

Guide to the Slit Spectra Reduction Task DOSLIT

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April 1992

ABSTRACT

Doslit extracts, sky subtracts, wavelength calibrates, and flux calibrates simple two dimensional slit spectra which have been processed to remove the detector characteristics; i.e. CCD images have been bias, dark count, and flat field corrected. It is primarily intended for spectrophotometry or radial velocities of stellar spectra with the spectra aligned with one of the image axes; i.e. the assumption is that extractions can be done by summing along image lines or columns. The alignment does not have to be precise but only close enough that the wavelength difference across the spectrum profiles is insignificant. The task is available in the **ctioslit**, **kpnoslit**, **kpnocoude**, and **specred** packages.

1. Introduction

Doslit extracts, sky subtracts, wavelength calibrates, and flux calibrates simple two dimensional slit spectra which have been processed to remove the detector characteristics; i.e. CCD images have been bias, dark count, and flat field corrected. It is primarily intended for spectrophotometry or radial velocities of stellar spectra with the spectra aligned with one of the image axes; i.e. the assumption is that extractions can be done by summing along image lines or columns. The alignment does not have to be precise but only close enough that the wavelength difference across the spectrum profiles is insignificant. Extended objects requiring accurate geometric alignment over many pixels are reduced using the **longslit** package.

The task is a command language script which collects and combines the functions and parameters of many general purpose tasks to provide a single, complete data reduction path and a degree of guidance, automation, and record keeping. In the following description and in the parameter section the various general tasks used are identified. Further information about those tasks and their parameters may be found in their documentation. **Doslit** also simplifies and consolidates parameters from those tasks and keeps track of previous processing to avoid duplications.

The general organization of the task is to do the interactive setup steps, such as the reference dispersion function determination, first using representative calibration data and then perform the majority of the reductions automatically, possibly as a background process, with reference to the setup data. In addition, the task determines which setup and processing operations have been completed in previous executions of the task and, contingent on the `redo` and `update` options, skip or repeat some or all the steps.

The description is divided into a quick usage outline followed by details of the parameters and algorithms. The usage outline is provided as a checklist and a refresher for those familiar with this task and the component tasks. It presents only the default or recommended usage since there are many variations possible.

2. Usage Outline

- [1] The images are first processed with **ccdproc** for overscan, zero level, dark count, and flat field corrections.
- [2] Set the **doslit** parameters with **eparam**. Specify the object images to be processed, one or more arc images, and one or more standard star images. If there are many object, arc, or standard star images you might prepare "@ files". Set the detector and data specific parameters. Select the processing options desired. Finally you might wish to review the `sparams` algorithm parameters though the defaults are probably

^{††}Operated by the Association of Universities for Research in Astronomy, Inc. under cooperative agreement with the National Science Foundation.

adequate.

- [3] Run the task. This may be repeated multiple times with different observations and the task will generally only do the setup steps once and only process new images. Queries presented during the execution for various interactive operations may be answered with "yes", "no", "YES", or "NO". The lower case responses apply just to that query while the upper case responses apply to all further such queries during the current execution and no further queries of that type will be made.
- [4] Apertures are defined for all the standard and object images. This is only done if there are no previous aperture definitions for the image. The highest peak is found and centered and the default aperture limits are set. If the resize option is set the aperture is resized by finding the level which is 5% (the default) of the peak above local background. If not using the quicklook option you now have the option of entering the aperture editing loop to check the aperture position, size, and background fitting parameters, and possibly add additional apertures. This step is highly recommended. It is important to check the background regions with the 'b' key. To exit the background mode and then to exit the review mode use 'q'.

The spectrum positions at a series of points along the dispersion are measured and a function is fit to these positions. If not using the quicklook option the traced positions may be examined interactively and the fitting parameters adjusted. To exit the interactive fitting type 'q'.
- [5] If dispersion correction is selected the first arc in the arc list is extracted. The dispersion function is defined using the task **identify**. Mark and identify a few arc lines with 'm' and use the 'l' line list identification command to automatically add additional lines and fit the dispersion function. Check the quality of the dispersion function fit with 'f'. When satisfied exit with 'q'.
- [6] If the flux calibration option is selected the standard star spectra are processed (if not done previously). The images are extracted and wavelength calibrated. The appropriate arc calibration spectra are extracted and the dispersion function refit using the arc reference spectrum as a starting point. The standard star fluxes through the calibration bandpasses are compiled. You are queried for the name of the standard star calibration data file.

After all the standard stars are processed a sensitivity function is determined using the interactive task **sensfunc**. Finally, the standard star spectra are extinction corrected and flux calibrated using the derived sensitivity function.
- [7] The object spectra are now automatically extracted, wavelength calibrated, and flux calibrated.
- [8] The option to examine the final spectra with **splot** may be given. To exit type 'q'. In quicklook mode the spectra are plotted noninteractively with **bplot**.
- [9] The final spectra will have the same name as the original 2D images with a ".ms" extension added.

3. Spectra and Data Files

The basic input consists of two dimensional slit object, standard star, and arc calibration spectra stored as IRAF images. The raw CCD images must be processed to remove overscan, bias, dark count, and flat field effects. This is generally done using the **ccdred** package. Lines of constant wavelength should be closely aligned with one of the image axes though a small amount of misalignment only causes a small loss of resolution. For large misalignments one may use the **rotate** task. More complex geometric problems and observations of extended objects should be handled by the **longslit** package.

The arc spectra are comparison arc lamp observations (they must all be of the same type). The assignment of arc calibration exposures to object exposures is generally done by selecting the nearest in time and interpolating. However, the optional *arc assignment table* may be used to explicitly assign arc images to specific objects. The format of this file is described in task **refspectra**.

The final reduced spectra are recorded in one, two or three dimensional IRAF images. The images have the same name as the original images with an added ".ms" extension. Each line in the reduced image is a one dimensional spectrum with associated aperture, wavelength, and identification information. With a single aperture the image will be one dimensional and with multiple apertures the image will be two dimensional. When the **extras** parameter is set the images will be three dimensional (regardless of the number of apertures) and the lines in the third dimension contain additional information (see **apsum** for further details). These spectral formats are accepted by the one dimensional spectroscopy tasks such as the plotting tasks **splot** and **specplot**.

4. Package Parameters

The package parameters, shown in Figure 1 for the **specred** package, set parameters which change infrequently and define the standard I/O functions.

Figure 1: Package Parameter Set for DOSLIT Packages

```

                                I R A F
                        Image Reduction and Analysis Facility
PACKAGE = imred
      TASK = specred

(extinct= onedstds$kpnoextinct.dat) Extinction file
(caldir = onedstds$spec16redcal/) Standard star calibration directory
(observa= observatory) Observatory of data
(interp =      poly5) Interpolation type
(dispxi=      2) Image axis for 2D images
(nsum    =      1) Number of lines/columns to sum for 2D images

(databas=      database) Database
(verbose=      no) Verbose output?
(logfile=      logfile) Log file
(plotfil=      ) Plot file

(records=      ) Record number extensions
(version= SPECRED V3: April 1992)
```

The extinction file is used for making extinction corrections and the standard star calibration directory is used for determining flux calibrations from standard star observations. The calibration directories contain data files with standard star fluxes and band passes. The available extinction files and flux calibration directories may be listed using the command:

```
cl> page onedstds$README
```

The extinction correction requires computation of an air mass using the task **setairmass**. The air mass computation needs information about the observation and, in particular, the latitude of the observatory. This is determined using the OBSERVAT image header keyword. If this keyword is not present the observatory parameter is used. See the task **observatory** for more on defining the observatory parameters.

The spectrum interpolation type is used whenever a spectrum needs to be resampled for linearization or performing operations between spectra with different sampling. The "sinc" interpolation may be of interest as an alternative but see the cautions given in **onedspec.package**.

The verbose parameter selects whether to print everything which goes into the log file on the terminal. It is useful for monitoring what the **doslit** task does. The log and plot files are useful for keeping a record of the processing. A log file is highly recommended. A plot file provides a record of the apertures, traces, and extracted spectra but can become quite large. The plotfile is most conveniently viewed and printed with **gkimosaic**.

5. Processing Parameters

The **doslit** parameters are shown in Figure 2.

Figure 2: Parameter Set for DOSLIT

```

                                I R A F
                        Image Reduction and Analysis Facility
PACKAGE = specred
    TASK = doslit

objects =                      List of object spectra
(arcs   =                      ) List of arc spectra
(arctabl=                      ) Arc assignment table (optional)
(standar=                      ) List of standard star spectra

(readnoi=      rdnoise) Read out noise sigma (photons)
(gain   =      gain) Photon gain (photons/data number)
(dispxi=  )_.dispaxis) Dispersion axis (1=along lines, 2=along columns)
(width  =      5.) Width of profiles (pixels)

(dispcor=      yes) Dispersion correct spectra?
(extcor =      no) Extinction correct spectra?
(fluxcal=      no) Flux calibrate spectra?
(resize =      no) Automatically resize apertures?
(clean  =      no) Detect and replace bad pixels?
(splot  =      no) Plot the final spectrum?
(redo   =      no) Redo operations if previously done?
(update =      no) Update spectra if cal data changes?
(quicklo=      no) Minimally interactive quick-look?
(batch  =      no) Extract objects in batch?
(listonl=      no) List steps but don't process?

(sparams=      ) Algorithm parameters

```

The input images are specified by image lists. The lists may be explicit comma separate image names, @ files, or image templates using pattern matching against file names in the directory. To allow wildcard image lists to be used safely and conveniently the image lists are checked to remove extracted images (the .ms images) and to automatically identify object and arc spectra. Object and arc images are identified by the keyword IMAGETYP with values of "object", "OBJECT", "comp", or "COMPARISON" (the current practice at NOAO). If arc images are found in the object list they are transferred to the arc list while if object images are found in the arc list they are ignored. All other image types, such as biases, darks, or flat fields, are ignored. This behavior allows simply specifying all images with a wildcard in the object list with automatic selections of arc spectra or a wildcard in the arc list to automatically find the arc spectra. If the data lack the identifying information it is up to the user to explicitly set the proper lists.

The arc assignment table is a file which may be used to assign specific arc spectra to specific object and standard star spectra. For more on this option see **refspectra**.

The next set of parameters describe the noise characteristics and spectrum characteristics. The read out noise and gain are used when "cleaning" cosmic rays and when using variance or optimal weighting. These parameters must be fairly accurate. Note that these are the effective parameters and must be adjusted if previous processing has modified the pixel values; such as with an unnormalized flat field.

The general direction in which the spectra run is specified by the dispersion axis parameter. Recall that ideally it is the direction of constant wavelength which should be aligned with an image axis and the dispersion direction may not be exactly aligned because atmospheric dispersion. The profile width should be approximately the full width at the profile base. This parameter is used for centering and tracing of the spectrum profiles.

The next set of parameters select the processing steps and options. The various calibration steps may be done simultaneously, that is at the same time as the basic extractions, or in separate executions of the task. Typically, all the desired operations are done at the same time. Dispersion correction requires at least one arc spectrum and flux calibration requires dispersion correction and at least one standard star observation.

The `resize` option resets the edges of the extraction aperture based on the profile for each object and standard star image. The default resizing is to the 5% point relative to the peak measured above the background. This allows

following changes in the seeing. However, one should consider the consequences of this if attempting to flux calibrate the observations. Except in quicklook mode, the apertures for each object and standard star observation may be reviewed graphically and adjustments made to the aperture width and background regions.

The `clean` option invokes a profile fitting and deviant point rejection algorithm as well as a variance weighting of points in the aperture. See the next section for more about requirements to use this option.

Generally once a spectrum has been processed it will not be reprocessed if specified as an input spectrum. However, changes to the underlying calibration data can cause such spectra to be reprocessed if the `update` flag is set. The changes which will cause an update are a new arc reference image and new standard stars. If all input spectra are to be processed regardless of previous processing the `redo` flag may be used. Note that reprocessing clobbers the previously processed output spectra.

The final step is to plot the spectra if the `splot` option is selected. In non-quicklook mode there is a query which may be answered either in lower or upper case. The plotting uses the interactive task **splot**. In quicklook mode the plot appears noninteractively using the task **bplot**.

The `quicklook` option provides a simpler, less interactive, mode. In quicklook mode a single aperture is defined using default parameters without interactive aperture review or trace fitting and the `splot` option selects a noninteractive plot to be shown at the end of processing of each object and standard star spectrum. While the algorithms used in quicklook mode are nearly the same as in non-quicklook mode and the final results may be the same it is recommended that the greater degree of monitoring and review in non-quicklook mode be used for careful final reductions.

The batch processing option allows object spectra to be processed as a background or batch job. This will occur only if the interactive `splot` option is not active; either not set, turned off during processing with "NO", or in quicklook mode. In batch processing the terminal output is suppressed.

The `listonly` option prints a summary of the processing steps which will be performed on the input spectra without actually doing anything. This is useful for verifying which spectra will be affected if the input list contains previously processed spectra. The listing does not include any arc spectra which may be extracted to dispersion calibrate an object spectrum.

The last parameter (excluding the task mode parameter) points to another parameter set for the algorithm parameters. The default parameter set is called `sparams`. The algorithm parameters are discussed further in the next section.

6. Algorithms and Algorithm Parameters

This section summarizes the various algorithms used by the **doslit** task and the parameters which control and modify the algorithms. The algorithm parameters available to you are collected in the parameter set **sparams**. These parameters are taken from the various general purpose tasks used by the **doslit** processing task. Additional information about these parameters and algorithms may be found in the help for the actual task executed. These tasks are identified below. The aim of this parameter set organization is to collect all the algorithm parameters in one place separate from the processing parameters and include only those which are relevant for slit data. The parameter values can be changed from the defaults by using the parameter editor,

```
cl> epar sparams
```

or simple typing `sparams`. The parameter editor can also be entered when editing the **doslit** parameters by typing `:e` when positioned at the `sparams` parameter. Figure 3 shows the parameter set.

Figure 3: Algorithm Parameter Set

```

                                I R A F
                        Image Reduction and Analysis Facility
PACKAGE = specred
    TASK = sparams

(line   =          INDEF) Default dispersion line
(nsum   =           10) Number of dispersion lines to sum
(extras =           no) Extract sky, sigma, etc.?

                                -- DEFAULT APERTURE LIMITS --
(lower  =          -3.) Lower aperture limit relative to center
(upper  =           3.) Upper aperture limit relative to center

                                -- AUTOMATIC APERTURE RESIZING PARAMETERS --
(ylevel =          0.05) Fraction of peak or intensity for resizing

                                -- TRACE PARAMETERS --
(t_step =           10) Tracing step
(t_func=       spline3) Trace fitting function
(t_order=           1) Trace fitting function order
(t_niter=           1) Trace rejection iterations
(t_low  =           3.) Trace lower rejection sigma
(t_high =           3.) Trace upper rejection sigma

                                -- APERTURE EXTRACTION PARAMETERS --
(weights=       none) Extraction weights (none|variance)
(pfit   =       fit1d) Profile fitting algorithm (fit1d|fit2d)
(lsigma =           3.) Lower rejection threshold
(usigma =           3.) Upper rejection threshold

                                -- BACKGROUND SUBTRACTION PARAMETERS --
(backgro=       fit) Background to subtract
(b_func=       legendre) Background function
(b_order=           1) Background function order
(b_sampl=  -10:-6,6:10) Background sample regions
(b_naver=       -100) Background average or median
(b_niter=           1) Background rejection iterations
(b_low  =           3.) Background lower rejection sigma
(b_high =           3.) Background upper rejection sigma

                                -- ARC DISPERSION FUNCTION PARAMETERS --
(coordli=linelists$idhenear.dat) Line list
(match  =           10.) Line list matching limit in Angstroms
(fwidth =           4.) Arc line widths in pixels
(cradius=          10.) Centering radius in pixels
(i_func=       spline3) Coordinate function
(i_order=           1) Order of dispersion function
(i_niter=           0) Rejection iterations
(i_low  =           3.) Lower rejection sigma
(i_high =           3.) Upper rejection sigma
(refit  =       yes) Refit coordinate function when reidentifying?
(addfeat=       no) Add features when reidentifying?

```

```

-- AUTOMATIC ARC ASSIGNMENT PARAMETERS --
(select =      interp) Selection method for reference spectra
(sort    =      jd) Sort key
(group   =     ljd) Group key
(time    =      no) Is sort key a time?
(timewra=    17.) Time wrap point for time sorting

-- DISPERSION CORRECTION PARAMETERS --
(lineari=     yes) Linearize (interpolate) spectra?
(log      =      no) Logarithmic wavelength scale?
(flux     =     yes) Conserve flux?

-- SENSITIVITY CALIBRATION PARAMETERS --
(s_func=    spline3) Fitting function
(s_order=      1) Order of sensitivity function
(fnu       =      no) Create spectra having units of FNU?
```

6.1. Aperture Definitions

The first operation is to define the extraction apertures, which include the aperture width, background regions, and position dependence with wavelength, for the input slit spectra and, if flux calibration is selected, the standard star spectra. This is done only for spectra which do not have previously defined apertures unless the `redo` option is set to force all definitions to be redone. Thus, apertures may be defined separately using the **apextract** tasks. This is particularly useful if one needs to use reference images to define apertures for very weak spectra which are not well centered or traced by themselves.

Initially a single spectrum is found and a default aperture defined automatically. If the `resize` parameter is set the aperture width is adjusted to a specified point on the spectrum profile (see **apresize**). If not in "quicklook" mode (set by the `quicklook` parameter) a query is printed to select whether to inspect and modify the aperture and background aperture definitions using the commands described for **apedit**. This option allows adding apertures for other objects on the slit and adjusting background regions to avoid contaminating objects. The query may be answered in lower case for a single spectrum or in upper case to permanently set the response for the duration of the task execution. This convention for query responses is used throughout the task. It is recommended that quicklook only be used for initial quick extractions and calibration and that for final reductions one at least review the aperture definitions and traces.

The initial spectrum finding and aperture definitions are done at a specified line or column. The positions of the spectrum at a set of other lines or columns is done next and a smooth function is fit to define the aperture centers at all points in the image. In non-quicklook mode the user has the option to review and adjust the function fitting parameters and delete bad position determinations. As with the initial aperture review there is a query which may be answered either in lower or upper case.

The above steps are all performed using tasks from the **apextract** package and parameters from the **sparams** parameters. As a quick summary, the dispersion direction of the spectra are determined from the **dispaxis** parameter if not defined in the image header. The default line or column for finding the object position on the slit and the number of image lines or columns to sum are set by the `line` and `nsum` parameters. A line of INDEF (the default) selects the middle of the image. The automatic finding algorithm is described for the task **apfind** and is basically finds the strongest peak. The default aperture size, background parameters, and resizing are described in the tasks **apdefault** and **apresize** and the parameters used are also described there. The tracing is done as described in **aptrace** and consists of stepping along the image using the specified `t_step` parameter. The function fitting uses the **icfit** commands with the other parameters from the tracing section.

6.2. Extraction

The actual extraction of the spectra is done by summing across the fixed width apertures at each point along the dispersion. The default is to simply sum the pixels using partial pixels at the ends. There is an option to weight the sum based on a Poisson variance model using the `readnoise` and `gain` detector parameters. Note that if the `clean` option is selected the variance weighted extraction is used regardless of the `weights` parameter. The sigma thresholds for cleaning are also set in the **sparams** parameters.

The cleaning and variance weighting options require knowing the effective (i.e. accounting for any image combining) read out noise and gain. These numbers need to be adjusted if the image has been processed such that the intensity

scale has a different origin (such as applying a separate background subtraction operation) or scaling (such as caused by unnormalized flat fielding). These options also require using background subtraction if the profile does not go to zero. For optimal extraction and cleaning to work it is recommended that any flat fielding be done using normalized flat fields (as is done in **ccdproc**) and using background subtraction if there is any appreciable sky. For further discussion of cleaning and variance weighted extraction see **apvariance** and **approfiles** as well as **apsum**.

Background sky subtraction is done during the extraction based on background regions and parameters defined by the default parameters or changed during the interactive setting of the apertures. The background subtraction options are to do no background subtraction, subtract the average, median, or minimum of the pixels in the background regions, or to fit a function and subtract the function from under the extracted object pixels. The background regions are specified in pixels from the aperture center and follow changes in center of the spectrum along the dispersion. The syntax is colon separated ranges with multiple ranges separated by a comma or space. The background fitting uses the **icfit** routines which include medians, iterative rejection of deviant points, and a choice of function types and orders. Note that it is important to use a method which rejects cosmic rays such as using either medians over all the background regions (`background = "median"`) or median samples during fitting (`b_naverage < -1`). The background subtraction algorithm and options are described in greater detail in **apsum** and **apbackground**.

6.3. Dispersion Correction

If dispersion correction is not selected, `dispcor=no`, then the object spectra are simply extracted. The extracted spectra may be plotted by setting the `splot` option. This produces a query and uses the interactive **splot** task in non-quicklook mode and uses the noninteractive **bplot** task in quicklook mode.

Dispersion corrections are applied to the extracted spectra if the `dispcor` processing parameter is set. There are three basic steps involved; determining the dispersion functions relating pixel position to wavelength, assigning the appropriate dispersion function to a particular observation, and either storing the nonlinear dispersion function in the image headers or resampling the spectra to evenly spaced pixels in wavelength.

The first arc spectrum in the arc list is used to define the reference dispersion solution. It is extracted at middle of the image with no tracing. Note extractions of arc spectra are not background subtracted. The interactive task **identify** is used to define the dispersion function. The idea is to mark some lines whose wavelengths are known (with the line list used to supply additional lines after the first few identifications define the approximate wavelengths) and to fit a function giving the wavelength from the pixel position.

The arc dispersion function parameters are for **identify** and it's related partner **reidentify**. The parameters define a line list for use in automatically assigning wavelengths to arc lines, a centering width (which should match the line widths at the base of the lines), the dispersion function type and orders, parameters to exclude bad lines from function fits, and defining whether to refit the dispersion function as opposed to simply determining a zero point shift. The defaults should generally be adequate and the dispersion function fitting parameters may be altered interactively. One should consult the help for the two tasks for additional details of these parameters and the interactive operation of **identify**.

The extracted reference arc spectrum is then dispersion corrected. If the spectra are to be linearized, as set by the `linearize` parameter, the default linear wavelength parameters are printed and you have the option to adjust them. The dispersion system defined at this point will be applied automatically to all other spectra as they are dispersion corrected.

Once the reference dispersion function is defined other arc spectra are extracted as required by the object spectra. The assignment of arcs is done either explicitly with an arc assignment table (parameter `arctable`) or based on a header parameter such as a time. This assignments are made by the task **refspectra**. When two arcs are assigned to an object spectrum an interpolation is done between the two dispersion functions. This makes an approximate correction for steady drifts in the dispersion.

The tasks **setjd** and **setairmass** are automatically run on all spectra. This computes and adds the header parameters for the Julian date (JD), the local Julian day number (LJD), the universal time (UTMIDDLE), and the air mass at the middle of the exposure. The default arc assignment is to use the Julian date grouped by the local Julian day number. The grouping allows multiple nights of data to be correctly assigned at the same time.

The assigned arc spectra are then extracted using the object aperture definitions (but without background subtraction or cleaning) so that the same pixels on the detector are used. The extracted arc spectra are then reidentified automatically against the reference arc spectrum. Some statistics of the reidentification are printed (if not in batch mode) and the user has the option of examining the lines and fits interactively if not in quicklook mode. The task which does the reidentification is called **reidentify**.

The last step of dispersion correction is setting the dispersion of the object image from the arc images. There are two choices here. If the `linearize` parameter is not set the nonlinear dispersion function is stored in the image

header. Other IRAF tasks interpret this information when dispersion coordinates are needed for plotting or analysis. This has the advantage of not requiring the spectra to be interpolated and the disadvantage that the dispersion information is only understood by IRAF tasks and cannot be readily exported to other analysis software.

If the `linearize` parameter is set then the spectra are resampled to a linear dispersion relation either in wavelength or the log of the wavelength using the dispersion coordinate system defined previously for the arc reference spectrum.

The linearization algorithm parameters allow selecting the interpolation function type, whether to conserve flux per pixel by integrating across the extent of the final pixel, and whether to linearize to equal linear or logarithmic intervals. The latter may be appropriate for radial velocity studies. The default is to use a fifth order polynomial for interpolation, to conserve flux, and to not use logarithmic wavelength bins. These parameters are described fully in the help for the task **dispcor** which performs the correction.

6.4. Flux Calibration

Flux calibration consists of an extinction correction and an instrumental sensitivity calibration. The extinction correction only depends on the extinction function defined by the package parameter `extinct` and determination of the airmass from the header parameters (the air mass is computed by **setairmass** as mentioned earlier). The sensitivity calibration depends on a sensitivity calibration spectrum determined from standard star observations for which there are tabulated absolute fluxes. The task that applies both the extinction correction and sensitivity calibration to each extracted object spectrum is **calibrate**. Consult the manual page for this task for more information.

Generation of the sensitivity calibration spectrum is done before processing any object spectra since it has two interactive steps and requires all the standard star observations. The first step is tabulating the observed fluxes over the same bandpasses as the calibrated absolute fluxes. The standard star tabulations are done after each standard star is extracted and dispersion corrected. You are asked for the name of the standard star as tabulated in the absolute flux data files in the directory `caldir` defined by the package parameters. The tabulation of the standard star observations over the standard bandpasses is done by the task **standard**. The tabulated data is stored in the file `std`. Note that if the `redo` flag is not set any new standard stars specified in subsequent executions of **doslit** are added to the previous data in the data file, otherwise the file is first deleted. Modification of the tabulated standard star data, such as by adding new stars, will cause any spectra in the input list which have been previously calibrated to be reprocessed if the `update` flag is set.

After the standard star calibration bandpass fluxes are tabulated the information from all the standard stars is combined to produce a sensitivity function for use by **calibrate**. The sensitivity function determination is interactive and uses the task **sensfunc**. This task allows fitting a smooth sensitivity function to the ratio of the observed to calibrated fluxes versus wavelength. The types of manipulations one needs to do include deleting bad observations, possibly removing variable extinction (for poor data), and possibly deriving a revised extinction function. This is a complex operation and one should consult the manual page for **sensfunc**. The sensitivity function is saved as a one dimensional spectrum with the name `sens`. Deletion of this image will also cause reprocessing to occur if the `update` flag is set.

7. References

7.1. IRAF Introductory References

Work is underway on a new introductory guide to IRAF. Currently, the work below is the primary introduction.

P. Shames and D. Tody, *A User's Introduction to the IRAF Command Language*, Central Computer Services, NOAO, 1986.

7.2. CCD Reductions

F. Valdes, *The IRAF CCD Reduction Package -- CCDRED*, Central Computer Services, NOAO, 1987.

F. Valdes, *User's Guide to the CCDRED Package*, Central Computer Services, NOAO, 1988. Also on-line as `help ccdred.guide`.

P. Massey, *A User's Guide to CCD Reductions with IRAF*, Central Computer Services, NOAO, 1989.

7.3. Aperture Extraction Package

F. Valdes, *The IRAF APEXTRACT Package*, Central Computer Services, NOAO, 1987 (out-of-date).

7.4. DOSLIT Task

P. Massey, *User's Guide to Slit Spectra Reductions*, Central Computer Services, NOAO, 1992.

7.5. Task Help References

Each task in the **specred** packages and tasks used by **doslit** have help pages describing the parameters and task in some detail. To get on-line help type

```
cl> help taskname
```

The output of this command can be piped to **lprint** to make a printed copy.

```
apall - Extract 1D spectra (all parameters in one task)
apdefault - Set the default aperture parameters and apidtable
apedit - Edit apertures interactively
apfind - Automatically find spectra and define apertures
apfit - Fit 2D spectra and output the fit, difference, or ratio
apflatten - Remove overall spectral and profile shapes from flat fields
apmask - Create and IRAF pixel list mask of the apertures
apnormalize - Normalize 2D apertures by 1D functions
aprecenter - Recenter apertures
apresize - Resize apertures
apscatter - Fit and subtract scattered light
apsum - Extract 1D spectra
aptrace - Trace positions of spectra

bplot - Batch plot of spectra with SPLOT
calibrate - Extinction and flux calibrate spectra
continuum - Fit the continuum in spectra
deredden - Apply interstellar extinction correction
dispcor - Dispersion correct spectra
dopcor - Doppler correct spectra
fitprofs - Fit gaussian profiles
identify - Identify features in spectrum for dispersion solution
msrespld - Create 1D response spectra from flat field and sky spectra
refspectra - Assign wavelength reference spectra to other spectra
reidentify - Automatically reidentify features in spectra
sapertures - Set or change aperture header information
sarith - Spectrum arithmetic
scombine - Combine spectra
scopy - Select and copy apertures in different spectral formats
sensfunc - Compute instrumental sensitivity from standard stars
setairmass - Compute effective airmass and middle UT for an exposure
setjd - Compute and set Julian dates in images
sfit - Fit spectra and output fit, ratio, or difference
skysub - Sky subtract extracted multispec spectra
slist - List spectrum header parameters
specplot - Scale, stack, and plot multiple spectra
splot - Preliminary spectral plot/analysis
standard - Tabulate standard star counts and fluxes

doslit - Process slit spectra
demos - Demonstrations and tests
```

Additional help topics

```
onedspec.package - Package parameters and general description of package
apextract.package - Package parameters and general description of package
approfiles - Profile determination algorithms
apvariance - Extractions, variance weighting, cleaning, and noise model
```

center1d - One dimensional centering algorithm
icfit - Interactive one dimensional curve fitting

Appendix A: DOSLIT Parameters

objects

List of object images to be processed. Previously processed spectra are ignored unless the `redo` flag is set or the `update` flag is set and dependent calibration data has changed. If the images contain the keyword `IMAGETYP` then only those with a value of "object" or "OBJECT" are used and those with a value of "comp" or "COMPARISON" are added to the list of arcs. Extracted spectra are ignored.

arcs = "" (at least one if dispersion correcting)

List of arc calibration spectra. These spectra are used to define the dispersion functions. The first spectrum is used to mark lines and set the dispersion function interactively and dispersion functions for all other arc spectra are derived from it. If the images contain the keyword `IMAGETYP` then only those with a value of "comp" or "COMPARISON" are used. All others are ignored as are extracted spectra.

arctable = "" (optional) (refspectra)

Table defining which arc spectra are to be assigned to which object spectra (see **refspectra**). If not specified an assignment based on a header parameter, `sparams.sort`, such as the Julian date is made.

standards = "" (at least one if flux calibrating)

List of standard star spectra. The standard stars must have entries in the calibration database (package parameter `caldir`).

readnoise = "rdnoise", gain = "gain" (apsum)

Read out noise in photons and detector gain in photons per data value. This parameter defines the minimum noise sigma and the conversion between photon Poisson statistics and the data number statistics. Image header keywords (case insensitive) may be specified to obtain the values from the image header.

dispaxis = "_dispaxis" (apextract)

Default dispersion axis. The dispersion axis is 1 for dispersion running along image lines and 2 for dispersion running along image columns. If the image header parameter `DISPAXIS` is defined it has precedence over this parameter. The default value defers to the package parameter of the same name.

width = 5. (apedit)

Approximate full width of the spectrum profiles. This parameter is used to define a width and error radius for the profile centering algorithm.

dispcor = yes

Dispersion correct spectra? This may involve either defining a nonlinear dispersion coordinate system in the image header or resampling the spectra to uniform linear wavelength coordinates as selected by the parameter `sparams.linearize`.

extcor = no

Extinction correct the spectra?

fluxcal = no

Flux calibrate the spectra using standard star observations?

resize = no (apresize)

Resize the default aperture for each object based on the spectrum profile?

clean = no (apsum)

Detect and correct for bad pixels during extraction? This is the same as the `clean` option in the **apextract** package. If yes this also implies variance weighted extraction.

splot = no

Plot the final spectra with the task **splot**? In quicklook mode this is automatic and in non-quicklook mode it is queried.

redo = no

Redo operations previously done? If no then previously processed spectra in the object list will not be processed unless required by the `update` option.

update = no

Update processing of previously processed spectra if the dispersion reference image or standard star calibration data are changed?

quicklook = no

Extract and calibrate spectra with minimal interaction? In quicklook mode only the initial dispersion function solution and standard star setup are done interactively. Normally the `splot` option is set in this mode to produce an automatic final spectrum plot for each object. It is recommended that this mode not be used for final reductions.

batch = yes

Process spectra as a background or batch job provided there are no interactive steps remaining.

listonly = no

List processing steps but don't process?

sparams = "" (pset)

Name of parameter set containing additional processing parameters. This parameter is only for indicating the link to the parameter set **sparams** and should not be given a value. The parameter set may be examined and modified in the usual ways (typically with "eparam sparams" or ":e sparams" from the parameter editor). The parameters are described below.

-- GENERAL PARAMETERS --

line = INDEF, nsum = 10

The dispersion line (line or column perpendicular to the dispersion axis) and number of adjacent lines (half before and half after unless at the end of the image) used in finding, resizing, editing, and tracing operations. A line of INDEF selects the middle of the image along the dispersion axis.

extras = no (apsum)

Include raw unweighted and uncleaned spectra, the background spectra, and the estimated sigmas in a three dimensional output image format. See the discussion in the **apextract** package for further information.

-- DEFAULT APERTURE LIMITS --

lower = -3., upper = 3. (apdefault)

Default lower and upper aperture limits relative to the aperture center. These limits are used when the apertures are first defined.

-- AUTOMATIC APERTURE RESIZING PARAMETERS --

ylevel = 0.05 (apresize)

Fraction of the peak to set aperture limits during automatic resizing.

-- TRACE PARAMETERS --

t_step = 10 (aptrace)

Step along the dispersion axis between determination of the spectrum positions. Note the `nsum` parameter is also used to enhance the signal-to-noise at each step.

t_function = "spline3", t_order = 1 (aptrace)

Default trace fitting function and order. The fitting function types are "chebyshev" polynomial, "legendre" polynomial, "spline1" linear spline, and "spline3" cubic spline. The order refers to the number of terms in the polynomial functions or the number of spline pieces in the spline functions.

t_niterate = 1, t_low = 3., t_high = 3. (aptrace)

Default number of rejection iterations and rejection sigma thresholds.

-- APERTURE EXTRACTION PARAMETERS --

weights = "none" (apsum) (none|variance)

Type of extraction weighting. Note that if the `clean` parameter is set then the weights used are "variance" regardless of the weights specified by this parameter. The choices are:

"none"

The pixels are summed without weights except for partial pixels at the ends.

"variance"

The extraction is weighted by the variance based on the data values and a poisson/ccd model using the `gain` and `readnoise` parameters.

pfit = "fit1d" (apsum and approfile) (fit1d|fit2d)

Type of profile fitting algorithm to use. The "fit1d" algorithm is preferred except in cases of extreme tilt.

lsigma = 3., usigma = 3. (apsum)

Lower and upper rejection thresholds, given as a number of times the estimated sigma of a pixel, for cleaning.

-- DEFAULT BACKGROUND PARAMETERS --

background = "fit" (apsum) (none|average|median|minimum|fit)

Type of background subtraction. The choices are "none" for no background subtraction, "average" to average the background within the background regions, "median" to use the median in the background regions, "minimum" to use the minimum in the background regions, or "fit" to fit across the dispersion using the background within the background regions. Note that the "average" option does not do any medianing or bad pixel checking, something which is recommended. The fitting option is slower than the other options and requires additional fitting parameter.

b_function = "legendre", b_order = 1 (apsum)

Default background fitting function and order. The fitting function types are "chebyshev" polynomial, "legendre" polynomial, "spline1" linear spline, and "spline3" cubic spline. The order refers to the number of terms in the polynomial functions or the number of spline pieces in the spline functions.

b_sample = "-10:-6,6:10" (apsum)

Default background sample. The sample is given by a set of colon separated ranges each separated by either whitespace or commas. The string "*" refers to all points. Note that the background coordinates are relative to the aperture center and not image pixel coordinates so the endpoints need not be integer. It is recommended that the background regions be examined and set interactively with the 'b' key in the interactive aperture definition mode. This requires `quicklook` to be no.

b_naverage = -100 (apsum)

Default number of points to average or median. Positive numbers average that number of sequential points to form a fitting point. Negative numbers median that number, in absolute value, of sequential points. A value of 1 does no averaging and each data point is used in the fit.

b_niterate = 1 (apsum)

Default number of rejection iterations. If greater than zero the fit is used to detect deviant fitting points and reject them before repeating the fit. The number of iterations of this process is given by this parameter.

b_low_reject = 3., b_high_reject = 3. (apsum)

Default background lower and upper rejection sigmas. If greater than zero points deviating from the fit below and above the fit by more than this number of times the sigma of the residuals are rejected before refitting.

-- ARC DISPERSION FUNCTION PARAMETERS --

coordlist = "linelists\$ldheneard.dat" (identify)

Arc line list consisting of an ordered list of wavelengths. Some standard line lists are available in the directory "linelists\$".

match = 10. (identify)

The maximum difference for a match between the dispersion function computed value and a wavelength in the coordinate list.

fwidth = 4. (identify)

Approximate full base width (in pixels) of arc lines.

cradius = 10. (reidentify)

Radius from previous position to reidentify arc line.

i_function = "spline3", i_order = 1 (identify)

The default function and order to be fit to the arc wavelengths as a function of the pixel coordinate. The functions choices are "chebyshev", "legendre", "spline1", or "spline3".

i_niterate = 0, i_low = 3.0, i_high = 3.0 (identify)

Number of rejection iterations and sigma thresholds for rejecting arc lines from the dispersion function fits.

refit = yes (reidentify)

Refit the dispersion function? If yes and there is more than 1 line and a dispersion function was defined in the initial arc reference then a new dispersion function of the same type as in the reference image is fit using the new pixel positions. Otherwise only a zero point shift is determined for the revised fitted coordinates without changing the form of the dispersion function.

addfeatures = no (reidentify)

Add new features from a line list during each reidentification? This option can be used to compensate for lost features from the reference solution. Care should be exercised that misidentified features are not introduced.

-- AUTOMATIC ARC ASSIGNMENT PARAMETERS --

select = "interp" (refspectra)

Selection method for assigning wavelength calibration spectra. Note that an arc assignment table may be used to override the selection method and explicitly assign arc spectra to object spectra. The automatic selection methods are:

average

Average two reference spectra without regard to any sort or group parameters. If only one reference spectrum is specified then it is assigned with a warning. If more than two reference spectra are specified then only the first two are used and a warning is given. There is no checking of the group values.

following

Select the nearest following spectrum in the reference list based on the sort and group parameters. If there is no following spectrum use the nearest preceding spectrum.

interp

Interpolate between the preceding and following spectra in the reference list based on the sort and group parameters. If there is no preceding and following spectrum use the nearest spectrum. The interpolation is weighted by the relative distances of the sorting parameter (see cautions in DESCRIPTION section).

match

Match each input spectrum with the reference spectrum list in order. This overrides any group values.

nearest

Select the nearest spectrum in the reference list based on the sort and group parameters.

preceding

Select the nearest preceding spectrum in the reference list based on the sort and group parameters. If there is no preceding spectrum use the nearest following spectrum.

sort = "jd" (setjd and refspectra)

Image header keyword to be used as the sorting parameter for selection based on order. The header parameter must be numeric but otherwise may be anything. Common sorting parameters are times or positions.

group = "ljd" (setjd and refspectra)

Image header keyword to be used to group spectra. For those selection methods which use the group parameter the reference and object spectra must have identical values for this keyword. This can be anything but it must be constant within a group. Common grouping parameters are the date of observation "date-obs" (provided it does not change over a night) or the local Julian day number.

time = no, timewrap = 17. (refspectra)

Is the sorting parameter a 24 hour time? If so then the time origin for the sorting is specified by the timewrap

parameter. This time should precede the first observation and follow the last observation in a 24 hour cycle.

-- DISPERSION CORRECTION PARAMETERS --

linearize = yes (dispcor)

Interpolate the spectra to a linear dispersion sampling? If yes the spectra will be interpolated to a linear or log linear sampling using the linear dispersion parameters specified by other parameters. If no the nonlinear dispersion function(s) from the dispersion function database are assigned to the input image world coordinate system and the spectral data is not interpolated. Note the interpolation function type is set by the package parameter `interp`.

log = no (dispcor)

Use linear logarithmic wavelength coordinates? Linear logarithmic wavelength coordinates have wavelength intervals which are constant in the logarithm of the wavelength.

flux = yes (dispcor)

Conserve the total flux during interpolation? If no the output spectrum is interpolated from the input spectrum at each output wavelength coordinate. If yes the input spectrum is integrated over the extent of each output pixel. This is slower than simple interpolation.

-- SENSITIVITY CALIBRATION PARAMETERS --

s_function = "spline3", s_order = 1 (sensfunc)

Function and order used to fit the sensitivity data. The function types are "chebyshev" polynomial, "legendre" polynomial, "spline3" cubic spline, and "spline1" linear spline. Order of the sensitivity fitting function. The value corresponds to the number of polynomial terms or the number of spline pieces. The default values may be changed interactively.

fnu = no (calibrate)

The default calibration is into units of F-lambda. If `fnu` = yes then the calibrated spectrum will be in units of F-nu.

PACKAGE PARAMETERS

The following package parameters are used by this task. The default values may vary depending on the package.

extinction (standard, sensfunc, calibrate)

Extinction file for a site. There are two extinction files in the NOAO standards library, `onedstds$`, for KPNO and CTIO. These extinction files are used for extinction and flux calibration.

caldir (standard)

Standard star calibration directory. A directory containing standard star data files. Note that the directory name must end with `'/'`. There are a number of standard star calibrations directories in the NOAO standards library, `onedstds$`.

observatory = "observatory" (observatory)

The default observatory to use for latitude dependent computations. If the `OBSERVAT` keyword in the image header it takes precedence over this parameter.

interp = "poly5" (nearest|linear|poly3|poly5|spline3|sinc) (dispcor)

Spectrum interpolation type used when spectra are resampled. The choices are:

- nearest - nearest neighbor
- linear - linear
- poly3 - 3rd order polynomial
- poly5 - 5th order polynomial
- spline3 - cubic spline
- sinc - sinc function

database = "database"

Database name used by various tasks. This is a directory which is created if necessary.

`verbose = no`

Verbose output? If set then almost all the information written to the logfile is also written to the terminal except when the task is a background or batch process.

`logfile = "logfile"`

If specified detailed text log information is written to this file.

`plotfile = ""`

If specified metacode plots are recorded in this file for later review. Since plot information can become large this should be used only if really desired.