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Starlink Project
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SPLAT-VO – A VO-enabled Spectral Analysis Tool

3.11-2

User's Manual

Abstract

SPLAT-VO is an extensible analysis and display tool for extracted spectra that can also interact with the Virtual Observatory.

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1 Overview

SPLAT-VO is a graphical tool for displaying, comparing, modifying and analysing astronomical spectra stored in FITS, VOTABLE, TEXT and NDF/NDX format. It can read spectra from local disk files, download them over the Internet, or it can interact with the Virtual Observatory and query servers for spectral data in a region of the sky.

SPLAT-VO can handle many spectra at the same time and will display these as line plots or point markers. Each display window, of which there can be many, can be used to view one or more spectra at the same time. Display windows can be interactively zoomed and scrolled, centred on specific wavelengths, provide continuous coordinate readout, annotated, produce printable hardcopy and be configured in many ways. They also provide the basis for interactive analysis facilities. All of these functions are displayed in a visually rich format, that is intended to appeal especially to casual users of spectral data analysis facilities.

The current set of analysis facilities include background estimation by the fitting of a polynomial to selected parts of a spectrum or by drawing an interpolated spline, the fitting of Gaussian, Lorentzian and Voigt profiles to emission and absorption lines, the filtering of spectra using average, median and line-shape window functions as well as wavelet denoising and the calculation of statistics. A database of laboratory line positions is available to aid in identification.

SPLAT-VO also supports a full range of coordinate systems for spectra, using the latest facilities of the Starlink AST library. This allows coordinates to be *displayed* and *aligned* in many different coordinate systems (wavelength, frequency, energy, velocity) and *transformed* between these and different standards of rest (topocentric, heliocentric, dynamic and kinematic local standards of rest *etc.*). It also uses AST to transform spectra between various flux systems, so that spectra from a wide variety of sources can be easily compared and used to produce a spectral energy distribution. This is an important feature for supporting the heterogeneous data supplied in the Virtual Observatory.

Finally, SPLAT-VO can interoperate with other VO enabled desktop tools using the SAMP protocol.

If you have any problems with SPLAT-VO then send these to the Starlink user support email list at:

`starlink@jiscmail.ac.uk`

or to the SPLAT-VO mailing-list at:

`http://lists.g-vo.org/cgi-bin/mailman/listinfo/splat-users`

where hopefully someone can help you.

2 Getting started

The next two sections describe how to display one spectrum and then how to display many spectra. If you're new to SPLAT-VO then do take the time to read these through and try out the example commands.

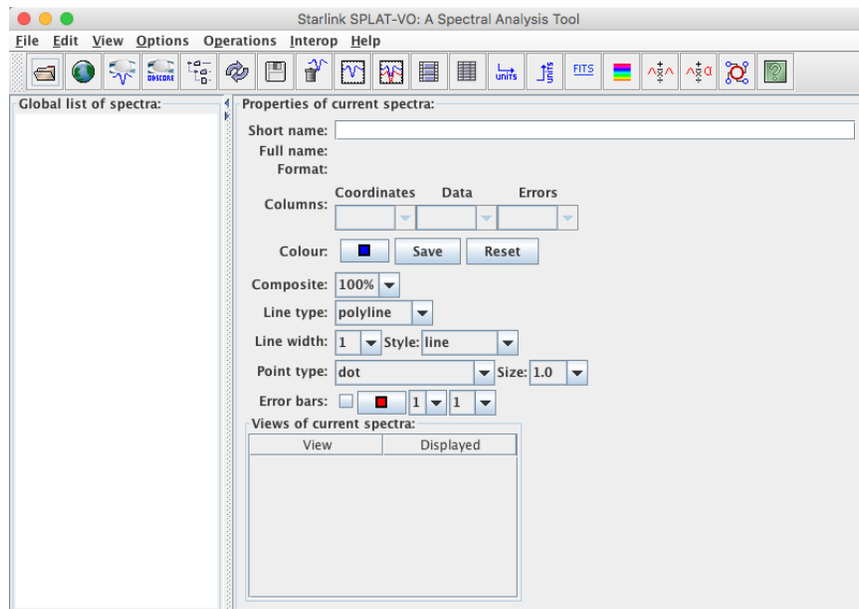
2.1 Displaying a spectrum

To start SPLAT-VO you should just need to type the command:

```
% splat &
```

This assumes that you have SPLAT-VO installed on your system as part of a standard Starlink installation. If not then you'll need follow any installation and pre-startup instructions that you have before using this command.

When SPLAT-VO appears it should look something like:



This is the main browser window. It has a list containing all the available spectra (in what we call the 'global list', but has been traditionally referred to as the 'stack'). Using this window you can add, save and remove spectra from the list, you can change how a spectrum draws itself, in all the places that it is shown and make a spectrum add itself to a plot – this is how you view spectra.

You can also do things like change the look and feel of the whole application (there are different colour and font schemes), perform some quite primitive mathematical operations on or between the spectra, run through an animated display sequence of some or all of the spectra, set the spectra coordinate systems and modify/inspect their values. So things that apply to each spectrum, wherever it is shown, or the program as a whole are generally done in this window.

To open your first spectrum select the **Open** item in the **File** menu. This will create a familiar open file dialogue window. Just navigate to your spectrum, select it, and press **Open** to proceed.

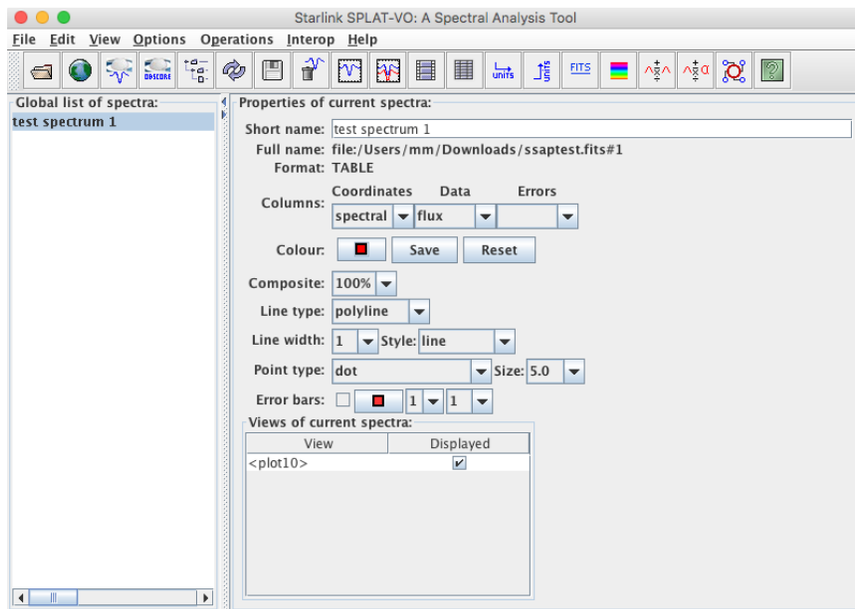
Alternatively you could have supplied the spectrum you want to use on the command-line:

```
% splat <spectrum_file> &
```

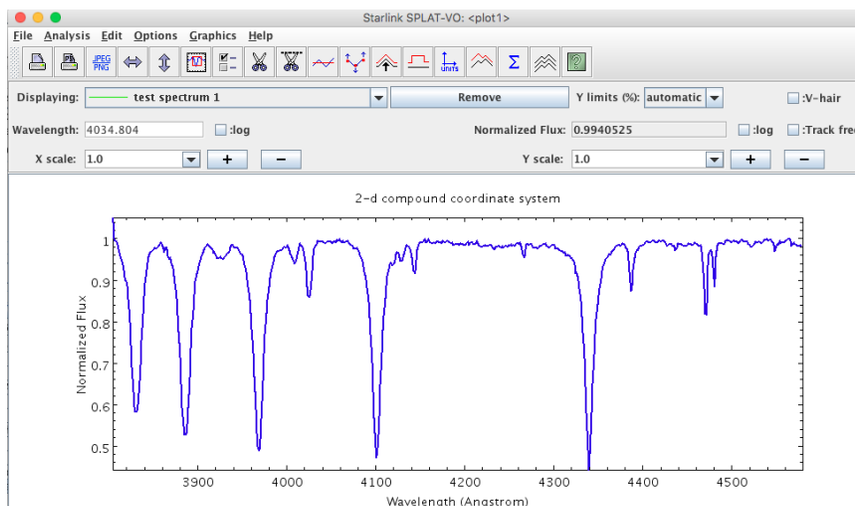
Which will read the spectrum stored in the named file. SPLAT-VO will read spectra stored in most astronomical formats (Starlink NDF, FITS spectra and tables, VOTables, simple TEXT files).

Now that you've actually read in a spectrum it will be immediately displayed in a 'plot'. Have a look at this plot window briefly, if you want, but close it before proceeding (to close it select the **Close** item in its **File** menu). You really need to understand how to display a spectrum in a plot yourself, if you're to gain complete control over the creation of plots and what they display.

The browser window (once again on its own) should now appear something like:



Now to re-create a plot and display a spectrum in it the hard way. Actually this couldn't be simpler, just 'double click' on the name in the **Global list of spectra**. Another way to achieve the same effect is to select the name (by just clicking on it once, you can tell it's selected when it becomes highlighted, as shown above) and choose the display selected spectra button in the toolbar. This is the button that looks like  (it's meant to represent a spectrum with axes drawn in a plot). If you're successful then a window looking like this:

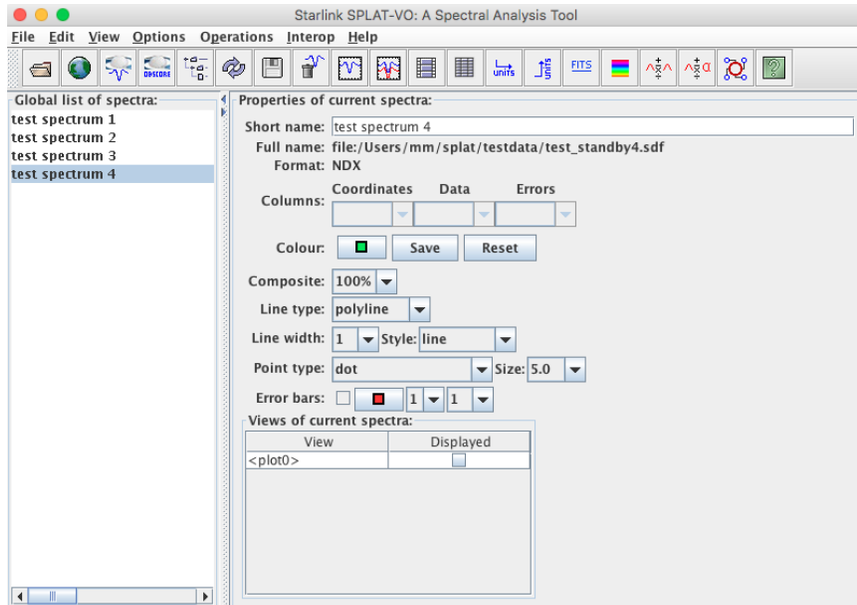


should appear as before.

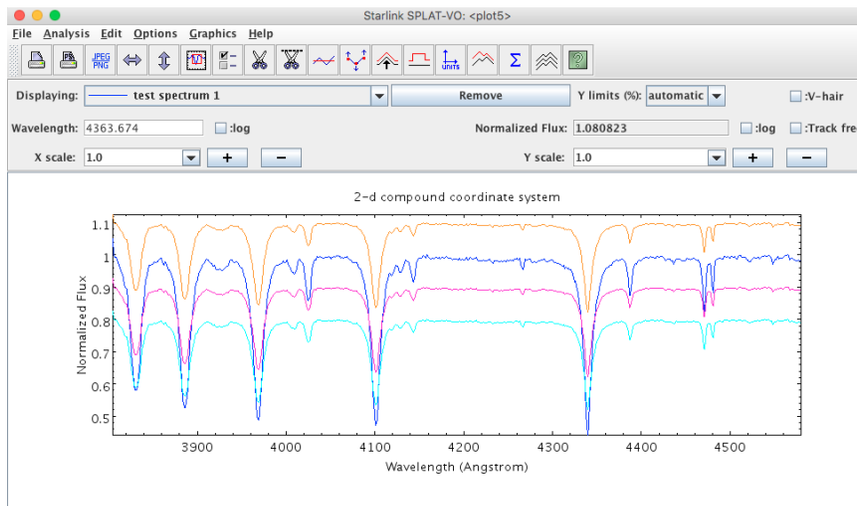
Since SPLAT-VO is meant to be capable of displaying many spectra (sometimes in the same plot as we'll see in the next section), you should confirm this by creating a second plot window, by just performing the same operation again (*i.e.* double click on the spectrum name).

2.2 Displaying more than one spectrum

Displaying many spectra at the same time isn't much more difficult than the single spectrum case. Just open up several spectra by giving their names on the command-line, or by opening them using the **File**→**Open** dialogue. This will produce a main window that looks something like:



If you've opened all the spectra at once (you can do this using the **File**→**Open** dialogue by selecting more than one file – just hold down the shift key while selecting each one) then they will have also all appeared displayed in a single plot, which should look something like:



Again you should now close the plot (or all the plots if you've opened the spectra one-by-one) so you can find out how to add spectra to plots manually.

There are three ways to display a set of spectra in a single plot, you can either open a plot just displaying one spectrum and then add the others one-by-one, by two different methods, or you can open a plot displaying a pre-selected list of spectra. Let's cover the latter option first.

The first thing to do is identify which spectra you want to display in the same plot, you do this by selecting their names in the **Global list of spectra:** list. You can select more than one name at a time by holding down the control key, while clicking on the name. It's also possible to select ranges of names by selecting the first name and then selecting the last name while holding down the shift key. Have a play to see the effects that you can get with combinations of these (hint: using the control key also allows you to remove single names from the selection). To select all the names you can also use the **Edit**→**Select all spectra** menu item.

Now that you've selected the spectra to display, just press the multi-display button:  and all the spectra will be displayed in a single plot. Note that if you inadvertently press the single display button: , then each spectrum will appear in a plot of its own (this can be a very bad idea if you have a very large number of spectra selected!).

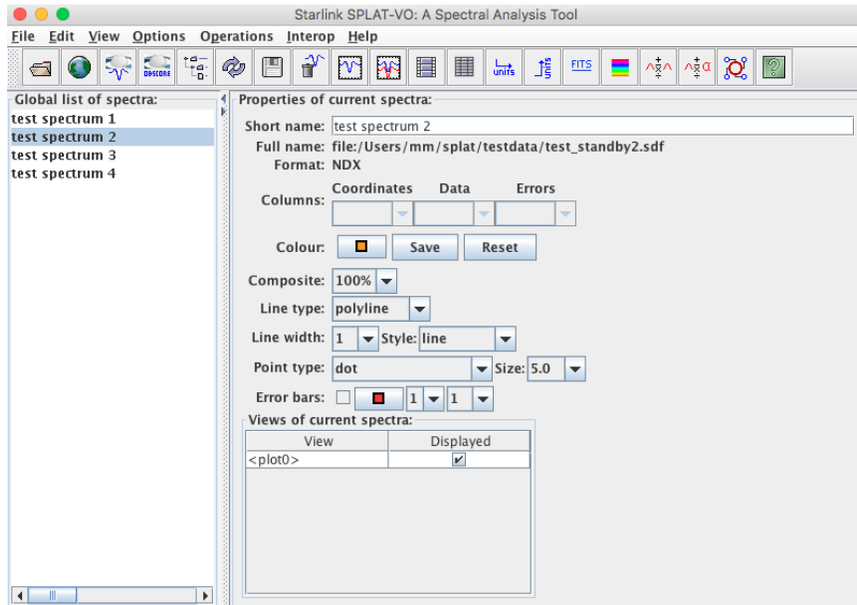
If you don't like the colours chosen for your spectra you can just press the auto-colour button:  to get a new set. Each press of this button should generate a new set of colours, so just keep clicking until you see a combination that you like.

The second method for adding more than one spectrum to a plot is the simplest, you can drag spectra from the **Global list of spectra:** and drop them into the display area of the plot. To drag a single spectrum just make sure that it is selected and then attempt to drag it from the **Global list of spectra:** into the plot (during dragging a document icon should be attached to your screen pointer). To drag more than one you just need to make all the spectra you want displayed selected and then attempt to drag any of the selected set.

The third method is actually also the method you'd use to remove a spectrum from a plot. The first thing to do is select a spectrum in the **Global list of spectra:**, this shows its properties in the **Properties of current spectra:** side of the browser window, including which views are being made (*i.e.* which plots are displaying it).

If you click on one of the displayed spectra on a plot, the respective entry in the **Global list of spectra:** will be highlighted. By clicking on an entry at the **Global list of spectra:**, the respective plotted spectrum will blink a few times (if it's plotted).

To add or remove a spectrum from a plot you need to click on the square shown in the **Displayed** column, so that it becomes ticked or unticked. See in the following image that test spectrum 2 is currently being displayed in <plot0>:



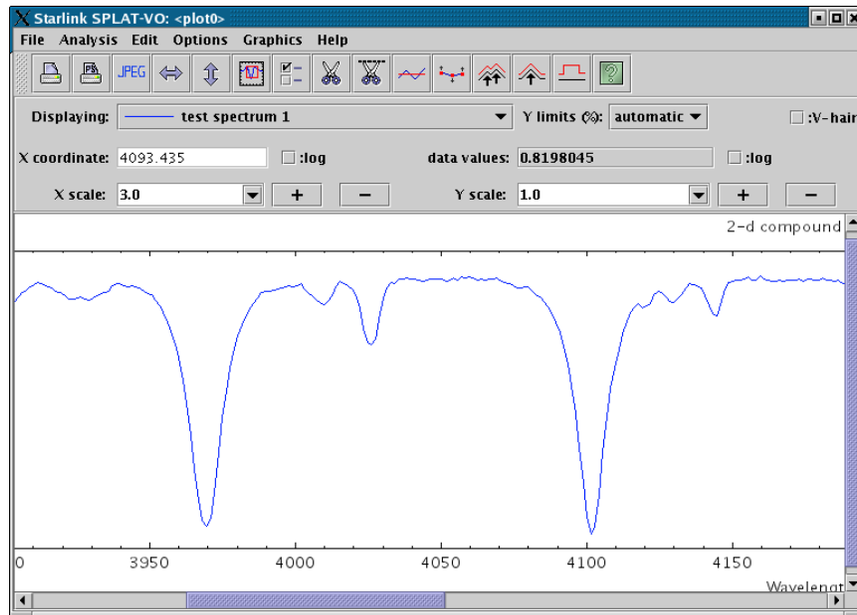
If any spectra were not displayed in this plot then they could be added to it using this mechanism. Naturally this also extends to the case when there are many plots available.

2.3 Basic control of a plot view

The final part of this introduction is to zoom and scroll the spectrum, so, for instance, you can inspect individual lines and read off accurate coordinates (this is after all the main purpose of SPLAT-VO).

Zooming of a plot is controlled by the **X scale:** and **Y scale:** controls. To adjust the zoom you may either select a value from the drop-down lists, or you may type in a value and press <Return>. Values may be non-integer, but should be one or more.

Once you have zoomed the plot scrollbars should appear at the bottom and maybe to the right (these appear when the size of the plot becomes larger than its visible size). Just drag the slider from side-to-side or up-and-down to view the part of the plot that you want to see.



A useful shortcut for zooming is to drag out a region using the middle mouse button. Just place your cursor on the plot, press the middle button and, keeping the button depressed, move the mouse until the rubber-band rectangle encloses the feature you want to zoom on. You can also zoom in by an increment of one in X and Y by pressing the middle button and just releasing it.

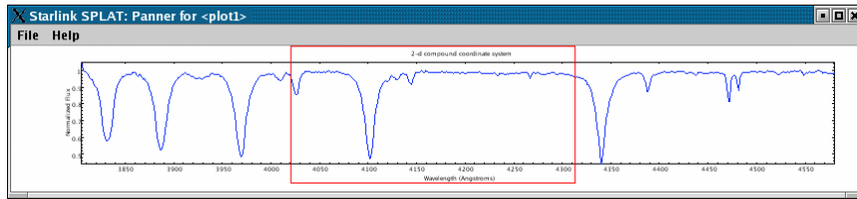
If you find that the figure is zoomed so much that you're having difficulty recognising the features, then there are several things that you can do.

The first is to resize the plot so that it has a scale of 1 in X or Y. That's what the toolbar buttons that appear as  and  do, make the plot scale its displayed spectra to fit the visible width or height. **Note:** These buttons are very useful when a plot seems to be displaying nothing or reports that the plot has no size etc., they are your best friends, press them a lot in such circumstances.

If what you really want to do is remain zoomed but quickly go to a given wavelength, then all you need to do is type in the wavelength in the X-coordinate readout entry field (labelled **Wavelength** above), press <Return> and the plot will be centred (or brought into the edge of the field of view).

If what you'd really like is a smaller view of the whole plot to work from, then press the  button, this will pop-up the 'panner'. This shows a red rectangle over a view of the full unzoomed plot (shown in the same aspect ratio as the full plot at the current scale factors). To move the rectangle, and hence the view shown in the plot, just point somewhere in the panner and press the left mouse button, or 'drag' the rectangle itself.

If the panner view seems very small, as can happen when you're using a high X scale factor, then try resizing the window to get a better fit to the aspect ratio (hint: make it taller).

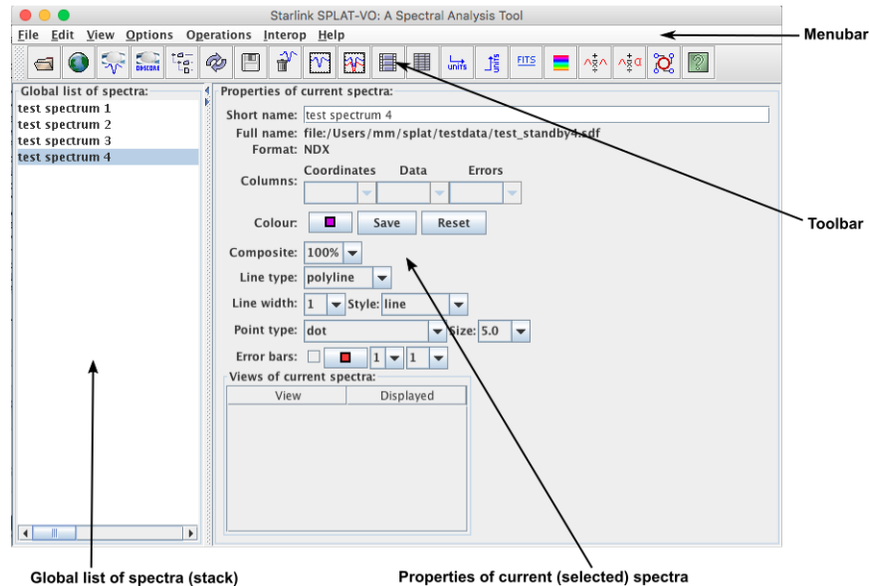


3 Help for individual windows

The following sections are descriptions of how to use the SPLAT-VO windows, hopefully these should only serve as reference material, the function of windows should be clear from their descriptions, balloon-help and context. Use a window's **Help** menu to view its related page on-line.

3.1 The browser window

The browser window has several main elements, each of which is shown in the following figure:



Global list of spectra

In the **Global list of spectra** short names for each 5 spectra are shown. The short names are intended to allow the quick identification of spectra and can be modified at any time by selecting the name and then typing a new name in the **Short name** field (remember to press <Return> to make the change happen). Initially short names are read from the **TITLE** or **OBJECT** properties of spectral data formats that support these (*i.e.* NDF and FITS respectively), otherwise they default to the file name.

Another important concept related to the global list, is using it to create a selected range or list of the spectra. Selection is important as this identifies which spectra you'd like to see properties for, or want to work with. There's a quick tutorial on selecting spectra in the introductory material (§2.2).

Properties of current spectra

In the properties area there are controls for changing how the currently selected spectra are displayed (a spectrum always uses the same display properties) and where they are displayed.

If more than one spectrum is selected then the properties shown are those of the first spectrum (*i.e.* highest in the global list). However, any changes that you then make are applied to all the selected spectra.

If you load a table into SPLAT-VO the **Columns:** controls will show the names of the columns selected. If these are incorrect then you can choose a different set.

The properties that you can change include the colour (press on this to get a swatch of selectable colours), the composite value (that is the transparency so you can see

lines or points through other line or points), the line type (simple connected polyline or histogram, or just a point marker). If you choose to draw lines then you can set the width and style (dashed *etc.*). If you choose to draw points then various types are available.

If the selected spectra have errors then you can display error bars by clicking on the **Error bars:** control. This is greyed out if errors are not available. The colour, number of sigma drawn and frequency of the error bars can be changed using the other controls on this line.

Views of current spectra

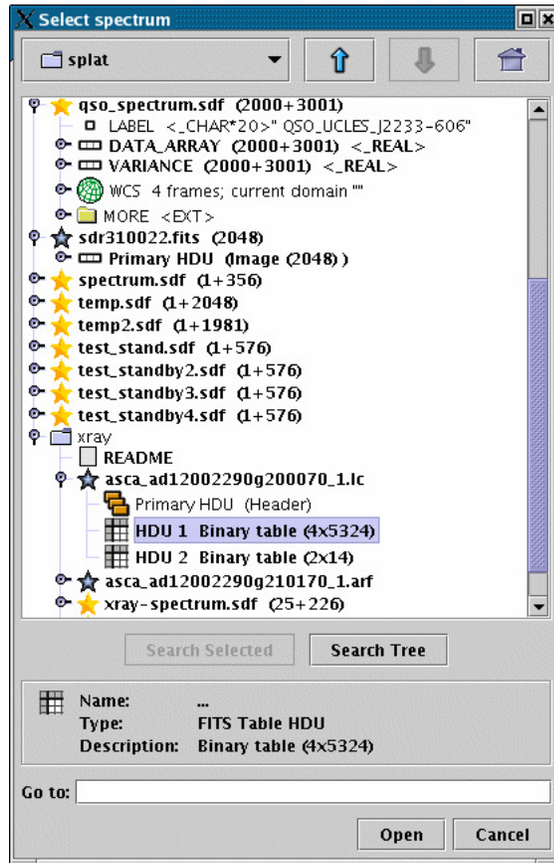
The views of the current spectra region shows the names of the plots that are displaying the currently selected spectra (this time if a plot is showing only some of the spectra, then it is highlighted in red). To add a spectrum or list of spectra to a plot, select their names in the global list and then click on the **Displayed** control (normally this results in a tick appearing). If a plot is already displaying some of the selected spectra then you force it to display them all by clicking twice – the first click removes the selected spectra and the second puts them all back, plus the ones not already shown.

To make a plot visible double click on its name. This should raise and deiconify it.

Toolbar and menus

The toolbar region has a series of buttons that act as short-cuts to many of the functions found in the main menus. Here's a quick break down of what they do:

-  Create an open file dialogue. You can use this to browse any local files and select a list of them for opening. When opened they are added to the global list and displayed in a new plot (unless you untick the **Display** control). You may open files that are already in the global list, this just creates a new entry. Normally the type of a spectrum is determined using the file extension, but using this dialogue you can select a type, or ask that it be guessed.
-  Download and display a spectrum from the Internet. If you have a URL for a spectrum to view then use this simple dialog.
-  Query any known Simple Spectral Access Protocol (SSAP) servers about spectra that they hold on a region of the sky and download these to SPLAT-VO. This tool provides access to the Virtual Observatory.
-  Query any known Observation Data Model Core (ObsCore) spectral services using the Table Access Protocol (TAP) and download these to SPLAT-VO. This tool provides access to the Virtual Observatory.
-  Browse a hierarchy of files. This dialogue can be used like the open file dialogue to choose a file containing a spectrum, but, it also allows you to look inside a file for any displayable components. This is useful for FITS files that contain extensions (spectral or table extensions can be opened), or HDS files that contain more than one NDF. The file type guessing used here is the same as that used when you set a type of guess (either on the command-line or by using the open file dialogue).



-  Re-open the selected spectra. If the files containing your spectra change on disk you can select them in the global list get them opened. Note that not all spectra are backed by a suitable disk-file, these types will be ignored.
-  Create a save file dialogue. Using this you can save a spectrum to disk-file. There are no restrictions on which file format you can save to, but information may be lost for certain types (*i.e.* TEXT format).
-  Remove the currently selected spectra from the global list. This does not modify disk files. Spectra that are only resident in memory will be lost, without warning.
-  Display the currently selected spectra in plots. Each of the selected spectra is displayed in a plot of its own. A similar effect is obtained by double-clicking a spectrum in the global list.
-  Display all the currently selected spectra in a single plot. Use this when you want to compare spectra.
-  Open a window to control the animation of some of the spectra. Using this window you can also create a JPEG or PNG sequence of the animation.
-  View and possibly modify the coordinates and data values of the selected spectra. This shows an editable table of values. It is also possible to make global changes using algebraic expressions to convert or create new values.
-  Open a window to set or remap the coordinate systems of any of the spectra. A variety of systems can be used covering wavelength, frequency, energy

and velocity. Unit information can be used to align spectra in a plot (so you can trivially compare spectra in say angstroms and nanometres, energy and frequency or even heliocentric and topocentric rest frames *etc.*).

-  Open a window to set the data units of any spectra. This can be useful when just setting the units displayed in a plot, but can also be used to convert spectra between flux systems when they are drawn (making them correctly aligned).
-  View the FITS headers of any selected spectra. A window displaying the headers will be opened for each of the selected spectra. Note that only 1D FITS and NDFs will generally have FITS headers other types, including FITS tables, will not.
-  Change the colour of all the spectra in the global list. Each time you press this button a new random seed is used to generate a different sequence of colours, so keep pressing until you see a sequence you like.
-  Perform simple maths (add, subtract, divide and multiply) between pairs of spectra.
-  Perform simple maths using a spectrum and a constant (add, subtract, divide and multiply).
-  Access the on-line help system.

Interesting menu items not found in the toolbar are a list of standard line position catalogues (for optical and some IR and submillimeter lines), these are found in **Options**→**Line identifiers**. Others are to do with controlling the selection of the spectra and plots (see the **Edit** menu, just like deleting spectra from the global list you can delete all the selected plots) and changing the ‘look and feel’ of SPLAT-VO. These options are found in the **Options**→**Look and Feel** and **Options**→**Colour Theme** sub-menus. Note that if you select the CDE/Motif look and feel you cannot change the colour and fonts used.

You can also save and restore the global stack of spectra to and from diskfile using a special data format that is only recognised by SPLAT-VO. Using this method allows you to save a working set of spectra and restart an analysis later. Properties such as the line colours are restored, but any relationship between disk files and the spectra in the global list is lost. Data saved in a stack are compressed.

The **Interop** menu contains items for communicating with other SAMP-aware desktop applications. This is described in the Tool interoperability section (§6).

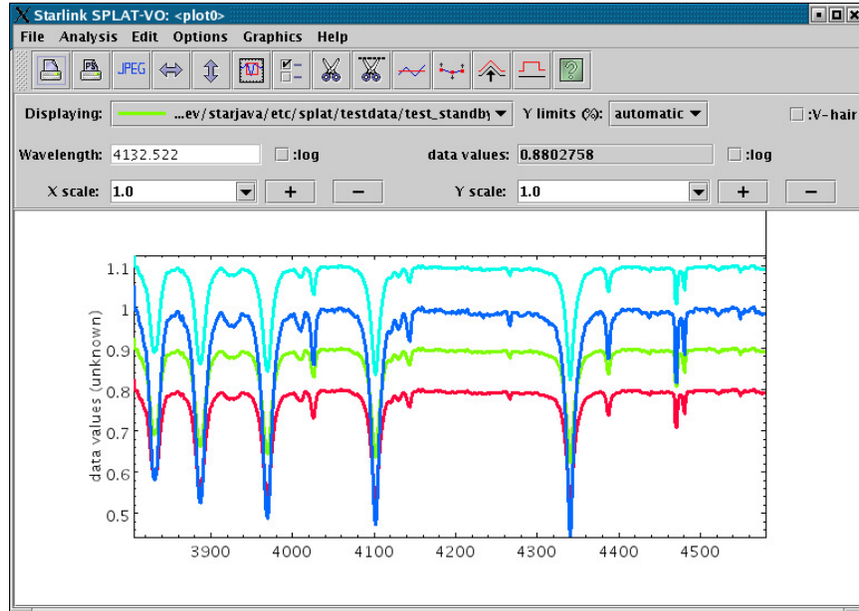
Finer control of the initial coordinate system picked by SPLAT-VO is given by the **Options**→**Find spectral coordinates** menu item. When this is selected (the default), SPLAT-VO will check for any spectral coordinate system in the input spectra and use that, if the default coordinate system isn’t spectral. It will also use any units it can locate to attempt a guess if all else fails. Unfortunately this means that if you intend to view another coordinate system (like GRID to see the channel numbers), it might fail, so switching this off gives you complete control. This option can also be enabled using the `-k, --keepcoords` switches on the command-line.

Accelerator keys

- **Control-o** Activate open file dialog.
- **Control-l** Activate location dialog.
- **Control-r** Re-open all spectra.
- **Control-w** Close window (and exit application).
- **Delete** Remove selected spectra.
- **Control-a** Select all spectra.
- **Control-d** Display selected spectra in plots.
- **Control-i** Display selected spectra in same plot.
- **Control-u** Re-colour all spectra.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.2 A plot window

The anatomy of a plot window is shown below:



It has three main regions, the menubar & toolbar, the control panel and the display area.

The display area

The display area shows a view of all the spectra that are currently associated with this plot. The colours, plotting style *etc.* of these spectra are determined by their global properties as set in the main browser window.

The plot title and axis labels are determined using the properties of the current spectrum, if possible. For instance if the current spectrum is an NDF then these values are determined from the NDF's title, units and axis components, or for more recent NDFs from the properties of its WCS component. Similarly for FITS files these values are determined using the world coordinates FITS keywords. To change these look in the **Configure Plot Options** window, which is activated by pressing the  button.

The size of the area used to display a spectrum is initially determined from an automatic match of its data and coordinate extents to the visible surface of the display area, you can change the size of the display surface using the zoom controls described below and/or you can change the actual data limits using the **Configure Plot Options** window. When you add or remove a spectrum the apparent size of the display region stays the same and all the spectra now displayed are scaled to fit within it. To return to a state where the visible surface matches the display area you need to press the  and  buttons.

The **Configure Plot Options** window also offers the ability change the colour of the display area background, to draw the lines using anti-aliasing (which reduces the

jagged effects, but at the cost of drawing speed), to modify the reserved area around the plot and do things like position the numeric labels, set the axis gaps *etc.*

The following interactions are provided in the display area.

- Continuous coordinate and data readout: when you move the mouse pointer over the display area the coordinates and data value of the nearest point of the current spectrum are shown in the control panel. If the readout is not updating you have probably moved outside of the extent of the current spectrum (the one selected in the **Displaying:** control).
If the vertical hair is enabled, then this also follows the mouse pointer.
- Region Zoom: you can zoom on a feature by dragging a rectangle over it using the middle mouse button.
- Incremental Zoom: you can increase the zoom factors by one simply by pressing the middle mouse button anywhere in the display area or pressing the `_ / =` keys. If you hold down the shift key these zoom in Y, otherwise in X.
- Scrollbars: when the X or Y zooms make the plot greater in size than the display area scrollbars appear to the bottom and right. Just drag the sliders to scroll.
- Keyboard Scroll: the arrows keys can be made to control the scroll, provided the keyboard focus is in the display area. Make sure of this by clicking the left mouse button while pointed in the display area.
- Interpolated data value: if the keyboard focus is in the display area and the vertical hair is shown, then pressing the space bar will display the interpolated position of the hair, rather than the X coordinate and data value of the nearest position in the spectrum.
- Interactive graphics control: if any interactive graphics figures are shown (such as those that define ranges for fitting backgrounds and spectral lines, or are produced by the controls in the **Graphics** menu), then these can be moved by selecting them (just click on one and little grips appear around the edges) and then dragging them and resized by dragging the edge grips.
- Drop target: you can drop any spectra dragged from the global list and they will be displayed.

The control panel

The control panel area provides controls for interacting with the plot. It also shows details of what spectra are displayed in the plot and a continuous readout of the X coordinate and corresponding data value.

The **Displaying:** drop down list shows the names and current rendering properties of any spectra that are displayed. If you choose a spectrum from this list it becomes the **current spectrum** and is used as the basis for the plot's coordinates (so the readouts now show the values of this spectrum, also the plot will be re-drawn if the coordinate system of the newly selected spectrum is different). The selected spectrum also becomes selected in the browser window (so you see its details more clearly and apply modifications). This spectrum is also the one used by any tools (such as background and line fitting) that work with only one spectrum. If you

remove this spectrum using the **Edit**→**Remove current spectrum** option then the next spectrum in the list becomes the current one.

The **Y limits (%)**: drop down list provides a series of quick cuts for setting the limits of the Y axis. Normally this is the full range, but using a percentage cut can usefully clip the range to reject extreme outlier data values. The data values used are restricted to the limits of then X coordinates (all when automatic ranging is used, the default).

The **Wavelength**: entry control has two functions. The first is to show the current wavelength. The second is to allow you to centre the display area on a specific coordinate. Just type in the value you want to see here and press <Return>. Note that the label of this area modifies to match the label of the X axis, so could say something different to **Wavelength**:

The **:V-hair** checkbox controls whether the ‘vertical hair’ is shown. The vertical hair is just a vertical line that follows the mouse pointer around the display area. It can be quite slow, so isn’t shown by default.

The **X scale**: and **Y scale**: controls change the zoom of the display area. The plus and minus controls increase and decrease the zoom by one, or you can choose a zoom from the drop-down lists, or you can type in a decimal value (which should be greater than equal to one) and press <Return>.

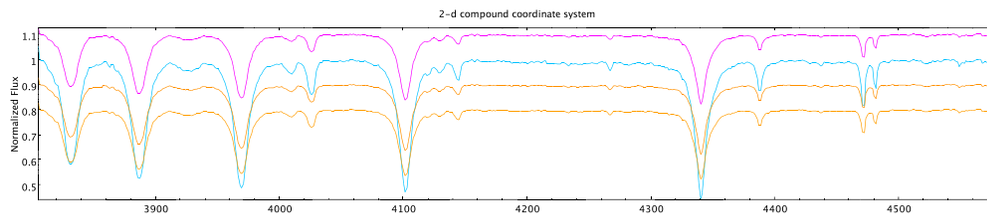
The **:log** controls determine if either of the axes are drawing using log spacing (this is not possible if either axis spans the value zero). Finer control of the log spacing and labelling can obtained using the plot configuration window.

The **Track free** checkbox determines what values are shown in the continuous coordinate readouts. When unchecked (the default) the values shown are the nearest ones of the current spectrum to the pointer position. When checked the readouts show the coordinates under the pointer.

Toolbar and menus

The toolbar region has a series of buttons that act as short-cuts to most of the functions found in the main menus. Some of these create new windows with complex actions that should be consulted for further help.

-  Print a copy of the display area to an installed printer or to a file. The output produced fits the output page format using the current aspect ratio of the display area, so if you’re zoomed by a large factor then you’ll get an output page that has a very small height. If you want a printout of just the visible part then cut it out and display it in a new plot first. Postscript output from this dialogue isn’t encapsulated. The next figure is an example of this output. Note that if you’re looking at a paper copy of this document you’re looking at the postscript output and if you’re looking at the on-line help it shows a scaled JPEG for on-line copies (see item after next).



-  Same as the print option, except, this only offers to write a postscript file and doesn't rely on having any printers installed on the local machine. The output can be in encapsulated form for inclusion in other documents.
-  Make a JPEG or PNG image version of the display area. Again the file produced fits the output graphic using the current aspect ratio of the display area. The default is to produce an image with a pixel-to-pixel correspondence to what you see on the screen (if it were possible to view the whole of the area). Alternatively you can scale the graphic to fit some dimensions that you enter, or just clip the graphic to these dimensions. (Hint: if you want anti-aliased lines drawn in the image select this plotting option first using the plot configuration window).
-  Make spectra fit the display area width. Use this to quickly return to an unzoomed coordinate state and to match the X axis width when you resize the plot window.
-  Make spectra fit the display area height. Use this to quickly return to an unzoomed data value state and to match the Y axis height when you resize the plot window.
-  Show the 'panner' window. You can use this to see a view of the whole extent of the currently displayed spectra (shown at the same aspect ratio as the display area). This also features a red rectangle that shows the extent of the display area, this can be moved to change the display area view.
-  Show the plot configuration window. This contains many controls for setting the display area data limits, changing the plot title and axis labels, drawing grid lines, defining how many ticks to display, changing the amount of space around the plot edges, setting the background colour and whether to draw text or text and lines using anti-aliasing. It also offers facilities to save these settings for future restoration.
-  Create a new spectrum (on the global list) that contains only the visible parts of the current spectrum. You can then display this new spectrum in this or a new plot. The new spectrum is given the name Cut <n> of <current_shortname>. This facility is useful to create printouts of just parts of a spectrum and to speed up processing of very large spectra.
-  Show the region cutter window. This window allows you to graphically define which regions you'd like to cut, delete or replace by linear interpolation in the current spectrum.
-  Show the toolbox for fitting a polynomial to defined regions of the current spectrum. This is used to define backgrounds prior to line fitting, or for subtracting and dividing backgrounds from spectra.

-  Show the toolbox for taking an interpolated curve (*i.e.* a spline drawn on the plot using the features of the **Graphics** menu) and creating a spectrum from it.
-  Show the toolbox for fitting emission and absorption lines. This measures the line position and equivalent width as well as fitting Gaussian, Lorentzian and Voigt profiles to selected wavelength ranges of the current spectrum. Remember to define a polynomial background before using this tool or have your data processed for fitting.
-  Show the toolbox for filtering the current spectrum. The filters offered are simple ones like the average and median over a windowed area, plus more complex ones like smoothing using any of the known line shapes and denoising using wavelet transformations.
-  Show the toolbox for selecting new coordinates or data units for the current spectrum (and therefore plot).
-  Show the toolbox for flipping, shifting or redshifting editable spectra shown in the plot. Also allows you create an editable copy of the current spectrum.
-  Show the toolbox for determining statistics (mean, standard deviation, *etc.*) of the whole or parts of the current spectrum.
-  Show the toolbox for apply an offsets to the displayed spectra so they can be viewed more easily or shown in some order, for instance time of observation.

The **Options** menu contains several control items whose values are preserved between invocations of SPLAT-VO.

- **Options**→**Match coordinates and/or fluxes**. When this is selected the plot will attempt to convert the spectral coordinates and fluxes of all displayed spectra into the systems of the current spectrum, so that they will be aligned when displayed.

Whether this works or not depends on whether the spectra that you have read in contain correct descriptions (or descriptions that SPLAT-VO can make good guesses about) of the spectral coordinates and data fluxes and whether the flux system is supported. If this doesn't work then check that your spectra have valid descriptions using the toolboxes in the main browser window for inspecting and setting spectral coordinates and data units ( and ).

In the case of line identifier catalogues, these will also be transformed to the rest frame of the observed source (*i.e.* they will be red or blue shifted if you have defined the source velocity as part of its coordinate system definition).

This option is switched on by default. If your spectra do not need aligning switch it off for extra performance.

- **Options**→**Match non-flux data units**. If your data units are not understood as fluxes it may still be possible to align them (dimensionally obvious unit conversions are possible for instance mK and K will be aligned correctly, as will eV and keV).
- **Options**→**Match sidebands** If your spectra are dual sideband then they can be aligned taking this into account, or they can be aligned just using the currently

selected sidebands. By default they are not aligned by taking the sideband into account. If you want to do otherwise select this item.

- **Options**→**Match origins** If you have defined origins for your spectra to display offset coordinate systems then selecting this will align the spectra in those coordinates not the absolute ones.
- **Options**→**Match using base system** When matching spectra it is usual to achieve this by transforming all coordinate systems into a common system, this system is wavelength by default. For some systems it may be better to use another common system so by selecting this option you can set the common system used to be that of the current spectrum. This option is most useful when working with velocities, when you want the alignment to happen in that system (because the spectra have different rest frequencies).
- **Options**→**Only display grid axes in visible area** Normally the labelled axes displaying the spectral and data coordinates are drawn to the full size of the display surface of the plot. This is the quickest mode as it reduces the need for re-drawing. If you select this option the axes will be only drawn within the the visible part of the plot, which makes it easier to see the coordinate labels, but means that the axes need to be re-drawn continually when scrolling.
- **Options**→**Only display spectra within axes** Select this option if you only want spectra (including line identifiers) to be drawn within the bounds of the axes. When combined with the previous option to only display axes in the visible area this will potentially slow down scroll operations a lot, so the default is off.
- **Options**→**Line identifiers**→**Load all matching line identifiers** If you select this option then all line identifiers known to SPLAT-VO (including any that you have opened yourself), and whose coordinate ranges fall within that of the plot, will be loaded into the global list and displayed in the plot. Note that this will not work if the spectra displayed in the plot do not have defined spectral coordinates.
- **Options**→**Line identifiers**→**Load all matching pre-loaded line identifiers** If you select this option then all line identifiers already loaded into the global list, and whose coordinate ranges fall within that of the plot, will be displayed in the plot. Note that this will not work if the spectra displayed in the plot do not have defined spectral coordinates.
- **Options**→**Line identifiers**→**Positions track current spectrum** If you select this option then line identifiers that do not have data positions (this is typically true) will be positioned above the current spectrum, rather than displayed in a line along the bottom of the plot.
- **Options**→**Line identifiers**→**Prefix name to labels** If you select this item then the short name of each line identifier spectrum displayed in the plot will be prefixed to the label. This allows you to discriminate between crowded labels. Note that the short name of a line identifier can be modified in the main window, once it is loaded into the global list. For legibility any names that are appended by `_lines`, will have that string removed and any underscores will be replaced by spaces.
- **Options**→**Line identifiers**→**Draw horizontal labels** Select this option if you want line identifier labels drawn horizontally rather than vertically.

- **Options**→**Line identifiers**→**Show dual sideband labels** Select this option if you want line identifier labels drawn for both sidebands of a dual sideband spectrum. The non-default sideband coordinates are displayed along the top edge of the plot and the matching labels will be displayed in red, with either **(USB)** or **(LSB)** postfixed to the label to avoid confusion.
- **Options**→**Error bar auto-ranging**. When auto-ranging a spectrum that has associated errors the limits do not include the extent of any error bars that will be drawn, if you want this to be otherwise select this option.
- **Options**→**Draw errors as spectrum**. If this option is selected and any of the displayed spectra have errors, then those spectra with errors will be drawn using the error value as each point on the line rather than the data values (and conversely the data values will be used to draw error bars, if that is enabled for any of the spectra). When displaying spectra without any errors this option has no effect.
- **Options**→**Auto-fit percentiles in Y**. When this option is selected choosing a percentile limit in Y results in the plot also being forced to fit the height of the plot.
- **Options**→**Short names in menu lists**. When this option is deselected the long names (usually file names) are shown in the drop-down menu.
- **Options**→**Display synopsis**. When this option is selected a description of various properties of the current spectrum is displayed overlaid on the plot. The description can be moved around by dragging.

The items possibly included in the synopsis are:

Name: the short name.

Telescope: the telescope, instrument and it's backend (TELESCOP/
INSTRUME/ BACKEND).

Object: The target, molecule and molecular transition being observed
(OBJECT/ MOLECULE/ TRANSITI).

Date obs: The date of the observation (DATE-OBS or Epoch in UTC).

Elevation: The observatory elevation (ELSTART or $0.5*(ELSTART+ELEND)$).

Exposure: Exposure time for the spectrum (EXTIME).

or

Exposure (median): Exposure time for the spectrum (EXP_TIME).

or

Exposure (elapsed): The time elapsed during exposure (INT_TIME
otherwise DATE-END-DATE-OBS).

Exposure (effective): Effective exposure time (EXEFFT).

Coord sys: The coordinate system code (System).

Spec position: The position of spectrum on the sky (EXRAX, EXDECX).

Src position: The position of the observation centre (EXRRA, EXRDEC).

or

Img centre: The centre of the originating image (EXRA, EXDEC).

Offset: The offset of spectrum from observation centre (EXRRAOF,
EXRDECOF or EXTRAOF, EXDECOF).

- Doppler RA, Dec:** The reference position (RefRA,RefDec).
SourceVel: The source velocity (SourceVel).
SourceVRF: The source velocity reference frame (SourceVRF).
SourceSys: The system of the source velocity (SourceSys).
StdOfRest: The standard of rest (StdOfRest).
RestFreq: The rest frequency (RestFreq).
ImagFreq: The image sideband equivalent of the rest frequency (ImagFreq).
Channel spacing: The channel spacing (derived value).
Number of channels: Number of coordinate positions.
TSYS: System temperature (TSYS).
or
TSYS (median): Median system temperature (MEDTSYS).
TSYS (est): System temperature (derived value, see region statistics window).
TRX: Receiver temperature (TRX).

These are somewhat tuned for submillimetre observations at the JAC and for spectra extracted by GAIA from cubes, so many may be missing for spectra from other sources. Names in parenthesis are either the FITS keywords or AST attributes used to obtain the values.

The Graphics menu

The **Graphics** menu provides methods for drawing text and a range of figures on the plot. This allows you to add annotations, but also provides the basis for the interactive graphics used by the various toolboxes.

To draw a figure on a plot you need to select an option from the **Drawing mode** sub-menu and then, depending on the figure type, either drag out a region or select fiducial points. When creating a figure using points these are completed using a double click on the last point.

In the **Drawing mode** menu the currently available figure types are:

-  a straight line,
-  a rectangle,
-  an ellipse,
-  a poly line (last point not connected to first),
-  a polygon (last point connected to first),
-  freehand curve,
-  a text string,
-  an interpolated curve (splines and special spectral shapes),
-  a rectangle which only moves along X (indicates a range of coordinates).

There are also two special modes:

-  select figure,

 edit figure.

When the mode is select (this is usually the default state), clicking on a figure selects it. You can then drag it around to move it, or drag one of the grips (the little squares) to change the shape. To select more than one figure you hold down the <Shift> while clicking.

The edit mode can be used to add new points to a polyline, polygon or interpolated curve. Just select this, click on the figure you want to edit and then start adding new points. You can also edit text strings, just choose the edit mode and click on the text, this produces a dialogue with the text for editing.

To change the properties of a figure (line width, fill/outline colour, font *etc.*) just select the figure and then choose the option you want to change from the various menus. You can define these before creating a figure too.

Interpolated curves are like polylines in that they are defined by a series of points, but they can only have increasing or decreasing X coordinates (so cannot double back on themselves). They are intended for drawing figures that are related to spectra, specifically spectral backgrounds and lines. In fact if you draw an interpolated curve you can convert it into a spectrum for removing from a real spectrum using the **Generate spectra from interpolated lines** toolbox. The types of interpolation scheme available are:

- Hermite, interpolation based on Hermite polynomials. The effect is supposed to construct reasonable analytic curves through discrete data points (*i.e.* like those a human would produce).
- Akima, interpolation based on Akima splines. Like Hermite this supposed to construct human-like analytic curves.
- Cubic, interpolation using natural cubic splines.
- Polynomial, interpolation using a simple polynomial of order one less than the current number of points.
- Linear, interpolation using straight-lines (like polyline, except it has sorted X coordinates).
- Gaussian, interpolation using a Gaussian line-profile. This can only have four reference positions (and is initialised from the first two indicated on the plot), these define the left extent, width, height/centre and right extent. The order will switch if these are dragged past each other (so dragging the width grip beyond the left extent grip will switch their meaning).
- Lorentz, interpolation using a Lorentzian line-profile. Works like the Gaussian curve.
- Voigt, interpolation using a Voigt line-profile. This also works like the Gaussian curve, but has five characteristic positions, left extent, Gaussian width, height/centre, Lorentzian width, right extent.

Once you have created your figures you may want to save them for restoration at a later time, this can be done using the **Graphics**→**Save/restore figures** dialogue. Figures are saved using the physical coordinates of the spectrum, so should be re-drawn at the correct wavelengths *etc.* on new plots.

Accelerator keys

- **Control-p** Activate print to printer dialog.
- **Control-t** Activate print to postscript file dialog.
- **Control-j** Activate print to JPEG file dialog.
- **Control-b** Scale spectrum to fit visible width.
- **Control-h** Scale spectrum to fit visible height.
- **Control-a** Show the panner window.
- **Control-o** Show the plot configuration window.
- **Control-w** Close the window.
- **Control-v** Cut out the current view of the spectrum.
- **Control-r** Activate cut out regions of spectrum window.
- **Control-y** Activate fit polynomial to background window.
- **Control-g** Activate generate spectrum from drawn curve window.
- **Control-l** Activate fit spectral lines window.
- **Control-f** Activate filter spectrum window.
- **Control-u** Activate quick change units window.
- **Control-l** Activate flip compare window.
- **Control-s** Activate statistics window.
- **Control-[1-9]** Select drawing mode for various graphics figures.
- **Delete** Delete any selected graphics figures.
- **Shift-Delete** Delete all graphics figures.
- **r** Raise any selected graphics figures.
- **l** Lower any selected graphics figures.
- **h** Hide or unhide all graphics figures (toggle).
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.3 The query VO for spectra window

This window provides access to the Virtual Observatory (VO). Using it you can search a region of sky for spectra or search for spectra associated with a specific astronomical object. Once you have done that you can then download all or some of these spectra for display and analysis in SPLAT-VO.

In the VO spectra are made available by SSAP servers (SSAP stands for Simple Spectral Access Protocol) and using this window you can also select which of these servers you want to query. That is important if you are only interested in spectra covering specific wavelengths or types of observations. These searches can be further refined using additional parameters like a range of wavelengths or times. Additional **Optional Parameters** are also provided for even more refinement, but note that these may not be supported by all SSAP servers.

The screenshot shows the 'Starlink SPLAT-VO: Query VO for Spectra' window. It is divided into three main sections:

- Service selection options:** Includes 'Observed data' and 'Theoretical data' radio buttons, 'Wave Band' (Radio, Optical, X-ray, Millimeter, UV, Gamma-ray) checkboxes, and 'Tags'.
- Input parameters:** Includes 'Search parameters' (Object: 60Cyg, Radius: 100.0, Band: MAXREC, Time: /), 'Query Format' (None), 'Wavelength calibration' (None), and 'Flux calibration' (None).
- Optional parameters:** A table with columns 'Use', 'Name', 'Value', and 'UCD'. Parameters include SPECRES, TARGETSPECTRY, VERSION (1.02), ARCHIVEID (ZCOSMOS_BRIG...), RETURNFORMAT (VOTable (SSA)), CAMERA, and IMAGE.
- SSAP Servers:** A list of servers with columns 'short name' and 'title'. Servers include TBL Narval, CALIFA DR2, Polarbase SSAP, HFA, CDPS SSA, etc.
- Query results:** A table with columns 'In...', 'object', 'coord_obs', 'min_wavelength', 'max_wavelength', 's_fov', 'tstart', 'tstop', and 'format'. It shows a list of 43 results for object 60Cyg.

The figure above shows the window after a query has been made. The window is divided in three main areas. The left area is the SSAP server selection area, where the current list of SSAP servers is shown and also options to select a subset of servers to be queried. The upper right area is the query parameters area, where query options can be entered. The right-lower part is the results area, showing the query results.

Server selection

You can select one or more servers from the **SSAP Servers** list by clicking on them. It's also possible to select a subset of servers based on their given metadata. Selection can be made according to data source (observation or theory data), and waveband. It's also possible to add tags to one or more servers, so only the tagged servers will be queried if the tag is selected. When right-clicking on a server in the **SSAP Servers** list you get a popup menu containing options to tag, to remove, or to see information on the service. To add a new SSAP server, which is currently unknown to the VO, to the current server list, click on the **Add New Server** button. A

new window will open where the information about the new server can be filled in. The **Query registry** allows you to query a VO registry for a list of all the SSAP servers it has registered.

SSAP query

To lookup the coordinates of an object just type in an identifier in the **Object** field and either press <Return> or the **Lookup** button. Alternatively if you know the coordinates (FK5/J2000) of the region of sky you can just enter the coordinates in the **RA** and **Dec** fields. It is also possible to query the servers about an object by name, useful for solar system work, in that case just enter a name and leave the **RA** and **Dec** fields empty.

The **Radius** value is a number of arcminutes to search about the position. The **Band** entries define a lower and upper limit, in meters, for the spectral bandpass (not all SSAP servers support this option, those that do may offer just an upper limit and allow you to set a wavelength to include by setting just a lower limit). The **Time** entries define either a time that should match the time the spectra were taken, or a range in which they should match (again not all servers support this option). The **MAXREC** field will define the maximum number of spectra that will be retrieved from each selected service. The file format, wavelength and flux calibration options can also be chosen. If the value is "None", the query will not contain these parameters.

Besides the mandatory parameters, some servers also support a set of other metadata parameters. The **Optional Parameters** panel shows the metadata parameter supported by at least one of the selected services. The checked parameters having a non empty value field will be added to the query.

Once you have identified the region to search or object press the green **SEND QUERY** button to contact the current list of SSAP servers. To make these queries it may be necessary for you to tell SPLAT-VO about your local web-proxy server (this is the same as that you use in your Internet browser), select the **Options**→**Configure proxy...** item to do this.

Query results

Once the SSAP query is complete lists of spectra that the servers hold will be shown in the **Query results:** region. There is a tab of results for each server. If no tab has been created for a selected server, then the server has no spectra for the selected region or is not available at the moment.

When right-clicking on a spectra, a popup menu will appear. From this menu you can download or display the spectrum, see its fields (and, if supported, see a preview), or copy the contents of the current table, line or cell to the clipboard.

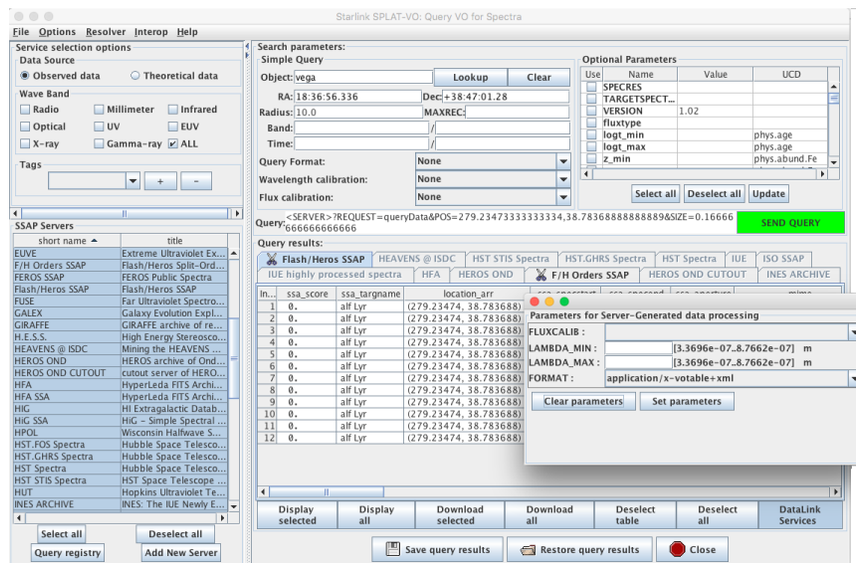
The next action is to select which spectra you want to download and view in SPLAT-VO. If you press **Display all** then all the spectra from all the servers will be downloaded and displayed in a single plot. Clearly this may take quite sometime if there are many spectra, so you can also download and display a subset. To do this select the rows that contain the spectra that you want to see (you can extend selections by using the control and shift keys while selecting a row and you can select spectra from more than one tab at a time) and then press the **Display selected** button.

Once the spectra have been downloaded they should appear together in a single plot. If this isn't the case then some of the spectra are probably invalid and you will need to add them to a plot yourself (see the section on displaying more than one spectrum). If you want SPLAT-VO to match the coordinate systems and fluxes of the spectra then make sure that the **Options**→**Match coordinates and/or fluxes** item is still switched on in the plot window.

To view a single spectrum quickly just double click on its row. This will download the spectrum and display it in its own plot window.

Server-side manipulation with DataLink

SPLAT-VO supports server-side manipulation through **DataLink**. After a SSAP Query is sent, SPLAT-VO checks the responding servers for **DataLink** parameters, and marks them with a symbol (✂). If at least one **DataLink** supporting server has been found, a **DataLink** button will appear below the query results. By clicking on it, **DataLink** will be activated, and all results tabs will be deactivated except the results from servers that support **DataLink**. When you choose one of the active services, a window will appear, where the respective parameters for sever-side manipulation can be entered. The parameters will be set by clicking on the **set parameters** button. While **dataLink** is activated and the parameters are set, the spectra you select and display or download will be manipulated using the chosen parameters. To deactivate **dataLink** and have retrieve the original spectra just click on the **dataLink** button again.



Implementation status

SSAP is a protocol being developed by the IVOA (International Virtual Observatory Alliance¹). Currently spectra returned from SSAP servers may only be in basic formats, that is simple FITS and VOTables (partially some SED data model serialisation). However, the SPLAT-VO data model includes errors as well as coordinates and data values.

Support for spectral coordinates is provided by the Starlink AST library (SUN/211) using an implementation of the FITS-WCS paper III *Representations of spectral coordinates in FITS* (Greisen, Valdes, Calabretta & Allen). AST also provides the flux and data value conversion utilities used by SPLAT-VO which is based on interpreting the units strings system described in FITS-WCS paper I *Representation of World Coordinate in FITS* by Greisen & Calabretta. Conversion between fluxes is currently restricted to flux per unit wavelength and flux per unit frequency is provided and does not require any 'dimensional analysis' information to be present, just correctly formed units strings.

¹<http://www.ivoa.net/>

Server metadata information is provided by the respective maintainers, and may not be complete or accurate.

Accelerator keys

- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.4 Query VO using ObsCore TAP

Besides SSAP, SPLAT can also query VO services supporting **ObsCore** (Observation Data Model Core) components via TAP (Table Access Protocol). These services are known as ObsTAP services. There are two ways of doing that in SPLAT-VO by using a cone search interface, like the SSAP query, or by sending an ADQL query. ADQL stands for Astronomical Data Query Language, based on SQL. There are introductions and tutorials for ADQL for instance at <http://docs.gvo.org/adql> (Markus Demleitner) or <http://is.gd/ADQLTutorial> (Simon Murphy).

Server Selection The server selection interface works in the same way as the server selection in the SSAP window. Just that the servers list contains ObsTap services instead of SSAP services, and there are no optional parameters, as all parameters can be queried using TAP.

Simple Search The simple search interface uses the same parameters as the SSAP cone search, with the same functionality.

The screenshot shows the SPLAT ObsCore Browser window. On the left, there is a 'Simple search' tab with 'Query Parameters' for Object (vega), RA (18:36:56.336), Dec (+38:47:01.28), Radius (10.0), and Band/Time fields. Below this is a list of 'Obscore Services' including GAVO DC TAP, HIPASS TAP, MACHO TAP, SkyMapper TAP, SUBARU_SUP, wgsdsstap, WiggleZ TAP, XCATDB, and XCATDB+. On the right, the 'Query results' tab shows a table with columns s_ra, s_dec, s_fov, and s_regio. The table contains 20 rows of data, all with s_regio values of 'Polygon ICRS UNKN...'. At the bottom, there are buttons for 'Display selected', 'Display all', 'Download selected', 'Download all', 'Deselect table', and 'Deselect all'.

ADQL Search The ADQL search is more flexible and more powerful than the simple cone search. The main part is a text area, which can be edited. The beginning of the query

```
SELECT TOP 10000 * from ivoa.Obscore WHERE dataproduct_type='spectrum'
```

will get the first 10000 spectra from the selected servers. The TOP value may be changed to another value, if necessary. This query should be then refined or completed, either by choosing the parameters and expressions from the options provided (which will create a simple query) or by editing directly the the text area. Please refer to the ObsCore standard for the available parameters. (<http://www.ivoa.net/documents/ObsCore/>).

The screenshot shows the SPLAT ObsCore Browser interface. On the left, there is a search panel with 'Simple search' and 'ADQL search' tabs. The ADQL search tab is active, showing a query: `SELECT TOP 10000 * from ivoa.Obscore WHERE dataproduct_type='spectrum'`. Below the query is a list of 'Obscore Services' with columns 'short name' and 'title'. The 'CADC Table Query (TAP) Service' is selected. On the right, the 'Query results' window displays a table with the following columns: `dataproduct_type`, `calib_level`, `obs_collection`, `facility_name`, and `instru`. The table contains 20 rows of data, all with `dataproduct_type` set to 'spectrum'. The `obs_collection` column lists 'GAVO DC TAP', 'XCATDB', and 'XCATDB+'. The `facility_name` column lists 'DAO 1.2-m' and 'DAO 1.8-m'. The `instru` column lists 'McKellar Spe' and 'Cassegrain S'. At the bottom of the results window, there are buttons for 'Display selected', 'Display all', 'Download selected', 'Download all', 'Deselect table', and 'Deselect all'.

The example below shows a query for spectra including 6500 angstroms, spectral resolution better than 15 angstroms, spatial resolution better than 2 arcseconds FWHM, and exposure time longer than 3600 seconds:

```
SELECT * from ivoa.Obscore
WHERE dataproduct_type='spectrum'
AND em_min < 650E-9
AND em_max > 650E-9
AND em_res_power > 6500/15.
AND s_resolution < 2
AND t_exptime > 3600
```

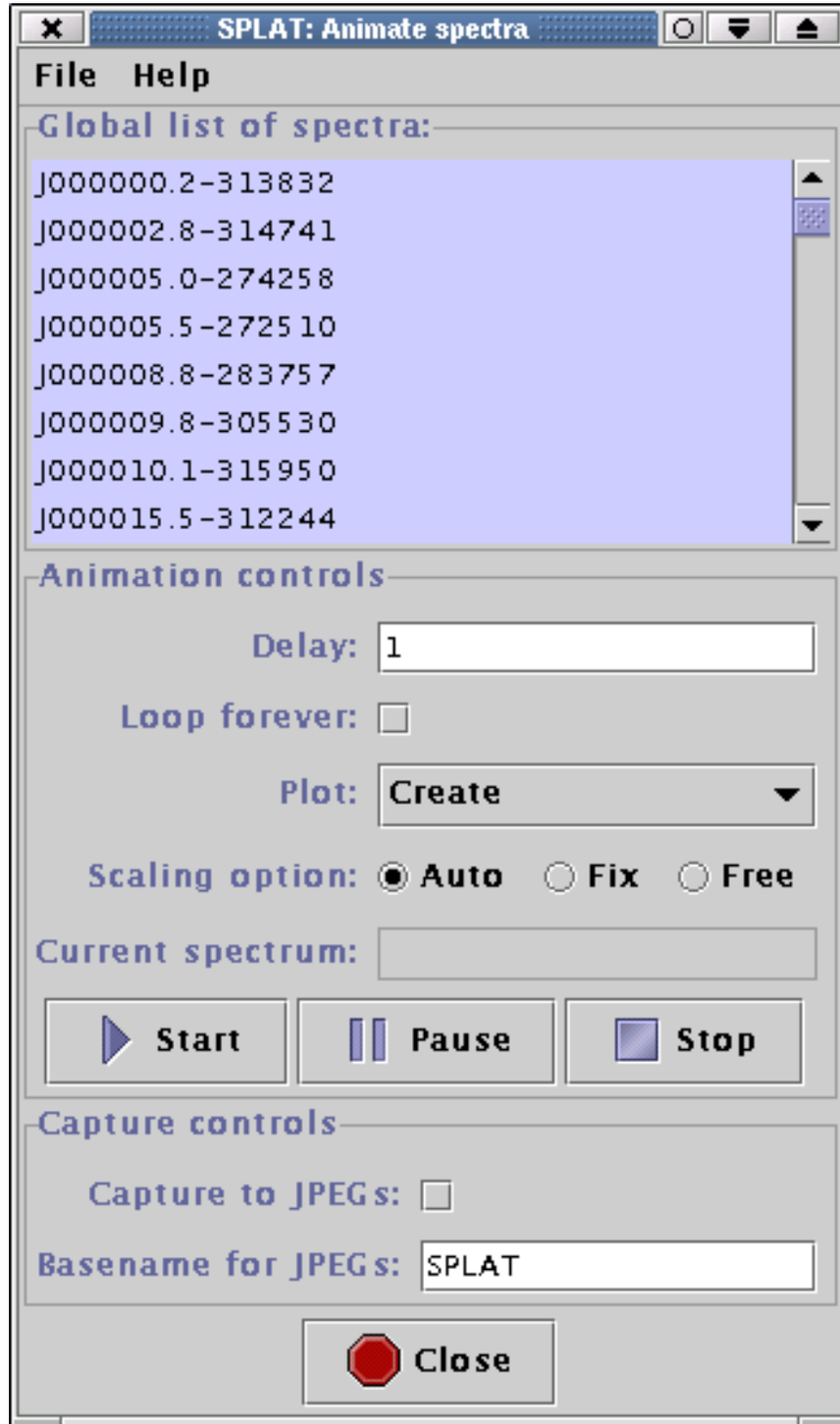
Implementation status

ObsCore and TAP are being developed by the IVOA (International Virtual Observatory Alliance²).

²<http://www.ivoa.net/>

3.5 The animation window

This window allows you to display a series of spectra from the global list, one after the other. It is possible to display these into an existing plot, that may already be displaying other spectra you'd like to use for comparison, or you can use a new plot.



With careful control of the plotting options you can display into a plot with fixed data ranges, or

see each spectrum plotted as autoscaled.

It is also possible to use this window to generate a series of JPEG or PNG images that you could combine into a movie of some kind.

Global list of spectra:

As in the browser window this is a view of the spectra currently stored in the global list. Initially all the spectra in the list are selected, but you may modify which spectra to animate by creating your own selection (reminder: the combination of shift-left-mouse click selects the range between the last selected row and the current position, and the combination control-left-mouse click, either adds the current row to the selection, or removes it, if already selected).

Animation controls:

The **Delay:** field accepts a decimal number of seconds to pause between displaying the selected spectra. You can modify this during an animated sequence by pressing the <Return> key.

The **Loop forever:** checkbox determines whether the animation sequence is restarted when all the selected spectra have been displayed. If set then you need to press the **Stop** button to stop the animation.

The **Plot:** drop-down list contains a list of all the currently available plots and the special entry **Create**. If the **Create** option is chosen then a new plot will be created to display the animation, otherwise the animation will take place in the selected plot.

The **Scaling option:** radio buttons control how a spectrum is scaled when it is displayed.

- **Auto** selecting this option (the default) makes each spectrum, and any other spectra already displayed in the target plot, fit themselves to the width and height of the plot display area.
- **Fix** selecting this option makes each spectrum respect the data limits already applied to the plot. So to make effective use of this option you need to display into an existing plot and have set the data limits (using the plot configuration window). It's expected that this will be most useful when comparing a list of spectra against a single pre-plotted spectrum, or when creating an animation sequence for a movie (so you need to also use the **Capture controls**).
- **Free** selecting this option makes the spectrum add itself without constraints to the plot. This is essentially the same as just adding a spectrum to a plot using the browser window controls (*i.e.* the data limits of the plot are produced by combining the data limits of any pre-plotted spectra and the one added, these are all scaled to fit).

The **Current spectrum** label just displays the name of the spectrum currently being added to the target plot. This information is repeated in the plot itself as the spectrum becomes the current one in the plot and is named in the **Displaying:** label.

The **Start**, **Pause** and **Stop** buttons do what is expected.

Capture controls

The capture controls are for creating a series of JPEG or PNG images of the plot as it displays each spectrum. To enable (after getting the required effect using the **Animation controls**) just click on the **Start capture:** check box. The output file names will start with what name you type into the **Basename for graphics files:** entry and have an integer number appended, plus the file type .jpg or .png. So a default sequence would be SPLAT0.jpg, SPLAT1.jpg, SPLAT2.jpg,... You'll then need to find some software to convert these into a movie. The ImageMagick convert command (when suitably configured) will convert these files into an MPEG or animated GIF. The image below was created with the command (note that the images were also scaled – view the html documentation or the pdf documentation in a viewer that supports animation such as Adobe Reader to see the animation.):

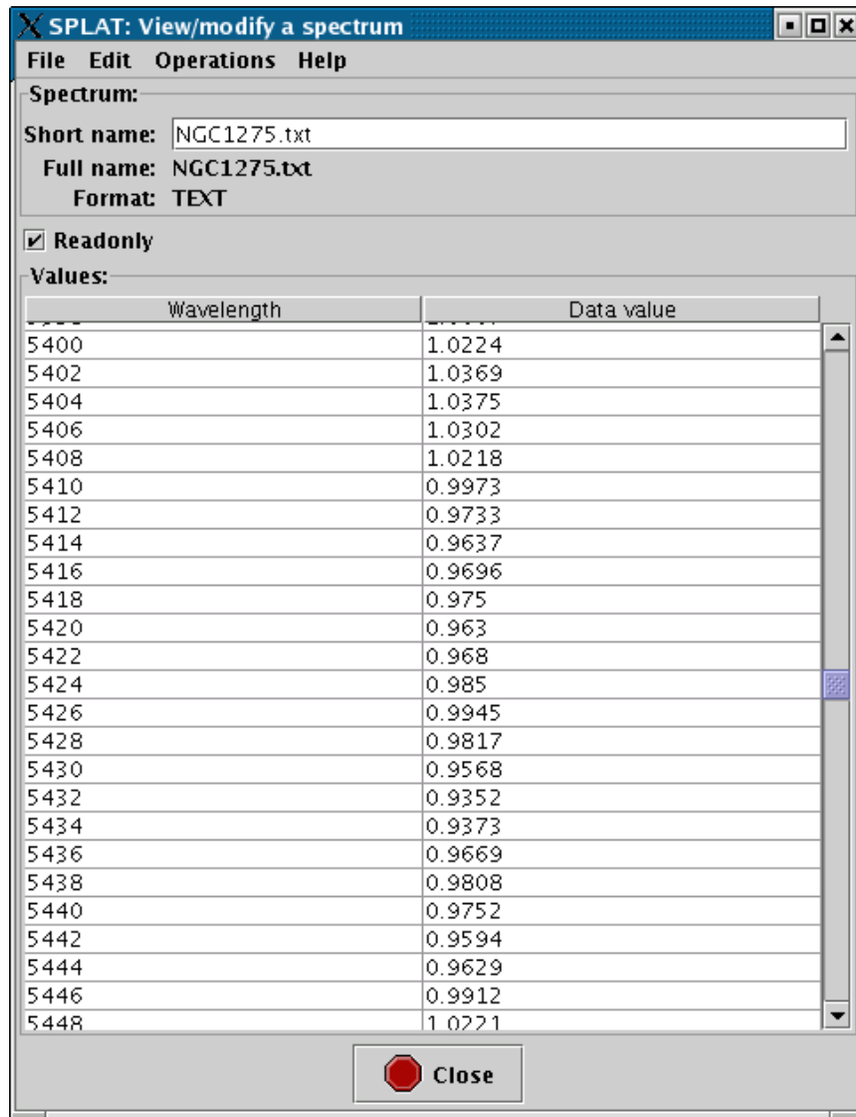
```
% convert -delay 50 -loop 0 SPLAT0.jpg SPLAT1.jpg SPLAT2.jpg \  
splatmovie.gif
```

Accelerator keys

- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.6 The view and modify spectral values window

The purpose of this window is to allow you to see the actual data values of a spectrum and to make modifications to them. The spectrum being shown has its basic details at the top of the window and the values of the spectrum are shown in a table area that you can scroll.



Modifications

If you want to modify any of the values, then you need to have a spectrum that isn't readonly. Spectra read from files always have readonly values, so this will generally be the case. The simplest way to get a modifiable spectrum is to untick the **Readonly** control. This will make a memory copy of your spectrum, enter this on the global list and then load it into this window replacing the previous spectrum. The new spectrum will be called 'Copy of <oldname>'.

Note that if you want to plot this new spectrum you'll need to go back to the main window and do this yourself (plots of the spectrum that this is a copy of are not going to show any modifications that you now make!).

Editing single values

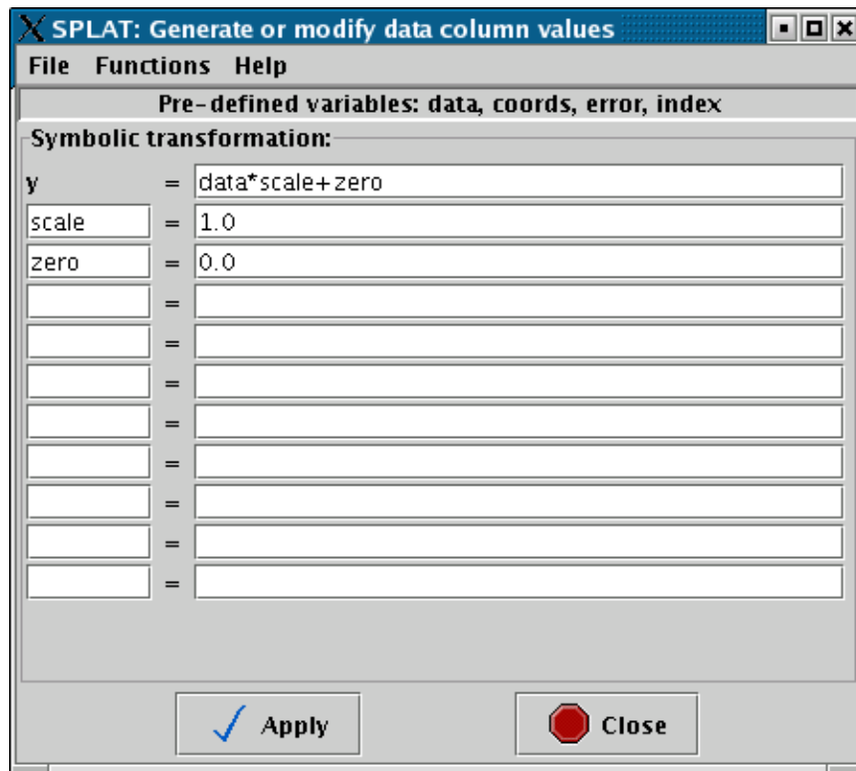
To change a single value of any kind you can now just select the table cell (usually by double clicking on it) and type in the new value. Press Return to make the edit happen (this should immediately be shown in any plots of this spectrum).

Editing whole columns

You can make global changes to a whole column of values using special dialogues that either offer special functions (linear transformations), or allow you to define an algebraic expression (in terms of the existing values or a simple index counter starting at 1 for the first row).

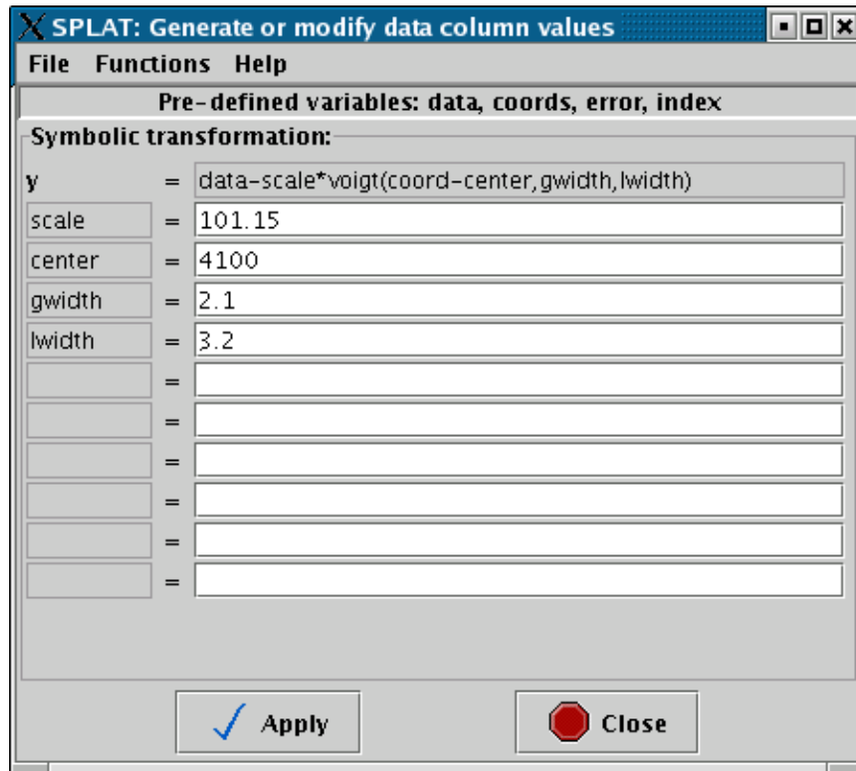
Editing the data values column

To apply a global edit to the data values column you need to select the **Operations**→**Modify data column** item. This should produce a new window.



In which you can see an example expression $data * scale + zero$, with the *scale* value set to 1.0 and the *zero* value to 0.0. A list of preset expressions that should be generally useful is found in the **Functions** menu.

Naturally if you applied this expression then the data values would stay the same. A more complex example is shown next.



This presents the possibility of subtracting a single Voigt profile (the top part of this is greyed out as Voigt is a built-in function that cannot be represented as an algebraic expression).

If you need to use an expression of your own you might like to view the full list of functions that are available in the MathMap section of the KAPPA SUN. A lot of C-like intrinsics are available (*abs, atan/sin/cos, tan/sin/cos, log, pow sqrt, etc.*), which can operate on the data columns *data, coord* and *error*, as well as the virtual *index* column.

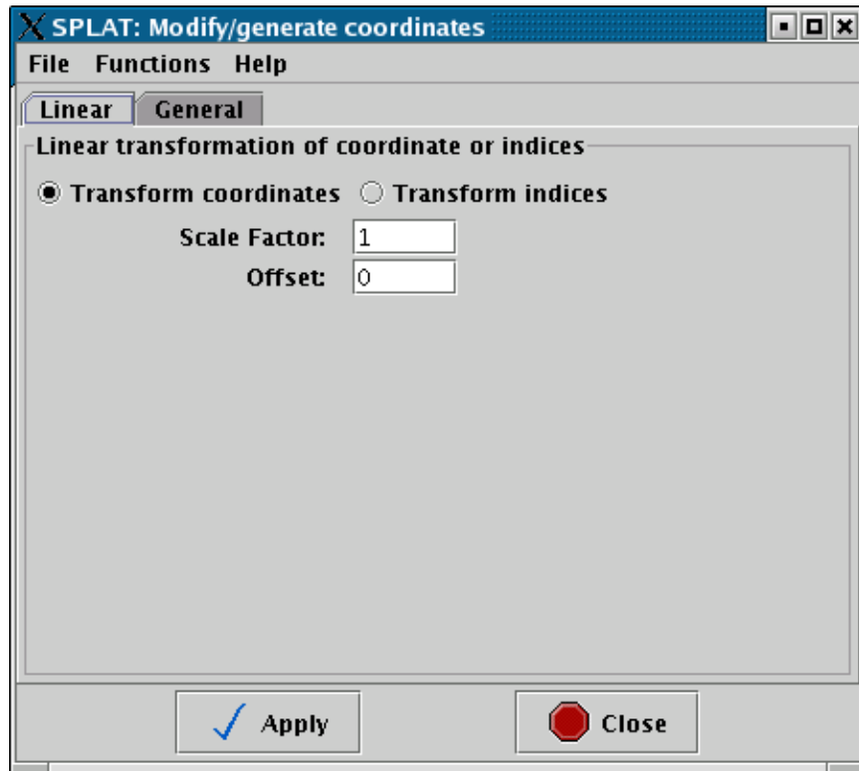
Editing the error values column

Editing the error column is much like editing the data column, except that you can actually create a new error column, see the **Operations**→**Modify error column** item.

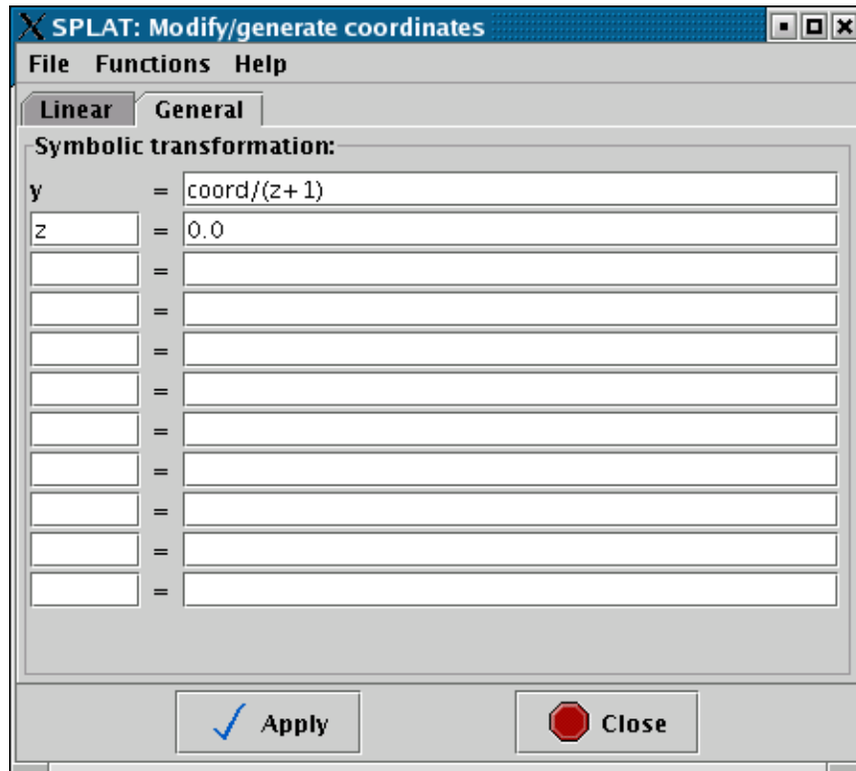
The error column can also be deleted using the **Edit**→**Delete error column** item.

Editing the coordinates column

Globally editing the coordinates can be done using the **Operations**→**Modify coordinate column** window.



A simple built-in option of applying a linear transformation is offered (useful for when the coordinates are in fact some index related to wavelength). Under the **General** tab you can use algebraic expressions like those for modifying the data columns.



In this case the pre-defined expression for blue-shifting a spectrum in some wavelength units is shown.

Note: when you modify coordinates either singly or as a column any coordinate system information will be lost. If you want to retain this you'll either have to re-enter this information or use the conversion facilities of the **Coordinate system attributes** window (using this other window you can transform between apparent place, Earth to Sun to Local Group *etc.* and to a source rest frame that has a velocity in a given direction, so you can correct for red/blue shift that way, in fact this is the recommended way).

Undoing and redoing changes

Using the **Edit** menu in the main view/modify window you can undo and redo any modifications that you make to the spectrum. The changes are propagated immediately so you can see these effects in any plots of the spectrum.

Accelerator keys

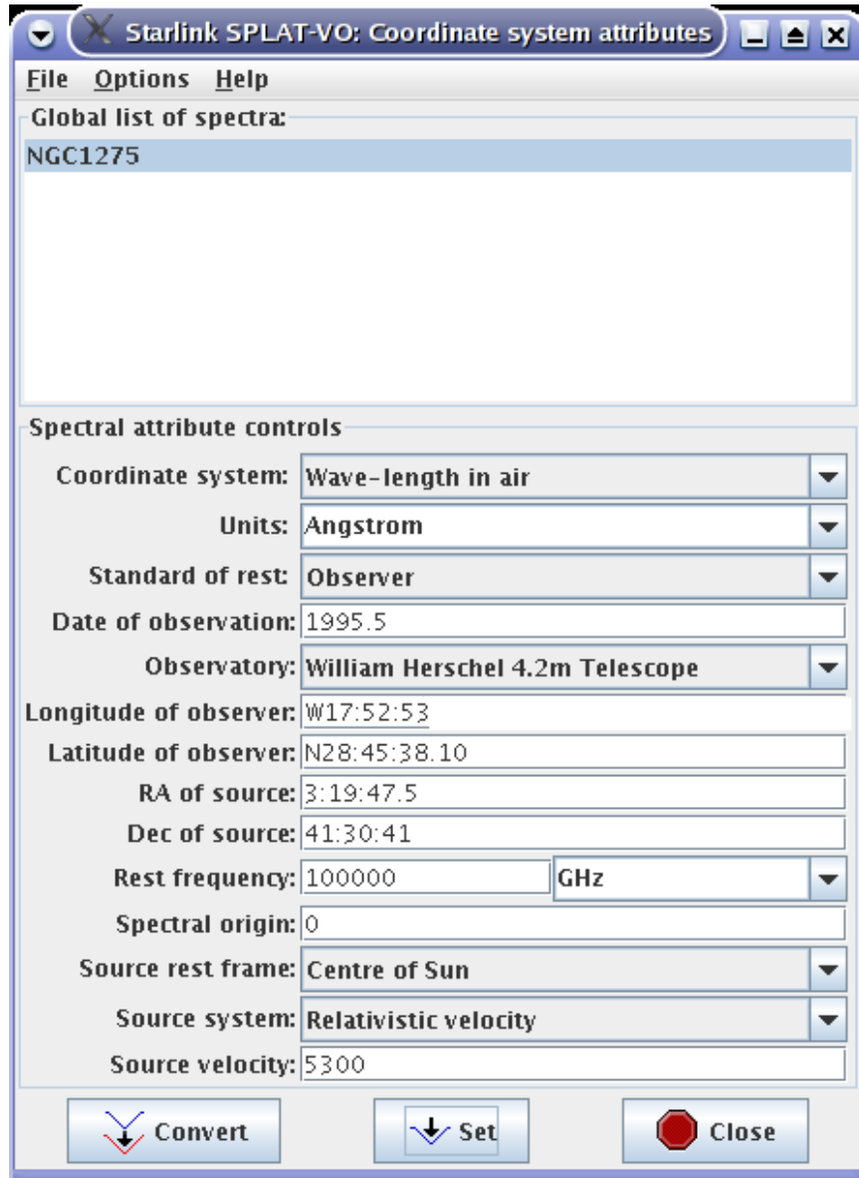
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.7 The spectral coordinate system attributes window

This window allows you to define the coordinate systems that describe your spectra. The most trivial use of these is simply to document what type of coordinate and units your spectra are measured in, examples are wavelength in air in angstroms, frequency in GHz, velocity in km/s, *etc.* When supplied these values will be used to annotate any plots, but can also be used to compare spectra in different coordinate systems and to transform a spectrum from one coordinate system to another.

A more complete description of the coordinate system also includes two further pieces of information, the time and place when the observation was made and the direction and speed of what was being observed. Taken together these define the 'rest frames' of the observation and the source.

In the following picture you can see that I have an observation (topocentric) of NGC1275 (RA: 03:19:47.49, Dec: 41:30:40.5 FK5 J2000) made in air, calibrated in angstroms, taken in June 1999 on the William Herschel Telescope. I've also entered that NGC1275 as observed from the centre of the Sun (heliocentric) has a recessional (relativistic) velocity of 5300km/s. Alternatively if you know the necessary redshift you can enter that value by selecting a **Source system of Redshift** (other possibilities are radio velocity, optical velocity and beta factor).



To apply this to a spectrum that has no coordinate information (or incorrect coordinate information) you need to press the **Set** button. Note that you can apply the same coordinates to more than one spectrum by selecting them all in the view of the global list shown in this window.

Now say that you wanted to compare this spectrum with a standard set of emission lines, obviously taken at rest in a laboratory. If you load the emission line catalogue and display it in the same plot (making sure that the **Options**→**Match coordinates and/or fluxes** item is switched on) the labels will be red-shifted to the rest frame of the source (*i.e.* where the lines from the galaxy originated). This is shown in action in the next figure where the lines before matching are shown in blue and red after matching.

3.8 The data units window

This window allows you to inspect, set or convert the units of the data values of your spectra. The most trivial use of these is simply to document what type of units your spectra are measured in. However, a more powerful use of data units can be made when displaying more than one spectrum in a plot. In that case you can request that the data values of all the spectra are transformed into the same system, so you can compare values in Janskys with those in other flux per unit frequency and per unit wavelength systems (note to activate this option you must have the **Options**→**Match coordinates and/or fluxes** menu item selected in the plot window). For instance all the following systems are understood as fluxes and can be intercompared:

Jy	(Jansky)
W/m ² /Hz	(W * m ⁻² * Hz ⁻¹)
W/m ² /Angstrom	(W * m ⁻² * Angstrom ⁻¹)
W/cm ² /um	(W * cm ⁻² * um ⁻¹)
erg/cm ² /s/Hz	(erg * cm ⁻² * s ⁻¹ * Hz ⁻¹)
erg/cm ² /s/Angstrom	(erg * cm ⁻² * s ⁻¹ * Angstrom ⁻¹)

Plus variations like *Joules* instead of *ergs* or maybe *Nm*, etc. When reading unit strings like W/m²/Hz you should say Watts per metre squared per Hertz, which is actually mathematically the same as the expressions shown in to the right in parentheses (W * m⁻² * Hz⁻¹ in this case). Note that dimensionally equivalent unit strings to those above maybe be recognised and re-formatted.

In addition to flux systems dimensionally similar ones can also be used and transformed between in a plot, things like temperature. SPLAT-VO understands that *K* and *mK* are temperatures in Kelvins and milliKelvins, same for units like *eV* and *keV*.

The units are described using strings that follow the conventions in FITS-WCS paper I *Representation of World Coordinate in FITS* by Greisen & Calabretta, and processed by the Starlink AST library (SUN/211). The following section is copied from SUN/211 for convenience. Not all units are relevant to data values (AST also provides the underlying coordinate transformations used in SPLAT-VO) and as noted above not all possible unit strings can be understood as fluxes (although support for more types is expected, for instance temperatures and magnitudes).

The adopted syntax is that described in FITS-WCS paper I *Representation of World Coordinate in FITS* by Greisen & Calabretta. We distinguish here between “basic” units and “derived” units: derived units are defined in terms of other units (either derived or basic), whereas basic units have no such definitions. Derived units may be represented by their own *symbol* (e.g. “Jy”—the Jansky) or by a *mathematical expression* which combines other symbols and constants to form a definition of the unit (e.g. “km/s”—kilometres per second). Unit symbols may be prefixed by a string representing a standard multiple or sub-multiple.

In addition to the unit symbols listed in FITS-WCS Paper I, any other arbitrary unit symbol may be used, with the proviso that it will not be possible to convert between spectra using such units. The exception to this is if both spectra refer to the same unknown unit string. For instance, an axis with unknown unit symbol “flop” *could* be converted to an axis with unit “Mflop” (Mega-flop).

Unit symbols (optionally prefixed with a multiple or sub-multiple) can be combined together using a limited range of mathematical operators and functions, to produce new units. Such expressions may also contain parentheses and numerical constants (these may optionally use “scientific” notation including an “E” character to represent the power of 10).

The following tables list the symbols for the basic and derived units which may be included in a units string, the standard prefixes for multiples and sub-multiples, and the strings which may be used to represent mathematical operators and functions.

Basic units		
Quantity	Symbol	Full Name
length	m	metre
mass	g	gram
time	s	second
plane angle	rad	radian
solid angle	sr	steradian
temperature	K	Kelvin
electric current	A	Ampere
amount of substance	mol	mole
luminous intensity	cd	candela

Prefixes for multiples & sub-multiples					
Sub-multiple	Name	Prefix	Sub-multiple	Name	Prefix
10^{-1}	deci	d	10	deca	da
10^{-2}	centi	c	10^2	hecto	h
10^{-3}	milli	m	10^3	kilo	k
10^{-6}	micro	u	10^6	mega	M
10^{-9}	nano	n	10^9	giga	G
10^{-12}	pico	p	10^{12}	tera	T
10^{-15}	femto	f	10^{15}	peta	P
10^{-18}	atto	a	10^{18}	exa	E
10^{-21}	zepto	z	10^{21}	zetta	Z
10^{-24}	yocto	y	10^{24}	yotta	Y

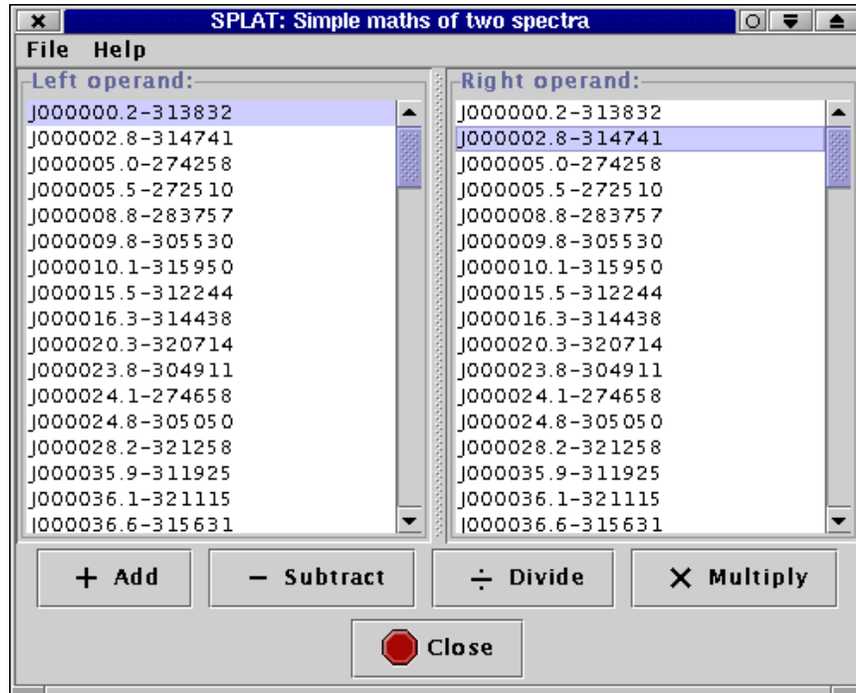
Mathematical operators & functions	
String	Meaning
sym1 sym2	multiplication (a space)
sym1*sym2	multiplication (an asterisk)
sym1.sym2	multiplication (a dot)
sym1/sym2	division
sym1**y	exponentiation (<i>y</i> must be a numerical constant)
sym1~y	exponentiation (<i>y</i> must be a numerical constant)
log(sym1)	common logarithm
ln(sym1)	natural logarithm
exp(sym1)	exponential
sqrt(sym1)	square root

Derived units			
Quantity	Symbol	Full Name	Definition
area	barn	barn	1.0E-28 m**2
area	pix	pixel	
area	pixel	pixel	
electric capacitance	F	Farad	C/V
electric charge	C	Coulomb	A s
electric conductance	S	Siemens	A/V
electric potential	V	Volt	J/C
electric resistance	Ohm	Ohm	V/A
energy	J	Joule	N m
energy	Ry	Rydberg	13.605692 eV
energy	eV	electron-Volt	1.60217733E-19 J
energy	erg	erg	1.0E-7 J
events	count	count	
events	ct	count	
events	ph	photon	
events	photon	photon	
flux density	Jy	Jansky	1.0E-26 W /m**2 /Hz
flux density	R	Rayleigh	1.0E10/(4*PI) photon.m**2 /s/sr
flux density	mag	magnitude	
force	N	Newton	kg m/s**2
frequency	Hz	Hertz	1/s
illuminance	lx	lux	lm/m**2
inductance	H	Henry	Wb/A
length	AU	astronomical unit	1.49598E11 m
length	Angstrom	Angstrom	1.0E-10 m
length	lyr	light year	9.460730E15 m
length	pc	parsec	3.0867E16 m
length	solRad	solar radius	6.9599E8 m
luminosity	solLum	solar luminosity	3.8268E26 W
luminous flux	lm	lumen	cd sr
magnetic field	G	Gauss	1.0E-4 T
magnetic flux	Wb	Weber	V s

Accelerator keys

- **Control-n** Convert from current data units to UI values.
- **Control-s** Set data units to UI values.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.9 The simple binary maths window



This window allows you to add, subtract, divide and multiply two spectra. To use it just select a spectrum in the **Left operand:** list and a spectrum in the **Right operand:** list. Now select the operation you want to apply between them. A new spectrum called:

- Sum: (short name of left operand) + (short name of right operand)
- Diff: (short name of left operand) - (short name of right operand)
- Ratio: (short name of left operand) ÷ (short name of right operand)
- Product: (short name of left operand) x (short name of right operand)

is created on the global list.

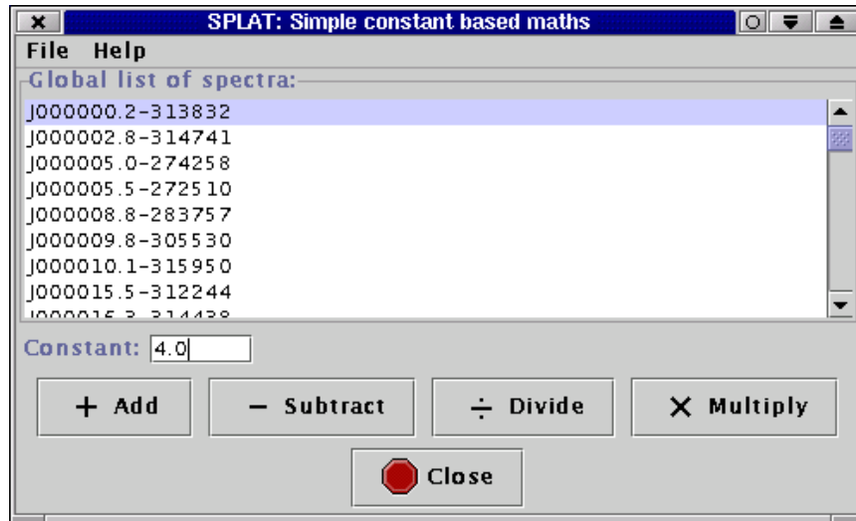
Note: the algorithm used to combine spectra that do not have the same wavelength positions is a simple interpolation. This does not preserve total flux, or errors, so this tool should only be used for quick qualitative arithmetic between real spectra. Artificial spectra (such as polynomial fits) do not have this restriction.

Accelerator keys

- **Control-d** Add the selected spectra from left and right.
- **Control-s** Subtract the selected spectra from left and right.
- **Control-m** Multiply the selected spectra from left and right.
- **Control-i** Divide the selected spectra from left and right.

- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.10 The simple constant maths window



This window allows you to add, subtract, divide and multiply a spectrum by a constant. To use it just select a spectrum in the **Global list of spectra:** and enter the value you want in the **Constant:** entry area. Now select the operation you want to apply. A new spectrum called:

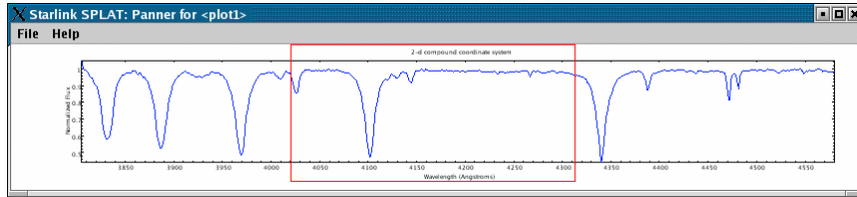
- Sum: (short name of spectrum) + (<constant>)
- Diff: (short name of spectrum) - (<constant>)
- Div: (short name of spectrum) ÷ (<constant>)
- Mult: (short name of spectrum) x (<constant>)

is created on the global list.

Accelerator keys

- **Control-d** Add the constant to the selected spectrum.
- **Control-s** Subtract the constant from the selected spectrum.
- **Control-m** Multiply the selected spectrum by the constant.
- **Control-i** Divide the selected spectrum by the constant.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.11 The panner window



This window always shows a full view of the spectra displayed in the related plot. It is designed to allow you to pan around the full sized view by moving the red rectangle.

The display is always drawn at the same aspect ratio as the full sized display, but is scaled to fit within the width or height of the panner window. Because of this there may be better widths or heights to fit any particular zoom. Resize the window to gain familiarity with this feature.

The following mouse interactions are provided:

- Centre display area, to centre the display area just put the mouse pointer over the panner and press the left mouse button.
- Scroll display area, to scroll the display area you need to drag the red rectangle. To do this just place the mouse pointer in the red rectangle and then press the left mouse button down, now move the mouse to drag the rectangle. When you're finished release the mouse button.
- Zoom display, to zoom the main display by a factor of 1 in X, just press the middle mouse button.
- Unzoom display, to unzoom the main display by a factor of 1 in X, just press the right mouse button.

Accelerator keys

- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.12 The plot configuration window

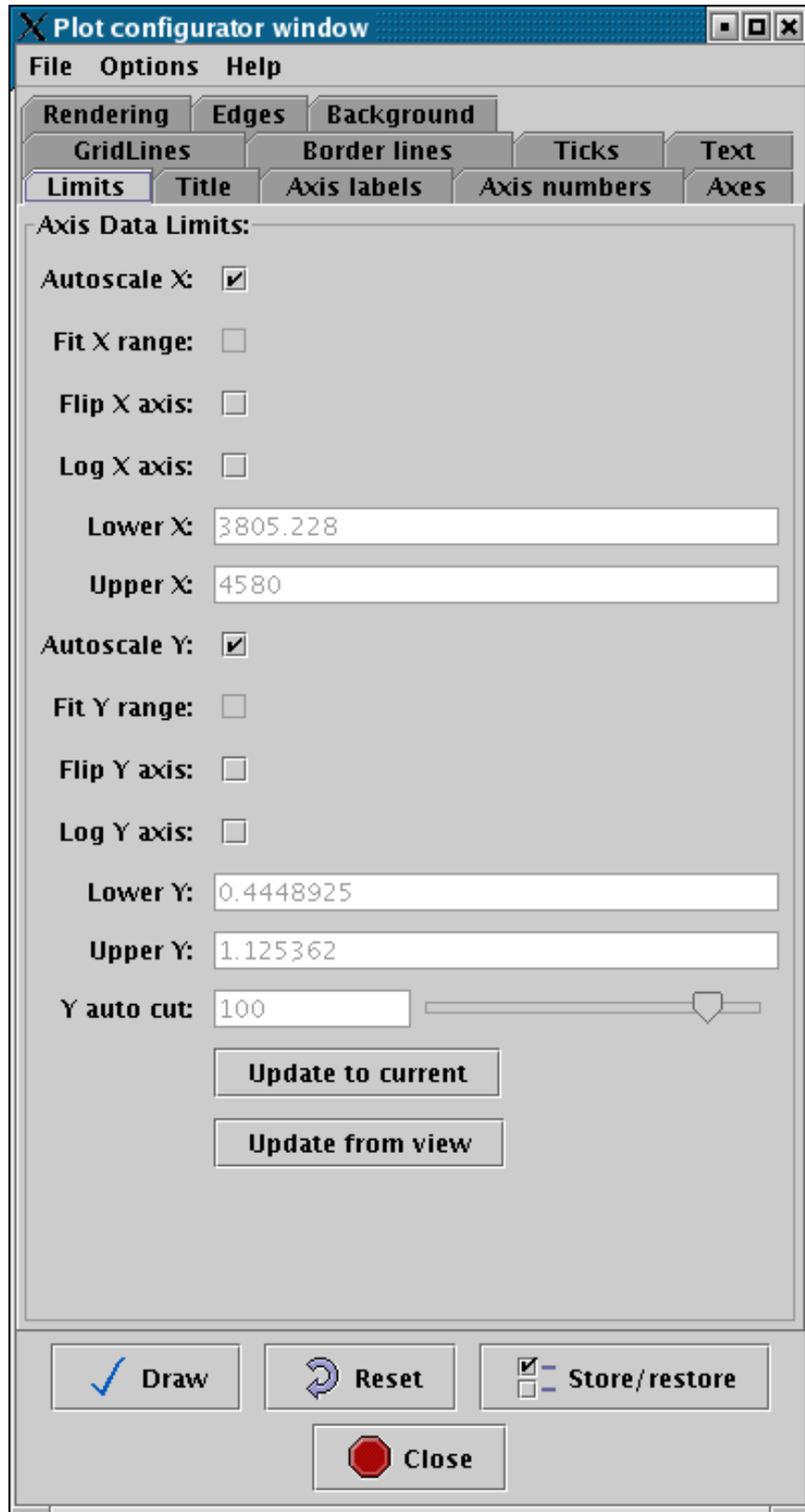
This window provides a series of paged controls for configuring the plot display area. This help section describes each page in turn. Global options that apply to the configuration window as a whole are the ability to **Auto-update** any changes to what the display area is showing and to save and restore configurations to and from an XML storage system.

Normally configuration changes happen immediately as **Auto-update** is switched on by default. If you prefer to wait for changes then you will need to press the **Draw** button. You also need to press the **Draw** button when making changes to the data limits (these may not be in a valid state until you've entered the complete coordinate value).

Accelerator keys

- **Control-d** Draw the plot with the current UI values.
- **Control-r** Reset values to their defaults.
- **Control-s** Activate the configuration store window.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

Axis Data Limits:



This page allows you to set the physical limits of the axis drawn in the display area. The default configuration auto-scales all spectra to fit within the currently available display area.

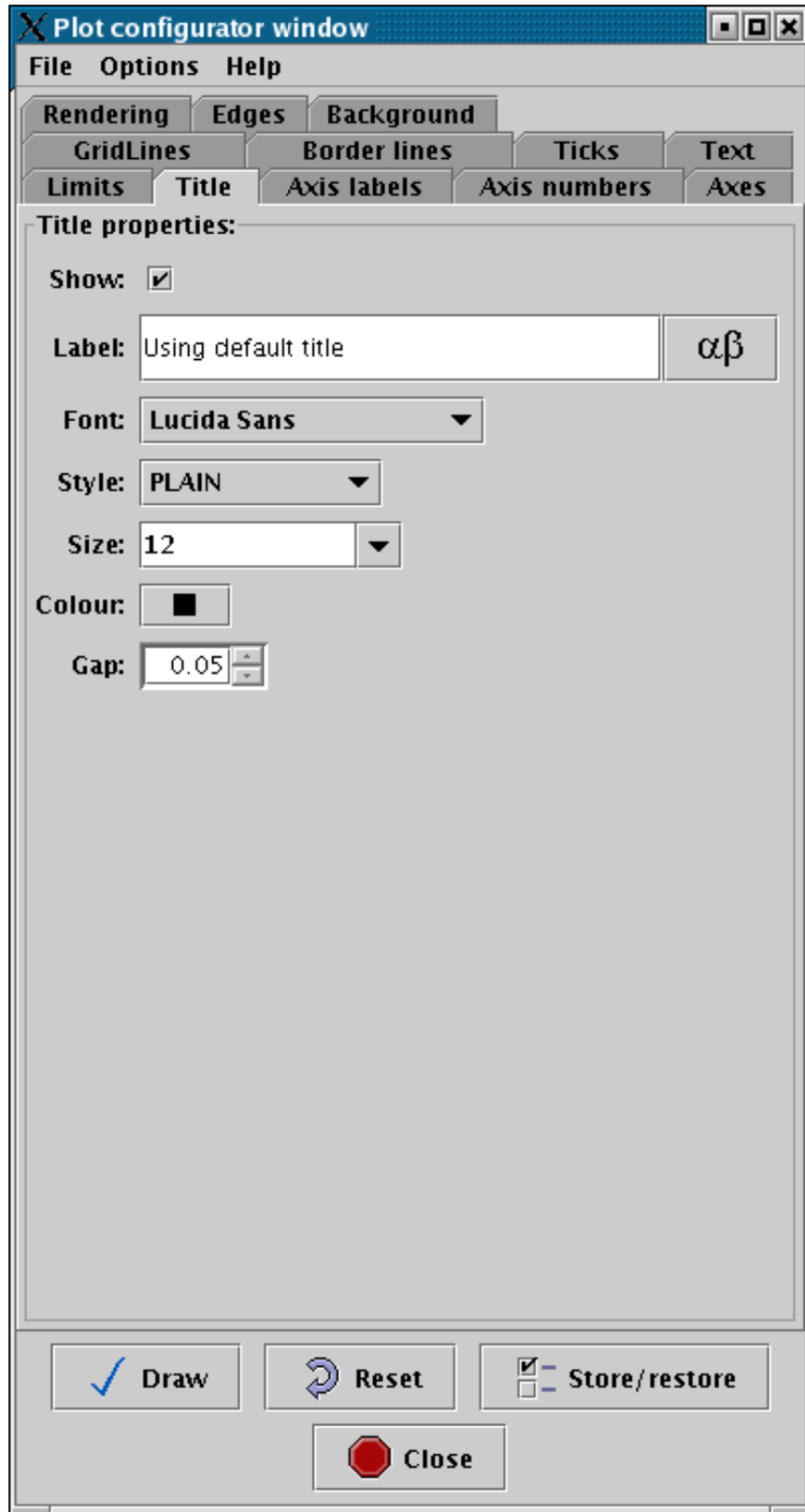
If you want to apply limits of your own then, untick the auto-scale checkboxes and enter your values in the lower and upper entry areas. Remember to press **Draw** to apply the limits.

You can also get either axis to display itself using logarithmic scaling (if possible, there are limitations, like not being able to contain the coordinate zero) and to display the axes ranges flipped top to bottom or left to right.

Other useful options are the checkboxes **Fit X range:** and **Fit Y range:** that also make the ranges that you supply fit the visible display area exactly (these are like the fitwidth and fitheight options in the plot window) and **Y auto cut:** which applies limits to the Y axis using percentile cuts (the data used to estimate the limits lies within the current X coordinate range).

The **Update to current** button sets the X and Y limits to those of the full plot (these are the values shown initially). The **Update from view** button sets the X and Y limits to those of the visible part of the display area. This is useful to just show coordinate axes for what you can see.

Title Properties:



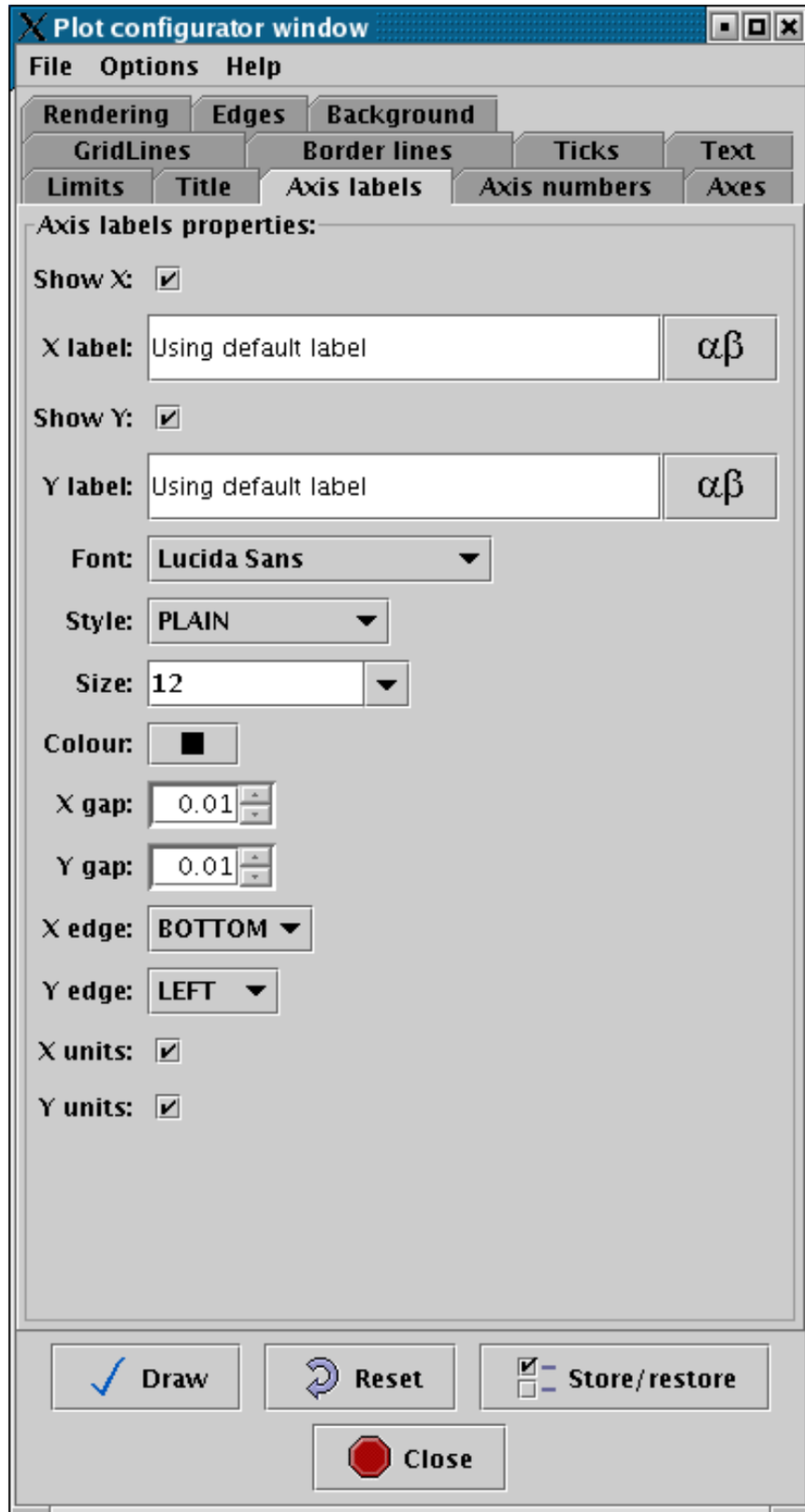
This page allows you to change the title shown in the plot. Using the controls you can define the font, style, size, colour and gap, from the nearest axis, of the title. You can also decide to have no title shown and insert any character from the chosen font (depends on the font, but usually these include the Å sign, press the $\alpha\beta$ button to see the character selection window).

The text shown in the **Label:** field is an accurate rendering of what is shown in the plot window. The default title can be restored by setting the text of this field to blank, or to the text `Using default title` (this text is used so that you can see the result of any changes you're about to make).

Superscript and subscripting are available using AST Escape sequences, for instance:

AST escape sequence	result
<code>super%^100+script%^+</code>	<i>super^{script}</i>
<code>sub%v100+script%v+</code>	<i>sub_{script}</i>
<code>%%</code>	<code>%</code>

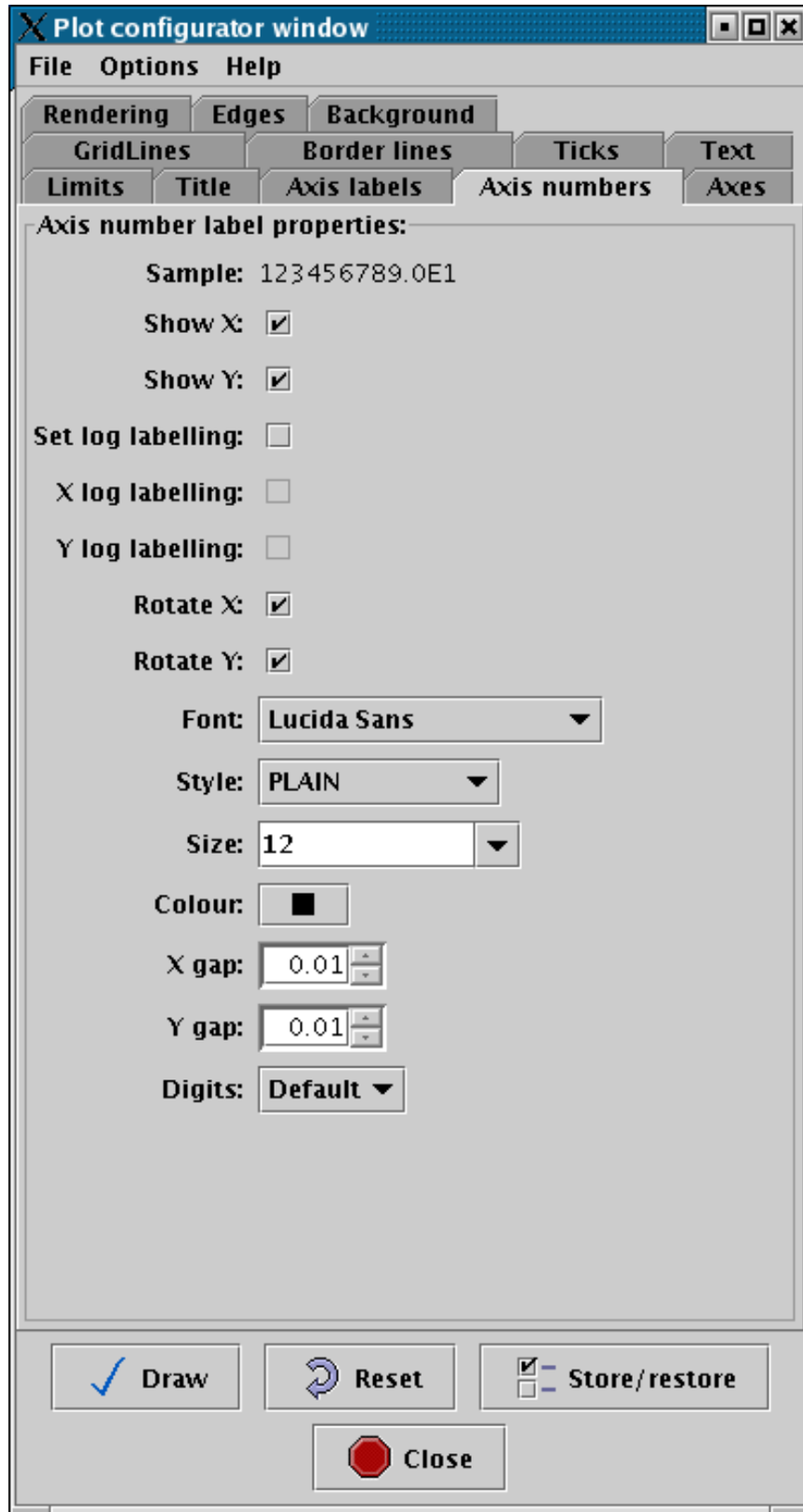
Axis Label Properties:



This page allows you to set the labels used on the X and Y axes. It also allows you to choose the font, style, size and colour used, as well as how far to position from the axis, which axis to attach to and whether to include any units (as a string in parenthesis). The text shown is an accurate rendering of what is shown in the plot window.

As with the title you can insert any characters from the font, and use AST Escape sequences to get effects like super and sub-scripting.

Axis Numbers:

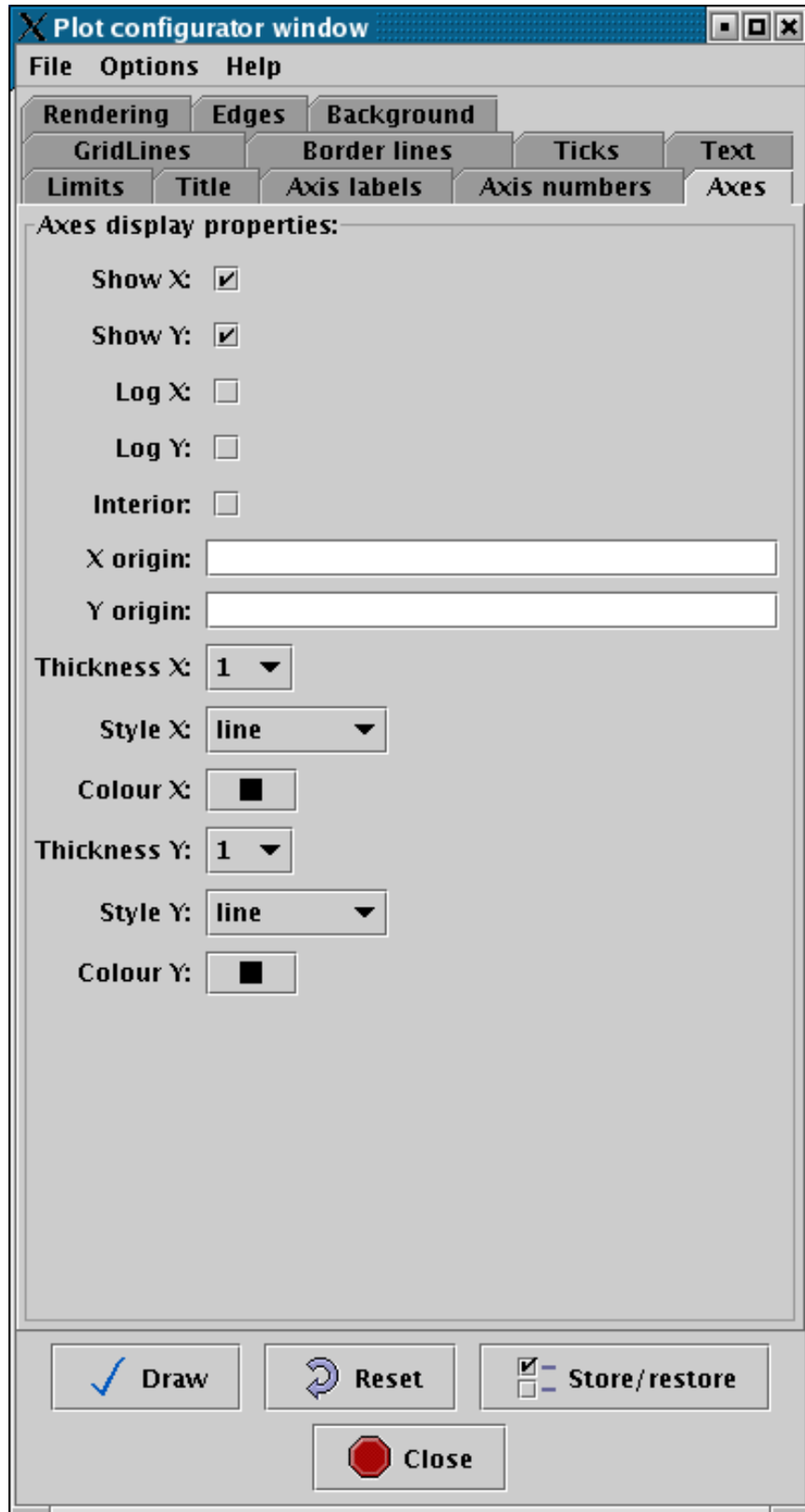


This page allows you to configure how the axis number labels are rendered. As with other pages you can define the font, style, size and colour used, as well as the separation between the actual axes and the labels. An accurate rendering of how the labels will be drawn is shown in the **Sample:** label.

The **Set log labelling:** option controls whether the numbers are drawn using an exponent notation. This is the default if log spacing has been selected for the axis (so you can also use this to not get exponent labelling when the axes are drawn using a log scale).

The **Digits:** drop down box allows you to override the default number of significant figures shown in the labels. Just choose a number that you want. However, note that the number of significant figures is used in the algorithm for determination how to space major ticks, so it may be necessary to increase this value when configuring the major tick separation.

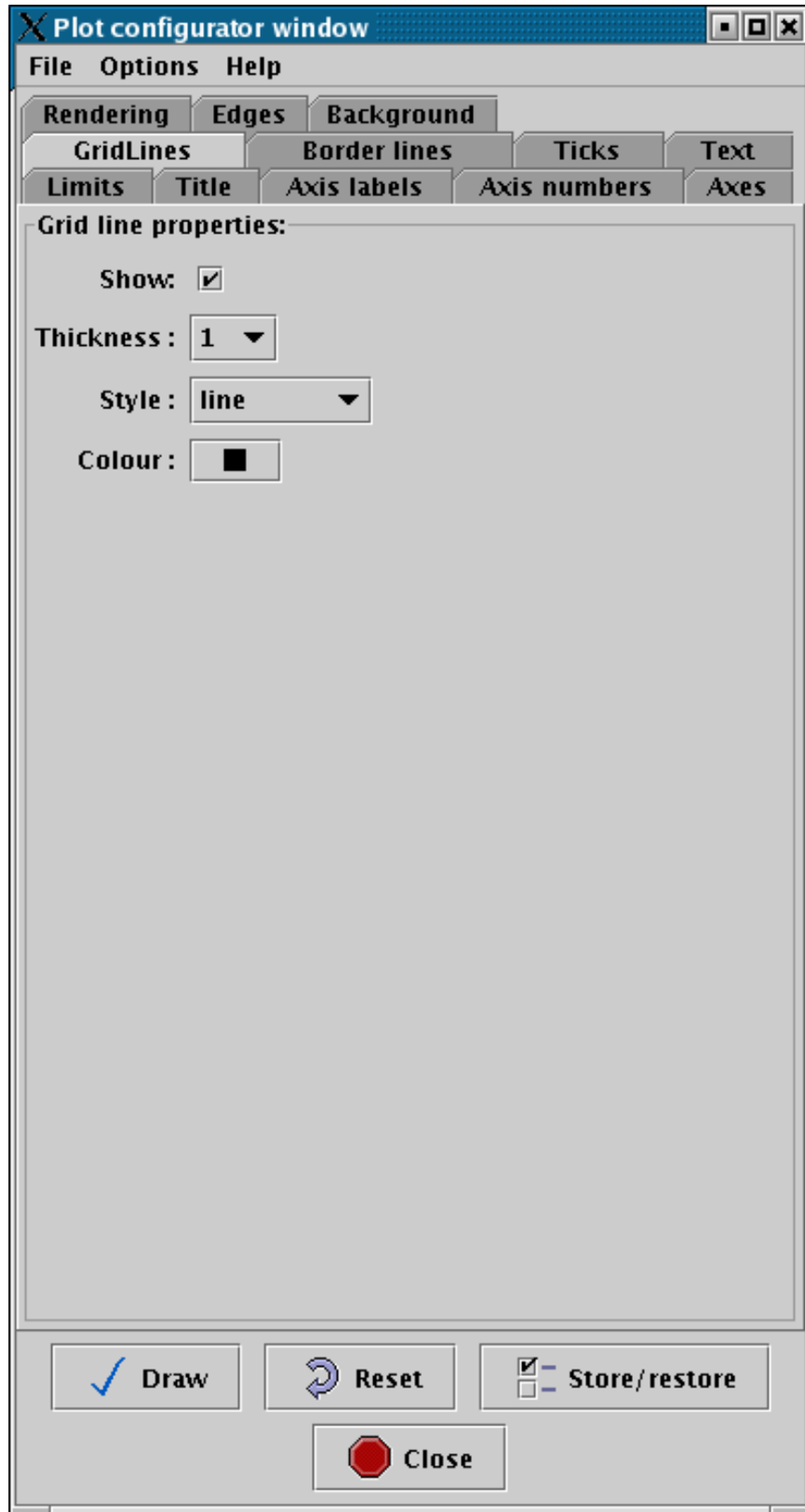
Axes:



This page allows you to control whether axes lines are drawn around the plot or not. In general this isn't very useful for spectral plots (when the outer border lines and axes are coincident), but it does allow you to draw the axes inside the display area and position where these are drawn, so you can define an arbitrary origin. Useful effects can then be obtained using grid lines.

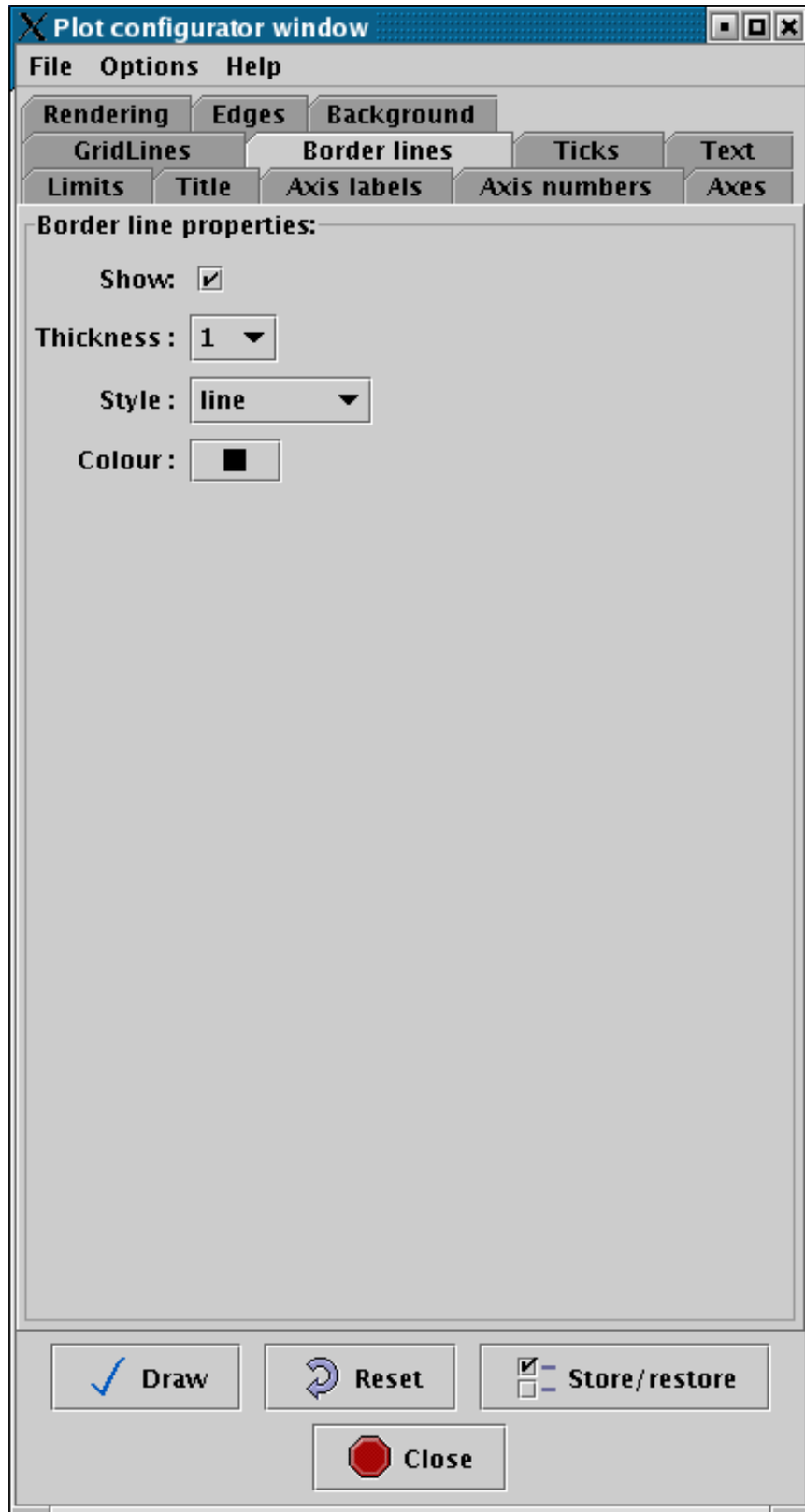
The **Log X:** and **Log Y:** options are the same as those shown in the **Limits** pane.

Grid Line Properties:



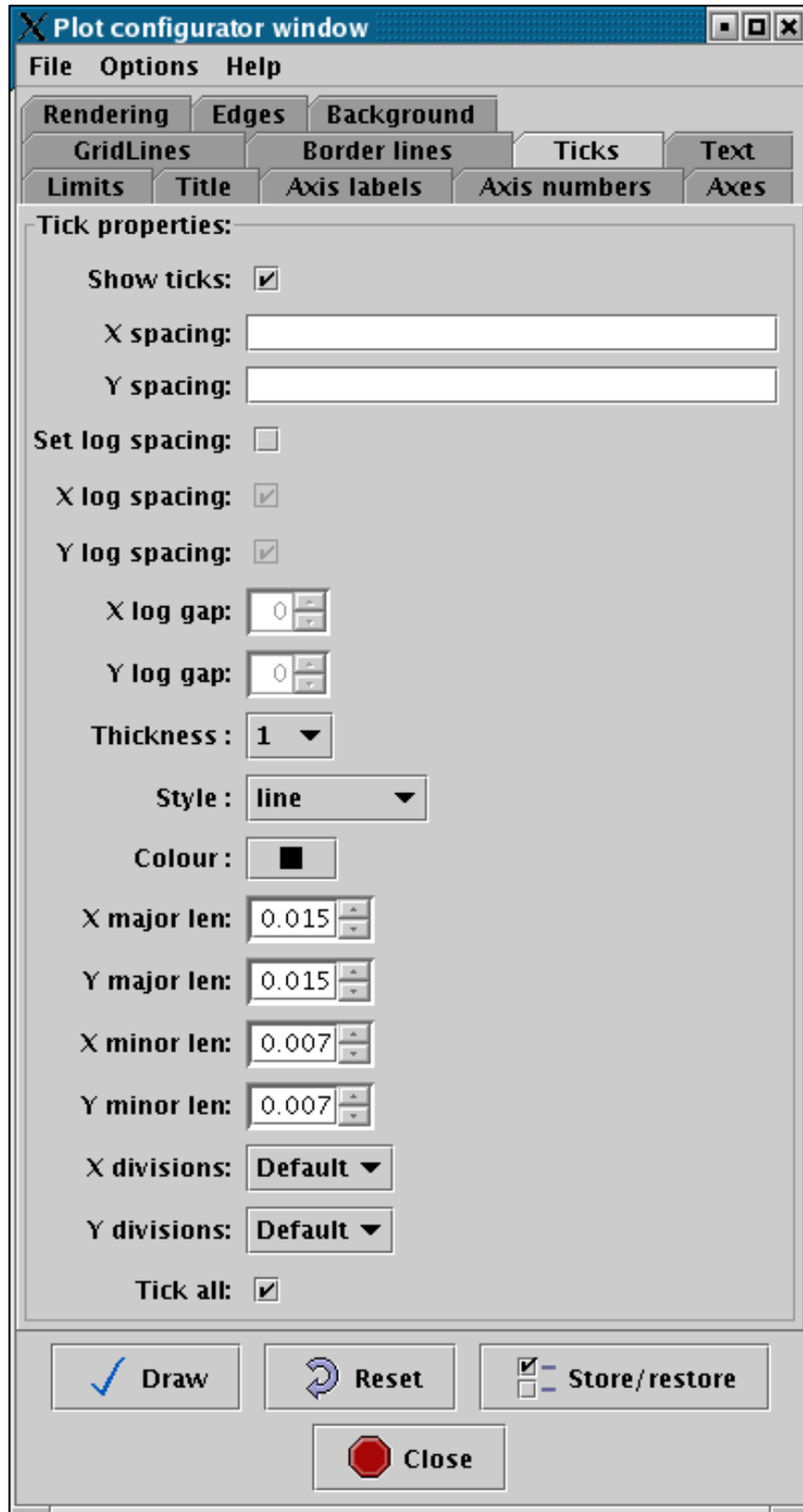
This page allows you to configure if grid lines are drawn in the plot, or not (not by default). Grid lines are drawn between the major ticks. You can also set the thickness, style and colour of the grid lines.

Border Line Properties:



This page allows you to configure if border lines are drawn in the plot, or not. Border lines are drawn at the data limits of the plot. You can also set the thickness, style and colour of the border lines.

Tick Properties:



This page allows you to define whether to show major and minor tick marks or not. It also allows you to define the spacing between major ticks and the number of divisions (minor ticks plus one) between major ticks, as well as the properties of the tick lines and their lengths.

The spacing used between major ticks may be dependent of the number of significant figures used in the axis number labels, so you may need to also change that.

The option **Set log spacing:** together with **X log spacing** and **Y log spacing** allow you to define if the spacing of major ticks along an axis are logarithmic. If the axis is scaled logarithmically (that is you've asked for log axes in the data limits tab) then log spacing is the default, otherwise major ticks are spaced linearly.

When spacing logarithmically you can define the separation of the major ticks using **X log gap:** and **Y log gap:**. These values are the power of 10 you require between adjacent ticks, so for instance, if you wanted ticks every factor of 10^2 you'd use 2.

Text Properties:

This window has no uses in SPLAT-VO.

Graphics Rendering Hints:

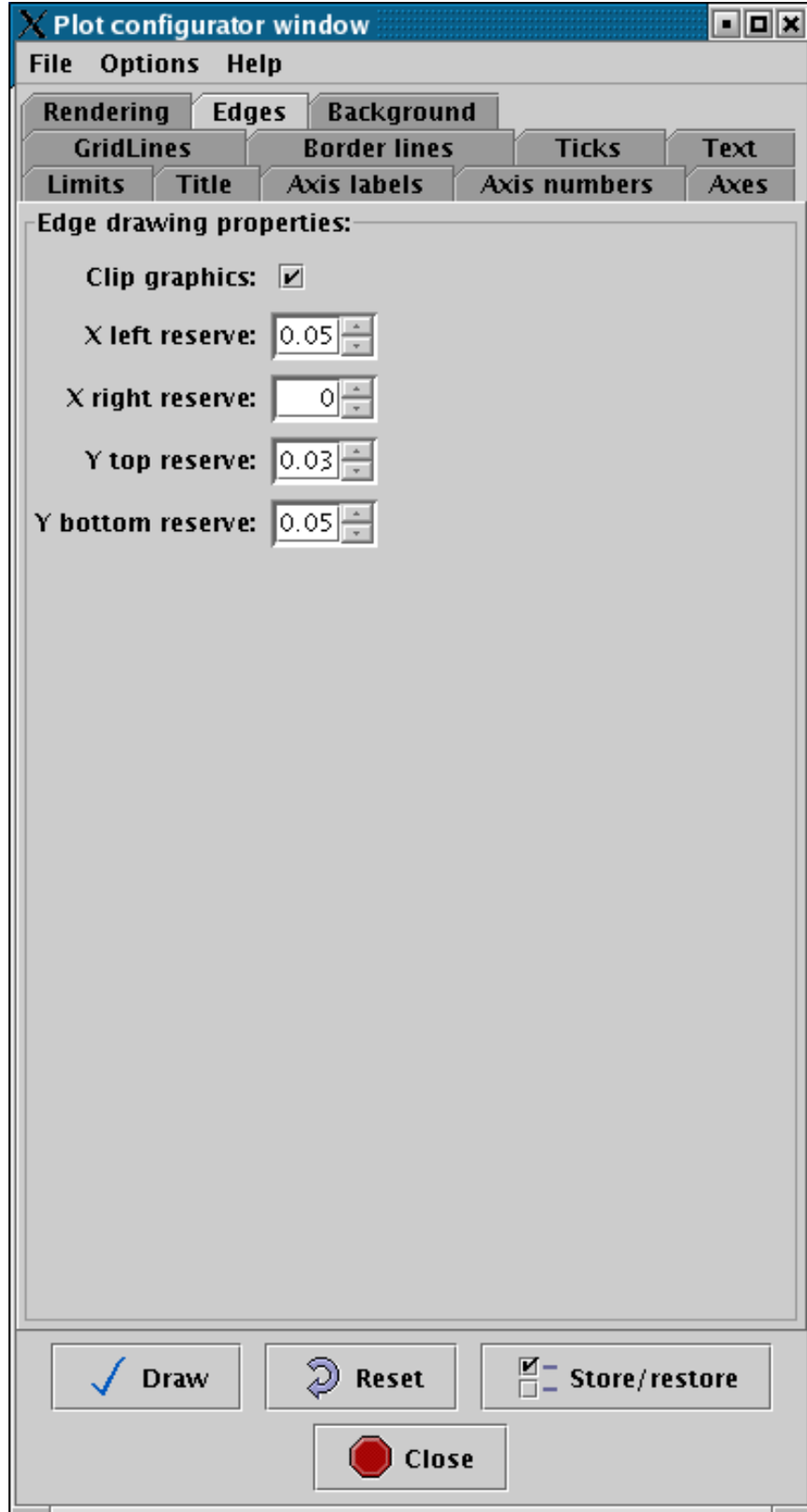


This page allows you to define how carefully the graphics are drawn in the plot display area. By default any text is drawn using anti-aliasing, which makes the graphics less jagged, and spectra are drawn without anti-aliasing. If you'd also like the spectra to be anti-aliased tick **Everything**:

It's not possible to anti-alias the spectra and not the text (so it's not just a perverse choice of options). Also note that any rendering options are applied to any captured JPEGs produced in the plot.

Finally anti-aliasing is a CPU expensive procedure so it can result in a significant performance degradation (so the default is switched off for spectra).

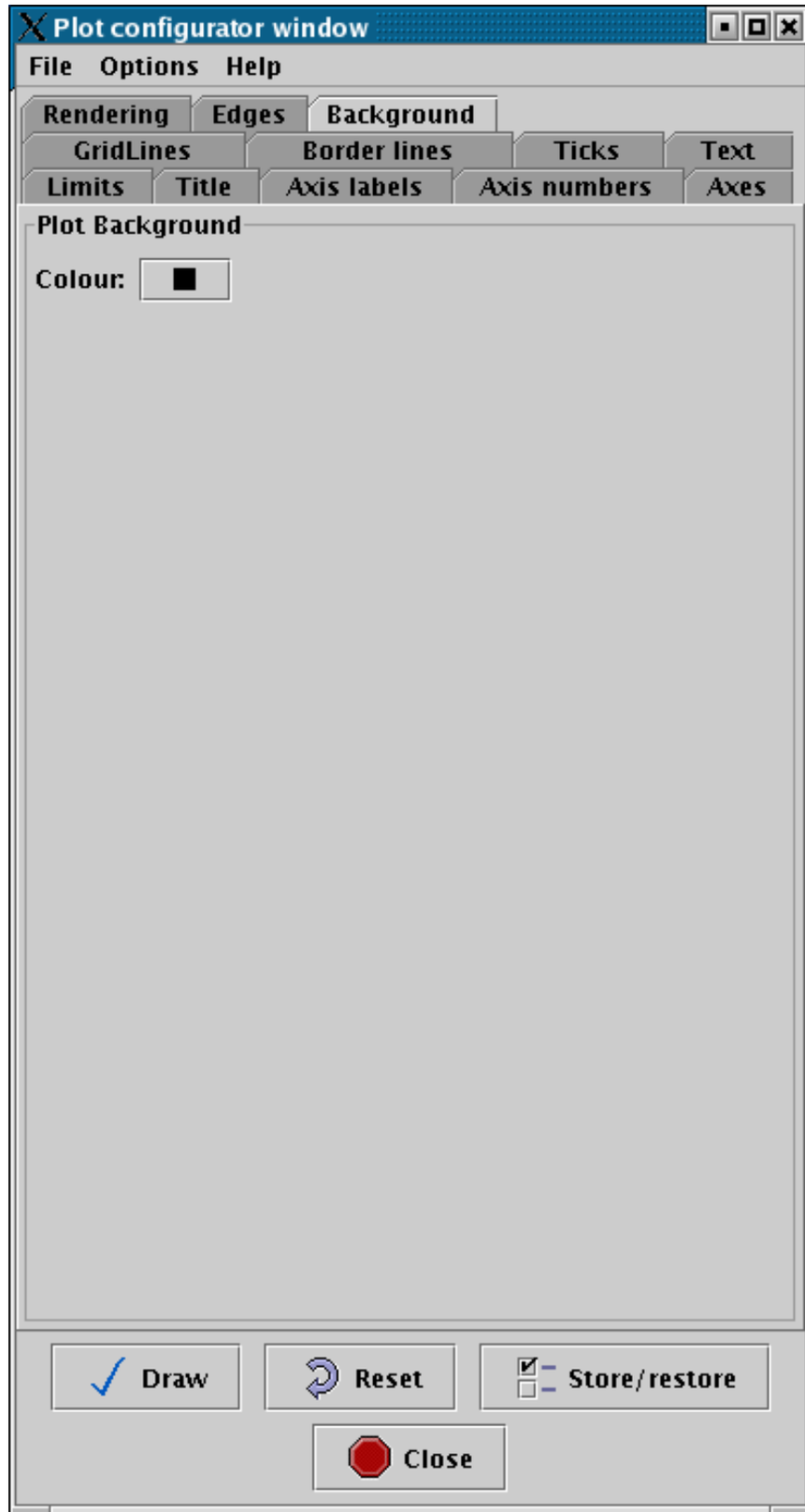
Edge Drawing Properties:



This page allows you to change the amount of space reserved around the plot for displaying titles and axis labels *etc.*

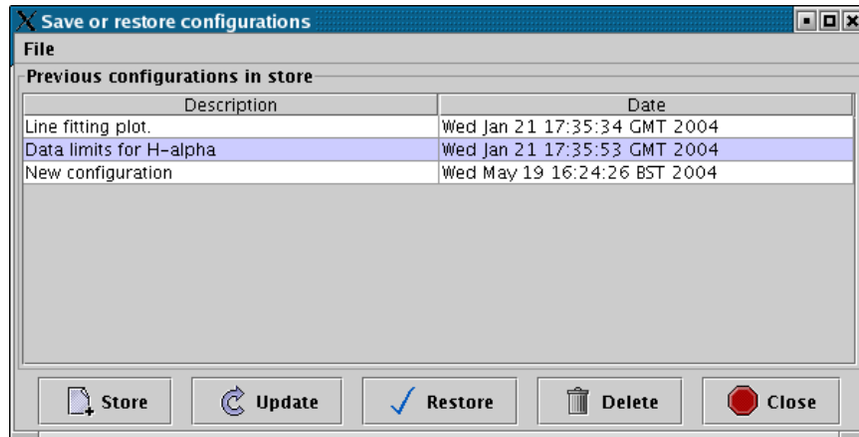
It also clips any points of any of the plotted spectra that lie outside the bounds of the data limits (this is not normally the case, unless you've defined data limits of your own).

Plot background:



Use this page to select a colour for the background of the plot display area.

Save or Restore Plot Configurations



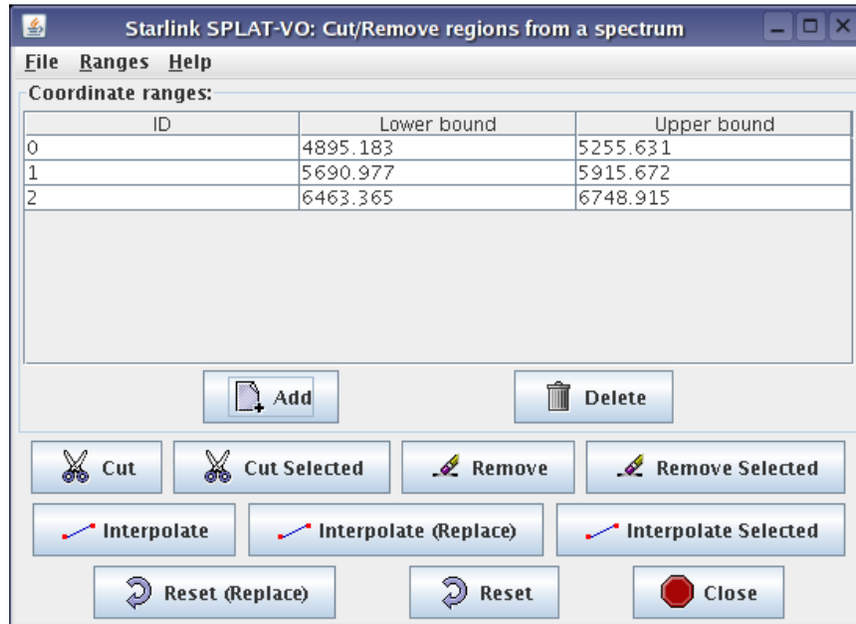
This additional window is activated by the **Store/restore** button. It's purpose is to allow you to create a database of configurations that you'd like to keep (after all if you've spent some time configuring the look of a plot you don't want to have to type it all in again when someone asks you to tweak the postscript).

The configurations are identified by two elements, a **Description** entered by you and the date when the configuration was created. To create a copy of the current configuration (note this doesn't include the spectra themselves) just press the **Store** button. This creates a new entry with the description `New configuration`. To change the description just double click on this text and type in your changes. Remember to press `<Return>` when you're done. To remove any un-needed configurations select their rows and press the **Delete** button.

To restore a configuration just select it and press the **Restore** button. The actual data is stored in the file: `$HOME/.splat/PlotConfigs.xml`. If things seem badly broken then try deleting this file.

3.13 The region extraction window

This window is used to cut out or remove sections from the current spectrum. The sections are defined as a series of graphics ranges that you draw on the plot display area.



To add a region press the **Add** button and then drag out a region in the display area of the plot window. This should result in the creation of a green rectangular figure.

You can interact with the figure, moving it side-to-side and resizing it. To do this point at the figure and press the left mouse button. This 'selects' the figure and adds grips to its exterior. Note that it also becomes the selected row in the **Coordinate ranges:** table. To move the figure just drag it and to resize it drag a grip (the little black squares). The associated coordinate range in the table updates with these changes.

To add a second range just press **Add** and repeat. The ranges can be overlapped or not.

To fine tune the ranges you can edit the values in the ranges table, just point at the cell you want to change and double click the left mouse button. This should enable the text editing cursor. Just make the modifications you want and press <Return> to make the changes permanent. (Note: if your spectra have sky coordinates shown for the X axis, then you should use the same format for your edits).

To read a set of ranges from a disk file choose the **File**→**Read ranges** menu item. The format of the input file is simple. It should have two fields separated by whitespace or commas. Comments are indicated by lines starting with a hash (#) and are ignored.

When you've got the ranges that you want to extract, remove or replace by linear interpolation, press one of various the **Cut**, **Remove** or **Interpolate** buttons. This creates and displays a new spectrum on the global list called Cut <n> of <shortname>, that you can save to disk, or display in another plot *etc.* To just extract, remove or interpolate, one or a subset of the ranges select the rows in the ranges table and press the **Cut Selected**, **Remove Selected** or **Interpolate Selected** buttons.

To clear the ranges press the **Reset** button, but note that this also clears any spectra that have been created on the global list. To keep these just close the window.

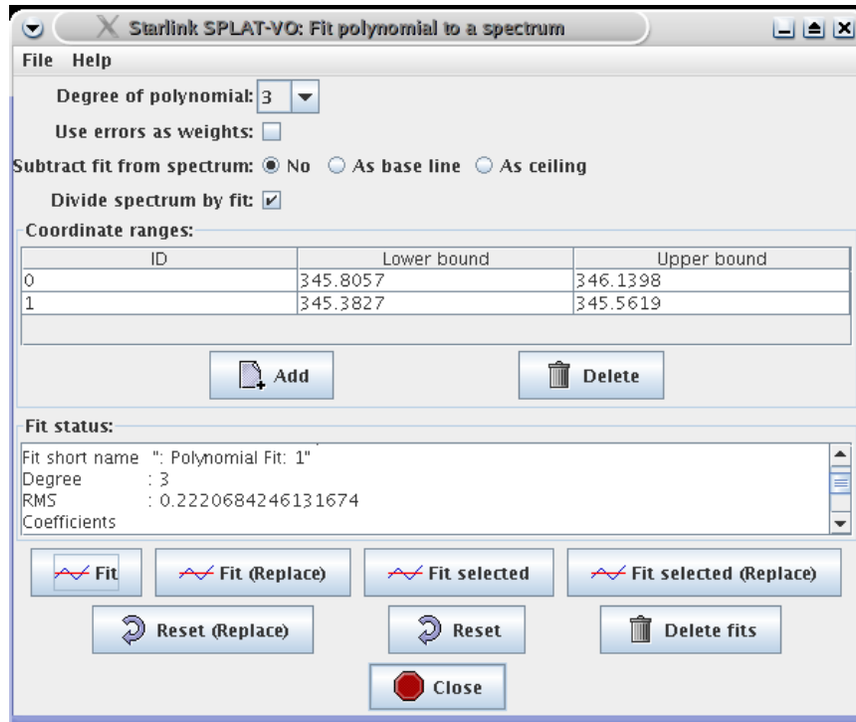
When interpolating you will often just want to see the resultant spectrum, so there is a special **Interpolate (Replace)** action. That performs the interpolation, removes the current spectrum and then displays the result of interpolation. To go back from this change use the **Reset (Replace)** button (only undoes one change).

Accelerator keys

- **Control-c** Cut out all regions from spectrum.
- **Control-s** Cut out selected regions from spectrum.
- **Control-m** Remove all regions from spectrum.
- **Control-o** Remove selected regions from spectrum.
- **Control-j** Interpolate all regions of the spectrum.
- **Control-k** Interpolate selected regions of the spectrum.
- **Control-r** Reset window by clearing ranges and any spectra.
- **Control-p** Reset an Interpolate (Replace) action.
- **Control-d** Add a coordinate range (interactive or non-interactive).
- **Control-e** Delete the selected coordinate ranges.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.14 The polynomial fitting window

The polynomial fitting window allows you to fit a polynomial of selected degree to various selected ranges of the current spectrum. You can select to use errors as weights for the fit and can save the fits on the global list as spectra in their own right. Fitting a polynomial is generally a pre-requisite to fitting spectral lines.



The **Degree of polynomial** drop-down list shows a series possible values for the degree of the polynomial that will be fitted. Obviously this should be chosen to match the expected complexity of the background.

The **Use errors as weights** checkbox is only effective if the spectrum being fitted has data errors.

The **Subtract fit from spectrum** series of radio buttons allows you to ask for the creation of a spectrum that is the difference between the spectrum and the polynomial fit. Use the **As a ceiling** option for absorption lines, and **As base line** for emission lines. These subtracted spectra appear on the global list and are added to the plot for display.

The **Divide spectrum by fit** checkbox allows you to ask for the creation of a new spectrum that is the result of dividing the spectrum by the polynomial fit. You will probably want to do this before attempting to fit absorption lines so that the continuum is normalized. The new spectrum appears on the global list (with the name `Ratio: <spectrum name> by <polynomial name>`) and can plotted in a new window before fitting the lines.

Coordinate ranges

This part of the window is used to define the regions of the spectra that you want to fit the polynomial to.

To add a region press the **Add** button and then drag out a region in the display area of the plot window. This should result in the creation of a green rectangular figure.

You can interact with the figure, moving it side-to-side and resizing it. To do this point at the figure and press the left mouse button. This 'selects' the figure and adds grips to its exterior. Note that it also becomes the selected row in the **Coordinate ranges:** table. To move the figure just drag it and to resize it drag a grip (the little black squares). The associated coordinate range in the table updates with these changes.

To add a second range just press **Add** and repeat. The ranges can be overlapped or not.

To fine tune the ranges you can edit the values in the ranges table, just point at the cell you want to change and double click the left mouse button. This should enable the text editing cursor. Just make the modifications you want and press <Return> to make the changes permanent. (Note: if your spectra have sky coordinates shown for the X axis, then you should use the same format for your edits).

To read a set of ranges from a disk file choose the **File**→**Read ranges** menu item. The format of the input file is simple. It should have two fields separated by whitespace or commas. Comments are indicated by lines starting with a hash (#) and are ignored. You can also save the ranges to disk (**File**→**Save ranges**).

When you've got the ranges that you want to fit, press the **Fit** button. This creates a new spectrum on the global list, called **Polynomial Fit: <n>** and displays it in the plot. You can save this to disk, or display in another plot *etc.*, just like any other spectrum. To just fit one or a subset of the ranges select the rows and then press the **Fit selected** button.

To clear any fits created by this window press the **Delete fits** button. To clear the ranges from the display area and delete the fits press the **Reset** button. To keep any fits just close the window.

Fit status

The **Fit status** text area shows some facts and figures about the fitting process. The RMS value is the root mean square difference between the fit and the spectra data *i.e.*:

$$\sqrt{\frac{\sum_{i=1}^N ((s_i - f_i) * (s_i - f_i))}{N}}$$

Where s_i and f_i are the data values of the spectrum and the fit (possibly interchanged) and N the number of positions in the spectrum.

Command Buttons

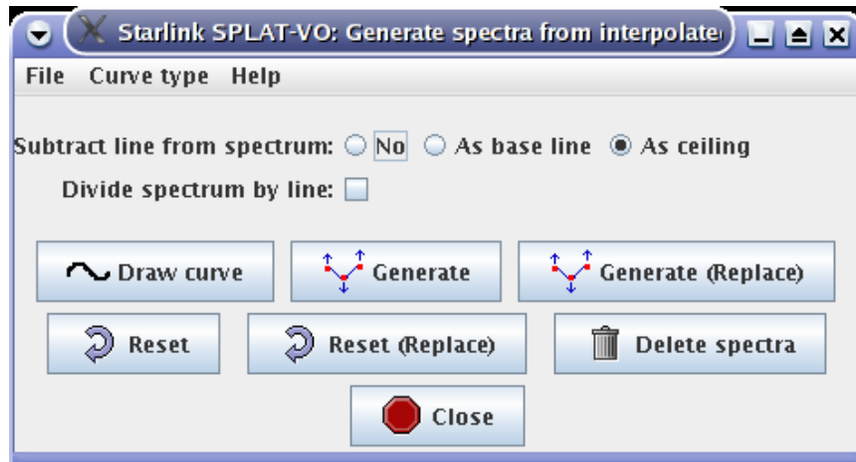
- **Fit** Performs a fit on all the regions you've defined and displays the fit and any subtracted or divided spectra in the plot.

- **Fit (Replace)** Same as **Fit**, except that if you've opted to subtract or divide the fit into the displayed spectrum, then that spectrum will be removed and you'll only see the subtracted and/or divided result. If you haven't opted to subtract or divide then the results are exactly the same as for **Fit**.
- **Fit selected** Performs a fit on only the currently selected regions. The results are displayed in the way as the **Fit** command.
- **Fit selected (Replace)** Same as **Fit (Replace)**, except only the selected regions are used for fitting.
- **Reset** Removes all fit spectra (from the plot and global list) and erases the ranges. The polynomial degree is set to the default (which will be the last value used in any polynomial toolbox).
- **Reset (Replace)** Also removes all fit spectra, but keeps the ranges and replaces the spectrum first removed by a **Fit (Replace)**. Effectively undoes a **Fit (Replace)**.

Accelerator keys

- **Control-f** Do fit to all ranges.
- **Control-l** Do fit to all ranges, only display any new spectra.
- **Control-s** Do fit to selected ranges.
- **Control-t** Do fit to selected ranges, only display any new spectra.
- **Control-p** Reset interface after a fit replace.
- **Control-r** Reset interface clearing all ranges and new spectra.
- **Control-i** Delete any new spectra (fits and corrected spectra).
- **Control-d** Add a coordinate range (interactive or non-interactive).
- **Control-e** Delete the selected coordinate ranges.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.15 The generate spectra from interpolated lines window



This window is for generating a spectrum that is equivalent to an interpolated line that you have drawn (or are about to draw) on the plot. Such lines can then be subtracted to or from, or divided into the current spectrum, thus providing a way to background correct a spectrum using a hand-drawn line.

Command Buttons

- **Draw Curve** Start an interaction in the plot window for drawing a curve. This is equivalent to selecting the **Drawing mode** → ~ option in the **Graphics** menu. Different types of interpolated lines can be selected first in the **Curve type** menu. The default is **hermite** a type of spline that attempts to connect points in a similar way to a hand-drawn curve.
- **Generate** Generates an actual spectrum from your interpolated curve and then displays it, along with any subtracted or divided spectra, in the plot.
- **Generate (Replace)** Same as **Generate**, except that if you've opted to subtract or divide the curve into the displayed spectrum, then that spectrum will be removed and you'll only see the subtracted and/or divided result. If you haven't opted to subtract or divide then the results are exactly the same as for **Generate**.
- **Reset** Removes any spectra that you've generated from the plot and global list.
- **Reset (Replace)** Same as **Reset**, but it also restores the spectrum removed in by the first **Generate (Replace)**. Effectively undoes a **Generate (Replace)**.

Accelerator keys

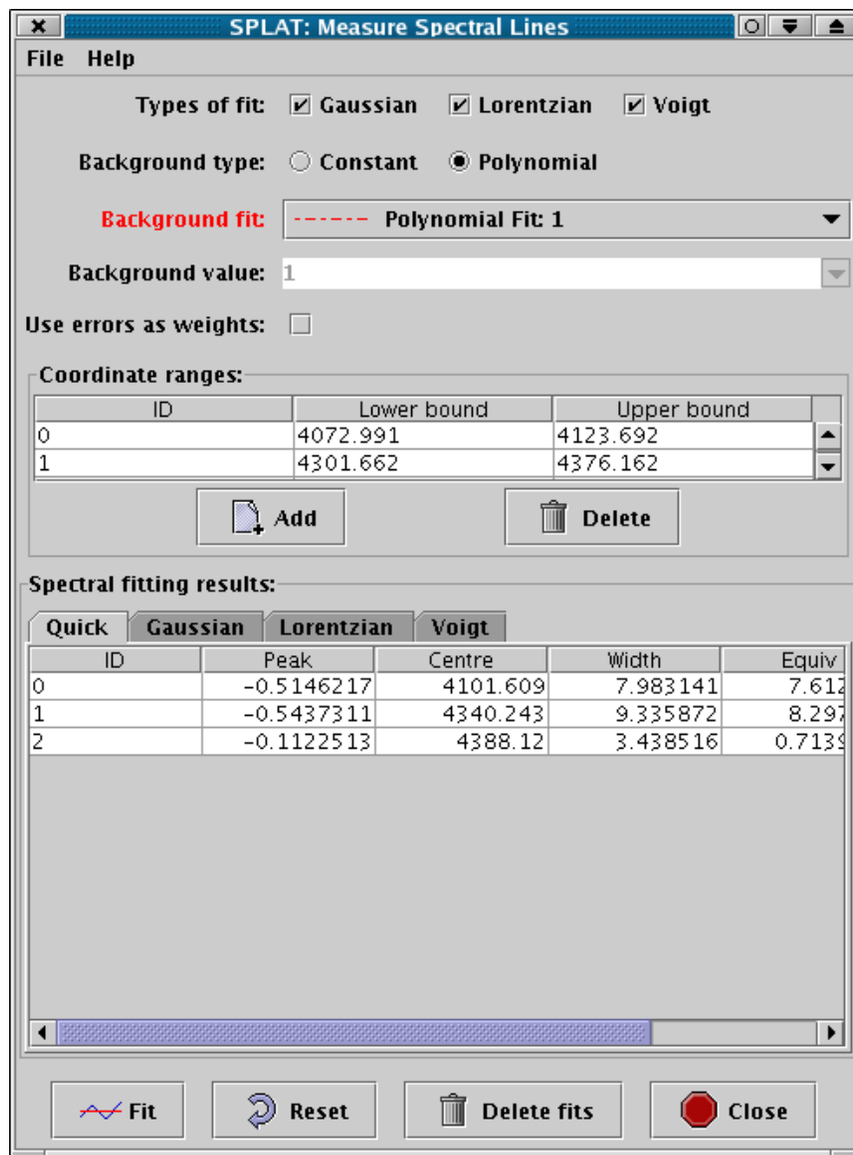
- **Control-d** Start interaction to draw a curve.
- **Control-g** Generate a spectrum from the drawn curve.
- **Control-p** Reset interface undoing a **Generate (Replace)**.
- **Control-r** Reset interface clearing all new spectra.

- **Control-i** Delete any new spectra.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.16 The spectral line fitting window

The spectral line fitting window allows you to measure the wavelength and equivalent width of emission or absorption lines and to fit various standard profiles to the lines.

The standard profiles offered are Gaussian, Lorentzian and Voigt, as well as a model independent intensity analysis measure.



Background type: Choose a type of background estimate to be subtracted from the spectrum before fitting. The choices are **Constant** and **Polynomial**. If you have already subtracted the background from the spectrum, or have applied a normalisation (so that the background is flat, usually with a value of 1) then you need to choose **Constant** and provide a suitable value. Otherwise choose **Polynomial** and select a spectrum from the list in **Background fit**.

Background fit: For emission lines you would normally define a background spectrum using the polynomial fitting tool. In this case select the **Polynomial** option and choose the background

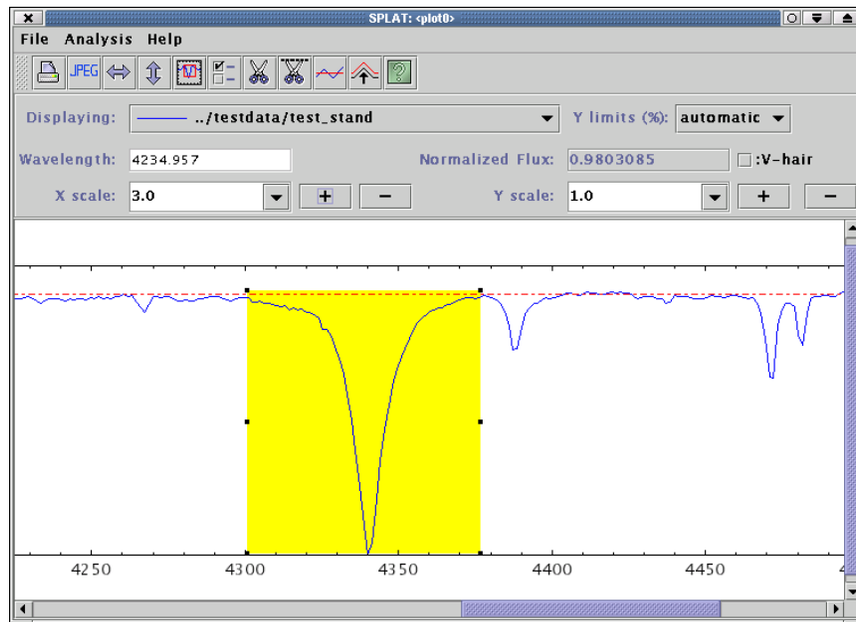
spectrum you have created. Note that you can actually use any spectrum from the global list as the background (so you are not restricted to what SPLAT-VO, provides) but if you choose the same spectrum as is being measured then a background of 0 is assumed.

The default background should be set to the first polynomial found on the global list.

Use error as weights: If your spectrum has associated error measurements then you can select this checkbox to use the errors as weights in the fitting procedure.

Coordinate ranges: This table shows the upper and lower coordinate range for each line that you want to measure.

To add a line to this list press the **Add** button. Now drag out a region on the plot display area that encompasses the spectral line. You should now see a yellow rectangle over the line:



Note that the rectangle is just defining a range, not an area, so you just need to worry about its width and position along the X axis, not its height.

To adjust the rectangle position select it and either drag it around, or use the grips on the exterior to resize it. You can see the physical extent of the region in the **Coordinate ranges:** table. To set these to a known value double click on the value you'd like to set, make the change, and then press <Return> to apply it.

The extents of lines can be saved to a simple text file and re-read. The format is simple. It should have two fields separated by whitespace or commas. Comments are indicated by lines starting with a hash (#) and are ignored.

Types of fit:

Once you have defined the extents of your spectral lines and chosen a background, you can proceed to make the measurements you want. Just select the types of fit you want from the list **Gaussian**, **Lorentzian** and **Voigt**. Note you can de-select all these and just get a **Quick** measurement.

All types except **Quick** use a non-linear least squares (Levenberg-Marquardt) minimisation stage to perform the actual fit of the model to the data (possibly with weighting, if the spectrum has any error measurements). So it is necessary to have some idea of the shape of the line, before attempting to fit it. This job is done by the **Quick** fit, which is compulsory. The description of each of the types and what their measurements represent is given below.

The error estimates are based on the variances produced by the fit. If you have errors for the data values, these will be used as weights and providing they are realistic the errors will be correct. In the absence of data errors the errors produced will be decided assuming that the fit is good (the chi-square statistic will be artificially forced to have a value of 1).

Quick: The default line measurement type. This is based on the ABLINE technique described in the FIGARO documentation. This just uses an analysis of the intensities in the selected region and the chosen background. The results you get are:

- **ID**, just a unique integer to label the line. This is the same for all fits.
- **Peak**, the peak, background subtracted, intensity in the region. Negative for absorption lines.
- **Centre**, the position of the median value *i.e.* that for which half of the total, background subtracted, area of the line, within the given region, lies to the left and right.
- **Width**, width of the line. For a Gaussian this would be ± 1 standard deviations wide.
- **Equiv**, the equivalent width of the line. This will be 0 if a zero background is given or assumed.
- **Flux**, the background subtracted integrated intensity of the line over the selected region. Requires a linear coordinate step.
- **Asym**, an asymmetry value for the line. This is a measure of the relative displacement of the centre of the line from the centre as measured by the upper and lower half widths.

Gaussian: The formula used for the Gaussian is:

$$y(r) = A * \exp \left(-0.5 * \left(\frac{x - c}{\sigma} \right)^2 \right)$$

where A is the scale height, r is the distance from the centre, c is the centre, x the distance from the origin and σ is the Gaussian sigma. The y values are background subtracted.

The values measured for this line shape are:

- **ID**, integer identifier (same as other fits).
- **Peak**, the Gaussian peak (A).
- **PeakErr**, an estimate of the error.
- **Centre**, the position of the Gaussian peak (c).
- **CentreErr**, an estimate of the error.

- Width, the Gaussian sigma (σ).
- WidthErr, an estimate of the error.
- FWHM, the full width of profile at half maximum.
- FWHMErr, an estimate of the error.
- Flux, the integrated flux of the Gaussian.
- FluxErr, an estimate of the error.
- Rms, the root mean square difference between the Gaussian and the line data.

Lorentzian: The formula used for the Lorentzian is:

$$y(r) = \frac{A}{1 + 0.5 * \left(\frac{x-c}{w}\right)^2}$$

where A is the scale height, r is the distance from the centre, c is the centre, x the distance from the origin and w the Lorentzian width. The y values are background subtracted. The full width half maximum ($fwhm$) of the curve is:

$$fwhm = w * 2\sqrt{2}$$

The values measured for this line shape are:

- ID, integer identifier (same as other fits).
- Peak, the Lorentzian peak (A).
- PeakErr, an estimate of the error.
- Centre, the position of the Lorentzian peak (c).
- CentreErr, an estimate of the error.
- Width, the Lorentzian width (w).
- WidthErr, an estimate of the error.
- FWHM, the full width of profile at half maximum.
- FWHMErr, an estimate of the error.
- Flux, the integrated flux of the Lorentzian.
- FluxErr, an estimate of the error.
- Rms, the root mean square difference between the Lorentzian and the line data.

Voigt: The formula for a Voigt profile (a convolution of a Gaussian and Lorentzian) is:

$$K(a, b) = \frac{b}{\pi} \int_{-\infty}^{\infty} \frac{\exp(-t^2)}{(a-t)^2 + b^2} dt$$

where:

$$a = \sqrt{\ln 2} \left(\frac{x - c}{\sigma} \right)$$

and

$$b = \sqrt{\ln 2} \left(\frac{w}{\sigma} \right)$$

In these equations c is the central wavelength, x the distance from the origin, σ is the Gaussian width and w the Lorentzian width. Which is complex to solve. However,

if we take a complex variable $z = a + ib$, then the Voigt function is, within a scale factor (A), also the real part of the complex error function:

$$w(z) = \exp(-z^2) \operatorname{erfc}(-iz)$$

which has been approximated by various numerical codes. Using this fact with a non-linear minimisation routine (plus some derivatives) is how SPLAT performs its Voigt fitting.

The values measured for this line shape are:

- ID, integer identifier (same as other fits).
- Peak, the Voigt peak (A).
- PeakErr, an estimate of the error.
- Centre, the position of the Voigt peak (c).
- CentreErr, an estimate of the error.
- Gwidth, the Gaussian width (σ).
- GwidthErr, an estimate of the error.
- Lwidth, the Lorentzian full width at half maximum (w).
- LwidthErr, an estimate of the error.
- FWHM, the full width of profile at half maximum (estimate).
- FWHMErr, an estimate of the error.
- Flux, the integrated flux of the Voigt profile.
- FluxErr, an estimate of the error.
- Rms, the root mean square difference between the Voigt and the line data.

3.16.1 Saving the results

The measurements, as shown in the **Spectral fitting results:** table can be written out to a text file. Just select the **File**→**Save line fits** item and either pick an existing file to overwrite, or enter a new file name. The format of this file is very simple, as shown by the following example (long lines are wrapped for presentation):

```
# File created by Starlink SPLAT-VO

# Quick results:
# ID Peak Centre Width Equiv Flux Asym
0 0.5280543 4101.737 9.664325 0 8.636 1.046

# Gaussian results:
# ID Peak PeakErr Centre CentreErr Width WidthErr FWHM FWHMErr
# Flux FluxErr Rms
0 0.47457 0.013518 4101.637 0.20611 6.266466 0.2061116 14.7564 0.4853556
7.454487 0.3243526 0.03108123

# Lorentzian results:
# ID Peak PeakErr Centre CentreErr Width WidthErr FWHM FWHMErr
```

```
# Flux FluxErr Rms
0 0.5306862 0.00502416 4101.669 0.05490553 4.108373 0.05540269 11.62023 0.1567025
9.686626 0.1596041 0.009050248

# Voigt results:
# ID Peak PeakErr Centre CentreErr Gwidth GwidthErr Lwidth
# LwidthErr FWHM FWHMErr Flux FluxErr Rms
0 0.5308469 0.0120453 4101.668 0.05545499 0.6207013 0.7700358 11.48657 0.3663733
11.68279 0.4136257 9.687501 0.1166889 0.009053993
```

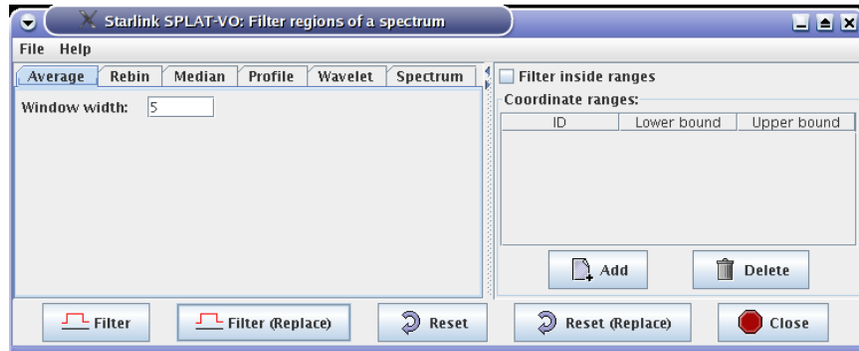
So each fit type has a section in which the results for each line are shown. Lines are uniquely identified by their ID value.

Accelerator keys

- **Control-f** Do line fits to all ranges.
- **Control-r** Reset interface clearing all ranges and fits.
- **Control-i** Delete any fits.
- **Control-d** Add a coordinate range (interactive or non-interactive).
- **Control-e** Delete the selected coordinate ranges.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.17 The filter spectrum window

This window provides the ability to filter the current spectrum displayed in a plot.



The various options are:

- **Average** take the running average value over a number of positions.
- **Rebin** rebin a spectrum so that one output value equals the mean of a number of input values.
- **Median** take the running median value over a number of positions.
- **Profile** smooth the values using a profile matched to typical line shapes, Gaussian, Lorentzian and Voigt and the standard Hanning, Hamming, Welch and Barlett filters.
- **Wavelet** use various wavelet filters to convert the spectrum into multi-resolution coefficients, removing a fraction of the coefficients, thus *denoising* the spectrum.
- **Spectrum** smooth the values using a profile defined in another spectrum (allows you to create kernels that match some shape, rather than just the standard profiles).

You can define ranges of the spectrum to apply the filter over (except for **Rebin**, that always applies to the whole spectrum, which is what happens for all filters if you've not defined any ranges). Depending on whether you've selected **Filter inside ranges** or not the ranges are used to define the parts of the spectrum to filter, or not to filter. Parts which are not filtered do not form part of the result spectrum.

Command Buttons

- **Filter** Filters the spectrum and displays the result in the plot.
- **Filter (Replace)** Same as **Filter**, except the current spectrum is removed so you only see the filtered one.
- **Reset** Removes all filtered spectra (from the plot and global list) and erases the ranges.
- **Reset (Replace)** Also removes all filtered spectra, but restores the spectrum first removed by **Filter (Replace)** Effectively undoes the first **Filter (Replace)**, since the last **Reset (Replace)**

Accelerator keys

- **Control-f** Apply current filter to spectrum.
- **Control-l** Apply current filter to spectrum and only display the new spectrum.
- **Control-p** Reset interface after a filter replace.
- **Control-r** Reset interface clearing any ranges and new spectra.
- **Control-d** Add a coordinate range (interactive or non-interactive).
- **Control-e** Delete the selected coordinate ranges.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.18 Quick change of coordinate or data units window

This window allows you to change the coordinate or data units of the current spectrum to a limited number of preset values. You can only change units when the current spectrum already has a correctly set coordinate system or data units.

If you need to set the coordinates or data units you must use the two toolboxes available in the main window. These also offer considerably more flexibility, particularly when defining spectral coordinates.

If you do not want to change the units of an axis, just leave the value set at **Not set or unrecognised**.

When you see the value **Not set or unrecognised**, that means that the current spectrum has no properly defined coordinates or data units for that axis (as appropriate), or, that the settings do not match one of the pre-defined set that you can switch to. Note that data units which match one of the preset values dimensionally will be displayed in the preset format, so may not match values shown elsewhere, say in the plot labels.

When the plot is displaying more than one spectrum, only the current one is actually modified. If you want the other spectra to be drawn as if in the same coordinates or data units, then you must make sure that the menu item **Options**→**Match coordinates and/or fluxes** is selected, but this will only work when all the displayed spectra have proper spectral coordinates and data units defined.

The **SideBand:** control is for switching between sidebands (upper and lower) when display spectra that are derived from a dual sideband instrument. This is not available when your spectra do not have this feature. When displaying a dual sideband spectrum the inactive sideband coordinates will be displayed along the unused X axis (usually the top). You can also use this control to display coordinates offset from the Local Oscillator value.

The **Origin:** control allows you to display offset coordinates. In this case you can select to have no offset (**None**), use the existing offset, if defined (**Default**, usually the same as **None**), or display coordinates offset from the rest frequency. Note for velocities this last option isn't useful as they are already displayed offset from this value. If you want to set another value as the origin then you need to use the advanced toolbox available in the main window.

The **Standard of rest:** control allows you to switch between the various supported rest frames. If the current spectrum doesn't support these the value will be set to **Not set or unrecognised**.

The **Rest Frequency:** field displays the current rest frequency in **GHz**. This can be changed in two ways, either by selecting a position along the plot X axis, which you start by pressing the **Pick pos** button and then clicking on the plot, or by selecting a line identifier, which you do by pressing the **Pick ID** button and clicking on the plot in the vicinity of the desired label. When picking a line identifier the rest frequency is set to that of the line, not of the position that it currently occupies on the plot (these will differ if the spectrum has a source velocity).

Accelerator keys

- **Control-a** Apply UI values converting from existing coordinates and/or data units to new ones.
- **Control-w** Close the window.

- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.19 Flip or shift or redshift a spectrum

This window allows you to do three related jobs.

- Create and optionally flip a copy of the current spectrum.
- Shift an editable spectrum.
- Redshift an editable spectrum.

Flipping and shifting a spectrum allows you to assess the symmetry of a line. Shifting alone to determine how well spectra are aligned and redshifting to transform a spectrum or set of line identifiers to guess or confirm the redshift (all line identifier lists are already editable).

The transformation is applied to the spectrum shown in the **Spectrum:** drop-down menu. Initially this will show any editable spectra displayed in the associated plot (like the line identifiers). If you want an editable copy of the current spectrum just press the **Copy** button.

One subtlety is that the offset value is either a redshift or a value in the coordinate units of the spectrum being modified, depending on whether the **Redshift** checkbox is selected, or not. The coordinate units of the spectrum may not be the same as those of the current spectrum in the plot, which can lead to surprising results.

When applying a redshift it is assumed that the spectrum being modified has a wavelength coordinate system, of some kind, unless it has a proper coordinate system defined. In this latter case the only restriction is that you cannot have a standard of rest set to 'Source' (as a transformation to and from Source is used to determine the redshift transform). In this latter case use the coordinate system toolbox to transform to another rest frame first.

The **Reset** button returns a spectrum to an offset of zero.

Visitor lists

It's often the case that you have a list of lines, in a spectrum, or many spectra, that you'd like to step through to check the radial velocity or redshift estimates, without the bother of reading each spectrum and line, positioning the plot *etc.* That's what a visitor list is designed for.

To use this create a simple text file containing either one, two, three or more whitespace separated columns. If one column is present then that is assumed to be the wavelength of the lines. If two columns are present then these are the wavelength and an associated spectrum. If three or more columns are present then the third columns contains the name of the associated spectrum and the second a label to be displayed at the position (when the label is missing it is set to the wavelength). Very short examples of the three types follow:

```
6562.8170
6583.6000
6716.4700
6730.8500
...
```

```
6562.8170 spectrum1.fits
6562.8170 spectrum2.fits
6562.8170 spectrum3.fits
6562.8170 spectrum4.fits
...
```

```

6562.8170 HALPHA    spectrum1.fits
6583.6000 [NII]    spectrum1.fits
6716.4700 [SII]6717 spectrum2.fits
6730.8500 [SII]6731 spectrum3.fits
...

```

Visitor lists are a type of line identifier spectrum, so can have fully specified spectral coordinate units written in a header section (see the line identifier lists in the SPLAT distribution themselves for details), so you are not restricted to having all your spectra in the same coordinate systems.

Once the visitor list is loaded **File**→**Read visitor line list**, you can start stepping through it using the **Visitor** controls, or jump to a particular line using the drop-down menu. The default behaviour of this tool, on each visit, is to create a copy of the current spectrum and then flip it centred on the line (the current zoom is also preserved). If you'd like to not create a copy of the spectrum for each visit, or indeed to flip it, then uncheck the **Flip** checkbox and select the menu item **Options**→**One spectrum for visitor** (these choices remain enforced between sessions).

Another way to use the visitor controls is to use a line identifier (either a built in one, or one of your own) as the visitor list. To do this load the line identifiers in the usual way (in the main window) and display them in the plot. You can then make this the visitor list by pressing **Set as visitor list**. Now using the visitor controls you can step between the lines (remember to de-select **Flip** and select **Options**→**One spectrum for visitor**, otherwise you'll get the line identifiers copied, flipped and maybe reused!).

SPEFO options

The original motivation for creating this tool (suggested by Petr Skoda, see the acknowledgements section) was really to do flip comparisons of lines to determine radial velocities in the same fashion as the SPEFO program. This has some additional requirements beyond those required for simple shifting in a linear sense, as well as in redshift, and are enabled by selecting the **Options**→**SPEFO options** menu item. This reveals two text areas (you need to resize the window to actually see them first time). The upper one shows the flip corrected offset (that's half the standard shift) and the radial velocity for that flipped shift (calculated as $c * Offset * 0.5 / (Flipcentre)$), assuming that the coordinate system is some wavelength.

There's also an editable area for writing some notes about the measurement, both these areas are written the file SPEFO.log by pressing the **Save to SPEFO.log file** button.

Accelerator keys

- **Control-r** Reset interface to defaults.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.20 Region statistics window

Using this window you can get simple constantly updated statistics on regions of the current spectrum, or fuller statistics for the whole spectrum, or collections of regions.

Constantly updated statistics for regions are:

- Mean
- Standard deviation
- Minimum
- Maximum
- Integrated flux
- TSYS

The integrated flux is a numerical estimate and does not require even coordinate spacing. It can be removed from this list by deselecting the **Options**→**Show flux integral** item. This can be used to quickly estimate the flux in a line, providing your spectra are background subtracted.

The TSYS value is a system temperature estimate and requires data from the JCMT, or you can set the necessary parameters by enabling the **Options**→**Set TSYS parameters** item, which reveals a set of controls not seen by default. Note that the effective exposure time is:

$$t_{eff} = t_{on} * t_{off} / (t_{on} + t_{off})$$

where t_{on} and t_{off} are the integration times spent on and off the source. The backend degradation factor is instrument specific.

When you use the buttons along the bottom of the window these statistics are extended to also include:

- Median
- Mode
- Sum
- Number of data points

Note that the Mode is determined as $3 * Median - 2 * Mean$, not from a distribution, so will not be correct for highly skewed distributions.

If you select the **Options**→**Show extra stats** menu item you can also see:

- Sum of squares
- RMS (square root of the mean square)
- Variance
- Standard error

- 25% and 75% quantiles
- Skew
- Kurtosis

and if you select **Error stats**→you can get basic statistics on the values that are being used as the spectrum errors (or variances).

To get statistics for the whole of the current spectrum just press the **Whole stats** button. If you want statistics for parts of the spectrum you will need to define the coordinate ranges you want to use (see the next part), and then press the **Selected stats** button to get statistics for the selected regions combined together (if none are selected then all regions are used), or **All stats** to get statistics for all the regions combined together.

Coordinate ranges

This part of the window is used to define the regions of the spectra that you want to see statistics for. Once defined you will see simple, continuously updated, statistics. You can also get fuller statistics using the **Selected stats** and **All stats** buttons.

To add a region press the **Add** button and then drag out a region in the display area of the plot window. This should result in the creation of a green rectangular figure.

You can interact with the figure, moving it side-to-side and resizing it. To do this point at the figure and press the left mouse button. This 'selects' the figure and adds grips to its exterior. Note that it also becomes the selected row in the **Coordinate ranges:** table. To move the figure just drag it and to resize it drag a grip (the little black squares). The associated coordinate range in the table updates with these changes.

To add a second range just press **Add** and repeat. The ranges can be overlapped or not.

To fine tune the ranges you can edit the values in the ranges table, just point at the coordinate you want to change and double click the left mouse button. This should enable the text editing cursor. Just make the modifications you want and press <Return> to make the changes permanent. (Note: if your spectra have sky coordinates shown for the X axis, then you should use the same format for your edits).

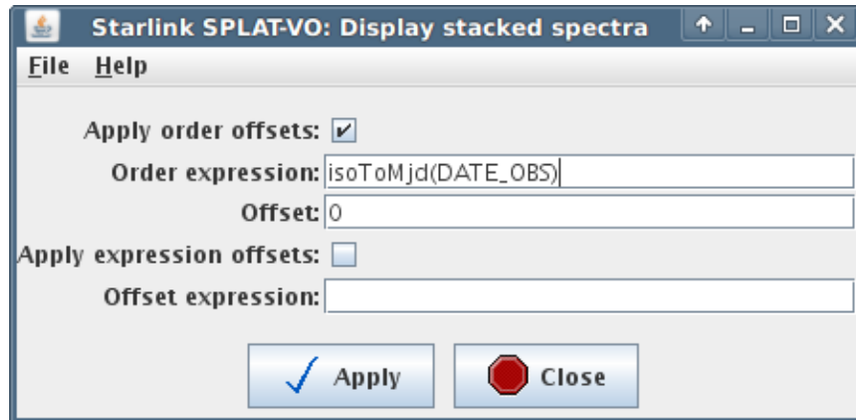
To read a set of ranges from a disk file choose the **File**→**Read ranges** menu item. The format of the input file is simple. It should have at least two fields separated by whitespace or commas. Comments are indicated by lines starting with a hash (#) and are ignored. You can also save the ranges and simple statistics to disk file using (**File**→**Save ranges**). It is also possible to append the contents of the **Full stats log:** region to the file `SPLATStats.log` by pressing the **Save to log file** button.

Accelerator keys

- **Control-s** Show full stats for all the selected ranges.
- **Control-l** Show full stats for all the ranges.

- **Control-h** Show full stats for the whole spectrum,
- **Control-d** Add a coordinate range (interactive or non-interactive).
- **Control-e** Delete the selected coordinate ranges.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

3.21 Display stacked spectra



Using this window you can change the display of multiple spectra so that an offset in data value is artificially applied so that the spectra can be viewed stacked one above the other. It can also be useful when say comparing spectra from different times or physical state, as the stacking order can be defined as an algebraic expression that can include time and dates or the actual offset as an expression of some other physical parameter, like period.

In both modes the offset is a value in the data units of the spectra, which must all be the same, no flux matching is supported.

If supplying an order expression it should evaluate to a number for each spectrum, higher values having their associated spectra placed above lower valued ones by the given offset.

If supplying a offset expression it should evaluate to a physical value, so scaling to appropriate values is often necessary.

Note that the values of any properties associated with the spectra are available for use as part of the expression, usually these are the values of the FITS headers and the keywords should be used.

The expressions are the same as used in the TOPCAT application (see SUN/253 or the TOPCAT builtin help), so include all the special functions it provides, for instance **isoToMjd** will convert an ISO 8601 date into an MJD, which is suitable as a order value. As an example the expression **isoToMjd(DATE_OBS)** will convert the **DATE-OBS** value into an MJD, provided the **DATE-OBS** value is in ISO 8601 format. Note that the minus sign in **DATE-OBS** has been changed into an underscore, that is deliberate. Any character in a keyword that cannot be a Java identifier will be converted into an underscore (allowed characters are any alpha-numeric, plus the underscore).

Accelerator keys

- **Control-a** Apply the current values.
- **Control-w** Close the window.
- **F1** Display help on SPLAT-VO.
- **Shift-F1** Display help on window.

4 Supported data formats

SPLAT-VO can read and write spectra stored in a wide variety of formats, either stored as 1D spectra or as columns in tables. 1D spectral formats supported are Starlink NDF & NDX, FITS and simple TEXT files. Table formats supported are FITS (binary and ascii), VOTable and more complex TEXT files. The 1D data formats currently have the best coordinate system support, so are recommended when a choice can be made. Some support is available for 2 and 3D data, see later).

Normally SPLAT-VO differentiates these types using a preset list of file extensions, for instance NDFs can only be stored in container files that have the extension `.sdf`, FITS files usually have one of the extensions `.fits` or `.fit` and TEXT files `.txt` or `.lis` and NDX format spectra, when stored on disk, will have extension `.xml`. However, you can choose a specific format when opening a file using the **File**→**Open** dialogue, so you can ask that a TEXT file is opened as a table, this allows you to select the columns. There is also a special type known as `guess`. When you choose that attempts to open the spectrum as different types are made until one succeeds, or the possibilities are exhausted.

You can specify a type for all the spectra opened on the command-line using the `--type` argument. Valid values for `--type` are:

```
fits
hds
text
hdx
table
line ids
guess
```

You free to truncate these values as long as they remain unique (so `--type line` will work for line identifiers).

Currently line identifier catalogues represent a special case and currently may have a single format - TEXT based in files with extension `.ids`.

1D spectra stored in TEXT files must have the format of one, two or three columns of data, separated by whitespace (that is spaces or tabs). Any comments in the file must be indicated by a hash sign, blank lines are also allowed. The three columns should have the following values:

```
[wavelength] count [error-in-count]
```

The wavelength and error columns being optional (if you don't have a wavelength column a simple row index is used, note you cannot have errors without wavelengths) . If you choose to open a TEXT file as a table you can select which columns should be used for these values, and the following rules apply:

- Each table row is represented by a single text line
- Anything after a '#' character on a line is ignored (except for optional heading line)

- Blank lines are ignored
- Within a line, fields are separated by one or more whitespace characters (space or tab)
- A field is either a sequence of non-whitespace characters or a sequence of characters between two matching quote characters (single (') or double (") quotes) - spaces are therefore allowed in quoted fields.
- In a quoted field any character preceded by a backslash is interpreted literally, so it is possible to insert quotes into a quoted string
- An empty string (two adjacent quotes) represents a null element
- All lines must contain exactly the same number of fields
- An optional heading line may be included: if the last comment line (line starting with a '#' before the first data line) contains the same number of fields as the columns in the table, these fields are interpreted as the headings (names) of the columns

The format of line identification catalogues is like that of simple TEXT spectral files, except the second column contains a string that identifies the line (with no spaces, *i.e.* H-alpha not H a.lpha). It is also possible to define the system of the coordinate values in a header section, see the SPLAT-VO built-in catalogues for examples of this.

NDFs stored within container files can be accessed, but only by entering the full HDS path on the command-line or in the **File**→**Open** dialogue. FITS extensions may also be open in this fashion (by appending the extension number in square brackets).

An alternative to specifying the component information is to use the component browser window , to see the contents of FITS and NDF files and choose the component to display directly. This uses the same techniques as the special guess type.

Internally all these formats are stored as double precision coordinates and data values. SPLAT-VO only requires that *coordinates increase or decrease monotonically* when performing any analyses.

4.1 Handling 2 and 3D data

If you have images or cubes that contain data that you want to view as spectra in some fashion, then SPLAT-VO will attempt to do this for you. There are three fundamental modes for doing this:

- Collapsing images into a spectrum or the planes of a cube into a set of spectra by adding together all values that lie perpendicular to the dispersion axis (so the spectra run along this axis, what a good spectral displaying tool should do).
- Extracting all spectra from each data line running along the dispersion axis of images and cubes.
- Vectorizing the whole data set into a single data line without any reference to its dimensionality (in effect mapping the whole image/cube as a single contiguous data line).

The default is to collapse. When collapsing cubes you have the option of which axis to step along selecting image planes, this is known as the `select` axis. To determine which option is used to handle your cubes or images you can either set values on the command-line, along with your spectra, or choose options in the open file dialogue. A command line that uses axis 3 as the dispersion axis and steps along axis 2 when choosing image planes might look like:

```
% splat --ndaction collapse --dispax 3 --selectax 2 <spectral_cube> &
```

or in shorthand:

```
% splat -n c -d 3 -s 2 <spectral_cube> &
```

If you don't choose a dispersion axis SPLAT-VO will make a good intelligent guess about this and step along the first axis that isn't the dispersion axis. The other possible values for `ndaction` are `extract` and `vectorize`. These are described as `extract` all spectra and open whole as 1D, respectively, in the open file dialogue.

5 Command-line control and tools

Remote control

The most useful of the remote control scripts are the commands:

- `splatdisp`
- `splatdispmany`

`splatdisp` takes the name of a spectrum and displays it. If a plot identifier is given after the spectrum name (this is the `n` in any names like `<plotn>` that you see) then you can add a spectrum to an existing plot.

`splatdispmany` takes a list of spectra and displays them all in a new plot, much like giving a list of spectra on the command-line when starting SPLAT-VO. The obvious uses of these commands are to have very basic remote control from any scripts that require display facilities.

Both of these commands are available as a standard part of SPLAT-VO *i.e.* you can just type in their names, just like the `splat` command. Command-line usage instructions are available either by typing in the command name, or by inspecting the comments in the file. This is true of all the following commands too.

More sophisticated control of how the spectrum, once plotted, can be displayed is provided by the following scripts:

- `$SPLAT_DIR/zoomandcentre`
- `$SPLAT_DIR/setcolour`
- `$SPLAT_DIR/setproperty`

`zoomandcentre` changes the zoom factor of the wavelength axis and optionally centres it on a given wavelength.

`setcolour` allows you to change the display colour of a spectrum.

`setProperty` is quite similar to `setcolour` but it also allows you to set the line thickness, type and style, as well as whether to display error bars.

Plugins

The one useful example plugin (*i.e.* code that is loaded into SPLAT-VO when it starts up) provides the ability to name a list of spectra in a directory that should be automatically loaded when SPLAT-VO starts up in that directory (this includes a special indicator to load all spectra).

To use this you need to do the following:

```
% setenv SPLAT_PLUGINS $SPLAT_DIR/example_plugin3.bsh
```

Now when SPLAT-VO starts it will look for a file `.splat_autoloads` in the current directory and if found it will read the lines (each of which are assumed to contain a file name) from it to construct a list of spectra to display. If the file only has one line `""` then all the NDFs are automatically loaded. If you'd like other types of spectra to be automatically loaded take a copy of `example_plugin3.bsh` and modify the `FILE_PATTERN` definition line. Now re-define `SPLAT_PLUGINS` to point at your copy.

Command-line SPLAT-VO

`$SPLAT_DIR` also contains several command-line scripts that only make use of SPLAT-VO classes, so don't need SPLAT-VO to be running. These are:

- `$SPLAT_DIR/fitgauss`
- `$SPLAT_DIR/fitgauss2`
- `$SPLAT_DIR/linepositions`

which as you might expect fit gaussians to lines and just locate accurate line positions from a initial list. Just run the commands without any arguments to get usage instructions. Look at the script headers for more details.

A one-off command that is currently only available as a script, but may become available as part of a proper toolbox is:

- `$SPLAT_DIR/deblend`

This fits a blend of spectral lines using a multi-component model, based on any of the three spectral line profiles supported by SPLAT-VO (Gaussian, Lorentz, Voigt). See the contents of this file for instructions on how to use it.

6 Using SAMP to exchange spectra

SPLAT-VO can communicate with other desktop tools using SAMP³ (Simple Applications Messaging Protocol).

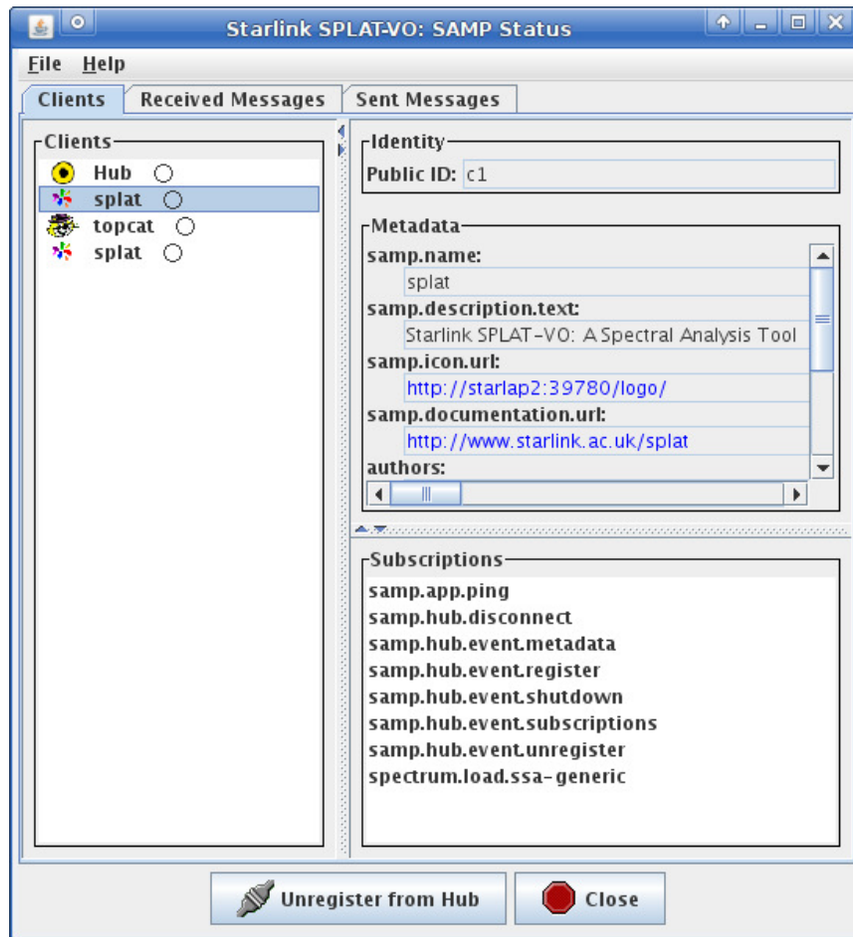
The main job that can be done using SAMP is to exchange spectra. If another tool sends a spectrum to SPLAT-VO, it will appear in the global list as if it had just been loaded. Conversely, SPLAT-VO can send a spectrum to another tool, which will probably respond in a similar way. To send from SPLAT-VO, select a spectrum in the global list, and then use the **Broadcast spectrum** or **Send spectrum to ...** options in the **Interop** menu. The Broadcast option sends to all listening tools, while the Send option sends only to a single selected tool; you choose which one from the submenu on the right. Only tools which (claim they) know what to do with spectral data will be listed; if none of the currently-registered tools is prepared to accept spectra, the Send option will be unavailable.

You can also send the results of an SSA query as a table from the **Query VO for spectra** window. Again, use the **Broadcast** and **Send** options from the **Interop** menu to do this.

The communication architecture of SAMP uses a central “Hub” process to pass messages between registered applications, so a hub must be running in order for the messaging to operate. If a hub is running when SPLAT-VO starts, or if one starts up while SPLAT-VO is in operation, it will connect to it automatically, unless you have started SPLAT-VO with the `--interop none` command-line option. Other tools have their own policies for connecting to the hub, but in general it is a good idea to start a hub first before starting up the tools which you want to talk to it. A hub can be started from within SPLAT-VO if required. The `--hub` or `--exthub` command-line flags start an “internal” or “external” hub respectively; the difference is that an internal hub will terminate when SPLAT-VO exits, while an external one continues running in its own process. Alternatively, a hub can be started from within the GUI.

The **SAMP control** button is visible in the browser window toolbar. Its appearance changes according to whether SPLAT-VO is currently registered:  or unregistered: . In either case, clicking the button will show the SAMP window.

³<http://www.ivoa.net/samp/>



The left hand panel shows which applications are registered, and the right hand panel gives you information about the selected one; it's not necessary to understand this in detail, though some of the information is self-explanatory. If no applications show up in the list, then SPLAT-VO is not registered, most likely because no hub is running. Clicking on the **(Un)Register with Hub** button will give you a chance to start one. The other tabs in the window provide information about messages sent to and from SPLAT-VO. They can be useful to see whether spectra which have been sent have arrived safely or not.

For those interested in the technical details, the domain-specific SAMP MTypes which SPLAT-VO knows about are:

- `spectrum.load.ssa-generic`: send and receive spectra
- `table.load.votable`, `table.load.fits`: send SSA query results

7 Acknowledgements

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- The JDOM Project (<http://www.jdom.org/>).
- The Apache Software Foundation (<http://www.apache.org/>).
- The DIVA project (<http://embedded.eecs.berkeley.edu/diva/>).
- The COLT distribution (<http://hoschek.home.cern.ch/hoschek/colt/>).
- JSci - A science API for Java (<http://jsci.sourceforge.net/>).
- TAMFITS - FITS data access library (<http://heasarc.gsfc.nasa.gov/docs/heasarc/fits/java>).
- EPSGRAPHICS - Java EPS Graphics2D package (GPL version) (<http://www.jibble.org/epsgraphics>).
- JETTY - Java Web server (<http://jetty.mortbay.org/>).

The BeanShell (remote control and scripting) is written by Pat Niemeyer (<http://www.beanshell.org>).

The flip toolbox and Hermitian interpolating polynomials are based on the similiar functionality in the SPEFO program, and were suggested by Petr Skoda (skoda@sunstel.asu.cas.cz, these features are particularly useful for hot star research, contact Petr for further details). Petr is also a major scientific advisor for the VO extensions added to SPLAT-VO.

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When using this software for research please consider citing it in your papers. The main refereed paper is "Spectroscopic analysis in the virtual observatory environment with SPLAT-VO", Škoda et al, *Astronomy & Computing*, 2014, doi:10.1016/j.ascom.2014.06.001.

The Astrophysics Source Code Library entry for SPLAT-VO is ascl:1402.008.

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Finally, many programmers working for Starlink have made contributions to the components used in SPLAT-VO, and many users have reported problems, as well as requests for useful features – thanks to them all.

⁴<http://java.sun.com>

⁵<http://www.sun.com/>