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# ROSS: Offline Spike Sorter V1.0.0

## Overview

Spike sorting is the process of extracting the activity of the single neurons from the extracellular recording, which is a vital step for investigating brain behavior in neuroscience. ROSS is a fast and accurate offline spike sorting software developed based on the methods described in [1], and is made freely available.

The software gets the raw extracellular recording as input and all steps including filtering, alignment, detection and sorting could be done in a fully-automated manner. Several configuration options are provided in each of the mentioned phases, enabling user to change the parameters of the algorithms. Moreover, efficient manual tools are provided along with the useful visualization methods in order to fine-tune the sorting results by user interference.

## Installation

Recommended System:

- MATLAB > 2018a
- 12GB RAM

Download the package to a local folder. Run matlab and navigate to the folder. Then you could easily start Offline Spike Sorter by running "main.mlapp" or typing "main" in the command window.

# File Menu

# Data types Definition & Supported file formats

Software input and output data are in the form of ".mat" files, which might include one or more of the following variables, depending on the processing level of the data.

- a. SpikeMat:
  - An MxN matrix, where M is the number of waveforms (or spikes), and N is the number of time samples in each of the waveforms, i.e. each row contains a spike waveform.
- b. SpikeTime:
  - An Mx1 vector, where the m'th element is the corresponding time of the occurrence of the m'th spike in the SpiekMat.
- c. optimal set:
  - A structure, containing the soring results. It has two fields: cluster\_index and typicality. cluster\_index contains the index of corresponding cluster of each waveform. Typicality is a criterion, which denotes the sample quality of being representative of its assigned cluster.

## d. tag:

A Cx1 vector, where C is the number of clusters. Each element is a user-defined label for the corresponding cluster (e.g. multi-unit, single-unit ...).

## Open

#### a. Raw Data

Opens Raw Data, which is the basic data file accepted by the software. It is a Mat file that must contain a vector of the extracellular recording signal. None of the above-defined variables is needed in Raw Data. In a case there exist multiple variables in your selected Mat File, a window will open which asks you to select the desired variable.

#### b. Detected Data

Opens Detected Data, which is a Mat file containing two variables: I) SpikeMat, and II) SpikeTime.

#### c. Sorted Data

Opens Sorted Data, which is a Mat file containing three variables: I) SpikeMat, II) SpikeTime, III) optimal set.

## Save

#### a. Detection Result

Saves detection results including the spike waveforms and spiking times in a Mat file with two variables, named SpikeMat, and SpikeTime, respectively.

### b. Sorting Result

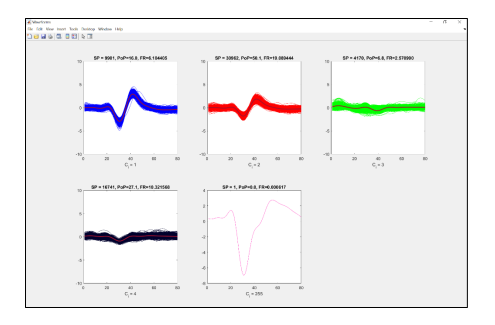
Saves sorting results including the spike waveforms, spiking times, sorting information and tags in a Mat file with four variables: SpikeMat, SpikeTime, optimal\_set, and tag.

# Visualization Menu

Visualization tools needed for manual analysis on spike detection and sorting results appear in this menu.

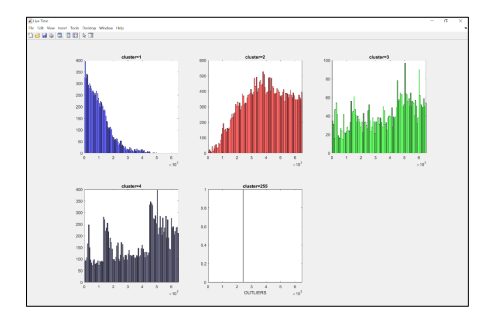
## **Plot**

When clicked, a window containing multiple time plots will open. Each plot represents waveforms assigned to one cluster versus time. In the bottom sample, the sorting result contains five clusters.



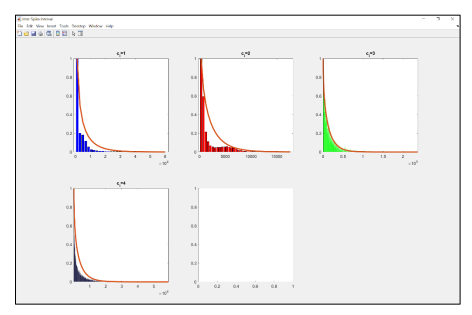
# Live Time

Clicking on Live Time will open a window containing multiple plots, each for one of the clusters. Each plot represents the histogram of the occurrence time of the spikes corresponding to that cluster. Live Time is useful for analysis on the firing rate of the neurons during the recording. A sample Live Time for five clusters is shown below.



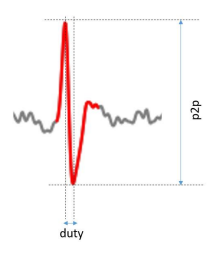
## ISI

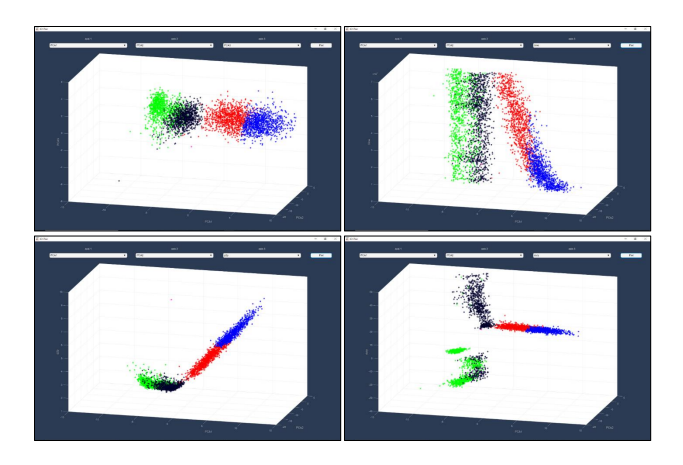
The distribution of inter-spike interval for each of the clusters will appear using this tool. Moreover a fitted gamma distribution is plotted for each cluster.



## 3D Plot

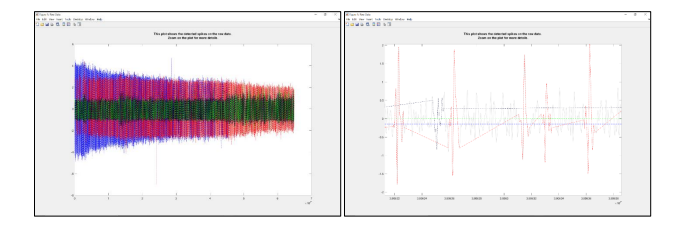
The tool provides different 3D plots to user. Each dimension of the 3D plots could be chosen arbitrary between the following statistics: "PCA1", "PCA2", "PCA3", "time" (time of the spike), "p2p" (amplitude distance between maximum and minimum of the spike) and "duty" (time distance between maximum and minimum of the spike). Sample 3D plots are shown below. User could also explore between different 2D views of the 3D plot by making the third axis "none".





# Raw Data

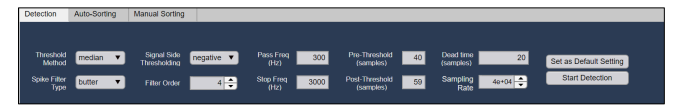
This option opens a window containing time plot of the raw data, where detected spikes could become visible by zooming.



# **Toolstrip**

## Detection

Detection tab in the toolstrip provides a fully automated and powerful spike detection tool described in [1]. The detection process is done in two steps: filtering, and thresholding. The following manual configurations are made available to user in the detection tab.



## **Filtering**

Filtering is the first step to separate spike activity from the LFP and background noise. A zero phase IIR filter is implemented by filtering the signal in forward and reverse orders in an offline mode. Configurable settings are as followed:

## Spike Filter Type

The filter type for zero phase filtering of raw data. Only "butterworth" type is supported in the current version.

#### Filter Order

The order of the IIR zero phase filter is selected using this option.

#### Pass Freq (Hz)

This option determines pass frequency of the band pass filter.

#### Stop Freq (Hz)

This option determines stop frequency of the band pass filter.

### Sampling Rate

The sampling rate of the input raw data is determined here.

## **Thresholding**

Spike detection is done by a thresholding method in this step. Settings of this section are:

## Threshold method

Threshold method must be chosen between three options: "median", "wavelet" or "plexon". This selection determines the method used to estimate the threshold. Each of the methods is described in the following.

Median: The standard deviation of noise  $(\sigma_n)$  is estimated according to Donoho's rule, then the spike detection threshold is set three times  $\sigma_n$ . The formula for "both" side threshoding would be:

$$\sigma_n = \frac{median(|r_F|)}{0.6745}$$
$$t_s = 3 \times \sigma_n$$

Where  $r_F$  denotes filtered signal, and  $t_S$  is the calculated threshold.

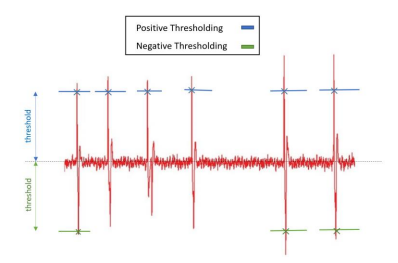
Wavelet: First the wavelet decomposition of unfiltered signal at level 2 using "db3" wavelet is calculated. Then the noise of wavelet coefficients are estimated. The detection threshold is defined as five times the estimated noise. "wavedec" and "wnoisest" matlab functions are used:

[c,l] = wavedec(Spike,2,'db3');
thr = 5\*wnoisest(c,l,2);

Plexon: First the samples with amplitudes higher than 2.7 times standard deviation of the full signal, are considered as outlier and would be ignored in threshold computation. Then the threshold is defined as four times the standard deviation of the filtered signal without considering the detected outliers.

### Signal Side Thresholding

This option is used to determine how the amplitude of the signal would be compared to the threshold. User could select "positive", "negative" or "two" amplitudes to be accounted for thresholding. Also by choosing "mean", an automatic decision between "positive" or "negative" options is made based on the sign of the signal mean.



## Pre-Threshold (samples)

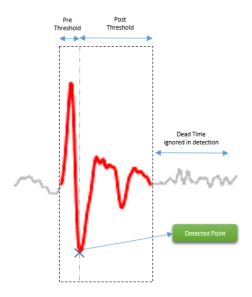
Number of samples before the detected (threshold crossing) point to be kept in the spike waveform.

## Post-Threshold (samples)

Number of samples after the detected (threshold crossing) point to be kept in the spike waveform.\(^{\)}

## Dead time (samples)

Number of samples after the detected spike waveform, which are ignored in detection.



## Set as default settings

Saves the current settings as the default.

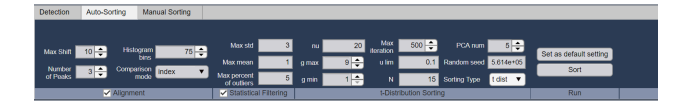
#### **Start Detection**

Starts the detection process.

## **Auto-Sorting**

Auto-Sorting tab provides useful tools for automatic spike sorting in four sections. Alignment tool is used to perform waveform alignment. Statistical Filtering is a noise removal process. The third section contains settings related to t-Distribution sorting or skew-t, and the last part includes Run commands.

<sup>&#</sup>x27;The total number of samples is "Pre-Threshold" + "Post-Threshold" + one.



## Alignment

Alignment is an effective step before clustering to improve results. Waveform alignment is implemented based on the method described in [1]. By checking "Alignment" the alignment method would run before the sorting process, otherwise, a simple alignment algorithm would be considered which aligns all spikes based on their minimum. Four settings are made available to user for configuration in the alignment algorithm:

#### Max Shift

Maximum number of points that the waveforms are allowed to be shifted in the alignment process.

## **Histogram Bins**

Number of histogram bins for plotting histogram of the waveform extremums.

#### Number of Peaks

Number of histogram peaks which are considered as aligning points.

#### Comparison Mode

Determines the mode for grouping the waveforms. If "magnitude" is selected, the waveforms are divided in two groups based on the amplitude of their extremums. (Samples with higher amplitude in maximum than minimum, are assigned to one group, and the remaining ones to the second group). Else, if "index" is selected, the waveforms are divided based on the indices of their extremums. (Samples with their maximum index lower than their minimum index, are assigned to one group and the remaining ones form the second group).

## Statistical filtering

An automatic noise removal process is designed in [1] based on data statistics. Settings related to this phase lie in this section. Detected noise samples would be assigned to cluster number 255 in the sorting phase.

### Max std

Magnitude of the threshold used for waveform standard deviation. Waveforms with higher standard deviation values would be considered as noise.

#### Max mean

Magnitude of the threshold used for waveform mean. Waveforms with higher absolute mean values will be considered as noise.

## Max percent of outliers

This parameter determines maximum percent of the waveforms allowed to be removed as outlier.

## Sorting

The clustering tool is implemented based on mixture of distributions modeling method, including t-distribution mixture modeling described in [2], and also skew-t mixture modeling presented in [1]. In addition, more three popular clustering methods are provided which are GMM, k-means and template matching. The provided manual configurations are as followed:

#### nu

This option is available in the case "t dist" or "skew-t dist" sorting type is selected, and determines initial value for nu ( $\nu$ ) parameter of t-distribution.

#### Max iterations

This option is available in all sorting types except "template matching". In the case "t dist" or "skew-t dist" sorting type is selected, this parameter determines maximum number of iterations for Expectation Maximization (EM) algorithm. For "GMM" sorting type, it indicates the maximum number of iterations for Gaussian mixture model fitting process. For "k-means" sorting type, it determines maximum number of iterations in k-means clustering algorithm.

### PCA num

Number of PCA components used as features for clustering. This option is available in all sorting types except "template matching".

## g max

Maximum number of clusters. This option is available in all sorting types except "template matching".

## g min

Minimum number of clusters. This option is available in all sorting types except "template matching".

#### u lim

This option is available in the case "t dist" sorting type is selected. "u lim" is the limit for determining waveforms typicality. The waveforms with typicality value lower than "u lim" will be accounted as not typical and assigned to cluster number 254.

#### error

This option is available in the case "skew-t dist" sorting type is selected. "error" is a threshold for defining another stop critera in addition to "Max iterations". The algorithm would stop in the case that maximum change among all parameters is smaller than "error".

#### tol

This option is available in the case "GMM" sorting type is selected. "tol" indicates the termination tolerance for the loglikelihood function value in fitting Gaussian mixture model to data.

#### Ν

Regularization parameter. This option is only available when "t dist" is selected as the sorting type.

#### Random seed

Random seed for MATLAB random generator engine.

## Matching mode

This option is available in the case "template matching" sorting type is selected and provides three options for template matching: "Euclidean", "Correlation" and "Chisquared".

- Euclidean: Euclidean distance between spikes and templates is employed for template matching.
- Correlation: Correlation between spikes and templates is employed for template matching.
- Chi-squared: In this method, first the subtraction of templates from spikes is calculated and considered as the residue. Then the distribution of the residue is compared to the distribution of raw data noise. This algorithm could not be performed when raw data does not exist in the system. Raw data is filtered by employing filtering parameters selected in Detection tab and the noise variance is estimated using the Donoho's rule referred to in Threshold method section. The noise in raw data is approximated as a Gaussian noise with mean of zero and the calculated variance value. Finally, the Chi-squared test is used to compare the residue distribution with the distribution of raw data noise with the given significance level (alpha). This test is performed for each spike in comparison to all templates. If the test is rejected (distributions are considered the same), that template is considered as a candidate template for the spike. Between all of the candidate templates, the one with the lowest residue variance is selected as the final cluster for that spike. In addition, in the case that no template is rejected in the Chi-squared test, again the template with the least amount of residue variance is selected.

### alpha

This option is available in the case "template matching" is selected as sorting type and "Chi-squared" is selected as matching mode. It determines significance level for Chi-squared test.

#### Combination

This option is available in the case "template matching" is selected as sorting type. By checking this checkbox, all two-member combinations of the existing templates will also be considered in the template matching process. The average of the two members is computed and considered as a new template.

## **Custom Templates**

This option is available in the case "template matching" is selected as sorting type. By pressing "Custom Templates" button a window will open asking user to select a ".mat" file containing custom templates. The selected file must include a variable named "templates" which is a matrix consisting of desired templates in each of the rows. After custom templates file is loaded successfully, a checkbox is enabled. Whenever the custom templates checkbox is checked, the user custom templates will be used in template matching, otherwise if there exist previous sorting results in the system, templates are extracted by computing the average of the existing clusters, else an initial GMM clustering is performed and then the templates are extracted from the created clusters.

## Sorting Type

There are five options for clustering method:

- 1. t dist: assumption of t distribution for clusters in the sorting algorithm. This method is implemented based on [2].
- 2. skew-t dist: assumption of skew-t distribution for clusters in the sorting algorithm. This method is implemented based on [1].
- 3. GMM: using Gaussian mixture models for data clustering. The optimal number of clusters is determined using the Calinski-Harabasz clustering evaluation criterion between "g min" and "g max".
- 4. K-means: using k-means for data clustering. The optimal number of clusters is determined using the Calinski-Harabasz clustering evaluation criterion between "g min" and "g max".
- 5. Template matching: three template matching methods are implemented which can be selected from Matching mode dropdown menu. There exist two modes for templates, either entered into the system by using Custom Templates option, or calculated by the software. In the second mode templates are extracted from the available clusters from a previous sorting process. If the sorting result does not exist in the system, an initial GMM sorting is performed and then the templates are extracted.

#### Run

### Set as default settings

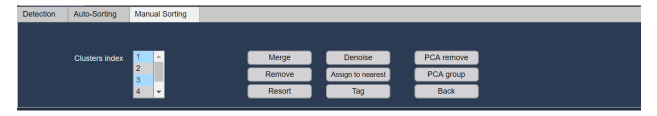
Saves the current configurations as default settings.

#### Run

Starts automatic sorting process.

## **Manual Sorting**

Manual Sorting tab provides several manual tools for modifying the auto-sorting results.



### **Clusters Index**

One or a number of clusters could be selected from this list for applying the manual changes.

## Merge

Using this button, the selected clusters in the "Clusters Index" list will be merged to form one cluster.

#### Remove

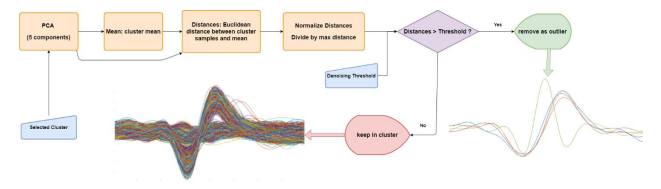
This option will remove the selected clusters in the "Clusters Index" list.

#### Resort

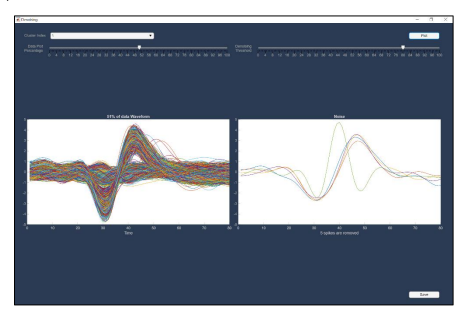
Performs auto-sorting only on the selected clusters in the "Clusters Index" list once again.

### Denoise

This option will open a window containing denoising tool. This tool removes the outliers by a thresholding method. The distance between each waveform and the average waveform of the cluster is normalized and compared to a threshold. The denoising algorithm is illustrated in the following figure.

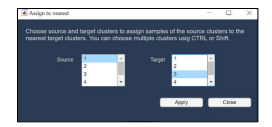


User can select the intended cluster from the "Cluster index" dropdown menu. The percentage of the random plotted data is set by changing "Data Plot Percentage", and the threshold for denoising algorithm is chosen in "Denoising Threshold". By clicking on "Plot", two figures will be plotted. The left figure displays random samples of the waveforms with the desired plot percentage. The right figure shows all of the detected noise waveforms. Clicking "save" will save the outliers under noise cluster (cluster index = 255).



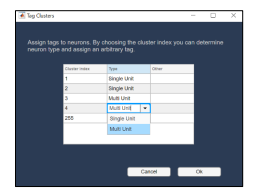
## Assign to nearest

This tool assigns all of the samples existing in the "source" cluster to the "target" clusters based on their distance from each of the targets.



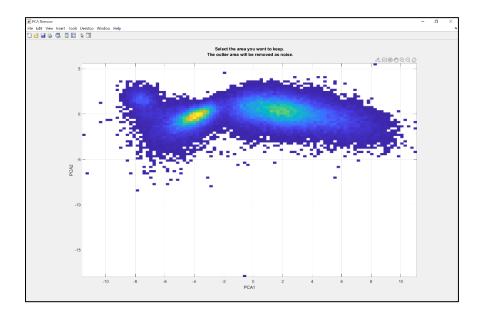
## Tag

User could define an arbitrary tag for each of the clusters, or choose the tag from the existing options. There exist a "Type" tag, including two options: "Single Unit" and "multi Unit". Also, there exist an "Other" tag, which could be any string.



## PCA remove

Samples could be removed manually from the feature space using this tool. A window containing 2D PCA plot of the samples is provided to user. User could manually select the desired region of the feature space and throw away the samples out of this region.



## PCA group

User could assign the desired regions in the feature space to a new cluster. A window similar to the previous figure is provided, and user could select the region of the new cluster.

#### Back

This button is used to undo the recent action. At most five last actions could be undone.

## **Batch Sorting**

Users could perform operations in batch mode using this tab. The desired process (detection, sorting or both) would be done on multiple files within a specified folder or determined in a file, and the results would be saved with the desired format.



### **Options**

This tab contains multiple settings for batch sorting.

#### Batch mode

Batch operation could be performed in two modes, which are selected by this option. In "path folder" mode, user selects a path containing the desired files (files can also exist in the sub-folders). In "batch file" mode, user imports a text file with a specific format that includes the required processes.

### **Process**

The process can be selected between "Detect", "Sort" or "Both" using this drop-down menu. This option is active when "path folder" is selected as the batch mode.

### Sub-folders level

The value of this option determines the level of the sub-folders to be considered for the batch process. This option is active when "path folder" is selected as the batch mode.

## **Browse folder**

Clicking on this button will open a window asking user to select a path. This option is active when "path folder" is selected as the batch mode.

## Browse file

Clicking on this button will open a window asking user to select a ".txt" batch file. This option is active when "batch file" is selected as the batch mode. The "batch file" must be in a format similar to the below example:

```
sort, path_to_file
detect, path_to_file, variable_name
both, path_to_file, variable_name
```

Each line of the text file specifies a process. In each line, the first word defines the type of process that could be one of these three options: "detect", "sort", or "both". If the process is "sort", that line must contain only two words, otherwise it must contain three words. The second word specifies the path of the input file. The third word determines the name of the variable in the raw data.

## **Regular Expression**

User can specify regular expression for names of the input files. This option is active when "path folder" is selected as the batch mode.

#### Variable Name

User can specify the variable name of the raw data matrix using this option. This option is active when "path folder" is selected as the batch mode.

#### Run

#### **Detect Save Format**

Saving file format of the output of the detection phase. The format can contain the input file name by using "%f" which will be replaced with the input name.

#### **Sort Save Format**

Saving file format of the output of the sorting phase. The format can contain the input file name by using "%f" which will be replaced with the input name.

#### Start

Starts the batch sorting process.

#### Stop

Stops the batch sorting process.

### Continue if Possible

This option can be used to continue a half-finished batch sorting process. By activating this checkbox, the desired operation is performed only for the input files for which the desired output is not already available.

### **Progress**

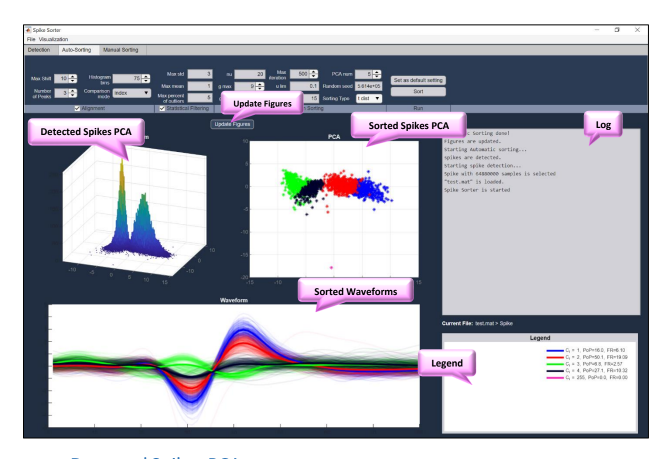
### **Processed**

The number of processed inputs while batch sorting appears here.

#### Left

The number of left inputs while batch sorting appears here.

## Main Window



**Detected Spikes PCA**: output of the detection phase, displaying histogram of 2D PCA of the detected spikes.

**Sorted Spikes PCA**: output of the auto-sorting phase, displaying PCA of the sorted spikes with different colors corresponding to different clusters. For each cluster, randomly selected samples are plotted.

**Sorted Waveforms**: output of the auto-sorting phase, displaying the waveforms of the sorted spikes with different colors for different clusters.

Log: keeps the history of the performed actions.

Legend: represents the legend for all visualizations.

Update figures: Updates all the plotted figures manually.

# Demo Video

Watch the demo video here.

# References

- [1] Toosi, Ramin, Mohammad Ali Akhaee, and Mohammadreza A. Dehaqani. "An Adaptive Detection for Automatic Spike Sorting Based on Mixture of Skew-t distributions." *bioRxiv* (2020).
- [2] Shoham, Shy, Matthew R. Fellows, and Richard A. Normann. "Robust, automatic spike sorting using mixtures of multivariate t-distributions." Journal of neuroscience methods 127.2 (2003): 111-122.