctrl-n (next,down) to search through only the history items that match what you have typed so far (min-match completion). If you use Ctrl-p or Ctrl-n at a blank prompt, they behave just like the normal arrow keys.

2. Search: Ctrl-r opens a search prompt. Begin typing and the system searches your history for lines that contain what you’ve typed so far, completing what it can. For example:

   ```plaintext
   CASA [37]: <CTRL-r>
   (reverse-i-search)’’:
   Typing anything after the colon will provide you with the last command matching the characters, for example, typing 'op' finds:
   (reverse-i-search)‘op’: im.open('ngc5921.ms')
   Subsequent hitting of Ctrl-r will search for the next command matching the characters.
   ```

## B.10 Macros

Macros can be made for easy re-execution of previous commands. For example to store the commands 13-15 to the macro 'example':

```plaintext
CASA [31]: macro example 13:16
Macro ‘example’ created. To execute, type its name (without quotes).
Macro contents:
x=1
y=3*x
z=x**2+y**2
CASA [32]: z
Out[32]: 6
CASA [33]: z=10
```
B.11 On-line editing

You can edit files on-line in two ways:

1. Using the shell access via '!vi'

2. Using the ed function; this will edit the file but upon closing, it will try to execute the file; using the 'script.py' example above:

```
CASA [13]: ed script.py # this will bring up the file in your chosen editor
    # when you are finished editing the file,
    # it will automatically
    # execute it (as though you had done a
    # execfile 'script.py'

    Editing... done. Executing edited code...
```

B.12 Executing Python scripts

Python scripts are simple text files containing lists of commands as if typed at the keyboard. Note: the auto-parentheses feature of IPython cannot be used in scripts, that is, you should make sure all function calls have any opening and closing parentheses.

```
# file is script.py
# My script to plot the observed visibilities
plotxy('ngc5921.ms','uvdist') #yaxis defaults to amplitude
```

This can be done by using the execfile command to execute this script. execfile will execute the script as though you had typed the lines at the CASA prompt.
CASA [5]: execfile 'script.py'
--------> execfile('script.py')

If you don’t want to launch CASA and execute your script from the command line, you can use the '-c' option:

unix$ casapy -c 'script.py'

B.13 How do I exit from CASA?

You can exit CASA by using the quit command. This will bring up the query

Do you really want to exit ([y]/n)?

to give you a chance in case you did not mean to exit. You can also quit using %exit or CTRL-D.

If you don’t want to see the question "Do you really want to exit [y]/n?", then just type Exit or exit followed by return, and CASA will stop right then and there.
Appendix C

Appendix: Models, Conventions, and Reference Frames

This appendix lists the available parameters, conventions, reference frames, and information on flux standards used in CASA.

C.1 Flux Density Models for setjy

`setjy` adds a source model given the source name, frequency, a standard (really, a set of models), and possibly a time. At cm wavelengths the flux density (FD) calibrators are typically one of several bright extragalactic sources. These objects are comparatively faint and less well characterized at shorter wavelengths, so for (sub)mm astronomy it is common to use Solar System objects.

Reliably setting the FD scale with astronomical calibrators requires that they be bright, not too resolved, and have simple dependencies on frequency and time. These criteria are somewhat mutually exclusive, so the number of calibrator sources supported by `setjy` is fairly small, although it could certainly be added to. This appendix is for describing the models that `setjy` uses. Choosing a FD calibrator of course has to be done before the observation and the observatory may provide additional information.

C.1.1 Long wavelength calibration

Synchrotron sources can vary over a light crossing time, so ones used as FD calibrators must have most of their emission coming from an extended region. The additional requirement that they be nearly unresolved therefore forces them to be distant, meaning that candidates which also have high apparent fluxes are quite rare. The following standards mostly share the same set of objects, and monitor their FDs every few years to account for variations. No interpolation is done between epochs, though - you are encouraged to choose the standard which observed your FD calibrator closest to the time you observed it at. The measurements are interpolated in frequency, however, using second to fourth degree polynomials of the frequency’s logarithm.
Table C.1: Recognized Flux Density Calibrators. Note that the VLA uses J2000 calibrator names. CASA accepts all strings that contain the names below. E.g. 'PKS 1934-638' will be recognized

<table>
<thead>
<tr>
<th>3C Name</th>
<th>B1950 Name</th>
<th>J2000 Name</th>
<th>Alt. J2000 Name</th>
<th>Standards</th>
</tr>
</thead>
<tbody>
<tr>
<td>3C48</td>
<td>0134+329</td>
<td>0137+331</td>
<td>J0137+3309</td>
<td>1,3,4,5,6,7</td>
</tr>
<tr>
<td>3C123</td>
<td>0433+295</td>
<td>0437+296</td>
<td>J0437+2940</td>
<td>2</td>
</tr>
<tr>
<td>3C138</td>
<td>0518+165</td>
<td>0521+166</td>
<td>J0521+1638</td>
<td>1,3,4,5,6</td>
</tr>
<tr>
<td>3C147</td>
<td>0538+498</td>
<td>0542+498</td>
<td>J0542+4951</td>
<td>1,3,4,5,6,7</td>
</tr>
<tr>
<td>3C196</td>
<td>0809+483</td>
<td>0813+482</td>
<td>J0813+4813</td>
<td>1,2,7</td>
</tr>
<tr>
<td>3C286</td>
<td>1328+307</td>
<td>1331+305</td>
<td>J1331+3030</td>
<td>1,2,3,4,5,6,7</td>
</tr>
<tr>
<td>3C295</td>
<td>1409+524</td>
<td>1411+522</td>
<td>J1411+5212</td>
<td>1,2,3,4,5,6,7</td>
</tr>
<tr>
<td></td>
<td>1934-638</td>
<td></td>
<td>J1939-6342</td>
<td>1,3,4,5,6</td>
</tr>
<tr>
<td>3C380</td>
<td>1828+487</td>
<td>1829+487</td>
<td>J1829+4845</td>
<td>7</td>
</tr>
</tbody>
</table>


C.1.1.1 Baars

The only standard to not have the year in the name. It is 1977.
The models are second order polynomials in log(ν), valid between 408 MHz and 15 GHz.

C.1.1.2 Perley 90


C.1.1.3 Perley-Taylor 95


C.1.1.4 Perley-Taylor 99

C.1.1.5 Perley-Butler 2010


C.1.1.6 Perley-Butler 2013

Flux scale for the constant flux sources 3C123, 3C196, 3C286, and 3C295. The models are time-dependent.


C.1.1.7 Scaife-Heald 2012

Low frequency, 30-300MHz, calibrators 3C48, 3C147, 3C196, 3C286, 3C295, and 3C380.


C.1.2 Short wavelength calibration

The usual approach in this regime is to use models that are, to first order, thermal sources in the Solar System. Their apparent brightness of course varies in time with their distance from the Earth (and Sun), and orientation if they are not perfect spheres with zero obliquity. However, most of them have almost constant surface properties, so once those properties are measured their apparent brightness distributions can in principle be predicted for any time, given an ephemeris. Planets, in particular, however, have more complex spectra and effects such as atmospheric lines, magnetic fields, seasons, polar caps and surface features need to be taken into account when they are available and significant. In CASA the Solar System objects supported by \texttt{setjy} are available through the ‘Butler-JPL-Horizons 2010’, and ‘Butler-JPL-Horizons 2012’ standards. The models are described in ALMA Memo 594 available on \texttt{https://science.nrao.edu/facilities/alma/aboutALMA/Technology/ALMA_Memo_Series/alma594/abs594}.

C.2 Velocity Reference Frames

CASA supported velocity frames are listed in Table C.2

C.2.1 Doppler Types

CASA supported Doppler types are listed in Table C.3
Table C.2: Velocity frames in CASA

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>REST</td>
<td>Laboratory</td>
</tr>
<tr>
<td>LSRK</td>
<td>local standard of rest (kinematic)</td>
</tr>
<tr>
<td>LSRD</td>
<td>local standard of rest (dynamic)</td>
</tr>
<tr>
<td>BARY</td>
<td>barycentric</td>
</tr>
<tr>
<td>GEO</td>
<td>geocentric</td>
</tr>
<tr>
<td>TOPO</td>
<td>topocentric</td>
</tr>
<tr>
<td>GALACTO</td>
<td>galactocentric</td>
</tr>
<tr>
<td>LGROUP</td>
<td>Local Group</td>
</tr>
<tr>
<td>CMB</td>
<td>cosmic microwave background dipole</td>
</tr>
<tr>
<td>Undefined</td>
<td>undefined frame</td>
</tr>
</tbody>
</table>

Table C.3: Doppler types in CASA

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>RADIO</td>
<td></td>
</tr>
<tr>
<td>Z</td>
<td></td>
</tr>
<tr>
<td>RATIO</td>
<td></td>
</tr>
<tr>
<td>BETA</td>
<td></td>
</tr>
<tr>
<td>GAMMA</td>
<td></td>
</tr>
<tr>
<td>OPTICAL</td>
<td></td>
</tr>
<tr>
<td>TRUE</td>
<td></td>
</tr>
<tr>
<td>RELATIVISTIC</td>
<td></td>
</tr>
</tbody>
</table>

C.3 Time Reference Frames

CASA supported time reference frames are listed in Table C.4.

C.4 Coordinate Frames

CASA supported time coordinate frames are listed in Table C.5.

Note that TOPO refers to a time stamp at a given observation date. If more than one observation is concatenated this may lead to vastly erroneous values. Any conversion from TOPO to other frames such as BARY and LSRK should be performed for each individual observation, prior to concatenation or simultaneous imaging.
Table C.4: Time reference frames in CASA

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>LAST</td>
<td></td>
</tr>
<tr>
<td>LMST</td>
<td></td>
</tr>
<tr>
<td>GMST1</td>
<td></td>
</tr>
<tr>
<td>GAST</td>
<td></td>
</tr>
<tr>
<td>UT1</td>
<td></td>
</tr>
<tr>
<td>UT2</td>
<td></td>
</tr>
<tr>
<td>UTC</td>
<td></td>
</tr>
<tr>
<td>TAI</td>
<td></td>
</tr>
<tr>
<td>TDT</td>
<td></td>
</tr>
<tr>
<td>TCG</td>
<td></td>
</tr>
<tr>
<td>TDB</td>
<td></td>
</tr>
<tr>
<td>TCB</td>
<td></td>
</tr>
<tr>
<td>IAT</td>
<td></td>
</tr>
<tr>
<td>GMST</td>
<td></td>
</tr>
<tr>
<td>TT</td>
<td></td>
</tr>
<tr>
<td>ET</td>
<td></td>
</tr>
<tr>
<td>UT</td>
<td></td>
</tr>
</tbody>
</table>

C.5 Physical Units

CASA also recognizes physical units. They are listed in Tables C.6, C.7, and C.8.

C.6 Physical Constants

The physical constants included in CASA can be found in Table C.9.
Table C.5: Coordinate frames in CASA

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>J2000</td>
<td>mean equator and equinox at J2000.0 (FK5)</td>
</tr>
<tr>
<td>JNAT</td>
<td>geocentric natural frame</td>
</tr>
<tr>
<td>JMEAN</td>
<td>mean equator and equinox at frame epoch</td>
</tr>
<tr>
<td>JTRUE</td>
<td>true equator and equinox at frame epoch</td>
</tr>
<tr>
<td>APP</td>
<td>apparent geocentric position</td>
</tr>
<tr>
<td>B1950</td>
<td>mean epoch and ecliptic at B1950.0</td>
</tr>
<tr>
<td>B1950_VLA</td>
<td>mean epoch(1979.9)) and ecliptic at B1950.0</td>
</tr>
<tr>
<td>BMEAN</td>
<td>mean equator and equinox at frame epoch</td>
</tr>
<tr>
<td>BTRUE</td>
<td>true equator and equinox at frame epoch</td>
</tr>
<tr>
<td>GALACTIC</td>
<td>Galactic coordinates</td>
</tr>
<tr>
<td>HADEC</td>
<td>topocentric HA and declination</td>
</tr>
<tr>
<td>AZEL</td>
<td>topocentric Azimuth and Elevation (N through E)</td>
</tr>
<tr>
<td>AZELSW</td>
<td>topocentric Azimuth and Elevation (S through W)</td>
</tr>
<tr>
<td>AZELNE</td>
<td>topocentric Azimuth and Elevation (N through E)</td>
</tr>
<tr>
<td>AZELGEO</td>
<td>geodetic Azimuth and Elevation (N through E)</td>
</tr>
<tr>
<td>AZELSWGEO</td>
<td>geodetic Azimuth and Elevation (S through W)</td>
</tr>
<tr>
<td>AZELNEGEO</td>
<td>geodetic Azimuth and Elevation (N through E)</td>
</tr>
<tr>
<td>ECLIPTIC</td>
<td>ecliptic for J2000 equator and equinox</td>
</tr>
<tr>
<td>MECLIPTIC</td>
<td>ecliptic for mean equator of date</td>
</tr>
<tr>
<td>TECLIPTIC</td>
<td>ecliptic for true equator of date</td>
</tr>
<tr>
<td>SUPERGAL</td>
<td>supergalactic coordinates</td>
</tr>
<tr>
<td>ITRF</td>
<td>coordinates wrt ITRF Earth frame</td>
</tr>
<tr>
<td>TOPO</td>
<td>apparent topocentric position</td>
</tr>
<tr>
<td>ICRS</td>
<td>International Celestial reference system</td>
</tr>
</tbody>
</table>
Table C.6: Prefixes

<table>
<thead>
<tr>
<th>Prefix</th>
<th>Name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y</td>
<td>(yotta)</td>
<td>$10^{24}$</td>
</tr>
<tr>
<td>Z</td>
<td>(zetta)</td>
<td>$10^{21}$</td>
</tr>
<tr>
<td>E</td>
<td>(exa)</td>
<td>$10^{18}$</td>
</tr>
<tr>
<td>P</td>
<td>(peta)</td>
<td>$10^{15}$</td>
</tr>
<tr>
<td>T</td>
<td>(tera)</td>
<td>$10^{12}$</td>
</tr>
<tr>
<td>G</td>
<td>(giga)</td>
<td>$10^{9}$</td>
</tr>
<tr>
<td>M</td>
<td>(mega)</td>
<td>$10^{6}$</td>
</tr>
<tr>
<td>k</td>
<td>(kilo)</td>
<td>$10^{3}$</td>
</tr>
<tr>
<td>h</td>
<td>(hecto)</td>
<td>$10^{2}$</td>
</tr>
<tr>
<td>da</td>
<td>(deka)</td>
<td>$10$</td>
</tr>
<tr>
<td>d</td>
<td>(deci)</td>
<td>$10^{-1}$</td>
</tr>
<tr>
<td>c</td>
<td>(centi)</td>
<td>$10^{-2}$</td>
</tr>
<tr>
<td>m</td>
<td>(milli)</td>
<td>$10^{-3}$</td>
</tr>
<tr>
<td>u</td>
<td>(micro)</td>
<td>$10^{-6}$</td>
</tr>
<tr>
<td>n</td>
<td>(nano)</td>
<td>$10^{-9}$</td>
</tr>
<tr>
<td>p</td>
<td>(pico)</td>
<td>$10^{-12}$</td>
</tr>
<tr>
<td>f</td>
<td>(femto)</td>
<td>$10^{-15}$</td>
</tr>
<tr>
<td>a</td>
<td>(atto)</td>
<td>$10^{-18}$</td>
</tr>
<tr>
<td>z</td>
<td>(zepto)</td>
<td>$10^{-21}$</td>
</tr>
<tr>
<td>y</td>
<td>(yocto)</td>
<td>$10^{-24}$</td>
</tr>
</tbody>
</table>
Table C.7: SI Units

<table>
<thead>
<tr>
<th>Unit (Name)</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$ (currency)</td>
<td>1 _</td>
</tr>
<tr>
<td>% (percent)</td>
<td>0.01</td>
</tr>
<tr>
<td>%% (permille)</td>
<td>0.001</td>
</tr>
<tr>
<td>A (ampere)</td>
<td>1 A</td>
</tr>
<tr>
<td>AE (astronomical unit)</td>
<td>149597870659 m</td>
</tr>
<tr>
<td>AU (astronomical unit)</td>
<td>149597870659 m</td>
</tr>
<tr>
<td>Bq (becquerel)</td>
<td>1 s⁻¹</td>
</tr>
<tr>
<td>C (coulomb)</td>
<td>1 s A</td>
</tr>
<tr>
<td>F (farad)</td>
<td>1 m⁻² kg⁻¹ s⁻⁴ A²</td>
</tr>
<tr>
<td>Gy (gray)</td>
<td>1 m² s⁻²</td>
</tr>
<tr>
<td>H (henry)</td>
<td>1 m² kg s⁻² A⁻²</td>
</tr>
<tr>
<td>Hz (hertz)</td>
<td>1 s⁻¹</td>
</tr>
<tr>
<td>J (joule)</td>
<td>1 m² kg s⁻²</td>
</tr>
<tr>
<td>Jy (jansky)</td>
<td>10⁻²⁶ kg s⁻²</td>
</tr>
<tr>
<td>K (kelvin)</td>
<td>1 K</td>
</tr>
<tr>
<td>L (litre)</td>
<td>0.001 m³</td>
</tr>
<tr>
<td>M₀ (solar mass)</td>
<td>1.98891944407×10³⁰ kg</td>
</tr>
<tr>
<td>N (newton)</td>
<td>1 m kg s⁻²</td>
</tr>
<tr>
<td>Ohm (ohm)</td>
<td>1 m² kg s⁻³ A⁻²</td>
</tr>
<tr>
<td>Pa (pascal)</td>
<td>1 m⁻¹ kg s⁻²</td>
</tr>
<tr>
<td>S (siemens)</td>
<td>1 m⁻² kg⁻¹ s⁻³ A²</td>
</tr>
<tr>
<td>S₀ (solar mass)</td>
<td>1.98891944407×10³⁰ kg</td>
</tr>
<tr>
<td>Sv (sievert)</td>
<td>1 m² s⁻²</td>
</tr>
<tr>
<td>T (tesla)</td>
<td>1 kg s⁻² A⁻¹</td>
</tr>
<tr>
<td>UA (astronomical unit)</td>
<td>149597870659 m</td>
</tr>
<tr>
<td>V (volt)</td>
<td>1 m² kg s⁻³ A⁻¹</td>
</tr>
<tr>
<td>W (watt)</td>
<td>1 m² kg s⁻³</td>
</tr>
<tr>
<td>Wb (weber)</td>
<td>1 m² kg s⁻² A⁻¹</td>
</tr>
<tr>
<td>_ (undimensioned)</td>
<td>1 _</td>
</tr>
</tbody>
</table>
Table C.7: SI Units – continued

<table>
<thead>
<tr>
<th>Unit</th>
<th>Name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>(year)</td>
<td>31557600 s</td>
</tr>
<tr>
<td>arcmin</td>
<td>(arcmin)</td>
<td>0.000290888208666 rad</td>
</tr>
<tr>
<td>arsec</td>
<td>(arcsec)</td>
<td>$4.8481368111 \times 10^{-6}$ rad</td>
</tr>
<tr>
<td>as</td>
<td>(arcsec)</td>
<td>$4.8481368111e \times 10^{-6}$ rad</td>
</tr>
<tr>
<td>cd</td>
<td>(candela)</td>
<td>1 cd</td>
</tr>
<tr>
<td>cy</td>
<td>(century)</td>
<td>3155760000 s</td>
</tr>
<tr>
<td>d</td>
<td>(day)</td>
<td>86400 s</td>
</tr>
<tr>
<td>deg</td>
<td>(degree)</td>
<td>0.0174532925199 rad</td>
</tr>
<tr>
<td>g</td>
<td>(gram)</td>
<td>0.001 kg</td>
</tr>
<tr>
<td>h</td>
<td>(hour)</td>
<td>3600 s</td>
</tr>
<tr>
<td>l</td>
<td>(litre)</td>
<td>0.001 m$^3$</td>
</tr>
<tr>
<td>lm</td>
<td>(lumen)</td>
<td>1 cd sr</td>
</tr>
<tr>
<td>lx</td>
<td>(lux)</td>
<td>1 m$^{-2}$ cd sr</td>
</tr>
<tr>
<td>m</td>
<td>(metre)</td>
<td>1 m</td>
</tr>
<tr>
<td>min</td>
<td>(minute)</td>
<td>60 s</td>
</tr>
<tr>
<td>mol</td>
<td>(mole)</td>
<td>1 mol</td>
</tr>
<tr>
<td>pc</td>
<td>(parsec)</td>
<td>$3.08567758065 \times 10^{16}$ m</td>
</tr>
<tr>
<td>rad</td>
<td>(radian)</td>
<td>1 rad</td>
</tr>
<tr>
<td>s</td>
<td>(second)</td>
<td>1 s</td>
</tr>
<tr>
<td>sr</td>
<td>(steradian)</td>
<td>1 sr</td>
</tr>
<tr>
<td>t</td>
<td>(tonne)</td>
<td>1000 kg</td>
</tr>
</tbody>
</table>
### Table C.8: Custom Units

<table>
<thead>
<tr>
<th>Unit</th>
<th>Name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;</td>
<td>(arcsec)</td>
<td>$4.8481368111 \times 10^{-6}$ rad</td>
</tr>
<tr>
<td>&quot;.2&quot;</td>
<td>(square arcsec)</td>
<td>$2.35044305391 \times 10^{-11}$ sr</td>
</tr>
<tr>
<td>&quot;&quot;</td>
<td>(arcmin)</td>
<td>$0.000290888208666$ rad</td>
</tr>
<tr>
<td>&quot;&quot;&quot;</td>
<td>(arcsec)</td>
<td>$4.8481368111 \times 10^{-6}$ rad</td>
</tr>
<tr>
<td>&quot;,2&quot;</td>
<td>(square arcsec)</td>
<td>$2.35044305391 \times 10^{-11}$ sr</td>
</tr>
<tr>
<td>&quot;,2&quot;</td>
<td>(square arcmin)</td>
<td>$8.46159499408 \times 10^{-8}$ sr</td>
</tr>
<tr>
<td>:</td>
<td>(hour)</td>
<td>$3600$ s</td>
</tr>
<tr>
<td>::</td>
<td>(minute)</td>
<td>$60$ s</td>
</tr>
<tr>
<td>::::</td>
<td>(second)</td>
<td>$1$ s</td>
</tr>
<tr>
<td>Ah</td>
<td>(ampere hour)</td>
<td>$3600$ s A</td>
</tr>
<tr>
<td>Angstrom</td>
<td>(angstrom)</td>
<td>$1e-10$ m</td>
</tr>
<tr>
<td>Btu</td>
<td>(British thermal unit (Int))</td>
<td>$1055.056$ m$^2$ kg s$^{-2}$</td>
</tr>
<tr>
<td>CM</td>
<td>(metric carat)</td>
<td>$0.0002$ kg</td>
</tr>
<tr>
<td>Cal</td>
<td>(large calorie (Int))</td>
<td>$4186.8$ m$^2$ kg s$^{-2}$</td>
</tr>
<tr>
<td>FU</td>
<td>(flux unit)</td>
<td>$10^{-26}$ kg s$^{-2}$</td>
</tr>
<tr>
<td>G</td>
<td>(gauss)</td>
<td>$0.0001$ kg m$^{-2}$ A$^{-1}$</td>
</tr>
<tr>
<td>Gal</td>
<td>(gal)</td>
<td>$0.01$ m s$^{-2}$</td>
</tr>
<tr>
<td>Gb</td>
<td>(gilbert)</td>
<td>$0.795774715459$ A</td>
</tr>
<tr>
<td>Mx</td>
<td>(maxwell)</td>
<td>$10^{-8}$ m$^2$ kg s$^{-2}$ A$^{-1}$</td>
</tr>
<tr>
<td>Oe</td>
<td>(oersted)</td>
<td>$79.5774715459$ m$^{-1}$ A</td>
</tr>
<tr>
<td>R</td>
<td>(mile)</td>
<td>$0.000258$ kg m$^{-1}$ A</td>
</tr>
<tr>
<td>St</td>
<td>(stokes)</td>
<td>$0.0001$ m$^2$ s$^{-1}$</td>
</tr>
<tr>
<td>Torr</td>
<td>(torr)</td>
<td>$133.322368421$ m$^{-1}$ kg s$^{-2}$</td>
</tr>
<tr>
<td>USfl_oz</td>
<td>(fluid ounce (US))</td>
<td>$2.95735295625 \times 10^{-5}$ m$^3$</td>
</tr>
<tr>
<td>USgal</td>
<td>(gallon (US))</td>
<td>$0.003785411784$ m$^3$</td>
</tr>
</tbody>
</table>
Table C.8: Custom Units – continued

<table>
<thead>
<tr>
<th>Unit</th>
<th>Name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>WU</td>
<td>(WSRT flux unit)</td>
<td>$5 \times 10^{-29} \text{kg s}^{-2}$</td>
</tr>
<tr>
<td>abA</td>
<td>(abampere)</td>
<td>10 A</td>
</tr>
<tr>
<td>abC</td>
<td>(abcoulomb)</td>
<td>10 s A</td>
</tr>
<tr>
<td>abF</td>
<td>(abfarad)</td>
<td>$10^9 \text{m}^{-2} \text{kg}^{-1} \text{s}^4 \text{A}^2$</td>
</tr>
<tr>
<td>abH</td>
<td>(abhenry)</td>
<td>$10^{-9} \text{m}^2 \text{kg}^{-2} \text{s}^{-2} \text{A}^{-2}$</td>
</tr>
<tr>
<td>abOhm</td>
<td>(abohm)</td>
<td>$10^{-9} \text{m}^2 \text{kg}^{-3} \text{A}^{-2}$</td>
</tr>
<tr>
<td>abV</td>
<td>(abvolt)</td>
<td>$10^{-8} \text{m}^2 \text{kg}^{-3} \text{A}^{-1}$</td>
</tr>
<tr>
<td>ac</td>
<td>(acre)</td>
<td>4046.8564224 m²</td>
</tr>
<tr>
<td>arcmin_2</td>
<td>(square arcmin)</td>
<td>$8.462159499408 \times 10^{-8}$ sr</td>
</tr>
<tr>
<td>arcsec_2</td>
<td>(square arcsec)</td>
<td>$2.35044305391 \times 10^{-11}$ sr</td>
</tr>
<tr>
<td>ata</td>
<td>(technical atmosphere)</td>
<td>98066.5 m⁻¹.kg.s⁻²</td>
</tr>
<tr>
<td>atm</td>
<td>(standard atmosphere)</td>
<td>101325 m⁻¹.kg.s⁻²</td>
</tr>
<tr>
<td>bar</td>
<td>(bar)</td>
<td>100000 m⁻¹.kg.s⁻²</td>
</tr>
<tr>
<td>beam</td>
<td>(undefined beam area)</td>
<td>1 -</td>
</tr>
<tr>
<td>cal</td>
<td>(calorie (Int))</td>
<td>4.1868 m² kg s⁻²</td>
</tr>
<tr>
<td>count</td>
<td>(count)</td>
<td>1 -</td>
</tr>
<tr>
<td>cwt</td>
<td>(hundredweight)</td>
<td>50.80234544 kg</td>
</tr>
<tr>
<td>deg_2</td>
<td>(square degree)</td>
<td>0.000304617419787 sr</td>
</tr>
<tr>
<td>dyn</td>
<td>(dyne)</td>
<td>$10^{-5}$ m kg s⁻²</td>
</tr>
<tr>
<td>eV</td>
<td>(electron volt)</td>
<td>$1.60217733 \times 10^{-19}$ m² kg s⁻²</td>
</tr>
<tr>
<td>erg</td>
<td>(erg)</td>
<td>$10^{-7}$ m² kg s⁻²</td>
</tr>
<tr>
<td>fl_oz</td>
<td>(fluid ounce (Imp))</td>
<td>$2.84130488996 \times 10^{-5}$ m³</td>
</tr>
<tr>
<td>ft</td>
<td>(foot)</td>
<td>0.3048 m</td>
</tr>
<tr>
<td>fu</td>
<td>(flux unit)</td>
<td>$10^{-26}$ kg s⁻²</td>
</tr>
<tr>
<td>fur</td>
<td>(furlong)</td>
<td>201.168 m</td>
</tr>
<tr>
<td>gal</td>
<td>(gallon (Imp))</td>
<td>0.00454608782394 m³</td>
</tr>
</tbody>
</table>
Table C.8: Custom Units – continued

<table>
<thead>
<tr>
<th>Unit</th>
<th>Name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>ha</td>
<td>(hectare)</td>
<td>10000 m²</td>
</tr>
<tr>
<td>hp</td>
<td>(horsepower)</td>
<td>745.7 m² kg s⁻³</td>
</tr>
<tr>
<td>in</td>
<td>(inch)</td>
<td>0.0254 m</td>
</tr>
<tr>
<td>kn</td>
<td>(knot (Imp))</td>
<td>0.514773333333 m s⁻¹</td>
</tr>
<tr>
<td>lambda</td>
<td>(lambda)</td>
<td>1</td>
</tr>
<tr>
<td>lb</td>
<td>(pound (avoirdupois))</td>
<td>0.45359237 kg</td>
</tr>
<tr>
<td>ly</td>
<td>(light year)</td>
<td>9.46073047 × 10¹⁵ m</td>
</tr>
<tr>
<td>mHg</td>
<td>(metre of mercury)</td>
<td>133322.387415 m⁻¹ kg s⁻²</td>
</tr>
<tr>
<td>mile</td>
<td>(mile)</td>
<td>1609.344 m</td>
</tr>
<tr>
<td>n_mile</td>
<td>(nautical mile (Imp))</td>
<td>1853.184 m</td>
</tr>
<tr>
<td>oz</td>
<td>(ounce (avoirdupois))</td>
<td>0.028349523125 kg</td>
</tr>
<tr>
<td>pixel</td>
<td>(pixel)</td>
<td>1</td>
</tr>
<tr>
<td>sb</td>
<td>(stilb)</td>
<td>10000 m⁻² cd</td>
</tr>
<tr>
<td>sq_arcmin</td>
<td>(square arcmin)</td>
<td>8.461594999408 × 10⁻⁸ sr</td>
</tr>
<tr>
<td>sq_arcsec</td>
<td>(square arcsec)</td>
<td>2.35044305391 × 10⁻¹¹ sr</td>
</tr>
<tr>
<td>sq_deg</td>
<td>(square degree)</td>
<td>0.000304617419787 sr</td>
</tr>
<tr>
<td>statA</td>
<td>(statampere)</td>
<td>3.33564095198 × 10⁻¹⁰ A</td>
</tr>
<tr>
<td>statC</td>
<td>(statcoulomb)</td>
<td>3.33564095198 × 10⁻¹⁰ s A</td>
</tr>
<tr>
<td>statF</td>
<td>(statfarad)</td>
<td>1.11188031733 × 10⁻¹² m⁻² kg⁻¹ s⁻¹ A²</td>
</tr>
<tr>
<td>statH</td>
<td>(stathenry)</td>
<td>899377374000 m² kg s⁻² A⁻²</td>
</tr>
<tr>
<td>statOhm</td>
<td>(statohm)</td>
<td>899377374000 m² kg s⁻³ A⁻²</td>
</tr>
<tr>
<td>statV</td>
<td>(statvolt)</td>
<td>299.792458 m² kg s⁻³ A⁻¹</td>
</tr>
<tr>
<td>u</td>
<td>(atomic mass unit)</td>
<td>1.661 × 10⁻²⁷ kg</td>
</tr>
<tr>
<td>yd</td>
<td>(yard)</td>
<td>0.9144 m</td>
</tr>
<tr>
<td>yr</td>
<td>(year)</td>
<td>31557600 s</td>
</tr>
</tbody>
</table>
## Table C.9: Physical Constants

<table>
<thead>
<tr>
<th>Constant</th>
<th>Name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>pi</td>
<td>pi</td>
<td>3.14159</td>
</tr>
<tr>
<td>ee</td>
<td>ee</td>
<td>2.71828</td>
</tr>
<tr>
<td>c</td>
<td>light vel.</td>
<td>(2.99792 \times 10^8 \text{ m s}^{-1})</td>
</tr>
<tr>
<td>G</td>
<td>grav. const</td>
<td>(6.67259 \times 10^{11} \text{ N m}^2 \text{kg}^{-2})</td>
</tr>
<tr>
<td>h</td>
<td>Planck const</td>
<td>(6.62608 \times 10^{-34} \text{ J s})</td>
</tr>
<tr>
<td>HI</td>
<td>HI line</td>
<td>1420.41 MHz</td>
</tr>
<tr>
<td>R</td>
<td>gas const</td>
<td>(8.31451 \text{ JK}^{-1} \text{mol}^{-1})</td>
</tr>
<tr>
<td>NA</td>
<td>Avogadro #</td>
<td>(6.02214 \times 10^{23} \text{mol}^{-1})</td>
</tr>
<tr>
<td>e</td>
<td>electron charge</td>
<td>(1.60218 \times 10^{-19} \text{ C})</td>
</tr>
<tr>
<td>mp</td>
<td>proton mass</td>
<td>(1.67262 \times 10^{-27} \text{ kg})</td>
</tr>
<tr>
<td>mp_me</td>
<td>mp/me</td>
<td>1836.15</td>
</tr>
<tr>
<td>mu0</td>
<td>permeability vac.</td>
<td>(1.25664 \times 10^{-6} \text{ H m}^{-1})</td>
</tr>
<tr>
<td>eps0</td>
<td>permittivity vac.</td>
<td>(1.60218 \times 10^{-19} \text{ C})</td>
</tr>
<tr>
<td>k</td>
<td>Boltzmann const</td>
<td>(1.38066 \times 10^{-23} \text{ JK}^{-1})</td>
</tr>
<tr>
<td>F</td>
<td>Faraday const</td>
<td>96485.3 \text{ C mol}^{-1})</td>
</tr>
<tr>
<td>me</td>
<td>electron mass</td>
<td>(9.10939 \times 10^{-31} \text{ kg})</td>
</tr>
<tr>
<td>re</td>
<td>electron radius</td>
<td>(2.8179 \times 10^{-15} \text{ m})</td>
</tr>
<tr>
<td>a0</td>
<td>Bohrs radius</td>
<td>(5.2918 \times 10^{-11} \text{ m})</td>
</tr>
<tr>
<td>R0</td>
<td>solar radius</td>
<td>(6.9599 \times 10^8 \text{ m})</td>
</tr>
<tr>
<td>k2</td>
<td>IAU grav. const^2</td>
<td>(0.000295912 \text{ AU}^3 \text{ d}^{-2} \text{ S0}^{-1})</td>
</tr>
</tbody>
</table>
Appendix D

Appendix: CASA Region File Format

The CASA region file format provides a flexible, easily edited set of region definitions which are accepted across CASA tasks. Region files may be written by hand or using the CASA viewer.

Alert: Whereas the region format is supported by all the data processing tasks, the viewer implementation is still limited to rectangles, ellipses, and some markers.

For a file to be recognized as a valid CASA region text file, the first line must contain the string:

```
#CRTF
```

"CRTF" stands for "CASA Region Text Format". One may also include an optional version number at the end of the string, so it reads #CRTFv0; this indicates the version of the format definition.

Region files have two different kinds of definitions, "regions" and "annotations", each of which is one line long. To indicate an annotation, a line must begin with "ann". Lines that begin with the comment character (#) are not considered for processing or display.

The second line of a file may define global parameters that are to be used for all regions and annotations in that file, in which case the line starts with the word "global". The parameters set here may also be overridden by keywords in a specific line, in which case the keywords pertain only to that one line.

- **Regions**: all regions are considered by tasks. They will be displayed by visualization tasks as well as used to create masks, etc., as appropriate.

- **Annotations**: these are used by display tasks, and are for visual reference only.

D.1 Region definitions

All regions lines will follow this general arrangement:
{shape} {additional parameter=value pairs}

The possible parameter/value pairs are described in more detail below. Note that most parameters beyond the shape and its coordinates can be defined globally.

Possible units for coordinates are:

- *sexagesimal*, e.g. 18h12m24s for right ascension or -03.47.27.1 for declination
- *decimal degrees*, e.g. 140.0342deg for both RA and Dec
- *radians*, e.g. 2.37666rad for both RA and Dec
- *pixels*, e.g. 204pix

Possible units of length are:

- *degrees*, e.g. 23deg
- *arcminutes*, e.g. 23arcmin
- *arcseconds*, e.g. 23arcsec
- *radians*, e.g. 0.00035rad
- *pixels*, e.g. 23pix

*Units must always be included when defining a region.* If any fractional pixels are covered, the entire pixel will be selected. For purely single-pixel work there are alternate methods which may be preferable to using regions, eg. ia.topixel, ia.toworld, ia.pixelvalue.

### D.2 Allowed shapes

- **Rectangular box**: the two coordinates are two opposite corners:

  \[
  \text{box}[[x_1, y_1], [x_2, y_2]]
  \]

- **Center box**: \([x, y]\) define the center point of the box and \([x_{\text{width}}, y_{\text{width}}]\) the width of the sides:

  \[
  \text{centerbox}[[x, y], [x_{\text{width}}, y_{\text{width}}]]
  \]

- **Rotated box**: \([x, y]\) define the center point of the box; \([x_{\text{width}}, y_{\text{width}}]\) the width of the sides; rotang the rotation angle:

  \[
  \text{rotbox}[[x, y], [x_{\text{width}}, y_{\text{width}}], \text{rotang}]
  \]
• **Polygon**: there could be many \([x, y]\) corners; note that the last point will connect with the first point to close the polygon:

\[\text{poly}[[x_1, y_1], [x_2, y_2], [x_3, y_3], \ldots]\]

• **Circle**: center of the circle \([x, y]\), \(r\) is the radius:

\[\text{circle}[[x, y], r]\]

• **Annulus**: center of the circle is \([x, y]\), \([r_1, r_2]\) are inner and outer radii:

\[\text{annulus}[[x, y], [r_1, r_2]]\]

• **Ellipse**: center of the ellipse is \([x, y]\); semi-major and semi-minor axes are \([b_{maj}, b_{min}]\); position angle of the major axis is \(p_a\):

\[\text{ellipse}[[x, y], [b_{maj}, b_{min}], p_a]\]

### D.3 Annotation definitions

In addition to the definitions for regions [above], the following are always treated as annotations:

• **Line**: coordinates define the end points of the line:

\[\text{line}[[x_1, y_1], [x_2, y_2]]\]

• **Vector**: coordinates define end points; second coordinate pair is location of tip of arrow:

\[\text{vector}[[x_1, y_1], [x_2, y_2]]\]

• **Text**: coordinates define leftmost point of text string:

\[\text{text}[[x, y], 'my text']\]

• **Symbol**: coordinates define location of symbol (see Sec. D.7.1 for a list of allowed symbols):

\[\text{symbol}[[x, y], \{\text{symbol}\}]\]
D.4 Global definitions

Definitions to be used throughout the region file are placed on a line beginning with ‘global’, usually at the top of the file. These definitions may also be used on any individual region or annotation line; in this case, the value defined on that line will override the predefined global (but only for that line). If a ‘global’ line occurs later in the file, subsequent lines will obey those definitions.

- **Coordinate reference frame:**
  - Possible values: J2000, JMEAN, JTRUE, APP, B1950, B1950_VLA, BMEAN, BTRUE, GALACTIC, HADEC, AZEL, AZELSW, AZELNE, AZELGEO, AZELSWGEO, AZELNEGEO, JNAT, ECLIPTIC, MECLIPTIC, TECLIPTIC, SUPERGAL, ITRF, TOPO, ICRS
  - Default: image value

  \[\text{coord} = \text{J2000}\]

- **Frequency/velocity axis:**
  - Possible values: REST, LSRK, LSRD, BARY, GEO, TOPO, GALACTO, LGROUP, CMB
  - Default: image value

  \[\text{frame} = \text{TOPO}\]

- **Frequency/velocity range:**
  - Possible units: GHz, MHz, kHz, km/s, Hz, channel, chan (=channel)
  - Default: image range

  \[\text{range} = [\text{min}, \text{max}]\]

- **Correlation axis:**
  - Possible values: I, Q, U, V, RR, RL, LR, LL, XX, XY, YX, YY, RX, RY, LX, LY, XR, XL, YR, YL, PP, PQ, QP, QQ, RCircular, LCircular, Linear, Ptotal, Plinear, PFtotal, PFlinear, Pangle
  - Default: all planes present in image

  \[\text{corr} = [X, Y]\]

- **Velocity calculation:**
  - Possible values: RADIO, OPTICAL, Z, BETA, GAMMA
- Default: image value
  
  veltype=RADIO

- **Rest frequency:**
  
  - Default: image value
  
  restfreq=1.42GHz

- **Line characteristics:**
  
  - Possible values: any line style recognized by matplotlib: ’-’=solid, ’--’=dashed, ‘:’=dotted
  - Default: linewidth=1, linestyle=’-’

    linewidth=1
    linestyle=’-’

- **Symbol characteristics:**
  
  - Symbol size and thickness:

    symsize = 1
    symthick = 1

- **Region, symbol, and text color:**
  
  - Possible values: any color recognized by matplotlib, including hex values
  - Default: color=green

    color=red

- **Text font characteristics:**
  
  - Possible values: see Sect. [D.7.2]
  
  - 'usetex' is a boolean parameter that determines whether or not the text line should be interpreted as LaTeX, and would require working LaTeX, dvipng, and Ghostscript installations (equivalent to the text.usetex parameter in matplotlib).

    font=Helvetica
    fontsize=10pt
    fontstyle=bold
    usetex=True/False
• *Label position:*
  
  – Possible values: 'left', 'right', 'top', 'bottom'
  – Default: 'top'
    
    `labelpos='right'`

• *Label color:*
  
  – Default: color of associated region.
  – Allowed values: same as values for region colors.
    
    `labelcolor='green'`

• *Label offset:*
  
  – Default: [0,0].
  – Allowed values: any positive or negative number, in units of pixels.
    
    `labeloff=[1, 1]`

### D.5 Allowed additional parameters

These must be defined per region line:

• *Labels:* text label for a region; should be placed so text does not overlap with region boundary
  
  `label='string'`

• *"OR/NOT" operators:* A "+" at the beginning of a line will flag it with a boolean "OR" (default), and a "-" will flag it with a boolean "NOT". Overlapping regions will be treated according to their sequence in the file; i.e., (((entireImage OR line1) OR line2) NOT line3) OR line4). This allows some flexibility in building "non-standard" regions. Note that a task (e.g., clean) will still consider all lines: if one wishes to remove a region from consideration, it should be commented out ("#.
  
  • Default: OR (+)

### D.6 Examples

A file with both global definitions and per-line definitions:

```plaintext
#CRTFv0
global coord=B1950_VLA, frame=BARY, corr=[I, Q], color=blue
```
APPENDIX D. APPENDIX: CASA REGION FILE FORMAT

# A simple circle region:
circle[[18h12m24s, -23d11m00s], 2.3arcsec]

# A box region, this one only for annotation:
ann box[[140.0342deg, -12.34243deg], [140.0360deg, -12.34320deg]]

# A rotated box region, for a particular range of velocities:
rotbox[[12h01m34.1s, 12d23m33s], [3arcmin, 1arcmin], 12deg], range=[-1240km/s, 1240km/s]

# An annular region, overriding some of the global defaults:
annulus[[17h51m03.2s, -45d17m50s], [0.10deg, 4.12deg]], corr=[I,Q,U,V], color=red, label='My label here'

# Cuts an ellipse out of the previous regions, but only for Q and a particular frequency range:
-ellipse[[17:51:03.2, -45.17.50], [0.25deg, 1.34deg], 45rad], range=[1.420GHz, 1.421GHz], corr=[Q], color=green, label='Removed this'

# A diamond marker, in J2000 coordinates:
symbol[[32.1423deg, 12.1412deg], D], linewidth=2, coord=J2000, symsize=2

D.7 Fonts and Symbols

D.7.1 Allowed symbols

'.', point marker
',', pixel marker
'o', circle marker
'v', triangle_down marker
')', triangle_up marker
'<', triangle_left marker
'>', triangle_right marker
'1', tri_down marker
'2', tri_up marker
'3', tri_left marker
'4', tri_right marker
's', square marker
'p', pentagon marker
'*', star marker
'h', hexagon1 marker
'H', hexagon2 marker
'+', plus marker
'x', x marker
'D', diamond marker
d', thin_diamond marker
'l', vline marker
D.7.2 Allowed fonts

D.7.2.1 Allowed fonts for Linux


D.7.2.2 Allowed fonts for MacOS X

APPENDIX D. APPENDIX: CASA REGION FILE FORMAT

Appendix E

The Measurement Equation and Calibration

The visibilities measured by an interferometer must be calibrated before formation of an image. This is because the wavefronts received and processed by the observational hardware have been corrupted by a variety of effects. These include (but are not exclusive to): the effects of transmission through the atmosphere, the imperfect details amplified electronic (digital) signal and transmission through the signal processing system, and the effects of formation of the cross-power spectra by a correlator. Calibration is the process of reversing these effects to arrive at corrected visibilities which resemble as closely as possible the visibilities that would have been measured in vacuum by a perfect system. The subject of this chapter is the determination of these effects by using the visibility data itself.

E.1 The HBS Measurement Equation

The relationship between the observed and ideal (desired) visibilities on the baseline between antennas i and j may be expressed by the Hamaker-Bregman-Sault Measurement Equation:

\[ \vec{V}_{ij} = J_{ij} \vec{V}_{ij}^{\text{IDEAL}} \]

where \( \vec{V}_{ij} \) represents the observed visibility, \( \vec{V}_{ij}^{\text{IDEAL}} \) represents the corresponding ideal visibilities, and \( J_{ij} \) represents the accumulation of all corruptions affecting baseline \( ij \). The visibilities are indicated as vectors spanning the four correlation combinations which can be formed from dual-polarization signals. These four correlations are related directly to the Stokes parameters which fully describe the radiation. The \( J_{ij} \) term is therefore a 4×4 matrix.

Most of the effects contained in \( J_{ij} \) (indeed, the most important of them) are antenna-based, i.e., they arise from measurable physical properties of (or above) individual antenna elements in a synthesis array. Thus, adequate calibration of an array of \( N_{\text{ant}} \) antennas forming \( N_{\text{ant}}(N_{\text{ant}} - 1)/2 \) baseline visibilities is usually achieved through the determination of only \( N_{\text{ant}} \) factors, such that

\[ J_{ij} = J_i \otimes J_j^* \]. For the rest of this chapter, we will usually assume that \( J_{ij} \) is factorable in this way, unless otherwise noted.

As implied above, \( J_{ij} \) may also be factored into the sequence of specific corrupting effects, each having their own particular (relative) importance and physical origin, which determines their unique algebra. Including the most commonly considered effects, the Measurement Equation can be written:

\[ \vec{V}_{ij} = M_{ij} B_{ij} G_{ij} D_{ij} E_{ij} P_{ij} T_{ij} \vec{V}_{ij}^{\text{IDEAL}} \]

where:

- \( T_{ij} = \) Polarization-independent multiplicative effects introduced by the troposphere, such as opacity and path-length variation.
- \( P_{ij} = \) Parallactic angle, which describes the orientation of the polarization coordinates on the plane of the sky. This term varies according to the type of the antenna mount.
- \( E_{ij} = \) Effects introduced by properties of the optical components of the telescopes, such as the collecting area’s dependence on elevation.
- \( D_{ij} = \) Instrumental polarization response. “D-terms” describe the polarization leakage between feeds (e.g. how much the R-polarized feed picked up L-polarized emission, and vice versa).
- \( G_{ij} = \) Electronic gain response due to components in the signal path between the feed and the correlator. This complex gain term \( G_{ij} \) includes the scale factor for absolute flux density calibration, and may include phase and amplitude corrections due to changes in the atmosphere (in lieu of \( T_{ij} \)). These gains are polarization-dependent.
- \( B_{ij} = \) Bandpass (frequency-dependent) response, such as that introduced by spectral filters in the electronic transmission system
- \( M_{ij} = \) Baseline-based correlator (non-closing) errors. By definition, these are not factorable into antenna-based parts.

Note that the terms are listed in the order in which they affect the incoming wavefront (\( G \) and \( B \) represent an arbitrary sequence of such terms depending upon the details of the particular electronic system). Note that \( M \) differs from all of the rest in that it is not antenna-based, and thus not factorable into terms for each antenna.

As written above, the measurement equation is very general; not all observations will require treatment of all effects, depending upon the desired dynamic range. E.g., bandpass need only be considered for continuum observations if observed in a channelized mode and very high dynamic range is desired. Similarly, instrumental polarization calibration can usually be omitted when observing (only) total intensity using circular feeds. Ultimately, however, each of these effects occurs at some level, and a complete treatment will yield the most accurate calibration. Modern high-sensitivity instruments such as ALMA and JVLA will likely require a more general calibration.
treatment for similar observations with older arrays in order to reach the advertised dynamic ranges on strong sources.

In practice, it is usually far too difficult to adequately measure most calibration effects absolutely (as if in the laboratory) for use in calibration. The effects are usually far too changeable. Instead, the calibration is achieved by making observations of calibrator sources on the appropriate timescales for the relevant effects, and solving the measurement equation for them using the fact that we have \( N_{\text{ant}}(N_{\text{ant}} - 1)/2 \) measurements and only \( N_{\text{ant}} \) factors to determine (except for \( M \) which is only sparingly used). (Note: By partitioning the calibration factors into a series of consecutive effects, it might appear that the number of free parameters is some multiple of \( N_{\text{ant}} \), but the relative algebra and timescales of the different effects, as well as the multiplicity of observed polarizations and channels compensate, and it can be shown that the problem remains well-determined until, perhaps, the effects are direction-dependent within the field of view. Limited solvers for such effects are under study; the calibrator tool currently only handles effects which may be assumed constant within the field of view. Corrections for the primary beam are handled in the imager tool.) Once determined, these terms are used to correct the visibilities measured for the scientific target. This procedure is known as cross-calibration (when only phase is considered, it is called phase-referencing).

The best calibrators are point sources at the phase center (constant visibility amplitude, zero phase), with sufficient flux density to determine the calibration factors with adequate SNR on the relevant timescale. The primary gain calibrator must be sufficiently close to the target on the sky so that its observations sample the same atmospheric effects. A bandpass calibrator usually must be sufficiently strong (or observed with sufficient duration) to provide adequate per-channel sensitivity for a useful calibration. In practice, several calibrators are usually observed, each with properties suitable for one or more of the required calibrations.

Synthesis calibration is inherently a bootstrapping process. First, the dominant calibration term is determined, and then, using this result, more subtle effects are solved for, until the full set of required calibration terms is available for application to the target field. The solutions for each successive term are relative to the previous terms. Occasionally, when the several calibration terms are not sufficiently orthogonal, it is useful to re-solve for earlier types using the results for later types, in effect, reducing the effect of the later terms on the solution for earlier ones, and thus better isolating them. This idea is a generalization of the traditional concept of self-calibration, where initial imaging of the target source supplies the visibility model for a re-solve of the gain calibration (\( G \) or \( T \)). Iteration tends toward convergence to a statistically optimal image. In general, the quality of each calibration and of the source model are mutually dependent. In principle, as long as the solution for any calibration component (or the source model itself) is likely to improve substantially through the use of new information (provided by other improved solutions), it is worthwhile to continue this process.

In practice, these concepts motivate certain patterns of calibration for different types of observation, and the calibrator tool in CASA is designed to accommodate these patterns in a general and flexible manner. For a spectral line total intensity observation, the pattern is usually:

1. Solve for \( G \) on the bandpass calibrator
2. Solve for \( B \) on the bandpass calibrator, using \( G \)
3. Solve for $G$ on the primary gain (near-target) and flux density calibrators, using $B$ solutions just obtained.

4. Scale $G$ solutions for the primary gain calibrator according to the flux density calibrator solutions.

5. Apply $G$ and $B$ solutions to the target data.

6. Image the calibrated target data.

If opacity and gain curve information are relevant and available, these types are incorporated in each of the steps (in future, an actual solve for opacity from appropriate data may be folded into this process):

1. Solve for $G$ on the bandpass calibrator, using $T$ (opacity) and $E$ (gain curve) solutions already derived.

2. Solve for $B$ on the bandpass calibrator, using $G$, $T$ (opacity), and $E$ (gain curve) solutions.

3. Solve for $G$ on primary gain (near-target) and flux density calibrators, using $B$, $T$ (opacity), and $E$ (gain curve) solutions.

4. Scale $G$ solutions for the primary gain calibrator according to the flux density calibrator solutions.

5. Apply $T$ (opacity), $E$ (gain curve), $G$, and $B$ solutions to the target data.

6. Image the calibrated target data.

For continuum polarimetry, the typical pattern is:

1. Solve for $G$ on the polarization calibrator, using (analytical) $P$ solutions.


3. Solve for $G$ on primary gain and flux density calibrators, using $P$ and $D$ solutions.

4. Scale $G$ solutions for the primary gain calibrator according to the flux density calibrator solutions.

5. Apply $P$, $D$, and $G$ solutions to target data.

6. Image the calibrated target data.

For a spectro-polarimetry observation, these two examples would be folded together.

In all cases the calibrator model must be adequate at each solve step. At high dynamic range and/or high resolution, many calibrators which are nominally assumed to be point sources become slightly resolved. If this has biased the calibration solutions, the offending calibrator may be imaged at any point in the process and the resulting model used to improve the calibration. Finally, if sufficiently strong, the target may be self-calibrated as well.
E.2 General Calibrater Mechanics

The calibrater tasks/tool are designed to solve and apply solutions for all of the solution types listed above (and more are in the works). This leads to a single basic sequence of execution for all solves, regardless of type:

1. Set the calibrator model visibilities
2. Select the visibility data which will be used to solve for a calibration type
3. Arrange to apply any already-known calibration types (the first time through, none may yet be available)
4. Arrange to solve for a specific calibration type, including specification of the solution timescale and other specifics
5. Execute the solve process
6. Repeat 1-4 for all required types, using each result, as it becomes available, in step 2, and perhaps repeating for some types to improve the solutions

By itself, this sequence doesn’t guarantee success; the data provided for the solve must have sufficient SNR on the appropriate timescale, and must provide sufficient leverage for the solution (e.g., D solutions require data taken over a sufficient range of parallactic angle in order to separate the source polarization contribution from the instrumental polarization).
Appendix F

Data Weights

F.1 Introduction

Visibility weight initialization and calibration has undergone several improvements in CASA 4.2.2 and (pending) CASA 4.3. This appendix briefly describes the formal weight definitions, and the changes occurring in these CASA versions. If data sets shall be combined that were reduced with different CASA versions, the weights may need to be adjusted accordingly. This can be achieved, e.g. by running the same version of `statwt` (§ 4.7.9) on all datasets before combination. The best option, however, is to use a single CASA version for all reductions, preferably 4.2.2 or later.

Note that post-calibration weights, e.g. imaging weights or tapers are not covered by this appendix.

F.2 SIGMA and WEIGHT columns

Formally, in CASA 4.2.2 and later, the SIGMA column in the measurement set will reflect the per-channel noise of the DATA as it depends on the channel bandwidth $\Delta \nu$ and the length of an integration $\Delta t$:

$$\text{SIGMA} = \frac{1}{\sqrt{2\Delta \nu \Delta t}}. \quad \text{(F.1)}$$

The factor of $\sqrt{2}$ is for cross-correlation only and auto-correlation data follows $\text{SIGMA} = \frac{1}{\sqrt{\Delta \nu \Delta t}}$.

SIGMA will only be updated if the time and channel widths are modified along with any DATA column manipulation, e.g. through averaging, binning, smoothing, etc. (tasks like `mstransform`, `cvel`, `split`, `exportuvfits`,...).

The WEIGHT column reflects how much weight each CORRECTED_DATA sample should receive when data are combined (e.g., in averaging). To start with, WEIGHT is initialized from the SIGMA column via:
Table F.1: Antenna-based WEIGHT calibration factor definitions for different CASA versions. For System Temperator and Bandpass, \( k \) is the channel index. ALMA has channelized \( T_{\text{sys}} \); EVLA does not.

\[
\text{WEIGHT} = \frac{1}{\text{SIGMA}^2} = 2\Delta \nu \Delta t \quad (\text{F.2})
\]

Data calibration by \texttt{applycal (§4.6.1)} with \texttt{calwt=T} will calculate and modify the WEIGHT values but not SIGMA. Calibration applies multiplicative factors and the WEIGHT of a visibility on a baseline between antennas \( i \) and \( j \) is calculated via

\[
\text{WEIGHT}_{ij} = \frac{\omega_i \omega_j}{\text{SIGMA}_{ij}^2} \quad (\text{F.3})
\]

where \( \omega_i \) and \( \omega_j \) are the antenna-based calibration factors derived by \texttt{applycal} (\( \omega_i = \omega_j \) for auto-correlation data). In Table F.1 we list the definitions of \textit{antenna-based} \( \omega \) for different calibration procedures and CASA versions. When more than one calibration is applied, the product of the relevant weight factors is used.

### F.2.1 Weights in CASA 4.2.1 and Earlier

The SIGMA and WEIGHT columns are initialized with values of “1”. Traditionally, this convention was adequate for datasets with uniform sampling in time in frequency; a global weight scale factor would not affect calibration and imaging fidelity. In data manipulation operations (e.g., \texttt{split}, etc.), SIGMA was treated as a per-channel value and WEIGHT as a per-spw (all channels) weight. Combined with unit initialization, this difference in definition could lead to incongruent weight scales for different spectral windows, in particular if bandwidth and channel count varied. CASA 4.2.1 is \textit{not} recommended for datasets which have variety in spectral window bandwidth and channelization and for which spectral windows are to be combined in imaging.
APPENDIX F. APPENDIX: DATA WEIGHTS

F.2.2 Weights in CASA 4.2.2

In CASA 4.2.2 the SIGMA and WEIGHT columns are properly initialized via the definition in Eqs. F.1 and F.2. Both are defined as per-channel values. Also, the weight calibration factors have been subtly updated to improve robustness, as indicated in Table F.1.

F.2.3 Weights in CASA 4.3

In CASA 4.3 frequency variations of the WEIGHT and SIGMA values are (optionally) captured in additional WEIGHT.SPECTRUM and SIGMA.SPECTRUM columns. This allows accommodation of variations of effective sensitivity on a channel by channel basis (e.g. band edges, atmospheric lines, spectral $T_{\text{sys}}$ variation etc.). WEIGHT.SPECTRUM will be recognized in the applycal task as well as in mstransform and clean. Calibration solvers, however, will not yet calculate and modify WEIGHT.SPECTRUM.

F.2.4 Weights in CASA 4.4 and later

Full support of WEIGHT.SPECTRUM. Also, statwt will be enhanced to permit calculating weights on a channel-dependent basis.
Appendix G

Annotated Example Scripts

The annotated scripts are available on the CASA homepage:

http://casa.nrao.edu/casa_scripts.shtml

These will be removed shortly, however, as the casaguides contain newer version of data reduction tutorials:

http://casaguides.nrao.edu
Appendix H

CASA Dictionaries

BETA ALERT: These tend to become out of date as we add new tasks or change names.

H.1 AIPS – CASA dictionary

In Table H.1 we provide a comparison of CASA and AIPS commands. The data reduction recipes and break-down of jobs in individual tasks, however, is not the same in both packages. Nevertheless, the table may give AIPS users a good start if they search for functionality in CASA.

H.2 MIRIAD – CASA dictionary

Table H.2 provides a list of common Miriad tasks, and their equivalent CASA tool or tool function names. The two packages differ in both their architecture and calibration and imaging models, and there is often not a direct correspondence. However, this index does provide a scientific user of CASA who is familiar with MIRIAD, with a simple translation table to map their existing data reduction knowledge to the new package.

H.3 CLIC – CASA dictionary

Table H.3 provides a list of common CLIC tasks, and their equivalent CASA tool or tool function names. The two packages are very similar since the CASA software to reduce IRAM data is based on the CLIC reduction procedures.
## Table H.1: AIPS – CASA dictionary

<table>
<thead>
<tr>
<th>AIPS Task</th>
<th>CASA task/tool</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>APROPOS</td>
<td>taskhelp</td>
<td>List tasks with a short description of their purposes</td>
</tr>
<tr>
<td>BLCAL</td>
<td>bical</td>
<td>Calculate a baseline-based gain calibration solution</td>
</tr>
<tr>
<td>BLCCHN</td>
<td>bical</td>
<td>Calculate a baseline-based bandpass calibration solution</td>
</tr>
<tr>
<td>BPASS</td>
<td>bandpass</td>
<td>Calibrate bandpasses</td>
</tr>
<tr>
<td>CALIB</td>
<td>gaincal</td>
<td>Calibrate gains (amplitudes and phases)</td>
</tr>
<tr>
<td>CLCAL</td>
<td>applycal</td>
<td>Apply calibration to data</td>
</tr>
<tr>
<td>COMB</td>
<td>immath</td>
<td>Combine images</td>
</tr>
<tr>
<td>CPASS</td>
<td>bandpass</td>
<td>Calibrate bandpasses by polynomial fitting</td>
</tr>
<tr>
<td>DBCON</td>
<td>concat</td>
<td>Concatenate u-v datasets</td>
</tr>
<tr>
<td>DEFAULT</td>
<td>default</td>
<td>Load a task with default parameters</td>
</tr>
<tr>
<td>FILLM</td>
<td>importvla</td>
<td>Import old-format VLA data</td>
</tr>
<tr>
<td>FITLD</td>
<td>importuvfits</td>
<td>Import a u-v dataset which is in FITS format</td>
</tr>
<tr>
<td>FITLD</td>
<td>importfits</td>
<td>Import an image which is in FITS format</td>
</tr>
<tr>
<td>FITTP</td>
<td>exportuvfits</td>
<td>Write a u-v dataset to FITS format</td>
</tr>
<tr>
<td>FITTP</td>
<td>exportfits</td>
<td>Write an image to FITS format</td>
</tr>
<tr>
<td>FRING</td>
<td>—</td>
<td>Calibrate group delays and phase rates.</td>
</tr>
<tr>
<td>GETJY</td>
<td>fluxscale</td>
<td>Determine flux densities for other cals</td>
</tr>
<tr>
<td>GO</td>
<td>go</td>
<td>Run a task</td>
</tr>
<tr>
<td>HELP</td>
<td>help</td>
<td>Display the help page for a task</td>
</tr>
<tr>
<td>IMAGR</td>
<td>clean</td>
<td>Image and deconvolve</td>
</tr>
<tr>
<td>IMFIT</td>
<td>imfit</td>
<td>Fit gaussian components to an image</td>
</tr>
<tr>
<td>IMHEAD</td>
<td>vishead</td>
<td>View header for u-v data</td>
</tr>
<tr>
<td>IMHEAD</td>
<td>imhead</td>
<td>View header for an image</td>
</tr>
<tr>
<td>IMLIN</td>
<td>imcontsub</td>
<td>Subtract continuum in image plane</td>
</tr>
<tr>
<td>IMSTAT</td>
<td>importfits</td>
<td>Import a FITS image</td>
</tr>
<tr>
<td>INF</td>
<td>inp</td>
<td>Measure statistics on an image</td>
</tr>
<tr>
<td>JMFIT</td>
<td>imfit</td>
<td>View task parameters</td>
</tr>
<tr>
<td>LISTR</td>
<td>listobs</td>
<td>Fit gaussian components to an image</td>
</tr>
<tr>
<td>MCAT</td>
<td>ls</td>
<td>Print basic data</td>
</tr>
<tr>
<td>MOMNT</td>
<td>immoments</td>
<td>List image data files</td>
</tr>
<tr>
<td>OHGEO</td>
<td>imregrid</td>
<td>Compute moments from an image</td>
</tr>
<tr>
<td>PBCOR</td>
<td>immath</td>
<td>Regrids an image onto another image’s geometry</td>
</tr>
<tr>
<td>PCAL</td>
<td>polcal</td>
<td>Correct an image for the primary beam</td>
</tr>
<tr>
<td>POSSM</td>
<td>plotcal</td>
<td>Calibrate polarization</td>
</tr>
<tr>
<td>POSSM</td>
<td>plotms</td>
<td>Plot bandpass calibration tables</td>
</tr>
<tr>
<td>PRTAN</td>
<td>listobs</td>
<td>Plot spectra</td>
</tr>
<tr>
<td>PRTAN</td>
<td>plotants</td>
<td>Print antenna locations</td>
</tr>
<tr>
<td>QUACK</td>
<td>flagdata</td>
<td>Plot antenna locations</td>
</tr>
<tr>
<td>RENAME</td>
<td>mv</td>
<td>Remove first integrations from scans</td>
</tr>
<tr>
<td>SETJY</td>
<td>setjy</td>
<td>Rename an image or dataset</td>
</tr>
<tr>
<td>SMOOTH</td>
<td>imsmooth</td>
<td>Set flux densities for flux cals</td>
</tr>
<tr>
<td>SNPLT</td>
<td>plotcl</td>
<td>Smooth an image</td>
</tr>
<tr>
<td>SPFLG</td>
<td>viewer</td>
<td>Plot gain calibration tables</td>
</tr>
<tr>
<td>SPLIT</td>
<td>split</td>
<td>Flag raster image of time v. channel</td>
</tr>
<tr>
<td>TASK</td>
<td>inp</td>
<td>Write out u-v files for individual sources</td>
</tr>
<tr>
<td>TGET</td>
<td>tget</td>
<td>Load a task with current parameters</td>
</tr>
<tr>
<td>TVALL</td>
<td>viewer</td>
<td>Load a task with parameters last used for that task</td>
</tr>
<tr>
<td>TVFLG</td>
<td>viewer</td>
<td>Display image</td>
</tr>
<tr>
<td>UCAT</td>
<td>ls</td>
<td>Flag raster image of time v. baseline</td>
</tr>
<tr>
<td>UFIX</td>
<td>fixvis</td>
<td>List u-v data files</td>
</tr>
<tr>
<td>UVFLG</td>
<td>flagdata</td>
<td>Compute u, v, and w coordinates</td>
</tr>
<tr>
<td>UVLIN</td>
<td>uvcontsub</td>
<td>Flag data</td>
</tr>
<tr>
<td>UVLSF</td>
<td>uvcontsub</td>
<td>Subtract continuum from u-v data</td>
</tr>
<tr>
<td>UVPLOT</td>
<td>plotms</td>
<td>Subtract continuum from u-v data</td>
</tr>
<tr>
<td>UVSUB</td>
<td>uvsms</td>
<td>Plot u-v data</td>
</tr>
<tr>
<td>WIPER</td>
<td>plotms</td>
<td>Subtracts model u-v data from corrected u-v data</td>
</tr>
<tr>
<td>ZAP</td>
<td>rmtables</td>
<td>Plot and flag u-v data</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Delete data files</td>
</tr>
</tbody>
</table>
### Table H.2: MIRIAD – CASA dictionary

<table>
<thead>
<tr>
<th>MIRIAD Task</th>
<th>Description</th>
<th>CASA task/tool</th>
</tr>
</thead>
<tbody>
<tr>
<td>blflag</td>
<td>Interactive baseline based editor/flagger</td>
<td>mp raster displays</td>
</tr>
<tr>
<td>egcurs</td>
<td>Interactive image analysis</td>
<td>viewer</td>
</tr>
<tr>
<td>egdisp</td>
<td>Image display, overlays</td>
<td>viewer</td>
</tr>
<tr>
<td>clean</td>
<td>Clean an image</td>
<td>clean</td>
</tr>
<tr>
<td>fits</td>
<td>FITS image filler</td>
<td>importfits, exportfits, importuvfits, exportuvfits</td>
</tr>
<tr>
<td>gpbolt</td>
<td>Set flux density scale</td>
<td>fluxscale</td>
</tr>
<tr>
<td>gpical</td>
<td>Polarization leakage and gain calibration</td>
<td>gaincal</td>
</tr>
<tr>
<td>gpcopy</td>
<td>copy calibration tables</td>
<td>not needed</td>
</tr>
<tr>
<td>gpplt</td>
<td>Plot calibration solutions</td>
<td>plotcal</td>
</tr>
<tr>
<td>imcomb</td>
<td>Image combination</td>
<td>immaths</td>
</tr>
<tr>
<td>imfit</td>
<td>Image-plane component fitter</td>
<td>imfit</td>
</tr>
<tr>
<td>impol</td>
<td>Create polarization images</td>
<td>clean</td>
</tr>
<tr>
<td>instat</td>
<td>Image statistics</td>
<td>instats</td>
</tr>
<tr>
<td>imsub</td>
<td>Extract sub-image</td>
<td>ia.subimage</td>
</tr>
<tr>
<td>invert</td>
<td>Synthesis imaging</td>
<td>clean</td>
</tr>
<tr>
<td>linmos</td>
<td>linear mosaic combination of images</td>
<td>clean</td>
</tr>
<tr>
<td>maths</td>
<td>Calculations involving images</td>
<td>immath</td>
</tr>
<tr>
<td>mfcal</td>
<td>Bandpass and gain calibration</td>
<td>bandpass</td>
</tr>
<tr>
<td>prthld</td>
<td>Print header of image or uvdata</td>
<td>imhead, listobs, vishead</td>
</tr>
<tr>
<td>restor</td>
<td>Restore a clean component model</td>
<td>clean</td>
</tr>
<tr>
<td>selical</td>
<td>selfcalibration of visibility data</td>
<td>clean, gaincal, etc.</td>
</tr>
</tbody>
</table>

### Table H.3: CLIC–CASA dictionary

<table>
<thead>
<tr>
<th>CLIC Function</th>
<th>Description</th>
<th>CASA task/tool</th>
</tr>
</thead>
<tbody>
<tr>
<td>load</td>
<td>Load data</td>
<td>importfits, importasdm, importuvfits</td>
</tr>
<tr>
<td>print</td>
<td>Print text summary of data</td>
<td>listobs</td>
</tr>
<tr>
<td>flag</td>
<td>Flag data</td>
<td>plotms, flagdata, viewer</td>
</tr>
<tr>
<td>phcor</td>
<td>Atmospheric phase correction</td>
<td>gaincal</td>
</tr>
<tr>
<td>rf</td>
<td>Radio frequency bandpass</td>
<td>bandpass</td>
</tr>
<tr>
<td>phase</td>
<td>Phase calibration</td>
<td>gaincal</td>
</tr>
<tr>
<td>flux</td>
<td>Absolute flux calibration</td>
<td>setjy, fluxscale</td>
</tr>
<tr>
<td>ampl</td>
<td>Amplitude calibration</td>
<td>gaincal</td>
</tr>
<tr>
<td>table</td>
<td>Split out calibrated data (uv table)</td>
<td>split</td>
</tr>
</tbody>
</table>
Appendix I

Writing Tasks

ALERT: This prescription for writing and incorporating tasks in CASA is for the power-user. This procedure is also likely to change in future releases.

It is possible to write your own task and have it appear in casapy. For example, if you want to create a task named “yourtask”, then must create two files, yourtask.xml and a task.yourtask.py. The .xml file is use to describe the interface to the task and the task.yourtask.py does the actual work. The argument names must be the same in both the yourtask.xml and task.yourtask.py file. The yourtask.xml file is used to generate all the interface files so yourtask will appear in the casapy system. It is easiest to start from one of the existing tasks when constructing these. You would make the name of the function in the yourtask.py be “yourtask” in this example.

We have provided the buildmytasks command in order to assemble your Python and XML into a loadable Python file. Thus, the steps you need to execute (again for an example task named “yourtask”):

- Create python code for task as task.yourtask.py
- Create xml for task as yourtask.xml
- Execute buildmytasks from the casapy prompt: !buildmytasks
- Initialize your new task inside casapy: execfile 'mytasks.py'

After this, you should see the help and inputs inside casapy, e.g. inp yourtask should work. Note that for the final step you invoke the file called mytasks.py, regardless of what you named the actual task. You now have a shiny new task yourtask that you can run and use in the same way as all other CASA tasks.

Note that if multiple custom tasks are stored in the same directory, they will all be built by !buildmytasks and will all be initialized by executing mytasks.py. To build and initialize only a single task, instead use ‘!buildmytasks taskname; you are then free to rename mytasks.py (e.g. load_taskname.py) and repeat this procedure for your other tasks. Our recommendation, for those of you who are managing multiple custom tasks, is to have each task live in its own directory.
mytasks.py file need not be in the current working directory to initialize your task, since you can provide the full path upon initialization (e.g. `execfile '/full_path_to_my_task/mytasks.py'`).

### I.1 The XML file

The key to getting your task into casapy is constructing a task interface description XML file.

Some XML basics, an xml element begins with `<element>` and ends with `</element>`. If an XML element contains no other XML element you may specify it via `<element/>`. An XML element may have zero or more attributes which are specified by attribute="attribute value". You must put the attribute value in quotes, i.e. `<element myattribute="attribute value">`.

All task xml files must start with this header information.

```xml
<?xml version="1.0" encoding="UTF-8"?>
<?xml-stylesheet type="text/xsl" ?>
<casaxml xmlns="http://casa.nrao.edu/schema/psetTypes.html"
xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance"
xsi:schemaLocation="http://casa.nrao.edu/schema/casa.xsd
file:///opt/casa/code/xmlcasa/xml/casa.xsd">

and the file must have the end tag

```xml
</casaxml>
```

Inside a `<task>` tags you will need to specify the following elements.

```xml
<task>
  Attributes
  type required, allowed value is "function"
  name required

  Subelements
  shortdescription required
  description required
  input optional
  output optional
  returns optional
  constraints optional

  <shortdescription> - required by <task>; A short one-line description describing your task

  Attributes
```
None
Subelements
None

<description> - required] by <task>, Also used by <param>a; A longer description describing your task with multiple lines

Attributes
None
Subelements
None

<input> - optional element used by <task>; An input block specifies which parameters are used for input

Attributes
None
Subelements
<param>, optional

<output> - optional  An output element that contains a list of parameters that are "returned" by the task.

Attributes
None
Subelements
<param>, optional

<returns> - optional Value returned by the task

Attributes

type optional; as specified in <param>

Subelements
<description>, optional

<constraints> - optional A constraints element that lets you constrain params based on the values of other params.

Attributes
None
Subelements
<when>, required.

<param> - optional The input and output elements consist of param elements.
Attributes

- **type**, required; allowed values are record, variant, string, int, double, bool, intArray, doubleArray, boolArray, stringArray
- **name**, required;
- **subparam**, optional; allowed values True, False, Yes or No.
- **kind**, optional;
- **mustexist**, optional; allowed values True, False, Yes or No.

All param elements require name and type attributes.

Subelements

- **<description>**, required;
- **<value>**, optional;
- **<allowed>**, optional;

**<value>** - optional Value returned by the task

Attributes

- **type**, required; as specified in **<param>** attributes.

Subelements

- **<value>**, optional

**<allowed>** - optional; Block of allowed values

Attributes

- **enum**, required; maybe enum or range. If specified as enum only specific values are allowed. If specified as range then the value tags may have min and max attributes.

Subelements

- **<value>**, optional

**<when>** - optional When blocks allow value specific handling for parameters

Attributes

- **param**, required; Specifies special handling for a **<param>**

Subelements

- **<equals>**, optional
- **<notequals>**, optional

**<equals>** - optional Reset parameters if equal to the specified value

Attributes

- **value**, required; the value of the parameter

Subelements

- **<default>**, required
\verb+<notequals> - optional+  Reset specified parameters if not equal to the specified value

\verb+Attributes+  
\verb+value , required; The value of the parameter+  
\verb+Subelements+  
\verb+<default> , optional+  

\verb+<default> - optional+  Resets default values for specified parameters

\verb+Attributes+  
\verb+param , required; Name of the <param> to be reset. +  
\verb+Subelements+  
\verb+<value> , required, the revised value of the <param>.+  

\verb+<example> - optional+  An example block, typically in python

\verb+Attributes+  
\verb+lang optional; specifies the language of the example, defaults to python. +  
\verb+Subelements+  
\verb+None+  

\section*{I.2 The \texttt{task.yourtask.py} file}

You must write the python code that does the actual work. The \texttt{task.*.py} file function call sequence must be the same as specified in the XML file. We may relax the requirement that the function call sequence exactly match the sequence in the XML file in a future release.

The \texttt{task.*.py} file should contain the following preamble

\begin{verbatim}
import os
from taskinit import *

plus any other global function imports you will need such as

import time

followed by the task function \texttt{def}. See Sect. I.3.2 for an example.

\section*{I.3 Example: The \texttt{clean} task}

Note that the following is for illustration only and does \textit{not} reflect the current implementation of \texttt{clean} – a task that is always developing further on a quick pace.
I.3.1 File clean.xml

Clean.xml gives a fairly comprehensive example of how to construct the XML file.

```xml
<?xml version="1.0" encoding="UTF-8"?>
<?xml-stylesheet type="text/xsl" ?>
<casaxml xmlns="http://casa.nrao.edu/schema/psetTypes.html"
xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance"
xsi:schemaLocation="http://casa.nrao.edu/schema/casa.xsd
file:///opt/casa/code/xmlcasa/xml/casa.xsd">

<!-- This is the param set for clean -->
<!-- This does the equivalent of -->
<!-- imgr:=imager('anyfile.ms'); -->
<!-- imgr.setdata(mode='channel',nchan=100,start=1,step=1,fieldid=1) -->
<!-- imgr.setimage(nx=512,ny=,cellx='1arcsec',celly='1arcsec',stokes='I',-->
<!-- mode='channel',start=35,step=1,nchan=40, -->
<!-- fieldid=[1]) -->
<!-- imgr.weight('natural'); -->
<!-- imgr.clean(algorithm='csclean',niter=500,model='field1') -->

<task type="function" name="clean">
  <shortdescription>Deconvolve an image with selected algorithm</shortdescription>
  <description>
  Form images from visibilities. Handles continuum and spectral line cubes.
  </description>

  <input>
    <param type="string" name="vis" kind="ms" mustexist="true">
      <description>name of input visibility file</description>
      <value></value>
    </param>
    <param type="string" name="imagename">
      <description>Pre-name of output images</description>
      <value></value>
    </param>
    <param type="string" name="field">
      <description>Field Name</description>
      <value></value>
    </param>

</task>
</casaxml>
```
<param type="any" name="spw">
   <description>Spectral windows:channels: \\
   is all </description>
   <any type="variant"></value>
   <value type="string"></value>
</param>

<param type="bool" name="selectdata">
   <description>Other data selection parameters</description>
   <value>False</value>
</param>

<param type="string" name="timerange" subparam="true">
   <description>Range of time to select from data</description>
   <value></value>
</param>

<param type="string" name="uvrange" subparam="true">
   <description>Select data within uvrange</description>
   <value></value>
</param>

<param type="string" name="antenna" subparam="true">
   <description>Select data based on antenna/baseline</description>
   <value></value>
</param>

<param type="string" name="scan" subparam="true">
   <description>Scan number range</description>
   <value></value>
</param>

<param type="string" name="mode">
   <description>Type of selection (mfs, channel, velocity, frequency)</description>
   <value>mfs</value>
   <allowed kind="enum">
      <value>mfs</value>
      <value>channel</value>
      <value>velocity</value>
      <value>frequency</value>
   </allowed>
</param>
<param type="int" name="niter">
<description>Maximum number of iterations</description>
[value]500[/value]
</param>

<param type="double" name="gain">
<description>Loop gain for cleaning</description>
[value]0.1[/value]
</param>

<param type="double" name="threshold" units="mJy">
<description>Flux level to stop cleaning. Must include units</description>
[value]0.0[/value]
</param>

<!-- Getting rid of this
<param type="bool" name="csclean">
<description>Use Cotton-Schwab style reconciliation with UV-data</description>
[value]False[/value]
</param>
-->

<param type="string" name="psfmode">
<description>method of PSF calculation to use during minor cycles</description>
[value]clark[/value]
<allowed kind="enum">
[value]clark[/value]
[value]hogbom[/value]
</allowed>
</param>

<param type="string" name="imagermode">
<description>Use csclean or mosaic. If "/", use psfmode</description>
[value][/value]
<allowed kind="enum">
[value][/value]
[value]csclean[/value]
[value]mosaic[/value]
</allowed>
</param>

<param type="string" name="ftmachine" subparam="true"/>
<description>Gridding method for the image</description>
<value>mosaic</value>
<allowed kind="enum">
  <value>mosaic</value>
  <value>ft</value>
  <value>sd</value>
  <value>both</value>
</allowed>

</param>

<param type="bool" name="mosweight" subparam="true">
  <description>Individually weight the fields of the mosaic</description>
  <value>False</value>
</param>

<param type="string" name="scaletype" subparam="true">
  <description>Controls scaling of pixels in the image plane.
    default='SAULT';
    example: scaletype='PBCOR'
    Options: 'PBCOR', 'SAULT'</description>
  <value>SAULT</value>
  <allowed kind="enum">
    <value>SAULT</value>
    <value>PBCOR</value>
  </allowed>
</param>

<param type="intArray" name="multiscale">
  <description>set deconvolution scales (pixels),
    default: multiscale=[] (standard CLEAN)</description>
  <value type="vector">
    <value></value>
  </value>
</param>

<param type="int" name="negcomponent" subparam="true">
  <description>
    Stop cleaning if the largest scale finds this number of neg components
  </description>
  <value>0</value>
</param>

<param type="bool" name="interactive">
  <description>use interactive clean (with GUI viewer)</description>
  <value>False</value>
</param>
</param>

<param type="any" name="mask">
  <description>cleanbox(es), mask image(s), and/or region(s) used in cleaning</description>
  <any type="variant"/>
  <value type="stringArray"></value>
</param>

<param type="int" name="nchan" subparam="true">
  <description>Number of channels (planes) in output image</description>
  <value>1</value>
</param>

<param type="any" name="start" subparam="true">
  <description>First channel in input to use</description>
  <any type="variant"/>
  <value type="int">0</value>
</param>

<param type="any" name="width" subparam="true">
  <description>Number of input channels to average</description>
  <any type="variant"/>
  <value type="int">1</value>
</param>

<param type="intArray" name="imsize">
  <description>x and y image size in pixels, symmetric for single value</description>
  <value type="vector">
    <value>256</value>
    <value>256</value>
  </value>
</param>

<param type="doubleArray" name="cell" units="arcsec">
  <description>x and y cell size. default unit arcsec</description>
  <value type="vector">
    <value>1.0</value>
    <value>1.0</value>
  </value>
</param>

<param type="any" name="phasecenter">
  <description>Image phase center: position or field index</description>
  <any type="variant"/>
  <value type="string"></value>
</param>
<param type="string" name="restfreq">
    <description>rest frequency to assign to image (see help)</description>
    <value></value>
</param>

<param type="string" name="stokes">
    <description>Stokes params to image (eg I, IV, QU, IQUV)</description>
    <value>I</value>
    <allowed kind="enum">
        <value>I</value>
        <value>IV</value>
        <value>QU</value>
        <value>IQUV</value>
        <value>RR</value>
        <value>LL</value>
        <value>RRLL</value>
        <value>XX</value>
        <value>YY</value>
        <value>XXYY</value>
    </allowed>
</param>

<param type="string" name="weighting">
    <description>Weighting to apply to visibilities</description>
    <value>natural</value>
    <allowed kind="enum">
        <value>natural</value>
        <value>uniform</value>
        <value>briggs</value>
        <value>briggsabs</value>
        <value>radial</value>
        <value>superuniform</value>
    </allowed>
</param>

<param type="double" name="robust" subparam='true'>
    <description>Briggs robustness parameter</description>
    <value>0.0</value>
</param>
<param type="bool" name="uvtaper">
  <description>Apply additional uv tapering of visibilities.</description>
  <value>False</value>
</param>

<param type="stringArray" name="outertaper" subparam="true">
  <description>uv-taper on outer baselines in uv-plane</description>
  <value type="vector">
    <value></value>
  </value>
</param>

<param type="stringArray" name="innertaper" subparam="true">
  <description>uv-taper in center of uv-plane</description>
  <value>1.0</value>
</param>

<param type="string" name="modelimage">
  <description>Name of model image(s) to initialize cleaning</description>
  <value></value>
</param>

<param type="stringArray" name="restoringbeam">
  <description>Output Gaussian restoring beam for CLEAN image</description>
  <value></value>
</param>

<param type="bool" name="pbcor">
  <description>Output primary beam-corrected image</description>
  <value>False</value>
</param>

<param type="double" name="minpb">
  <description>Minimum PB level to use</description>
  <value>0.1</value>
</param>

<param type="any" name="noise" subparam='true'
  <description>noise parameter for briggs abs mode weighting</description>
  <any type="variant"/>
<param type="int" name="npercycle" subparam='true'>
   <description>Number of iterations before interactive prompt</description>
   <value>100</value>
</param>

<param type="double" name="cyclefactor" subparam='true'>
   <description>Change depth in between of csclean cycle</description>
   <value>1.5</value>
</param>

<param type="int" name="cyclespeedup" subparam='true'>
   <description>Cycle threshold doubles in this number of iteration</description>
   <value>-1</value>
</param>

<constraints>
   <when param="selectdata">
      <equals type="bool" value="False"/>
      <equals type="bool" value="True">
         <default param="timerange"><value type="string"></value>
      </default>
      <default param="uvrange"><value type="string"></value>
      <default param="antenna"><value type="string"></value>
      <default param="scan"><value type="string"></value>
   </equals>
</when>

   <when param="multiscale">
      <notequals type="vector" value="[]" >
         <default param="negcomponent"><value>-1</value>
      </default>
   </when>
</constraints>
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<notequals>

</when>

<when param="mode">
  <equals value="mfs"/>
  <equals value="channel">
    <default param="nchan"><value>1</value></default>
    <default param="start"><value>0</value>
      <description>first input channel to use</description>
    </default>
    <default param="width"><value>1</value></default>
  </equals>
</when>

<when param="weighting">
  <equals value="natural"/>
  <equals value="uniform"/>
  <equals value="briggs">
    <default param="robust"><value>0.0</value></default>
    <default param="npixels"><value>0</value>
      <description>number of pixels to determine uv-cell size
      0=&gt; field of view</description>
    </default>
  </equals>
</when>
<equals value="briggsabs">
  <default param="robust"><value>0.0</value></default>
  <default param="noise"><value type="string">1.0Jy</value></default>
  <default param="npixels"><value>0</value>
    <description>number of pixels to determine uv-cell size</description>
    0=&gt; field of view
  </default>
</equals>

<equals value="superuniform">
  <default param="npixels"><value>0</value>
    <description>number of pixels to determine uv-cell size</description>
    0=&gt; +/-3pixels
  </default>
</equals>

<when param="uvtaper">
  <equals type="bool" value="False"/>
  <equals type="bool" value="True">
    <default param="outertaper"><value type="vector"></value></default>
    <default param="innertaper"><value type="vector"></value></default>
  </equals>
</when>

<when param="interactive">
  <equals type="bool" value="False"/>
  <equals type="bool" value="True">
    <default param="npercycle"><value>100</value></default>
  </equals>
</when>

<when param="imagermode">
  <equals value=""/>
  <equals value="csclean">
    <default param="cyclefactor"><value>1.5</value></default>
    <default param="cyclespeedup"><value>-1</value></default>
  </equals>
  <equals value="mosaic">
    <default param="mosweight"><value>False</value></default>
    <default param="ftmachine"><value type="string">mosaic</value></default>
    <default param="scaletype"><value type="string">SAULT</value></default>
    <default param="cyclefactor"><value>1.5</value></default>
    <default param="cyclespeedup"><value>-1</value></default>
  </equals>
</when>

<!--Get rid of that soon-->
The main clean deconvolution task. It contains many functions

1) Make 'dirty' image and 'dirty' beam (psf)
2) Multi-frequency-continuum images or spectral channel imaging
3) Full Stokes imaging
4) Mosaicking of several pointings
5) Multi-scale cleaning
6) Interactive clean boxing
7) Initial starting model

vis -- Name of input visibility file
default: none; example: vis='ngc5921.ms'

imagename -- Pre-name of output images:
default: none; example: imagename='m2'
output images are:
m2.image; cleaned and restored image
With or without primary beam correction
m2.psf; point-spread function (dirty beam)
m2.flux; relative sky sensitivity over field
m2.model; image of clean components
m2.residual; image of residuals
m2.interactive.mask; image containing clean regions

field -- Select fields in mosaic. Use field id(s) or field name(s).
[ 'go listobs' to obtain the list id's or names]
default: ''=all fields
If field string is a non-negative integer, it is assumed to
be a field index otherwise, it is assumed to be a
field name
field='0~2'; field ids 0,1,2
field='0,4,5~7'; field ids 0,4,5,6,7
field='3C286,3C295'; field named 3C286 and 3C295
field = '3,4C*'; field id 3, all names starting with 4C

spw -- Select spectral window/channels
NOTE: This selects the data passed as the INPUT to mode
default: ''=all spectral windows and channels
spw='0~2,4'; spectral windows 0,1,2,4 (all channels)
spw='0:5~61'; spw 0, channels 5 to 61
spw='&lt;2'; spectral windows less than 2 (i.e. 0,1)
spw='0,10,3:3~45'; spw 0,10 all channels, spw 3, channels 3 to 45.
spw='0~2:2~6'; spw 0,1,2 with channels 2 through 6 in each.
spw='0~10,15~60'; spectral window 0 with channels 0-10,15-60
spw='0~10,1:20~30,2:1;2:3'; spw 0, channels 0-10, spw 1, channels 20-30, and spw 2, channels 1,2 and 3

selectdata -- Other data selection parameters
default: True

&spw; expandable parameters
See help par.selectdata for more on these

timerange -- Select data based on time range:
default = '' (all); examples,
timerange = 'YYYY/MM/DD/hh:mm:ss~YYYY/MM/DD/hh:mm:ss'
Note: if YYYY/MM/DD is missing date defaults to first
day in data set
timerange='09:14:00~09:54:00' picks 40 min on first day
timerange = '25:00:00~27:30:00' picks 1 hr to 3 hr
30min on NEXT day
timerange='09:44:00' pick data within one integration
of time
timerange='&gt;10:24:00' data after this time

uvrange -- Select data within uvrange (default units meters)
default: '' (all); example:
uvrange='0~1000klambda'; uvrange from 0-1000 kilo-lambda
uvrange='&gt;4klambda'; uvranges greater than 4 kilo lambda

antenna -- Select data based on antenna/baseline
default: '' (all)
If antenna string is a non-negative integer, it is
assumed to be an antenna index, otherwise, it is
considered an antenna name.
antenna='5&amp;6'; baseline between antenna index 5 and
index 6.
antenna='VA05&amp;VA06'; baseline between VLA antenna 5
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and 6.
antenna='5&6;7&8'; baselines 5-6 and 7-8
antenna='5'; all baselines with antenna index 5
antenna='05'; all baselines with antenna number 05
   (VLA old name)
antenna='5,6,9'; all baselines with antennas 5,6,9
   index numbers

scan -- Scan number range.
default: '' (all)
example: scan='1~5'
Check 'go listobs' to insure the scan numbers are in order.

mode -- Frequency Specification:
NOTE: See examples below:
default: 'mfs'
mode = 'mfs' means produce one image from all specified data.
mode = 'channel'; Use with nchan, start, width to specify output image cube. See examples below
mode = 'velocity', means channels are specified in velocity.
mode = 'frequency', means channels are specified in frequency.

mode expandable parameters (for modes other than 'mfs')
Start, width are given in units of channels, frequency or velocity as indicated by mode, but only channel is complete.

nchan -- Number of channels (planes) in output image
default: 1; example: nchan=3

start -- Start input channel (relative-0)
default=0; example: start=5

width -- Output channel width in units of the input channel width (&gt;1 indicates channel averaging)
default=1; example: width=4

examples:
spw = '0,1'; mode = 'mfs'
will produce one image made from all channels in spw 0 and 1
spw='0:5^28~2'; mode = 'mfs'
will produce one image made with channels (5,7,9,...,25,27)
spw = '0'; mode = 'channel': nchan=3; start=5; width=4
will produce an image with 3 output planes
plane 1 contains data from channels (5+6+7+8)
plane 2 contains data from channels (9+10+11+12)
plane 3 contains data from channels (13+14+15+16)
spw = '0:0.63^3'; mode='channel'; nchan=21; start = 0;
width = 1
will produce an image with 20 output planes
Plane 1 contains data from channel 0
Plane 2 contains data from channel 2
Plane 21 contains data from channel 61
spw = '0:0.40^2'; mode = 'channel'; nchan = 3; start = 5; width = 4
will produce an image with three output planes
plane 1 contains channels (5,7)
plane 2 contains channels (13,15)
plane 3 contains channels (21,23)

psfmode -- method of PSF calculation to use during minor cycles:
default: 'clark': Options: 'clark', 'hogbom'
'clark' use smaller beam (faster, usually good enough)
'hogbom' full-width of image (slower, better for poor
uv-coverage)
Note: psfmode will be used to clean if imagermode = ''
imagermode -- Advanced imaging e.g. mosaic or Cotton-Schwab clean
default: imagermode=''': Options: '', 'csclean', 'mosaic'
default '' =&gt; psfmode cleaning algorithm used
Image as a mosaic of the different pointings (uses csclean
style too)
mosweight -- Individually weight the fields of the mosaic
default: False; example: mosweight=True
This can be useful if some of your fields are more
sensitive than others (i.e. due to time spent
on-source); this parameter will give more weight to
higher sensitivity fields in the overlap regions.
ftmachine -- Gridding method for the image;
Options: ft (standard interferometric gridding), sd
(standard single dish) both (ft and sd as appropriate),
mosaic (gridding use PB as convolution function)
default: 'mosaic'; example: ftmachine='ft'
scaletype -- Controls scaling of pixels in the image plane.
(Not fully implemented...for now only controls
what is seen if interactive=True...but in the future will
control the image on which clean components are searched)
default='SAULT'; example: scaletype='PBCOR'
Options: 'PBCOR','SAULT'
'SAULT' when interactive=True shows the residual
with constant noise across the mosaic. If
pbcor=False, the final output image is NOT
corrected for the PB pattern, and therefore is not "flux correct". Division of SAULT &lt;imagename&gt;.image by the &lt;imagename&gt;.flux image will produce a "flux correct image", can also be achieved by setting pbcor=True.

'PBCOR' uses the SAULT scaling scheme for deconvolution, but if interactive=True shows the primary beam corrected image; the final PBCOR image is "flux correct" if pbcor=True.

imagermode='csclean' expandable parameter(s): Image using the Cotton-Schwab algorithm in between major cycles
cyclefactor -- Change the threshold at which the deconvolution cycle will stop, degrid and subtract from the visibilities. For poor PSFs, reconcile often (cyclefactor=4 or 5); For good PSFs, use cyclefactor 1.5 to 2.0. Note: threshold = cyclefactor * max sidelobe * max residual.
default: 1.5; example: cyclefactor=4
cyclespeedup -- Cycle threshold doubles in this number of iterations default: -1; example: cyclespeedup=3
multiscale -- set of scales to use in deconvolution. If set, cleans with several resolutions using hobgom clean. The scale sizes are in units of cellsize. So if cell='2arcsec', a multiscale scale=10 = 20arcsec. First scale should always be 0 (point), we suggest second on the order of synthesized beam, third 3-5 times synthesized beam, etc. For example if synthesized beam is 10" and cell=2", try multscale = [0,5,15]. Note, multiscale is currently a bit slow.
default: multiscale=[] (standard CLEAN using psfmode algorithm, no multi-scale). Example: multscale = [0,5,15]
negcomponent -- Stop component search when the largest scale has found this number of negative components; -1 means continue component search even if the largest component is negative. default: -1; example: negcomponent=50
imsize -- Image pixel size (x,y)
default = [256,256]; example: imsize=[350,350]
imsize = 500 is equivalent to [500,500]

multiscale expandable parameter(s): negcomponent -- Stop component search when the largest scale has found this number of negative components; -1 means continue component search even if the largest component is negative. default: -1; example: negcomponent=50
imsize -- Image pixel size (x,y)
default = [256,256]; example: imsize=[350,350]
imsize = 500 is equivalent to [500,500]
cell -- Cell size (x,y)
default= '1.0arcsec'; example: cell=['0.5arcsec','0.5arcsec'] or
cell=['1arcmin', '1arcmin']
cell = '1arcsec' is equivalent to ['1arcsec', '1arcsec']

NOTE: cell = 2.0 => ['2arcsec', '2arcsec']

phasecenter -- direction measure or fieldid for the mosaic center
default: '' => first field selected ; example: phasecenter=6
or phasecenter='J2000 19h30m00 -40d00m00'

restfreq -- Specify rest frequency to use for output image
default=''; Occasionally it is necessary to set this (for
example some VLA spectral line data). For example for
NH_3 (1,1) put restfreq='23.694496GHz'

stokes -- Stokes parameters to image
default='I'; example: stokes='IQUV';
Options: 'I','IV''QU','IQUV','RR','LL','XX','YY','RRLL','XXYY'

niter -- Maximum number iterations,
if niter=0, then no CLEANing is done ("invert" only)
default: 500; example: niter=5000

gain -- Loop gain for CLEANing
default: 0.1; example: gain=0.5

threshold -- Flux level at which to stop CLEANing
default: '0.0mJy';
example: threshold='2.3mJy' (always include units)
threshold = '0.0023Jy'
threshold = '0.0023Jy/beam' (okay also)

interactive -- use interactive clean (with GUI viewer)
default: interactive=False
example: interactive=True

interactive clean allows the user to build the cleaning
mask interactively using the viewer. The viewer will
appear every npercycle iteration, but modify as needed
The final interactive mask is saved in the file
imagename_interactive.mask. The initial masks use the
union of mask and cleanbox (see below)

fft=interactive=True expandable parameter npercycle -- this is the
number of iterations between each clean to update mask
interactively. Set to about niter/5, but can also be
changed interactively.

mask -- Specification of cleanbox(es), mask image(s), and/or
region(s) to be used for CLEANing. As long as the image has
the same shape (size), mask images from a previous
interactive session can be used for a new execution. NOTE:
the initial clean mask actually used is the union of what
is specified in mask and &lt;imagename&gt;.mask default: [] (no
masking); Possible specification types: (a) Explicit
cleanbox pixel ranges example: mask=[110,110,150,145] clean
region with blc=110,100; trc=150,145 (pixel values) (b)
Filename with cleanbox pixel values with ascii format:
example: mask='mycleanbox.txt' &lt;fieldid blc-x blc-y
trc-x trc-y&gt; on each line
1 45 66 123 124
2 23 100 300 340
(c) Filename for image mask example: mask='myimage.mask'
(d) Filename for region specification (e.g. from viewer)
exampel: mask='myregion.rgn' (e) Combinations of any of the
above example: mask=[[110,110,150,145],'mycleanbox.txt',
'myimage.mask','myregion.rgn']

uv taper -- Apply additional uv tapering of the visibilities.
default: uvtaper=False; example: uvtaper=True

outertaper -- uv-taper on outer baselines in uv-plane
[bmaj, bmin, bpa] taper Gaussian scale in uv or
angular units. NOTE: uv taper in (klambda) is roughly on-sky
FWHM(arcsec/200)
default: outertaper=[]; no outer taper applied
example: outertaper=['5klambda'] circular taper
FWHM=5 kilo-lambda
outertaper=['5klambda','3klambda','45.0deg']
outertaper=['10arcsec'] on-sky FWHM 10"
outertaper=['300.0'] default units are meters
in aperture plane

innertaper -- uv-taper in center of uv-plane
[bmaj,bmin,bpa] Gaussian scale at which taper falls to
zero at uv=0
default: innertaper=[]; no inner taper applied
NOT YET IMPLEMENTED

model image -- Name of model image(s) to initialize cleaning. If
multiple images, then these will be added together to
form initial staring model NOTE: these are in addition
to any initial model in the &lt;imagename&gt;.model image file
default: '' (none); example: modelimage='orion.model'
modelimage=['orion.model','sdorion.image'] Note: if the
units in the image are Jy/beam as in a single-dish
image, then it will be converted to Jy/pixel as in a
model image, using the restoring beam in the image
header

weighting -- Weighting to apply to visibilities:
default='natural'; example: weighting='uniform';
Options: 'natural','uniform','briggs',
'superuniform','briggsabs','radial'

Weighting expandable parameters
For weighting='briggs' and 'briggsabs'
robust -- Brigg's robustness parameter
   default=0.0; example: robust=0.5;
   Options: -2.0 to 2.0; -2 (uniform)/+2 (natural)
For weighting='briggsabs'
   noise -- noise parameter to use for Briggs "abs"
   weighting
eexample noise='1.0mJy'
For superuniform/briggs/briggsabs weighting
   npixels -- number of pixels to determine uv-cell size
   for weight calculation
eexample npixels=7
restoringbeam -- Output Gaussian restoring beam for CLEAN image
   [bmaj, bmin, bpa] elliptical Gaussian restoring beam
   default units are in arc-seconds for bmaj,bmin, degrees
   for bpa default: restoringbeam=[]; Use PSF calculated
   from dirty beam.
eexample: restoringbeam=['10arcsec'] circular Gaussian
   FWHM 10" example:
   restoringbeam=['10.0','5.0','45.0deg'] 10"x5"
at 45 degrees
pbcor -- Output primary beam-corrected image
   default: pbcor=False; output un-corrected image
   example: pbcor=True; output pb-corrected image (masked outside
   minpb) Note: if you set pbcor=False, you can later
   recover the pbcor image by dividing by the .flux image
   (e.g. using immath)
minpb -- Minimum PB level to use default=0.1; example:
   minpb=0.01 Note: this minpb is always in effect
   (regardless of pbcor=True/False)

</example>
</task>
</casaxml>

I.3.2 File task_clean.py

Task clean implementation file.

import os
from taskinit import *
from cleanhelper import *
def clean(vis, imagename, field, spw, selectdata, timerange, uvrangle, antenna, scan, mode, niter, gain, threshold, psfmode, imagermode, ftmachine, mosweight, scaletype, multiscale, negcomponent, interactive, mask, nchan, start, width, imsize, cell, phasecenter, restfreq, stokes, weighting, robust, uvtaper, outertaper, inner taper, modelimage, restoringbeam, pbcor, minpb, noise, npixels, npercycle, cyclespeedup):

# Python script

casalog.origin('clean')

maskimage=''
if((mask==[]) or (mask=='')):
    mask=['']
if (interactive):
    if( (mask=='') or (mask==['']) or (mask==[]):
        # try:
        if(1):
            imCln=imtool.create()
            imset=cleanhelper(imCln, vis)

            if((len(imagename)==0) or (imagename.isspace())):
                raise Exception, 'Cannot proceed with blank imagename'
            casalog.origin('clean')

            imset.defineimages(imsize=imsize, cell=cell, stokes=stokes, mode=mode, spw=spw, nchan=nchan, start=start, width=width, restfreq=restfreq, field=field, phasecenter=phasecenter)

            imset.datselweightfilter(field=field, spw=spw, timerange=timerange, uvrangle=uvrange, antenna=antenna, scan=scan, wgttype=weighting, robust=robust, noise=noise, npixels=npixels, mosweight=mosweight, inner taper=innertaper, outertaper=outertaper)
if(maskimage==''):
    maskimage=imagename+'.mask'
imset.makemaskimage(outputmask=maskimage,imagename=imagename,
                        maskobject=mask)

###define clean alg
alg=psfmode
if(multiscale==[0]):
    multiscale=[]
if((type(multiscale)==list) and (len(multiscale)>0)):
    alg='multiscale'
imCln.setscales(scalemethod='uservector',
                   uservector=multiscale)
if(imagermode=='csclean'):
    alg='mf'+alg
if(imagermode=='mosaic'):
    if(alg.count('mf')<1):
        alg='mf'+alg
    imCln.setoptions(ftmachine=ftmachine, padding=1.0)
imCln.setvp(dovp=True)

###PBCOR or not
sclt='SAULT'
if((scaletype=='pbcor') or (scaletype=='pbcor')):
    sclt='NONE'
imCln.setvp(dovp=True)
else:
    if(imagermode != 'mosaic'):
        ##make a pb for flux scale
        imCln.setvp(dovp=True)
imCln.makeimage(type='pb', image=imagename+'.flux')
imCln.setvp(dovp=False)

###restoring
imset.setrestoringbeam(restoringbeam)

###model image
imset.convertmodelimage(modelimages=modelimage,
                        outputmodel=imagename+'.model')

###after all the mask shenanigans...make sure to use the
###last mask
maskimage=imset.outputmask
if((imagermode=='mosaic')):
    imCln.setmfcontrol(stoplargenegatives=negcomponent,scaletype=sclt,
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minpb=minpb,cyclefactor=cyclefactor,
cyclespeedup=cyclespeedup,
fluxscale=[imagename+’.flux’])
else:
imCln.setmfcontrol(stoplargenegatives=negcomponent,
cyclefactor=cyclefactor, cyclespeedup=cyclespeedup)
imCln.clean(algorithm=alg,niter=niter,gain=gain,
threshold=qa.quantity(threshold,’mJy’),
model=[imagename+’.model’],
residual=[imagename+’.residual’],
image=[imagename+’.image’],
psfimage=[imagename+’.psf’],
mask=maskimage, interactive=interactive,
npercycle=npercycle)
imCln.close()
presdir=os.path.realpath(’.’)
newimage=imagename
if(imagename.count(’/’) > 0):
newimage=os.path.basename(imagename)
os.chdir(os.path.dirname(imagename))
result
= ’\’’ + newimage + ’.image’ + ’\’’;
fluxscale_image = ’\’’ + newimage + ’.flux’ + ’\’’;
if (pbcor):
if(sclt != ’NONE’):
##otherwise its already divided
ia.open(newimage+’.image’)
pixmask = fluxscale_image+’>’+str(minpb);
ia.calcmask(pixmask,asdefault=True);
pixels=’iif(’+ fluxscale_image+’>’+str(minpb)+’,’
+ result+’/’+fluxscale_image+’, 0)’
ia.calc(pixels=pixels)
ia.close()
else:
## people has imaged the fluxed corrected image
## but want the
## final image to be non-fluxed corrected
if(sclt==’NONE’):
ia.open(newimage+’.image’)
result=newimage+’.image’
fluxscale_image=newimage+’.flux’
pixels=result+’*’+fluxscale_image
ia.calc(pixels=pixels)


ia.close()

os.chdir(presdir)

del imCln

# except Exception, instance:
#     print '*** Error *** ',instance
#     raise Exception, instance