Wave Tracking Code Document

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February 29, 2016

$\underline{\mathbf{History}}$

Version 1 - 03/2016 - Basic operating information

1 Introduction

If you spot any mistakes or suggestions please email richard.morton@northumbria.ac.uk .

2 Selecting data sets

The automated wave measurement tools are designed to analyse features in time-distance (TD) diagrams. There are three tools available to create time-distance diagrams, diag_slit.pro, track_rout.pro and spline_slit.pro.

2.1 diag_slit

The diag_slit.pro routine is a simple point and click routine that lets you pick two data points, calculates a straight line between the two points and extracts the intensity values along the line via cubic interpolation, doing this for all time-frames in the data cube.

A typical calling procedure for diag_slit is:

 $IDL > diag_slit, data_cube, slit$

where *slit* is the outputted time-distance diagram. In Figure 1 a typical example of data selection is given and the output shown is shown in Figure 2.

2.2 track_rout

2.3 spline_slit

This routine probably the better of the three, but is slightly more difficult to use. A full description of the routine is given in an alternate document.

3 Wave tracking

The basic premise behind the wave tracking routine is just a simple feature finding routine. The routine identifies peaks in intensity throughout the TD diagram of interest. A peak is defined as the local maximum intensity value in a search box, whose size can be set by the user. The gradient of the slopes either side of the local maximum are used to determine whether the found local maximum is identified as a peak or not. The local maximum is required to have a positive and negative slope on the left and right hand side respectively. The user also has to select the minimum value of the gradient of the slope, i.e., a cut-off value. This value is not the same for different data sets and the user should test different values for the gradient cut-off. A gradient too large selects no points, a gradient cut-off that is too small ends up picking up the noise in the data, as well as the signal. The routine essentially scans through the TD diagram, identifying all peaks that meet the criteria. NOTE: only positive peaks are found - so if your interested in features that have lower intensities than the background (e.g., fibrils) you will need to invert the intensities! (It is recommended to multiply the TD by -1).

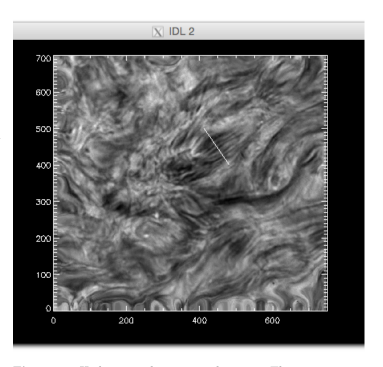


Figure 1: Hydrogen α line centre data set. The units are in pixels and the white slit shows the location of the slit from which the data for the time-distance diagram is extracted using $diag_slit$. The slit is placed across a large-scale feature composed of many fine-scale dark fibrils.

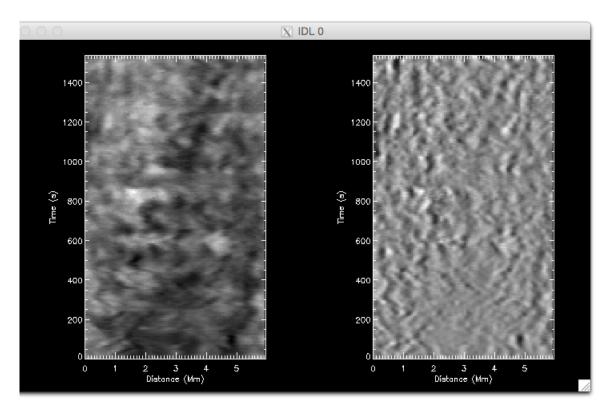


Figure 2: Time-distance-diagrams. The left hand panel shows the intensity time-distance diagram from the slit in Figure 1. The right hand panel is an unsharp mask version of the same data, revealing more clearly the swaying of the dark fibrils.

The routine has two modes for finding the location of the peaks, one more accurate than the other. The first mode just takes the given location of maximum intensity at pixel level accuracy, and are given an error that is ± 0.5 pixel. This is a clearly a very coarse method.

The second mode is based on the least-squares fitting of a Gaussian function to the peak in intensity, allowing for a sub-pixel estimate of the location of the intensity peak.

3.1 quick_plot_ft

The various wave tracking routines are made up of a number of smaller sub-routines that perform various tasks. The routine *quick_plot_ft* is essentially a quick look routine, to test out how various input parameters influence the location and fitting of intensity peaks.

The basic call is: IDL>quick_plot_ft,td

where td is the time-distance diagram to process. This call uses the basic method for peak finding. The user will be prompted to input a gradient cut-off value. A suitable value of the gradient for different data sets will require some trial and error. In Figure 3 we show the output from the routine (default plot to screen), the result is shown for the case when the cut-off is far to restrictive, and no peaks have been selected. A better selection of the gradient finds the structure, but doesn't introduce too much noise into the found peaks, an example of which is shown in Figure 4.

In Figure 4 there is clearly the influence of noise. The influence of noise can be reduced, e.g., by averaging, smoothing, or a high-frequency noise filtering technique (Figure 5).

To test the second fitting method, the call is:

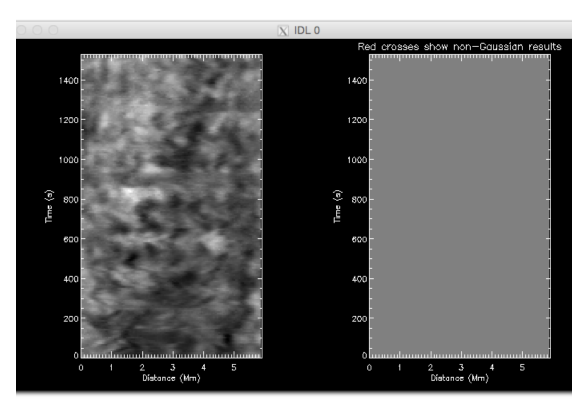


Figure 3: Peak finding. The left hand panel shows the intensity time-distance diagram. The right hand panel shows the results of a harsh gradient constraint - nothing is found!

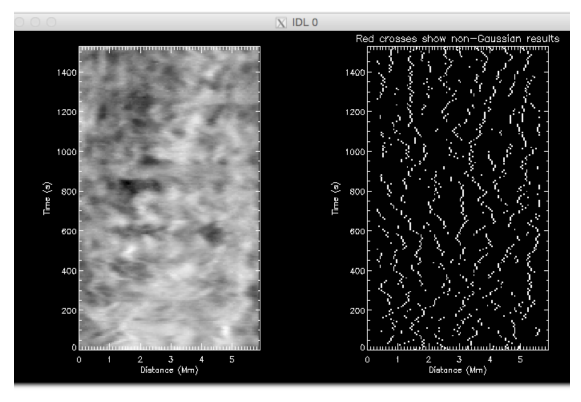


Figure 4: Peak finding. The left hand panel shows the intensity time-distance diagram. The right hand panel shows the results of the peak location.

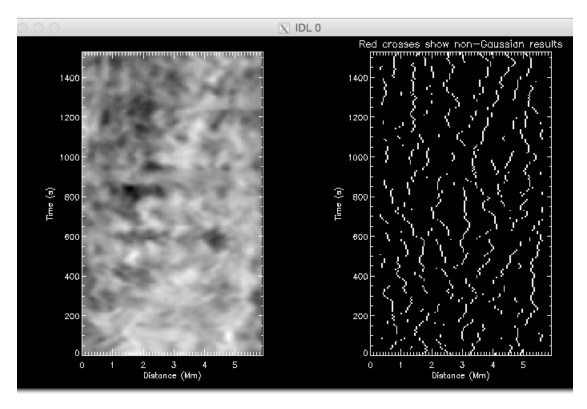


Figure 5: Peak finding. The left hand panel shows the intensity time-distance diagram. The right hand panel shows the results of the peak location, with a smoothing of the TD diagram in the time-direction.

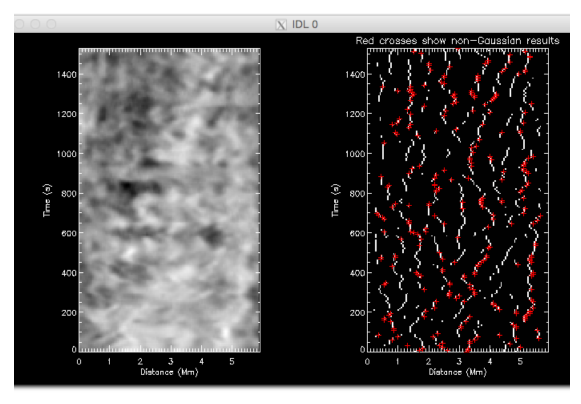


Figure 6: Peak finding. The left hand panel shows the intensity time-distance diagram. The right hand panel shows the results of the peak location, using smoothed TD and the Gaussian peak location. The red crosses show where the Gaussian fit was rejected and the basic peak location is used.

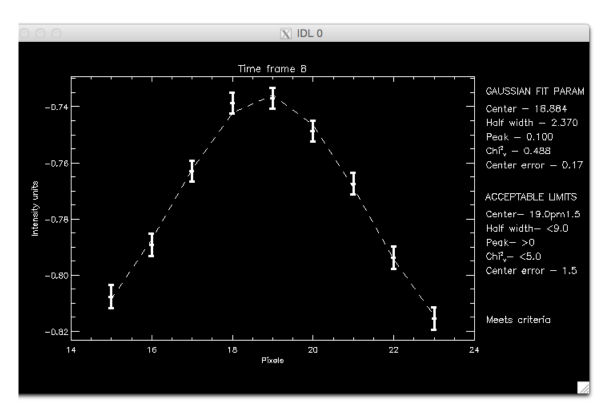


Figure 7: Example of fitted fibril cross-section.

 $IDL>quick_plot_ft,td,/gauss,error=td_er$

Here, the td_err is the uncertainties on the intensity values in the time-distance diagram. It should be the same dimensions and size as td. The uncertainties are used to weight the Gaussian fit. They are required for the code to run as it makes judgements about the goodness of fit of the Gaussian to the peak. If the peak doesn't meet the goodness of fit criteria, then the location of the peak is recorded using the basic properties, of the first mode.

In Figure 6 the output window is shown for the Gaussian fitting. The red crosses show the locations where the fit of the Gaussian failed to meet a list of criteria (see Section). For this example the assumed uncertainties of 0.5% of the intensity value. NOTE: careful analysis of data uncertainties is highly recommended! Otherwise results will not accurately reflect the uncertainties of the peak location.

There are a number of other keywords that lets you play with different parameters.

$3.2 \quad track_rout$

The track_rout routine uses the same feature finding as discussed for quick_plot_ft. Once the peaks have been identified, an another sub-routine then 'stitches' together the peaks by crawling through the data. The routine starts at a peak and searches forward in time for the next 'local' peak. The search box is 5 pixels in space by 4 in time. If no pixel is found in the box, the thread is considered to be finished. The means small jumps in time are allowed for.

The basic calls for the routine are:

 $IDL > track_rout, data = td$

or

 $IDL > track_rout, data = td, /gauss, error = td_er$

Here, data can be either one or many TD diagrams, in an array that is ordered as [x,t,n]. The user will be prompted to enter a value for the gradient cut-off and also a minimum thread length. The routine will then reject any thread whose length is less than that value. Further, any thread that has more than half of its value set to 0 will be rejected. A window will be plotted to screen that shows the TD and the found peaks - there is no indication at this stage of how many peak fittings failed for the Gaussian fit. Note, all peaks found are plotted - this image does not reflect the threads found.

Once the fitting and stitching is complete, the user will be prompted as to whether they want to fit the threads. If y, then the it takes you to an interactive fitting process, the details of which are given in the next sub-sub-section. Answering n skips the fitting.

After the fitting stage, user is then asked whether they want the results to be saved. If y is chosen the you will be prompted to enter a location to save the results. Once you have entered the information the routine will either quit (if only one TD was input) or repeat (for many TD).

On opening the saved file, the user will find the outputs generated by the routine are saved as structures. Depending on whether the threads where fitted or not, the saved outputs will either be threads or threads and threads_fit. The threads contains two extensions, threads.pos and threads.err_pos, which give the measured location of the peaks and the uncertainty on that position. Each array has the same time length as the original TD. The time frames where no peak is found contain a -1. The remaining either contain the position value or a 0. The 0 corresponds to time-frames that were jumped over at the crawling stage, i.e., no peak has been found between the time frames.

Note, it is up to the user to decide which found threads are worth using for further science.

If the user has used the internal fitting routine, then there will also be a structure called *threads_fit* in the idl save file. The saved data has three extensions, *threads_fit.pos*, *threads_fit.err_pos*, and *threads_fit.fit_result*. The first two extensions contain similar information to *threads* (although slightly modified - see wave fitting section for details). The extension *threads_fit.fit_result*, predictably, contains the details related to the fitting (again, see fitting section for more details).

3.3 Wave fitting

At present, the wave fitting part of the routine is quite rigid. The user has access to three functional forms:

$$f(t) = A_0 + A_1 \sin\left(\frac{2\pi t}{A_2} - A_3\right) + tA_4,\tag{1}$$

$$f(t) = A_0 + A_1 \sin\left(\frac{2\pi t}{A_2} - A_3\right) \exp(-A_5 t) + tA_4,\tag{2}$$

$$f(t) = A_0 + A_1 \sin\left(\frac{2.\pi t}{A_2(1 + A_6 t)} - A_3\right) \exp(-A_5 t) + tA_4.$$
(3)

The function Eq. 1 is the default fitting option, the others are selected by keywords when starting the routine via keywords, i.e.,

 $IDL > track_rout.data = td. damped$

or

 $IDL > track_rout, data = td, tdp$

However, this means that which ever function is chosen at the start of the routine is used for fitting all threads found! (This is obviously restrictive and we are looking to improve this aspect).

Once the fitting section of the routine has been entered, the routine performs a gap-filling. Any threads that have zero's in it's entries are filled. The default option for the filling is to give the blank entry the same value as found for the previous entry in the array and assigning it an error of ± 1 pixel. The large value of the error than essentially

gives the filled location less weight in the weighted lease squares. It is these filled arrays that are then saved in threads_fit.pos and threads_fit.err_pos. This fill assumes that the wave doesn't move more than 1 pixel between time-frames - how reasonable this is depends on what type of events you are measuring.

The routine will then show you which thread you are fitting, printing some text to the terminal and plotting the thread to screen. From here, there are a number of options for user interaction via the terminal. You will first be asked whether you would like to fit this thread. Answering n moves on to the next thread. Answering y, you will then be asked if you want to change the start and end positions of the thread. This allows you focus on fitting certain portions of the thread. Answering n moves on to the next step. Answering y shows you the current values of time-frames it is fitting between and ask you to enter new t and t1 values, e.g.,

```
Fit this thread? - y/n y
Change start end of thread? - y/n y
Current value of t=1 and t1=20
Enter t value: 2
Enter t1 value: 17
```

After this, you will then be prompted about whether you want to subtract a polynomial trend line. Answering n will skip this section. Selecting y will then prompt you to enter the degree of the polynomial you wish to remove. After entering a number, a polynomial is fit to the thread and removed, with the window updated with the new thread. The χ^2 value of the fit is printed to screen for reference.

```
Change polynomial degree? - y/n y
Fit trend - enter polynomial degree: 2
Chi^2 of fit 10.7489
Is the fit good? y/n
```

If you are happy with the subtracted polynomial then answer y. Answering n repeats the polynomial selection again. Note: the removal of a polynomial trend before the fitting of the sinusoid will impact upon the estimated uncertainties on the fitted parameters of the sinusoid, usually underestimating the uncertainties. In reality, the polynomial terms should be fit with the sinusoid, rather than individually.

The next section of the routine begins the fitting of the selected function (Eq. 1-3). The fitting is performed with a Levenberg-Marquadt algorithm (mpfit.pro - cite), which requires initial guesses for the parameters to be fit. The routine has a default parameter range set, but this will likely be unsuitable for most cases. The user will be asked whether the initial variables should be changed, selecting n tries the default initial values while selecting n lets you change the start values for amplitude and period (Note, in our experience we have found that the initial values for period and amplitude have the greatest impact on the fit, hence they are the only variables allowed to change.):

```
Initial variables: Constant=43.0000 Amplitude=1.00000 Period=20.0000 Phase=0.500000 Linear=1.00000 Change initial variables? y/n y

Enter amplitude: 2.

Enter period: 15.

%

Fitted variables 3.33458 68.1474 75.1892 -2.55123 4.85452

Error on fits 126.310 320.733 122.048 0.929392 17.6052

Chi^2 75.1915

Repeat fitting procedure? - y/n
```

The fit is overplotted in the window. Should you be unhappy with the fit, selecting y starts the process again from the selection of time-frames. This process can be repeated as many times as desired. Selecting n then ends the fitting of the current thread. You are then prompted whether to save the results of the fitting. Selecting y will record the fit parameters in $threads_fit.fit_result$, selecting n will not record the values. The routine then moves onto the next thread.

As mentioned, the results of the fit are stored in threads_fit.fit_result, with the following positions:

```
fit_result/0/=constant
fit_result/1 = amplitude
fit_result/2|=period
fit_result/3/=phase
fit_result/4/=linear coefficient
fit_result/5/=error on constant
fit_result/6/=error on amplitude
fit_result/7/=error on period
fit_result/8/=error on phase
fit_result/9/=error on linear coef
fit_result/10/=chi^2 for fit
fit_result[11]=start time from start of time-distance diagram
fit_result/12/=end time
fit_result[13]=pre-fitting used; (1)-non (2,3....)- polynomial degree
fit_result[14]=damping term (if /damped used)
fit_result/15]=error on damping term
fit_result[16]=time-dependent period term (if /tdp used)
fit_result[17]=error on time-dependent period term
```

4 To do

The following is a list of future upgrades that may be implemented. This list is not exclusive and will be updated. If you find bugs or have suggestions, they can be added to the list. Further, if anybody would like to work on any of these issues please feel free!! We will be running the code through Git, so version control will be in place.

- Update χ^2 cut-off for Gaussian fitting of cross-section At present, reduced χ^2 is used and set to a hard value of 5. This should be either set by a critical value for rejection. However, reduced χ^2 is not an appropriate measure for goodness of fit for non-linear functions (cite), so implementing a Kolmogorov-Smirnov test may provide greater confidence.
- Make the fitting section more flexible option for user to input own function for each fit.
- Make a GUI version!
- Fully automated routine removing the hand fitting section and replacing with something. Will likely form a separate routine to this one.
- Add consistent full Gaussian fit to cross-sections, returning intensity and structure width. This routine is almost complete but there are some issues regarding the rejected fits and how to provide a consistent value for intensity.
- Full documentation the future plan for the documentation is to include a discussion of the theory of least-squares fitting, this is being done by Krishna and should be available at the end of 2017.