SAFT - Semi-automatic fluorescence trace analysis

The purpose of SAFT is to automate some steps in analysis of recordings of synaptic activity, namely fluorescence time series corresponding to the emission of a neurotransmitter indicator. The steps that can be automated are background subtraction and peak finding. The user can tune both of these algorithms, and check the consequences interactively. The main workflow is directed towards recordings that come from different 'regions of interest' ('ROIs') over the same neuron that is stimulated in a regular fashion. The stimulus protocol is not known, but is inferred from the statistics of the mean responses. This makes analysis possible in a coordinated and efficient way over multiple conditions using the same parameters.

Installing

SAFT is written in Python, but basic Python is not enough. SAFT has quite a few dependencies, meaning external Python libraries that must be downloaded. These are for calculations (numpy and scipy), data management (pandas) and the GUI (PySide2 and pyqtgraph). You need the versions below at time of writing. Unfortunately, some unclear incompatibilities mean that some more modern versions of the relevant packages do not work.

Getting all these packages aligned is not easy. You can use anaconda, pip or compile from source. It is best to start with a fresh environment.

Conda currently provides Qt 5.9.7 and this makes it convenient to use conda to set up the package environment. To make getting the right environment in place easy, we include a conda environment .yml file in the distribution (SAFTenv.yml). To use this, you must have Anaconda or Miniconda. If you don't have Anaconda already, just get [Miniconda](# <u>https://docs.conda.io/projects/continuumio-conda/en/latest/user-guide/install/macos.html</u>).

To install the packages needed for SAFT into a fresh environment with Miniconda, go to the SAFT directory and enter the following in the Terminal,

conda env create --file SAFTenv.yml

This will create an environment called 'SAFT' with a working set of packages. As conda will tell you, to switch to this environment, enter the following,

conda activate SAFT

then navigate to the SAFT directory and execute the commands as above to launch SAFT.

If you happen to already have an environment named SAFT, you would need to remove it first:

conda env remove --name SAFT

Very important : Qt versions after 5.9 are not open source (breaking the conditions of our license), and coincidentally have serious bugs on macOS. We need Qt for the high-performance scientific plotting library pyqtgraph. PySide2 versions up to 5.15 seem to run happily with Qt 5.9.7 but it may be difficult to avoid a concurrent Qt update, unless you have a separate version of Qt compiled from open source and know how to point PySide2 at it. The latter solution is not stable.

Dependencies current as of 2020-11-12

```
Python 3.7.9
qt==5.9.7
PySide2>=5.9.0a1
numpy==1.19.2 scipy==1.5.2
pyqtgraph==0.11.0
pandas==0.24.0
matplotlib==3.3.2
xlrd==1.2.0
```

openpyxl==3.0.5
pyobjc-framework-Cocoa==6.2.2
python.app 2

To get SAFT as intended, launch it with the command from the directory where you unpacked it:

pythonw SAFT.py

After launching SAFT, you are confronted with the main window, 'Semi-automatic Fluorescence Trace analysis'. A few commands are in the menu bar, but the advanced controls are presented directly in the interface.



SAFT main window with data file loaded.



SAFT expects a set of time series as input. Current input format accepted is Excel. You can export from LibreOffice if you do not like Microsoft.

SAFT has four windows for analysing the peak responses.

- 1. Main window (SAFT.py) for loading trace data and identifying the position of peaks, removing baseline. This window also shows histograms of the peak data.
- 2. Peak extraction dialog (extractPeakResponses.py, opened by 'Extract peaks from all ROIs' button in the main window) for extracting peak data from known timepoints across a large number of ROI and sets. Automatic baseline correction is done. On completion and return to the main window, the peaks can be seen (as well as failures), edited and compared with the baseline subtracted traces.
- 3. Group peaks dialog (processGroupedPeaks.py called by the "*Extract group responses*" button in the main window) where you can get statistics for repetitive responses, for example, if you repetitively stimulate with trains, you can get the average and SD of the 2nd, 3rd, nth response in the train. Data, including automatically generated graphs, can be saved here. This kind of analysis is quite easy to do in Excel for example, but takes a lot of cutting and pasting, which is time consuming and leads to errors.
- 4. Histogram fitting (histogramFitDialog.py which is opened by the 'Launch Histogram Fit' button) in which you can fit histograms with binomial and poisson models of vesicle release, if they represent quantal responses. Global fitting of models across conditions can also be done. No allowance is made for reporter characteristics, such as non-linearity.

To get started, load the file SAFT_Data.xlsx using the File > Open File command. These traces were automatically extracted from movies in imageJ using the SNFR_ROI_Finder.ijm script (written by Benjamin König). This script is also in the repository.

Traces load and are automatically colour-coded. Three panels show the entire traces for each condition with the baseline correction plotted separately ('Traces and background subtraction', bottom left), a zoom into a chosen trace ('zoom', the scope is indicated by the blue translucent region in the bottom left panel) and histograms of automatically found peak responses (upper right).

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On the left hand side of the main window, a small graph shows all the traces overlaid and the automatically subtracted baselines. You can see here how "wobbly" the subtraction is. If you don't want to subtract the steadystate response of the indicator, it should be smooth.

There is also a panel that shows you the live histograms of the peak responses for each trace in the current ROI.



Traces and background subtract

In the centre, the zoom graph displays the trace (select which one in the "Peak finding and editing panel"). You can adjust the peak finding controls to collect the most peaks automatically - these are overlaid as square points on the trace, then switch to manual mode. A left click will add a peak, or remove it if you click on an existing one. You can look at any ROI but SAFT will use the "Mean" ROI for a reference (you can choose which condition in "Data display and processing").

When you are happy that all the peak locations are accurately specified, you can launch "Extract peaks from all ROIs".

On the right side of the main window, a set of panels gives you control over analysis and data display. These controls are divided into four sections:

'Data display and processing'

The dataset takes the name of the file, and you can select which ROI to view. Two automatically generated virtual ROIs, the mean and the variance are also viewable.

Here, a panel of some buttons give easy access to downstream analysis windows, saving data and you can choose which trace from the virtual ROI 'Mean' you want to use as the reference

'Automatic baseline cleanup'

Baseline removal can be switched on ('Auto'), off ('None') or locked with fixed parameters ('Lock'). Datasets generated following analysis will automatically turn off baseline removal. The baseline removal can be tuned with the p and lambda (length) scale controls. The lower the value of lambda, the shorter the region of a steady state response that will be removed. The baseline that is subtracted is drawn in the bottom left graph panel.

Savitsky-Golay smoothing is also implemented but did not yet work well in out hands.

'Peak finding and editing'

Select which condition to show in the zoom graph panel (here it's the '4 mM' condition). You can switch between auto peak finding, and manual editing of peaks with the mouse. If you switch to manual editing, a crosshair appears and you can click to mark peaks (squares). Click any square to remove it. If *Auto find peaks* is enabled, you

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Data display and processing	Data display and processing					
Load traces Dataset	SAFT_Data					
Select ROI	Mean					
• Combined traces	Split traces					
Reference condition for patt	ern extraction 4 mM ᅌ					
Extract peaks from all ROIs	Extract grouped responses					
Quantal Histogram Fit						
Automatic baseline cleanup						
Baseline removal? Auto						
Save baselined ROI traces	lambda					
Savitzky-Golay smoothing	Off \diamond Window 15 \diamondsuit					
Peak finding and editing						
Show 4 mM ᅌ	in zoom.					
Auto find peaks	 Edit peaks with mouse 					
Auto-find peaks with	wavelet 💠 algorithm.					
Prominence / SNR	1.3					
Width (wavelet only)	6					
Ignore peaks <	30 % 🗘 Of largest.					
Show current peak data	Save peak data					
Histogram options						
Show histograms Separated						
dF/F max 1	Fit Histograms					
No. of bins 100 🗘	No. of Gaussians 5					

SAFT main window control panel

can choose between a simple threshold method and the wavelet algorithms. Parameters like *Prominence* (for the simple algorithm) or *SNR* and *width* for the wavelet algorithm can be adjusted. You can optionally ignore any small peaks too.

There is a button ('Show current peak data') to give a quick popup of the underlying values.

'Histogram options'

There is an option to sum the histograms, which makes sense if the quantal size is consistent across conditions. The scale of the histogram display can be controlled here. There is also the option to fit the histograms with simple functions.

Extract Peaks according to a reference pattern

•	Extract peaks according to reference pattern				
	Extracting 46 peal over the sets nam				
	Skipping 0 traces out of 84 for low SNR.			3 0	
	Search range around peak (data points)			3	
	Treat peaks as failures when < SD x			1.5	
No peaks set to be failures.					
	Extract respon	nses	Cancel	Accept and Return	

The number of peaks in each trace, the number of traces (ROIs) and the theoretical maximum total of peaks to find over the entire set of conditions is shown. You can choose to skip individual traces on the basis of the signal to noise. You can adjust the range in each trace to search for a maximum around the peak location.

Extract peaks according to reference pattern					
Extracting 46 peaks from [28 28 28] ROIs (total 3864) over the sets named 0.5 mM, 2 mM, 4 mM					
Skipping 18 traces out of 84 for low SNR.	5 0				
Search range around peak (data points)	3				
\checkmark Treat peaks as failures when < SD x	1 0				
451 peaks set as failures (11.7% of total).					
Extract responses Cancel	Accept and Return				

Once you have clicked "Extract responses", you have the option to treat detected peaks as "failures", that is, set their amplitude to zero, when they are less than the SD x factor. The lower this value, the more peaks you will discard. The dialog tells you the number that were set to failures and the fraction.

When you click "Accept and Return", you go back to the main window and a new dataset corresponding to the detected peaks (with the baseline-subtracted traces) is created. Some controls are disabled for this dataset. You can use it for analysing the responses according to repetitive groups and the statistics of the peak amplitudes in "*Histogram Fitting with Quantal parameters*"

Extract grouped responses

-		
	Group analysis	
Number of respons	ses in group	5 0
46 peaks in each t	race and so 9 repeats.	
Grouping peaks fro over the condition	om combit 50601 dataset: [30, 30, 30] ROIs is 0.5 mM, 2 mM, 4 mM	
	Extract grouped data	
	Save grouped peak data	
Sa	ve graphs of grouped peaks	Done

If each trace corresponds to repetitive grouped responses, for example trains of 5 action potentials, you can analyse these groups and get the mean of the first response in the group, second response in the group etc as well as their standard deviation. This can be useful for comparing to synaptic transmission measured with electrophysiology.

You select how many responses are in each group and the dialog will then indicate how many repeats are found.

You can only save the results of this analysis directly from the "Group analysis" dialog

Histogram Fitting with Quantal parameters

One goal of optical reporting is to gain access to quantal parameters from a large number of synapses at once. To do this, the peak amplitude distributions can be fitted with different models of vesicle release.



The histograms of peak responses for one region of interest are shown. The X- and Y-axis scales are automatically the same for each. You can reset these with the mouse or the histogram controls. You can also choose to display the histograms summed into a single histogram of peaks or separated by condition.

Binomial statistics assume a finite (small) pool of independent vesicles, and from this, you can estimate the release probability (P_r). Poisson statistics assume an infinite pool, but in doing so, one less parameter is needed to describe the shape of the distribution. However, the "rate" of release that is obtained from Poisson fit is harder to interpret than the release probability.

SAFT also offers the opportunity to fit the summed histogram, to find the width (w) and quantal spacing (called 'dF(q)' in the interface). You can also fix the Gaussian width, w, to be the same as the SD for the baseline of the trace (peaks are automatically blanked for this calculation. For some as yet incomprehensible reason, using the SD for the baseline seems to give Gaussian functions that are wider than expected.

When you do a fit, you choose the number of Gaussian components - this is not optimised. If you choose a global fit, the quantal spacing and width is taken to be the same for each condition. The width can be optimised.

Fit results are autosaved to a file HFtemp.xlsx in the SAFT directory every time you change ROI. If you click "save", you get to pick a filename. Tick the box to save the fitted curves in the same excel file in a separate sheet.