# Plotting and Analysis for Neural Database-Oriented Research Applications (PANDORA) Toolbox — User's and Programmer's Manual

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#### 1 Introduction

#### 1.1 What is the PANDORA Toolbox?

The PANDORA Toolbox is a software package which consists of a collection of MAT-LAB object-oriented classes and script functions for creating, analyzing and visualizing databases based on data from electrophysiological neuron simulations and recordings.

#### 1.2 Why did you make it?

Motivations to create this software were:

- Analyze data generated by brute-force and other parameter search methods.
- Analyze subsets of parameter spaces and special cases.
- Evaluate robustness of model neurons.
- Find functional roles of specific conductances.

#### 1.3 How is it implemented?

A custom database management system (DBMS) is written from scratch in the MAT-LAB language. The toolbox design follows object-oriented programming principles. It uses functions from the statistics and signal processing toolboxes of MATLAB, but they are not strictly necessary. It does not use MATLAB's database (DB) toolbox.

#### 1.4 How can I use it?

The PANDORA Toolbox uses an object-oriented approach to provide maximal flexibility for interactive use on the MATLAB command-line. Objects can be created, modified, analyzed, and visualized interactively in few steps. It is straightforward to save and load binary representations of these objects into files. Scripts can be made to programmatically repeat these procedures. Existing object classes are designed with the prospect of future extension, to accommodate new types of data and analyses.

#### 1.5 Who is it made for?

PANDORA Toolbox is customized for neuroscientific research. However, the concepts of a complex dataset, extraction of multiple observations from each item of the dataset, and analysis of multi-dimensional parameter spaces are universal. In its current form the database and dataset classes can be used for data other than electrophysiologic sources. As this toolbox is designed for flexible extensibility, one can add extensions that deal with different types of data and analyses.

<sup>&</sup>lt;sup>1</sup>At the time of initial design, the author did not have access to the DB toolbox. Future versions may support the DB toolbox.

<sup>&</sup>lt;sup>2</sup>This version of the toolbox does not yet have a general graphical user interface (GUI). The author prefers to have a flexible command-line interface than to maintain a limited GUI. However, once commonly used functions can be conveniently placed within a GUI, it will be added to the toolbox.

#### 1.6 Finding your way around

The source code uses MATLAB's documentation system, therefore all methods and classes are documented. To get help about all classes, issue the

>> help djlab

at the MATLAB prompt. This should give you an overview of available classes. Then, to learn about a specific class, ask for the documentation for the constructor method. For instance, for the trace class, issuing

>> help trace

gives you the documentation for the constructor together with an overview for the class. Sometimes, if there are multiple methods with the same name under different classes, you may get the wrong documentation. In that case, you can specify the class from which to take the method by prepending the class name to the method, such as in

>> help trace/spikes

In order to learn all methods available for a class, you can use MATLAB's methods command. For the trace class, do

>> methods(trace)

However, some documentation may be outdated or simply wrong. Please report these to the author via e-mail to cgunay@emory.edu.

#### 1.7 Overview of this document

Next, Section 2 guides the reader through the installation of the package and other dependencies. You can skip this section if you already have a running software environment. Section 3 introduces the essential components of the software and talks about their design decisions. You can also skip this part if you're not interested in the guts of the system and you are in favor of a quick start. The recipes in Section 5 provide a tutorial for some common tasks. It may be easier for some readers to follow these recipes to jump-start using the software. However, it is recommended that you familiarize yourself with the basic organization of the classes before proceeding into more complex tasks. Section 6 takes the tutorial approach to describe common visualization tasks. Finally, Section A points to the list of individual methods provided by the software. These methods are documented in detail using the MATLAB online help system.

# 2 Installation

Download the latest package file from:

http://userwww.service.emory.edu/~cgunay/pandora.

Unpack the archive anywhere in your system, using

\$ tar xzf pandora-xyz.tar.gz

and follow the instructions in the README file. Basically it involves pointing your MATLAB installation to look at the pandora/ subdirectory for loading the PANDORA files.

This can be achieved either by copying the pandora/directory under the toolbox/directory in your MATLAB installation, or by adding the path to your MATLAB session. While the former is a permanent solution, the latter method requires repeated application for each new session. To avoid repetition, you can have a startup script, startup.m, in the directory that you run MATLAB with the following commands:

```
%--- startup.m for matlab
addpath /my/download/directory/pandora-1.0b/pandora
%--- end startup.m
```

This will be loaded everytime you run MATLAB from this directory. in UN\*X systems, this can be improved further by placing the command in the file \$HOME/matlab/startup.m, which is executed no matter from where Matlab is called, especially if you are running Matlab from different or unknown places each time.

# Part I

# Software Architecture

# 3 Toolbox Components

An overview of the toolbox functionality is shown in Figure 1. In the figure, boxes represent objects that can be created with the toolbox. Flow starts from the dataset object on the top left which represents the collection of raw data files. The raw data is loaded using information in the dataset to create intermediate objects that, for instance, contain data traces. These objects define electrophysiological measurements to be entered into the data matrix of the database object on the top right. The database object allows filtering and querying to refine its contents. From the database object, one can always go back to the dataset and find the raw data that results from a query. The arrows going to bottom objects and corresponding plots show the types of possible analyses that can be done on a database object. These analyses are typically for displaying statistical information. The red arrow is a speacial analysis for searching and matching rows between different databases. The match is done by taking a row from a database created with data from real neurons and finding best matching model neurons from a simulation database.

The objects in the figure are instances of classes that define their properties in the object-oriented framework. Each class comes with a hierarchy of of subclasses that specialize to specific functions. Subsequent sections describe each of these class hierarchies that make up the main components of the toolbox.

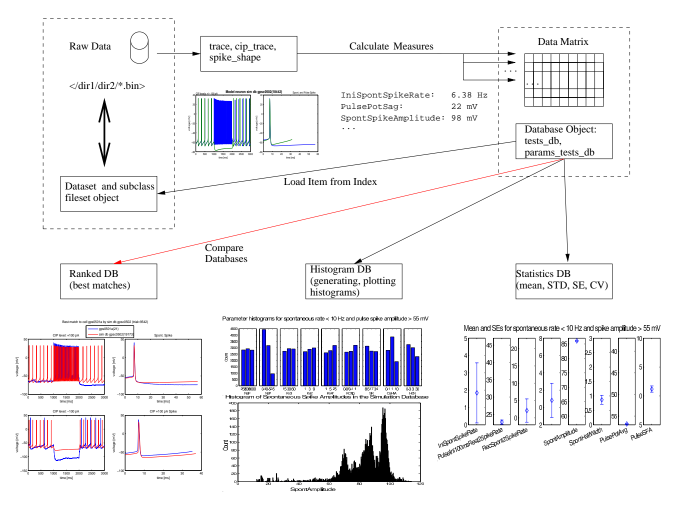


Figure 1: Schematic overview of the database software.

#### 3.1 Databases hold all the information

The database object is at the center of this toolbox (see Figure 1). It holds a data matrix with rows as observations and columns as attributes. The rows would normally correspond to results from individual data traces, or simply neurons. The columns hold values of separate measurements, statistical data, or parameter values.

A database object can be created from any of the classes in the hierarchy of Figure 2. The top-level database class is tests\_db which contains a two-dimensional data matrix of real numbers and some metadata. The metadata consists of column labels (e.g., measure names), a dataset label, and data properties (e.g., time resolution). The subclasses are specialized for different tasks.

If the database object is created using a dataset object, this maintains a connection from the elements of the database (e.g., neurons) to the raw data. This allows raw data associated with database contents to be visualized during analysis. However, a database can be created from any data matrix given in the proper format.

Some specialized subclasses of tests\_db are as follows:

- params\_tests\_db The first num\_params columns are reserved for parameters that were changed between different rows. It contains methods that treat these columns specially. Parameters can be simulation parameters, or pharmacological applications to experiments.
- tests\_3D\_db Contains a three-dimensional data matrix that has additional dimension for pages of information. This is mainly used to look at change in measurements with a parameter using the invarParam method of params\_tests\_db. Three dimensional databases can be useful for other purposes as well.
- **stats\_db** Contains few rows that describe the statistics obtained possibly from another database. It can contain the mean and standard deviation or error, or in some cases, the minimal and maximal values of columns in a database. It contains special plotting functions. There are methods that use the statistics collected by this class.
- ranked\_db Contains distances that resulted from a comparison of a database with a criterion. Its rows are ranked and sorted according to this distance value. Each row would point to a row in dex into the original database. Contains methods to generate reports from information about matching neurons.
- **spikes\_db** Contains results from individual spike shapes of a trace object. It can be obtained using the trace/analyzeSpikesInPeriod method.
- **histogram\_db** Each row corresponds to a histogram bin. Contains plotting methods.
- corrcoefs\_db Each row corresponds to a correlation coefficient. Contains plotting methods.
- **cluster db** Each row corresponds to a cluster centroid. Contains plotting methods.

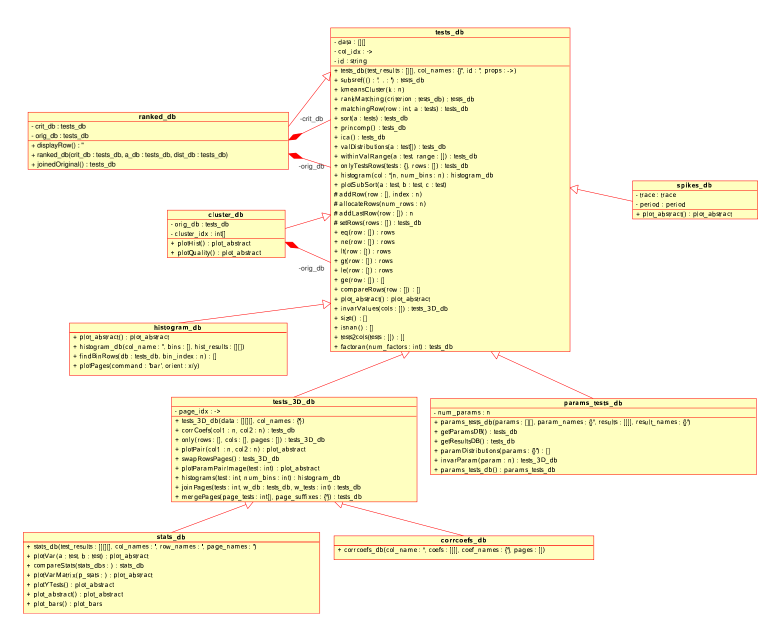


Figure 2: Database class hierarchy.

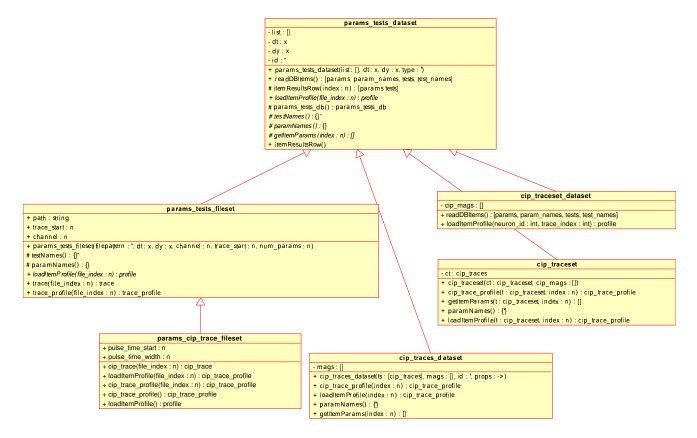


Figure 3: Dataset class hierarchy.

#### 3.2 Datasets create the databases

The dataset object is responsible for creating the database objects (see Figure 1). It defines where the raw data is stored and what parameters are used to load and analyze it. It knows that raw data has parameters associated which individual raw data traces and how and which measures will be generated. This information is used to automatically generate a database from the dataset. It also allows reaching back the raw data from rows of an analyzed database.

Figure 3 shows the hierarchy for the dataset classes. The top-level dataset class is params\_tests\_dataset which is an incomplete class. That is, this class defines general utilities that can work for a variety of dataset subclasses, but one cannot make a object from the params\_tests\_dataset class directly. Instead, one of its subclasses must be chosen and used. Some of these specialized subclasses are as follows:

params\_tests\_fileset This class assumes each raw data item resides in a file and all of these files are in the same directory. The parameter names and values are obtained from each file name itself. This class is mostly useful for simulation filesets.

- params\_cip\_trace\_fileset This is a subclass of params\_tests\_fileset,
   therefore it inherits the notion of one file per data item. The files must conform
   to the current-pulse injection experiments and have a starting time and duration
   for the pulses. The pulse magnitude is read from the pAcip parameter. This
   class is mostly useful for simulation filesets.
- **physiol\_cip\_traceset** This is a subclass of params\_tests\_dataset. It is designed to load a set of physiology traces from a single file generated by the PCDX stimulation and acquisition software.
- physiol\_cip\_traceset\_fileset This is a subclass of params\_tests\_dataset.
   It is designed to load traces from multiple PCDX data files. It uses the physiol\_cip\_traceset class for this purpose.
- cip\_traces\_dataset, cip\_traceset, cip\_traceset\_dataset These
   are obsolete classes that allow loading physiology traces from older MATLAB
   formatted objects.

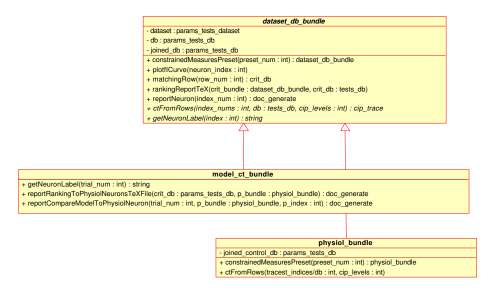


Figure 4: Bundle class hierarchy.

# 3.3 Bundling the database and dataset together

Since dataset and database objects are related and work together for some operations, it is convenient to have another object that bundles them together. There are several analysis routines that start from the database, retrieve raw data traces and other related information from the dataset and create a result. For instance, matching neurons from one database to another requires first comparing the measurements to find match candidates, and then comparing raw traces to visually represent the match quality.

The top-level dataset\_db\_bundle class in Figure 4 fulfills this purpose by bundling a dataset with the raw database, db, created from it, and with the reduced database, joined\_db, that contains a one-row-per-neuron representation. Although being a virtual class that cannot be instantiated, it contains general methods and prototype methods that must be implemented in subclasses. This way, it provides guidelines for defining subclasses. Its two subclasses provide specialize methods for model and physiology databases, respectively.

model\_ct\_bundle Contains methods to name and visualize neurons in the model database. It has methods to compare real neurons to model neurons to find best matching candidates.

physiol\_bundle Contains methods to name and visualize neurons in the physiology database. It contains a new attribute, <code>joined\_control\_db</code>, that holds only the neurons recorded without any pharmacological treatments.

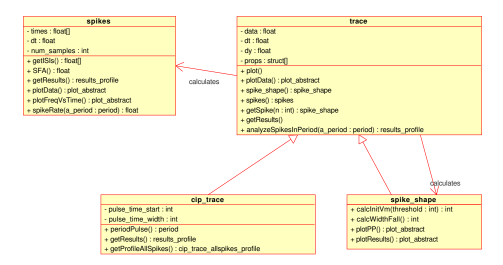


Figure 5: Data wrapper class hierarchy.

#### 3.4 Wrapper classes hold raw data

Wrapper classes are designed to hold data and provide simple methods that operate on them. They can either hold raw data, or intermediate processed forms of data being byproducts of analysis routines. In the overall schema of Figure 1, the raw traces obtained from the dataset object are kept in data wrapper objects.

Figure 5 shows the hierarchy for the data wrapper classes. The most basic data wrapper class in this toolbox is the trace class, which holds raw voltage or current traces. The spikes object contains the spike times obtained by analyzing a trace object.

A data wrapper class does more than just holding the data. It defines a set of operations in terms of method functions that can work on the data held by the class. As a rule of thumb, if one needs to add some new functionality into the toolbox, it should be added as a method into a class holding the data on which to operate.

Some of the data wrapper classes are as follows:

- **trace** Generic object that holds a vector of data that changes over time. It has a time resolution and y-axis resolution. Contains simple analysis routines such as finding average values within different periods, or finding spikes given a threshold.
- **cip\_trace** A subclass of trace class for current-injection recording protocols. It defines an initial spontaneous period, followed by a current-injection period, and final recovery period. It contains period-specific analyses that apply to the experimental protocol.
- **spike\_shape** A subclass of trace that holds the shape of a single spike. It contains spike shape measurements.

**spikes** A generic class to hold the event times for spikes. It contains methods for making measurements based on spike times, such as rate and ISI calculations.

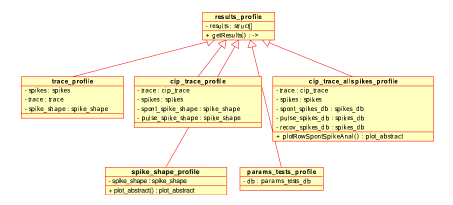


Figure 6: Profile class hierarchy.

#### 3.5 Profiles hold results of measurements

Profile classes are designed to hold results of analysis and measurements on the data wrapper or database objects. The data and results are separated into different classes for added flexibility of saving data and results separately. Yet, the profiles normally keep a copy of the data wrapper object from which they obtained the measurements. The intention is to save the measurement results for possible visualization or later inspection, without having to repeat the analyses.

In Figure 6, the top-level results\_profile class contains a simple MATLAB structure variable, results, that holds a set of name-value pairs. These are names of measurements and their corresponding values. Most of the subclasses are simplistic, and they exist only for organizational reasons. Some of them may implement specialized plotting methods that make use of the saved measurements. These subclasses can be briefly described as follows:

- **trace\_profile** Holds measurements from a trace object. It contains the trace object and the spikes found in it, and averaged spike\_shape object.
- cip\_trace\_profile Holds measurements from a cip\_trace object with a currentinjection period. It contains the original cip\_trace object and the spikes
  found in it. In addition, it holds averaged spike\_shape objects from the spontaneous and current-injection periods.
- cip\_trace\_allspikes\_profile Extended version of cip\_trace\_profile. Instead of single averaged spike shapes, it contains spike databases from the spontaneous, current-injection and recovery periods. These databases only retain measurements made from individual spikes, but not their shapes.
- spike\_shape\_profile Holds measurements made from a spike\_shape object.
- params\_tests\_profile Holds analysis results from a params\_tests\_db object.

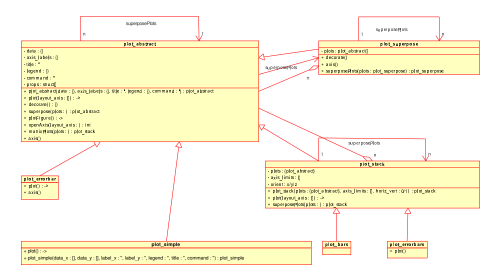


Figure 7: Plot classes hierarchy.

### 3.6 Integrated plotting for easy visualization

To integrate visualization into each class, common MATLAB plotting features are implemented in the supporting classes seen in Figure 7. These bring an object-oriented approach to plot generation in MATLAB. Plots can be generated as objects, saved, modified and included as subplots in larger plots.

The main plotting classes are plot\_abstract, plot\_superpose, and plot\_stack. The most general plotting template class, and the top-level class in the hierarchy, is plot\_abstract, which plots an axis using a single MATLAB command, like plot or bar. Multiple plot\_abstract objects that use the same command can be superposed and still act as a single plot\_abstract object. If they require different plotting commands (e.g., mixing plot and text labels), a plot\_superpose object must be used that is composed of an array of plot\_abstract objects. Multiple plot\_abstract objects or any of the subclass objects can be composed together in a horizontal or vertical stack using the plot\_stack class. Since plot\_stack is itself a subclass of plot\_abstract, it can be stacked as well. This allows creating virtually any complex structured figure using the three classes. Each of these classes have several properties that control the layout and details of placement and looks.

The rest of the classes in the hierarchy create typical types of plots for convenience:

**plot\_bars** Multi-axis bar plot with extending errorbars using a combination of the bar, errorbar, and text commands.

plot\_errorbar Single-axis errorbar plot using the errorbar command.

plot\_errorbars Multi-axis errorbar plot using the errorbar command.

plot\_simple Simplified single-axis, single command plot.

#### 3.7 Miscellaneous classes

These are miscellaneous classes that do not fit into any of the above categories:

- **period** Defines a period composed of a start and end time for operations on traces, etc.
- **script\_array** Defines a looping construct that can be extended. It defines an initialization routine, a job that needs to be repeated, and a finalization routine.
- script\_array\_for\_cluster Subclass of script\_array, it can submit the
  array job to run in parallel on a computing cluster that supports the Sun Grid
  Engine (SGE) commands.
- script\_factory Factory class to generate an enumerated array of script files to be
  distributed on several machines and run in parallel. It also defines a final function
  to gather results. It is recommended to use script\_array\_for\_cluster
  instead.

# 4 Programming Conventions

# 4.1 Using property structures for passing optional arguments to methods

For flexibility in passing optional arguments to methods, this toolbox adopted using property structures. A MATLAB structure, mostly called props, is passed to a method as the last argument:

```
>> props.optionalParam = 1
>> myFunc('hello', props)
```

Each method defines a list of accepted arguments that can be defined as fields in the structure, but should be able to execute without them by substituting defaults. Using a property structure is advantageous over using the varargin keyword for variable number of arguments, because properties allow adding and deleting arguments in methods without causing changes to the contents of the method. Since arguments are addressed by names rather than positional arguments, missing arguments do not affect the other arguments.

Most objects keep a property structure that define custom attributes passed at time of construction. These classes define a setProp method to modify properties after being created.

#### 4.2 Overloaded operators for transparent access to object contents

The simplistic implementation of object-oriented programming features in Matlab impose several strict limitations. MATLAB's powerful and flexible operator overloading feature helps overcome these limitations.

PANDORA Toolbox uses MATLAB operator overloading to facilitate manipulation of local and parent object fields. In MATLAB, object fields can only be accessed from the object's private methods. This means one cannot access the object fields using the dot operator. To give an example, the trace object has a dt field for time resolution. The following command fails:

```
>> mytrace.dt = 1e-4;
??? Object fields can only be accessed within methods.
```

Everytime object contents need to be addressed, a method must be called. The recommended way to do this is by defining separate getter/setter methods for each field of the object. For instance, writing getDt and setDt methods for accessing the dt field. This creates a lot of burden for the programmer not just creating a class, but also maintaining it later. Although this probably was intended for strictness in building object-oriented constructs, it is highly inconvenient for command-line manipulations. Therefore our toolbox objects offer generic get and set methods that can read or write the value of any of its fields:

```
>> mytrace = set(mytrace, 'dt', 1e-4)
>> get(mytrace, 'dt')
ans = 1e-04
```

These methods are almost identical across different classes. In addition to this, defining the special subsref method for objects allow overloading the dot (.), parenthesis (()), and curly brace ({}) operators. Most<sup>3</sup> of the objects in the toolbox allows using the dot operator to read or write to fields. Overloading these operators also help with the limitation of accessing parent object fields, a problem not found in other object-oriented languages such as JAVA. For example without any overloading, from the subclass class cip\_trace one needs to first address the parent class name, and then dt:

```
>> myciptrace.trace.dt
ans = 1e-4
```

After defining the overloaded operator that call parent methods, one get reach dt directly:

```
>> myciptrace.dt
ans = 1e-4
```

Some classes overload indexing operators to allow accessing special functions. For instance the main database class, tests\_db, overloads parenthesized indexing to access cells in the database matrix. Some classes define the special subsasgn method to overload the assignment operations when the object is on the left-hand-side of the operation. This allows the command:

```
>>  mytrace.dt = 1e-4;
```

which would otherwise need to be done the following way:

```
>> mytrace = set(mytrace, 'dt', 1e-4);
```

<sup>&</sup>lt;sup>3</sup>May not be implemented for all objects.

#### 4.3 Troubleshooting errors

For debugging problems with methods, one can turn on the verbosity of information display during execution with:

```
>> warning on verbose
>> warning on backtrace
```

#### 4.4 Creating a new class

To get the benefit of overloading, the top-level class must have the generic subsref and subsasgn methods. These methods can be copied from any of the other top-level classes. Any subclasses should have the generic get and set methods in place.

# Part II

# **User's Manual**

# 5 Recipes for Common Tasks

#### 5.1 Loading a database

A database can be created directly from a data matrix, or indirectly by loading a dataset. For the latter, first a dataset object must be created that point to the data sources. There are different dataset classes that allow using different data sources. For instance, physiology and simulation data require different operations. In physiology data, one can record information about the treatments and other conditions, whereas in simulations one can keep track of changing parameters.

Once the dataset object is obtained, the database object can be created with

```
>> mydb = params_tests_db(dataset)
```

which initiates the loading of files. This operation is the same no matter what type of dataset or fileset object is used. The following commands reduce the verbosity of output during this long process:

```
>> warning off verbose
>> warning off backtrace
>> warning off calcInitVm:info
```

#### 5.1.1 Creating a dataset for physiology data

Physiology data can be obtained from multiple sources.

**Loading data by specifying tracesets in a text file** The preferred way to load physiology traces is to first create a text file where each line specifies traces to load from a single data source (e.g., a PCDX file). The format of this text file is explained in the

PANDORA Toolbox: Prog.'s Mahraft v805 on 2007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

help of the physiol\_cip\_traceset\_fileset class. The physiology fileset can be created from the text file with a command such as follows:

This command reads the cell\_traces.txt file and records the tracesets to read from each file. The structure passed indicates to use the getProfileAllSpikes method to calculate the measurements on the traces.

The fileset can then be used to generate the database, as shown above, using its params\_tests\_db method. The fileset object holds within, a separate physiol\_cip\_traceset object for each line in the text file.

The cip\_list optional parameter must be used with caution. To determine actual CIP-levels, the current channel of the trace is analyzed. cip\_list entries are used to discretize the noisy current channel. Current levels will coerce to the nearest entry from cip\_list. In the above example, all current levels below -200 pA will be assumed to be -200 pA. The default cip\_list resides in the physiol\_cip\_traceset/CIPform method.

**Loading data from existing cip\_traces objects** The now obsolete cip\_traces Matlab objects have been used to hold some earlier physiological data. Each object holds a set of traces with varying CIP levels applied to the cell. The following command creates a dataset object from a cell array ct\_list of cip\_traces objects by choosing only the traces with 100 pA and -100 pA CIP levels

#### 5.1.2 Creating a dataset for simulation data

First a dataset or a fileset must be created. An example to load GENESIS .bin files would be

The explanation of arguments can be obtained by issuing a

PANDORA Toolbox: Prog.'s MaBuraft v805 on32007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

```
>> help params_cip_trace_fileset
```

in MATLAB. In this example, all GENESIS files were created with the same characteristics:  $\mathrm{d}t=10^{-4},\,\mathrm{d}V=10^{-3},\,\mathrm{pulse}$  during samples [20001, 30000]. Optional properties (the last argument) indicates that the first 10000 samples should be discarded and that the data should be prescaled to yield the  $\mathrm{d}V$  indicated. Note that, using an absolute path to refer to data files ensures that they can be reached from different directories after the fileset object is saved.

Loading heterogeneous set of simulation files 
Sometimes not all data files in a simulation set would have the same length, or CIP start time. The brute-force simulation set is such an example, where the spontaneous trace and different CIP level traces are in different files. I have a special superclass that contains multiple fileset objects to automatically handle this kind of data. It resides not in the general distribution directory, but in my personal directory /djlab/shared/matlab/classes/cengiz. This class is an example of how to create composite fileset objects. An instance of this class can be created with:

which will find all the files in the given directory and put them in separate pre-defined fileset objects according to their <code>\_pAcip\_</code> suffixes. Parameter range definition and value files are used to read parameter values for each simulation. A single database object can be loaded using the <code>params\_tests\_db</code> method on the <code>m\_filesetall</code> object above. The help on this method explains how to load only certain filesets at a time. This helps to load different filesets in parallel, since databases can be concatanated easily afterwards.

#### 5.2 Finding constrained subsets in a database

Once a database with more-than-sufficient number of measures is available, subsets of this database can be extracted easily for other tasks. New databases can be formed by filtering rows, columns or pages of an existing database. For choosing any of these dimensions, the user can specify an array of indices, or a logical array. For instance,

```
>> db2 = db1(1:10, :);
```

creates a database object db2 by the first ten rows of db1 and all its columns. For three-dimensional databases, a third parameter can be specified, as in

```
>> db2 = db1(1:10, :, [1 3]);
```

which will take only the first and third page from db1.

For measures, columns can also be specified as a single string value, or a cell array of strings, as in

```
>> db2 = db1(1:10, 'pAcip');
```

which chooses only the pAcip column of the first 10 rows of db1 or

PANDORA Toolbox: Prog.'s Mahraft v805 on32007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

```
>> db2 = db1(1:10, {'pAcip', 'IniSpontSpikeRate'});
```

which chooses two columns. Finally, composite queries can be formed when cell arrays are used for addressing:

```
>> db2 = db1(1:10, {1:10, 'IniSpontSpikeRate', 234});
```

which will select the first ten measures, the spontaneous spike rate, and the measure number 234.

Rows of the database signify neurons or simulation runs. Therefore it is important to find subset of neurons that match a certain criteria. This can be done specifying a list of rows that is the result of a logical operation. A logical operation finds rows that satisfy constraints on a single parameter or measure of a database. For instance,

```
>> rows = db1(:, 'IniSpontSpikeRate') > 10;
```

gives a logical array that contains a true value for all rows in db1 that has spontaneous firing faster than 10 Hz. If this is used as the row specifier in a subset operation, a new database with only these neurons can be obtained by

```
>> db2 = db1 (rows, :);
```

If we want more constrains it is straightforward to use any logical operation such as AND (&), OR (|) and NOT( $\sim$ ) on these logical arrays such as

```
>> db2 = db1((rows | rows2) & rows3, :);
```

which says choose all rows from db1 where either the tests rows and rows2 are satisfied and while rows3 is always satisfied. All these operations can be specified in-place such as in

which will create a database of neurons that spontaneously fire between 10-20 Hz.

#### 5.2.1 Complex Queries

Complex queries can be constructed using results of queries in nested fashion. The following example shows an example of finding all neurons that match any of the neurons in another database and then find the ones that match certain criteria:

```
>> sub_phys_es2 = phys_joined_db(phys_joined_db(:, 'NeuronId') == es2(:, 'NeuronId'), :);
>> displayRows(sub_phys_es2(sub_phys_es2(:, 'Apamin') > 0, 'NeuronId'))
```

Here, the first query returns all rows that match the NeuronIds from the es3 database. The inner term in the second query finds among these neurons the ones for which apamin blocker data is present. The final enclosing block uses these rows to get a subset of the phys\_joined\_db with only the NeuronId column. This type query has equivalent computational power to using nested SELECT statements in the SQL language.

#### 5.3 Sorting the database according to a measure

First, all rows where the desired measure value is NaN should be eliminated:

```
>> ampDecayTau_nonNaN_db =
    dball(~isnan(dball(:, 'PulseSpikeAmpDecayTau')), :)
```

This finds all rows in dball that the PulseSpikeAmpDecayTau measure is not NaN and creates a new DB object ampDecayTau\_nonNaN\_db, which includes these rows with all measures from dball. Notice that the newly created DB contains less rows than the original DB. The number of rows in the new DB can be obtained by just typing the name of the DB and pressing enter at the MATLAB prompt.

Then, one can sort the new database using:

This will create DB which is sorted with increasing values of the PulseSpikeAmpDecayTau measure. Displaying the first few rows gives lowest values:

```
>> displayRows(ampDecayTau_sorted_db, 1:3)
ans =
                                            1000] [ 250] [ 250]
0.5000] [ 0.5000] [ 2.5000]
60] [ 15] [ 30]
     'NaF'
     'NaP'
                                      Γ
     'Kv3'
                                     [
     'Kv2'
                                                   9] [
                                                                    3] [
                                                                                       31
                                     Γ
                                                   5] [ 1] [
     'Kv4f'
                                     ]
                                                                                      251
                                  [ 0.1000] [ 0.0100] [ 0.1000] [ 8.5000]
     'KCNQ'
     'SK'
                                            10] [ 0.1000] [
     'CaHVA'
                                   [
                                                                                     10]
                                   [
[
     'HCN'
                                                30] [ 0.3000]
                                                                           [
                                                                                      301
                                              100] [ 100]
     'pAcip'
                                                                           [
                                                                                     1001
     'pAcip' [ 100] [ 100] [ 100] 
'IniSpontISICV' [3.9448e-04] [ 0.0051] [ 0.0452] 
'IniSpontPotAvg' [ -64.9161] [-52.6859] [-41.5876] 
'IniSpontSpikeRate' [ 14.0014] [ 18.0018] [ 81.0081] 
'PulseISICV' [ 0.0226] [ 0] [ 0.0366]
                 SpikeRate' [ 14.0014] [ 18.0018] [ 81.0081] [ 0.0226] [ 0] [ 0.0366] [ 10.0081SICV' [ 0.0541] [ 0] [ 0.0814] [ 1827 char] [ 28.8953] [ 0] [ 88.9086] [ 1827 char] [ 26.6785] [ 0] [ 86.7052]
     'PulseIni100msISICV'
                                            30] [ 20] [
                  [1x22 char] [
                                                                                     1001
                 [1x25 char] [ 29.4118] [166.6667] [ 96.0512]
     'PulsePotAvg' [ -61.0044] [-32.7775] [-34.1518]
     'PulsePotMin'
                        [ NaN] [
[ NaN] [
[ 1.1254] [
                                   [ NaN] [ NaN] [ NaN]
     'PulsePotSag'
                                                                NaN]
                                                                           [
     'PulseSFA'
                                                                NaN] [ 1.3571]
     'PulseSpikeAmpDecayDelta' [ 4.2764] [ 9.2041] [ 17.8389] 
'PulseSpikeAmpDecayTau' [ 0.2000] [ 0.2000] [ 0.3000] 
'PulseSpikeRate' [ 28.0028] [ 2.0002] [ 89.0089]
```

Displaying the last few rows gives the highest values:

>> displayRows(ampDecayTau_s	or	ted_db, 1	L3879:	13881)		
ans =						
'NaF'	[	250]	[	250]	]	1000]
'NaP'	[	2.5000]	[	2.5000]	]	2.5000]
'Kv3'	[	15]	[	15]	]	60]
'Kv2'	[	3]	[	3]	]	9]
'Kv4f'	[	5]	[	5]	]	25]
'KCNQ'	[	0.1000]	[	0.1000]	]	0.0100]
'SK'	[	8.5000]	[	8.5000]	[	17]
'CaHVA'	[	10]	[	10]	[	10]
'HCN'	[	30]	[	3]	]	30]
'pAcip'	[	-100]	[	-100]	]	100]
'IniSpontISICV'	[	0.0027]	[	0.0027]	[ 9	.1376e-04]
'IniSpontPotAvg'	[	-28.5685]	[	-28.5687]	1	-67.7567]
'IniSpontSpikeRate'	[	69.0069]	[	69.0069]	[	14.0014]
'PulseISICV'	[	0.0046]	[	0.0046]	[	0.0091]
'PulseIni100msISICV'	[	0.0080]	[	0.0080]	1	0]
[1x27 char]	[	71.1269]	[	71.1269]	1	24.4499]
[1x27 char]	[	71.1427]	[	71.1427]	Ĺ	26.6785]
[1x22 char]	[	80]	[	80]	[	20]
[1x25 char]	[	70.6357]	[	70.6357]	1	25.4453]
'PulsePotAvg'	[	-30.1705]	[	-30.1718]	1	-65.2721]
'PulsePotMin'	[	NaN]	[	NaN]	1	NaN]
'PulsePotSag'	[	NaN]	[	NaN]	[	NaN]
'PulseSFA'	[	0.9792]	[	0.9792]	1	1.0407]
'PulseSpikeAmpDecayDelta	[	-1.2791]	[	-1.2868]	1	1.2201]
'PulseSpikeAmpDecayTau'	[	999.6000]	]	999.6000]	Ĺ	1000]
'PulseSpikeRate'	[	72.0072]	]	72.0072]	[	25.0025]

# 5.4 Preprocessing a raw (physiology) database by elimination and averaging

Mostly, raw physiology databases are subject to redundancies and unwanted recordings. We usually apply the following steps before we start analyzing a raw physiology database. Similar steps may be employed for simulation databases, too.

#### 5.4.1 Limiting range of bias currents

Recordings with high bias current are undesirable. We commonly filter-out high bias currents with:

which will limit the bias current,  $i_b$ , to  $-30 \,\mathrm{pA} < i_b < 30 \,\mathrm{pA}$ .

#### 5.4.2 Choosing few current levels

To get a profile for a neuron, usually both hyperpolarizing and depolarizing CIP-levels need to be included. Moreover, to capture the spiking frequency vs. current response of the neuron, multiple depolarizing CIP-level information may need to be included.

There are two counterparts to selecting which CIP-levels to include in a DB. First, one can select what CIP-levels are available in the raw data and what discretization levels should be used while loading the database. This is done with the <code>cip\_list</code> optional parameter described in Section 5.1.1. GP recordings prior to mid-2005 have current channel data which are too noisy to be quantized to levels of 10 pA. Instead, at least a step size of 20 pA needs to be used. Later recordings have both better recordings, and feature 20 pA steps in the experimental protocol anyway.

Second, after the database is loaded, one can filter-out unwanted CIP-level traces:

This operation can be simplified to take advantage of complex query form:

```
>> phys_dball_limitedcip =
    phys_dball_big(phys_dball_big(:, 'pAcip') == [-100; 0; 50; 100; 200])
```

which will choose rows with current levels matching any of the given values.

#### 5.4.3 Adding new columns calculated from existing measures

Some measures can be deduced from measures collected from raw data. These do not need to be calculated at time of loading the raw data, but rather can be added to the database later. Some measures must be added later because they may be composed of measurements from multiple traces or averages. Here is an example for adding a new measure:

# 5.4.4 Averaging multiple traces of same neuron with same CIP-level and (pharmacological) parameters

In making a one-row-per-CIP-level database, it is essential to include all available information from the raw database. Especially in physiology datasets, there may be multiple traces of a neuron where the same CIP-level and the same pharmacological conditions were applied. These rows can be averaged to obtain a single row for each CIP-level of a neuron.

Before doing this, the parameters of the raw database should only include parameters that uniquely distinguish neurons. The averaging operation tries to find each distinct set of parameters and then averages all rows that has this combination. For example, the NeuronId and pAcip parameters need to be distinct. However, the pAbias parameter does not need to be distinct for each neuron. The non-unique parameters need to be filtered-out before the averaging process. The following shows all the parameters of a raw physiology database:

```
>> phys_db
params_tests_db, tracesets from .../cip_traces_all_axoclamp.txt
ans =
   num_params: 16
        props: [0x0 struct]
     tests_db: [1x1 tests_db]
Optional properties of params_tests_db:
ans =
0x0 struct array with no fields.
tests_db, tracesets from .../cip_traces_all_axoclamp.txt
1527 rows in database with 182 columns, and 1 pages.
Column names:
          'pulseOn'
    [ 1]
          'pulseOff'
    [ 2]
          'traceEnd'
      3]
    Γ
          'pAcip'
      4]
    'pAbias'
      5]
           'Cadmium'
       6]
      7]
            'PicroTx'
            'Apamin'
      8]
      91
            'Glycine'
    [ 10]
            'KynAcid'
            'TTX'
    [ 11]
    [ 12]
            'XE991'
    [ 13]
            'drug_4AP'
            'EBIO'
    [ 14]
    [ 15]
            'NeuronId'
    [ 16]
            'TracesetIndex'
```

One can choose which parameters need to be distinct for each row by specifying as the second argument in the call to the meanDuplicateRows method, whereas the third argument specifies the measures to be averaged:

```
>> phys_mean_db = meanDuplicateRows(phys_db, [4 6:15], [17:161])
params_tests_db, averaged tracesets from .../cip_traces_all.txt
ans =
    num_params: 13
        props: [0x0 struct]
        tests_db: [1x1 tests_db]

Optional properties of params_tests_db:
ans =
0x0 struct array with no fields.
```

```
tests_db, averaged tracesets from .../cip_traces_all.txt
690 rows in database with 158 columns, and 1 pages.
Column names:
    [ 1]
             'pAcip'
    [ 2]
             'Cadmium'
    [ 3]
             'PicroTx'
    [ 4]
             'Apamin'
             'Glycine'
       51
             'KynAcid'
    Γ
      61
             'TTX'
       7]
    ſ
             'XE991'
       8]
             'drug_4AP'
      9]
    [ 10]
             'EBIO'
    [ 11]
             'NeuronId'
             'NumDuplicates'
    [ 12]
    [ 13]
             'RowIndex'
    [ 14]
             'IniSpontISICV'
```

This command ignores the pulse time information and the bias current, but includes all the pharmacological parameters, in distinguising the unique traces. It also eliminates some measures, such as the ItemIndex, from averaging.

# 5.5 Making a database by merging multiple rows from another database

A simple example for making a new database out of multiple rows in an existing database is combining multiple traces from the same neuron with different current pulse injection (CIP) levels. The initial database contains a row for each CIP level with redundant information, such as spontaneous period measurements.

#### 5.5.1 Making a one-row-per-neuron DB from multiple CIP-level rows

Measures of same cell obtained with multiple CIP-levels can be merged to make a single row. Note that, different pharmacological conditions applied to one cell must be kept in a different rows. The following command selects measures from each of the  $\{-100, 0, 40, 100, 200\}$  CIP levels to be included in the merged database:

This command operates on the previously averaged database, phys\_mean\_db, where each CIP level only occurs once for each distinct pharmacological setting for each neu-

ron. Note that, we filter-out columns 14 and 15 from the averaged DB while supplying the first argument to mergeMultipleCIPsInOne, which are artifacts of the averaging process and need not be included in the merged database. The second argument is a cell array of pairs of a suffix string and a corresponding list of measures for each of the CIP levels in phys\_mean\_db, in increasing order. The merged phys\_joined\_db looks like this:

```
params_tests_db, averaged tracesets from .../cip_traces_all_axoclamp.txt mult CIP
ans =
   num_params: 12
     tests_db: [1x1 tests_db]
tests_db, averaged tracesets from .../cip_traces_all_axoclamp.txt mult CIP
179 rows in database with 258 columns, and 1 pages.
Column names:
    [ 1]
[ 2]
            'Cadmium'
            'PicroTx'
    [ 3]
            'Apamin'
            'Glycine'
    [ 4]
    [ 5] 'KynAcid'
    [ 6] 'TTX'
          'XE991'
    [7]
          'drug 4AP'
    [ 8]
           'EBIO'
    [ 9]
            'Gabazine'
    [ 10]
            'NeuronId'
    [ 11]
            'TracesetIndex'
    [ 12]
           'PulseISICV_H100pA'
    [ 13]
            'PulseIni100msISICV_H100pA'
    [ 14]
    [ 15]
            'PulseIni100msRest1SpikeRate_H100pA'
            'PulseIni100msRest2SpikeRate_H100pA'
    [ 16]
    [ 17]
            'PulseIni100msSpikeRate_H100pA'
    [ 18] 'PulseIni100msSpikeRateISI_H100pA'
    [255]
            'PulseSpikeRiseTimeMean_D200pA'
    [256]
            'PulseSpikeRiseTimeMode_D200pA'
    [257]
            'PulseSpikeRiseTimeSTD_D200pA'
    [258]
            'PulseSpontAmpRatio_D200pA'
```

Note how each measure suffix indicate the CIP-level it belongs.

# 5.5.2 Making a one-row-per-neuron DB from dual CIP-level rows

The following statement uses the params\_tests\_db/getDualCIPdb method to merge rows of depolarizing and hyperpolarizing CIP-levels:

Here, the cell array variables depol\_tests and hyper\_tests hold the names of measures to be selected from depolarizing CIP and hyperpolarizing CIP, respectively.

The last two arguments define the suffixes to be applied to distinguish the measures from each CIP. The original DB is

```
>> dball
dball
params_tests_db, sim dataset gpsc0501
ans =
    num_params: 10
        props: [0x0 struct]
      tests_db: [1x1 tests_db]
Optional properties of params_tests_db:
ans =
0x0 struct array with no fields.
tests_db, sim dataset gpsc0501
39366 rows in database with 62 columns, and 1 pages.
Column names:
    [ 1]
           'NaF'
    [2]
           'NaP'
          'Kv3'
    [3]
           'Kv2'
    [ 4]
           'Kv4f'
    [5]
           'KCNQ'
    [6]
           'SK'
    [7]
    [8]
           'CaHVA'
           'HCN'
    [ 9]
    [10]
           'pAcip'
           'IniSpontISICV'
    [11]
           'IniSpontPotAvg'
    [12]
    [13]
           'IniSpontSpikeRate'
    [14]
           'PulseISICV'
    [15]
           'PulseIni100msISICV'
           'PulseIni100msRest1SpikeRate'
    [16]
            'PulseIni100msRest2SpikeRate'
    [17]
            'PulseIni100msSpikeRate'
    [18]
            'PulseIni100msSpikeRateISI'
    [19]
    [20]
            'PulsePotAvg'
    [21]
            'PulsePotMin'
            'PulsePotSag'
    [22]
    [23]
            'PulseSFA'
            'PulseSpikeAmpDecayDelta'
    [24]
    [25]
            'PulseSpikeAmpDecayTau'
    [26]
            'PulseSpikeRate'
    [27]
            'PulseSpikeRateISI'
    [28]
            'RecIniSpontPotRatio'
            'RecIniSpontRateRatio'
    [29]
    [30]
            'RecSpont1SpikeRate'
    [31]
            'RecSpont2SpikeRate'
            'RecSpontISICV'
    [32]
            'RecSpontPotAvg'
    [33]
    [34]
            'RecSpontSpikeRate'
```

```
[35]
            'SpontAmplitude'
            'SpontBaseWidth'
    [36]
    [37]
            'SpontDAHPMag'
    [38]
            'SpontFallTime'
    [39]
            'SpontHalfVm'
    [40]
            'SpontHalfWidth'
    [41]
            'SpontInitTime'
            'SpontInitVm'
    [42]
            'SpontMaxAHP'
    [43]
            'SpontMinTime'
    [44]
            'SpontMinVm'
    [45]
            'SpontPeakVm'
    [46]
            'SpontRiseTime'
    [47]
    [48]
            'PulseAmplitude'
    [49]
            'PulseBaseWidth'
    [50]
            'PulseDAHPMag'
    [51]
           'PulseFallTime'
    [52]
           'PulseHalfVm'
    [53]
           'PulseHalfWidth'
    [54]
           'PulseInitTime'
           'PulseInitVm'
    [55]
           'PulseMaxAHP'
    [56]
           'PulseMinTime'
    [57]
           'PulseMinVm'
    [58]
            'PulsePeakVm'
    [59]
            'PulseRiseTime'
    [60]
    [61]
            'PulseSpontAmpRatio'
    [62]
            'ItemIndex'
Optional properties of tests_db:
ans =
0x0 struct array with no fields.
           After merging, it becomes
>> sdball
sdball
params_tests_db, sim dataset gpsc0501 dual cip
ans =
    num_params: 9
       props: [0x0 struct]
      tests_db: [1x1 tests_db]
Optional properties of params_tests_db:
0x0 struct array with no fields.
tests_db, sim dataset gpsc0501 dual cip
19683 rows in database with 50 columns, and 1 pages.
Column names:
    [1]
           'NaF'
           'NaP'
    [2]
           'Kv3'
    [3]
           'Kv2'
    [4]
```

# 5 RECIPES FOR COMMON TASKS

```
[5]
            'Kv4f'
    [ 6]
            'KCNO'
    [7]
            'SK'
    [8]
            'CaHVA'
    [ 9]
            'HCN'
    [10]
            'RecIniSpontPotRatioHyp100pA'
    [11]
            'RecIniSpontRateRatioHyp100pA'
            'RecSpont1SpikeRateHyp100pA'
    [12]
            'RecSpont2SpikeRateHyp100pA'
    [13]
            'RecSpontISICVHyp100pA'
    [14]
            'RecSpontPotAvgHyp100pA'
    [15]
            'ItemIndexHyp100pA'
    [16]
            'IniSpontSpikeRate'
    [17]
    [18]
            'PulseIni100msSpikeRate'
            'PulseIni100msSpikeRateISI'
    [19]
    [20]
            'PulseIni100msISICV'
    [21]
            'PulseIni100msRest1SpikeRate'
    [22]
            'PulseIni100msRest2SpikeRate'
    [23]
            'RecSpont1SpikeRate'
    [24]
            'RecSpont2SpikeRate'
    [25]
            'RecIniSpontRateRatio'
            'IniSpontISICV'
    [26]
            'PulseISICV'
    [27]
            'RecSpontISICV'
    [28]
            'PulseSFA'
    [29]
    [30]
            'PulseSpikeAmpDecayTau'
    [31]
            'PulseSpikeAmpDecayDelta'
    [32]
            'IniSpontPotAvg'
    [33]
            'PulsePotAvg'
    [34]
            'RecSpontPotAvg'
    [35]
            'RecIniSpontPotRatio'
    [36]
            'SpontInitVm'
    [37]
            'SpontAmplitude'
    [38]
            'SpontMaxAHP'
            'SpontDAHPMag'
    [39]
            'SpontRiseTime'
    [40]
            'SpontFallTime'
    [41]
            'SpontHalfWidth'
    [42]
    [43]
            'PulseInitVm'
    [44]
            'PulseAmplitude'
    [45]
            'PulseMaxAHP'
    [46]
            'PulseDAHPMag'
    [47]
            'PulseRiseTime'
    [48]
            'PulseFallTime'
    [49]
            'PulseHalfWidth'
    [50]
            'ItemIndex'
Optional properties of tests_db:
0x0 struct array with no fields.
```

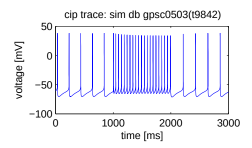


Figure 8: Example trace plot.

# 6 Visualization

By default, MATLAB prints figures in portrait orientation with a 8x6 aspect ratio, ignoring their size on the screen. The command;

>> orient tall

changes that behavior to print a full page in portrait orientation. The command;

>> orient landscape

does the same but rotates the figure to landscape orientation. If you want the printed figure to reflect its current screen size, issue the command;

```
>> set(figurenum, 'PaperPositionMode', 'auto')
```

Figures generated by the plotting system of the PANDORA Toolbox has a special resizing capability. Everytime the figure is resized, it will be drawn from scratch after calculating proper spacing between subplots according to font size. However, in some conditions this may cause other problems, such as crashing MATLAB in figure editing mode or causing loss of manual changes to the figure. To disable the auto-resize function, issue the following command after creating the figure;

```
>> set(figurenum, 'ResizeFcn', ")
```

Finally, to print the figure, consult your Matlab manual or issue a command such as;

```
>> print -depsc2 figurename.eps
```

See below for specific types of figures you can create.

# **6.1** Visualizing traces

The trace, and its subclasses such as cip\_trace, has the plot command overloaded to draw the raw trace in Figure 8:

```
>> a_ct = ctFromRows(mbundle, 9842, 100);
>> plot(a_ct)
```

Note that the plot in the figure has been created with a more precise control:

```
>> plotFigure(plotData(a_ct, ", struct('PaperPosition', [0 0 3 2])))
```

# **6.2** Displaying database contents

The displayRows method of tests\_db can be used to display rows of a database:

```
>> displayRows(sdball, 1:3)
ans =
   'NaF'
                       [
                            2501
                                  [
                                      250] [
                                                   10001
                       [ 2.5000] [ 2.5000] [ 15] [
                                                 2.5000]
   'NaP'
   'Kv3'
                                                  601
                                         3] [
   'Kv2'
                       [
                             31
                                [
                                                      91
                                     5] [
                             5] [
   'Kv4f'
                                                     251
                       Γ
                       [ 0.1000] [ 0.1000] [ 0.0100]
[ 8.5000] [ 8.5000] [ 17]
   'KCNO'
   'SK'
   'CaHVA'
                          10] [ 10] [
                                                    101
   'HCN'
                      [
                             30] [
                                        3] [
                                                     30]
                                                  100]
   'pAcip'
                      [ -100] [
                                      -100] [
   'IniSpontISICV' [ 0.0027] [ 0.0027] [9.1376e-04]
```

Note that, in the output, database rows appear as columns, and database columns appear as rows. See in above Section 5.3 for more example outputs from displayRows.

displayRows returns a cell array of column names justapoxed to a matrix of values. This cell array is intended for display on the screen and for generating reports. The displayRowsTeX method uses output from displayRows to generate a LATeX table that can be printed or converted to PDF:

With this the LATEX code to generate a table with the given caption is saved in a text file called fast\_spiking.tex. This file can then be included from a regular LATEX document to generate PDF output. See a LATEX manual on how to do that.

An alternative to displayRows is using the tests\_db/rows2Struct method:

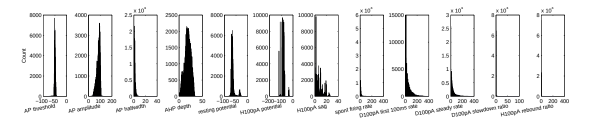


Figure 9: Example measure distribution plot.

```
IniSpontSpikeRate: 0
IniSpontSpikeRateISI: 0
PulseISICV: 0.0265
PulseIni100msISICV: 0.0584
PulseIni100msRest1SpikeRate: 40.0089
PulseIni100msRest2SpikeRate: 40.0178
PulseIni100msSpikeRate: 50
PulseIni100msSpikeRateISI: 43.5256
PulsePotAvg: -57.3737
PulsePotMin: NaN
PulsePotSag: NaN
PulseSFA: 1.1698
```

. . .

This method returns the database contents as a structure array. It is more natural for programming interfaces to use the database contents in a structure array than a cell array. The database columns become field names in the the structure. If multiple rows are requested, the displayed output would not contain the values. The desired row can be reached via indexing (e.g., s(1)). For instance, analysis in <code>cip\_trace/getProfileAllSpikes</code> method is done using this method for getting statistics from the <code>spikes\_db</code> databases.

### **6.3** Plotting all measure histograms

For plotting all measure histograms in a DB, the following method of tests\_db creates the horizontal stack plot in Figure 9:

# 6.4 Plotting all parameter histograms

For plotting all parameter histograms in a DB, the following method of params\_tests\_db creates the horizontal stack plot in Figure 10:

```
>> plotFigure(plotParamsHists(sdball));
```

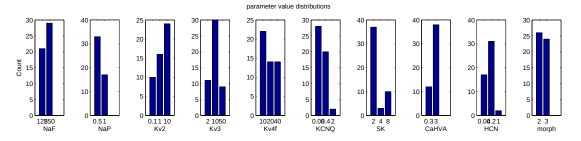


Figure 10: Example parameter distribution plot.

# 6.5 Plotting database statistics

The stats\_db object allows keeping statistical information obtained from a database. Statistics are calculated using one of the tests\_db converter methods, such as statsAll, statsMeanStd, etc.:

```
>> my_stats = statsMeanStd(my_db(:, {'IniSpontSpikeRate', 'PulseSpikeRate'}));
```

Then, the statistics can be plotted with diamonds indicating the mean and symmetric errorbars indicating upper and lower extensions (SE or Std):

```
>> plot(my_stats);
```

which is equivalent to:

>> plotFigure(plot\_abstract(my\_stats));

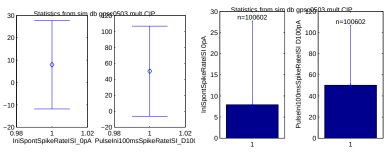
An alternative plotting form is using filled bars with extending errorbars:

```
>> plotFigure(plot_bars(my_stats));
```

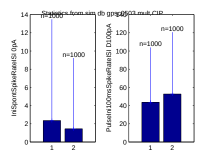
which are both seen in Figure 11 (a) and (b). The plot\_bars method is better suited for comparing statistics from multiple stats\_dbs coming from different sources. In this case, we take the first 1000 rows from the DB as a subset, and compare it with the statistics from the second 1000 rows:

The combined statistics object can then be fed into plot\_bars as seen in Figure 11 (c):

```
>> plotFigure(plot_bars(compared_two_subset_stats));
```



(a) Default visualization for stats\_db. (b) Optional plot\_bars visualization for stats\_db.



(c) Comparing multiple  ${\tt stats\_dbs}$  with  ${\tt plot\_bars}.$ 

Figure 11: Plotting statistics for two selected measures.

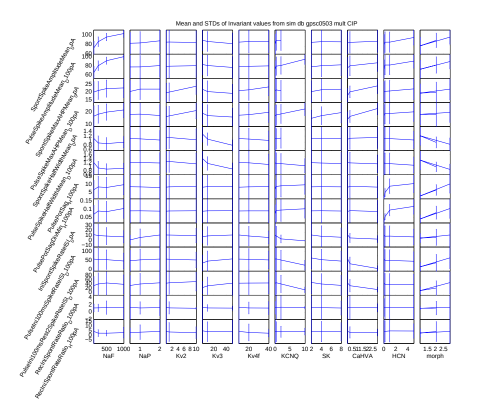


Figure 12: Example parameter-measure variation statistics plot.

# 6.6 Plotting all parameter-measure variations

To plot all parameter-measure variations, the plotVarMatix method of params\_tests\_db can be used (see Figure 12). plotVarMatix requires the DB object and the p\_stats array of stats\_db objects that hold the mean and standard error information (or possibly other statistics) for each of the possible parameter-measure combinations. The p\_stats array can be created using the paramsTestsHistsStats method, which in turn requires the p\_t3ds array of 3-dimensional databases each of which contain effects of a parameter invariant of other parameters. The p\_t3ds array can be created using the invarParams method. The sequence of commands<sup>4</sup> is then becomes:

```
>> p_t3ds = invarParams(noNaNRows(sdball))
>> [pt_hists, p_stats] = paramsTestsHistsStats(p_t3ds)
>> ap = plotVarMatrix(p_stats)
>> plotFigure(ap)
```

This will create a matrix plot with as many columns as parameters and as many rows as measures in the sdball object. It may be difficult to read if sdball contains large number of measures. One can divide the measures into two plots with the following sequence of commands

```
>> sdb1 = sdball(:, [1:9, 10:35])
>> sdb2 = sdball(:, [1:9, 36:49])
```

by choosing all parameters in both DBs, but only some measures for each. Then, the plots can be created for each DB by issuing

```
>> p1_t3ds = invarParams(sdb1)
>> p2_t3ds = invarParams(sdb2)
>> [pt1_hists, p1_stats] = paramsTestsHistsStats(p1_t3ds)
>> [pt2_hists, p2_stats] = paramsTestsHistsStats(p2_t3ds)
>> ap1 = plotVarMatrix(p1_stats)
>> plotFigure(ap1)
>> ap2 = plotVarMatrix(p2_stats)
>> plotFigure(ap2)
```

# 6.7 Insets

See Figure 13 which was created with the following set of commands.

```
>> im_p = plot_abstract({50 * rand(5)}, {}, ", {}, 'image');
>> plotFigure(plot_absolute([im_p, im_p], [0 0 1 1; 0.5 0.5 0.3 0.3]))
```

<sup>&</sup>lt;sup>4</sup>The noNaNRows function is required to filter out any rows containing NaN values in measurements before running statistic functions. Otherwise, the statistics functions (such as mean and std) will eliminate NaN values within each database column automatically, scrambling the row order and loosing the association with parameters.

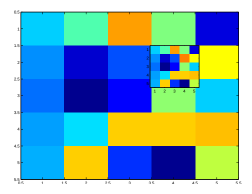


Figure 13: Creating insets in figures.

# 6.8 Generating a report comparing two databases

Most commonly a database of physiology neurons need to be compared to a database of a large body of simulation neurons and find best matches. One may need to see the match quality for a number of best matching candidates for each of the physiology neurons. Assuming multiple plots and tables are required to do a fair job of comparing a neuron to a thousands of simulation neurons, it becomes a difficult job to do this manually. An automatic report generation system has been built into the system for this purpose.

Currently a LaTeX document is created that needs to be included in either a proper LaTeX document, or included in a LayX document. The including document should provide the context in which the included part becomes meaningful. The report contains a set of tables and figures with proper captions. The table of contents, list of figures and list of tables facilities of LaTeX becomes useful to make the automatically generated document eaisly readable.

The report can be created with the command

where the simulation database sdb is searched for best matches to each row of physiology database phys\_sdb. The dataset for each is provided for the report to contain raw data associated with best matches. The obtained report contained in tex\_string can be saved as a ASCII LATEX file with

```
>> string2File(tex_string, 'myreport.tex');
```

<sup>&</sup>lt;sup>5</sup>This document is prepared using the LyX document preparation system [Ettrich et al., 2003] which uses the LaTeX  $2\varepsilon$  typesetting language [Lamport, 1994]. LyX is copyrighted by Matthias Ettrich and covered by the terms of the GNU General Public License (GPL), and LaTeX  $2\varepsilon$  is copyrighted by D. E. Knuth and the Free Software Foundation, Inc. and is covered by both the TeX copyright and the GNU GPL.

# References

Matthias Ettrich et al. LyX. A document preparation system that allows the author to concentrate on content, rather than typesetting., 2003. URL http://www.lyx.org.

Leslie Lamport. *ETeX: A document preparation system.* Addison-Wesley, Reading, Massachusetts, second edition, 1994. ISBN 0-201-52983-1.

# **Appendices**

# A Function Reference

See Section 1.6 on how to get online help about the software within MATLAB.

# A.1 Class chans\_db

# A.1.1 Constructor chans\_db/chans\_db

**Summary:** A database of channel activation and kinetics.

**Usage:** 

```
a_chans_db = chans_db(data, col_names, orig_db, crit_db, id, props)
```

**Description:** This is a subclass of tests\_db. Channel tables can be imported from Genesis using the utils/chanTables2DB script.

#### **Parameters:**

data: Database contents.

col names: The channel variable names.

channel\_info: Structure that holds scalar data elements such as Gbar.

id: An identifying string.

props: A structure with any optional properties.

### Returns a structure object with the following fields:

```
tests_db, channel_info, props.
```

See also: tests\_db (p. 217), chanTables2DB (p. ??)

# A.1.2 Method chans\_db/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.1.3 Method chans\_db/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

### A.1.4 Method chans\_db/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.1.5 Method chans\_db/plotInf

Summary: Plot the product of minf variables and the gmax of the given channel.

#### **Usage:**

```
a_plot = plotInf(a_chans_db, chan_name, gate_subnames)
```

# **Parameters:**

**a\_chans\_db:** A chans\_db describing channel variables.

**chan\_name:** Name of channel that make up the stem of variable names.

gate subnames: Gate names of the channel.

**props:** A structure with any optional properties.

(rest passed to plot\_abstract.)

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153)

# A.1.6 Method chans\_db/plotGateVars

**Summary:** Plot given channel gate variables of the same channel superposed.

# Usage:

```
a_plot = plotGateVars(a_chans_db, chan_name, gate_subnames)
```

#### **Parameters:**

```
a_chans_db: A chans_db describing channel variables.
```

**chan\_name:** Name of channel that make up the stem of variable names.

gate\_subnames: Gate names of the channel.

**props:** A structure with any optional properties.

**usePowers:** Use the gate powers, Luke. (rest passed to plot\_abstract.)

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2007/07/01

# A.1.7 Method chans\_db/plotAllInf

**Summary:** Plots the steady-state (infinity) response of all channels.

#### **Usage:**

```
a_plot = plotAllInf(a_chans_db, title_str, props)
```

### **Parameters:**

```
a_chans_db: a chans_db
```

title\_str: Plot title.

**props:** A structure with any optional properties.

(rest passed to matrixPlots.)

# **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153)

# A.1.8 Method chans\_db/plotAllVars

**Summary:** Plot all channel variables by grouping activation and time constant curves per channel.

#### **Usage:**

```
a_plot = plotAllVars(a_chans_db, title_str, props)
```

#### **Parameters:**

```
a_chans_db: A chans_db describing channel variables.
```

**id:** String that identify the source of the tables structure.

**props:** A structure with any optional properties.

(rest passed to plot\_abstract.)

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2007/03/05

# A.2 Class cip\_trace

### A.2.1 Constructor cip\_trace/cip\_trace

**Summary:** A trace with a current injection pulse (CIP).

#### Usage:

```
obj = cip_trace(datasrc, dt, dy, pulse_time_start, pulse_time_width, id, props)
```

#### **Parameters:**

datasrc: A vector of data points containing the spike shape.

dt: Time resolution [s].

**dy:** y-axis resolution [ISI (V, A, etc.)]

pulse\_time\_start, pulse\_time\_width: Start and width of the pulse
 [dt]

id: Identification string.

**props:** A structure with any optional properties, such as:

**trace\_time\_start:** Samples in the beginning to discard [dt] (see trace for more)

# Returns a structure object with the following fields:

trace, pulse\_time\_start, pulse\_time\_width, props.

PANDORA Toolbox: Prog.'s MaBuraft v805 on52007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

```
See also: trace (p. 255), spikes (p. 195), spike_shape (p. 183), period (p. 137)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

# A.2.2 Method cip\_trace/periodPulseIni50ms

Summary: Returns the first 50ms of the CIP period of cip\_trace, t.

Usage:

the\_period = periodPulseIni50ms(t)

#### Parameters:

t: A trace object.

#### **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

# A.2.3 Method cip\_trace/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.2.4 Method cip\_trace/periodRecSpontRestPeriod

Usage:

the\_period = periodRecSpont(t)

# **Parameters:**

t: A trace object.

**iniPeriod:** the time following pulse offset that is ignored. The rest of the time is kept

# **Returns:**

the\_period: A period object.

**See also:** period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>,Tom Sangrey 2006/01/26

# A.2.5 Method cip\_trace/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.2.6 Method cip\_trace/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.2.7 Method cip\_trace/plotData

**Summary:** Plots a trace by calling trace/plotData but also adds optionaldecorations.

Usage:

```
a_plot = plotData(t, title_str, props)
```

**Description:** If t is a vector of traces, returns a vector of plot objects.

#### **Parameters:**

t: A trace object.

title\_str: (Optional) String to append to plot title.

**props:** A structure with any optional properties.

**stimBar:** If true, put a bar indicating the CIP duration. (rest passed to trace/plotData)

### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/17

# A.2.8 Method cip\_trace/getBurstResults

Summary: Calculate test results related to Burst behavior.

Usage:

```
results = getRateResults(a_cip_trace, a_spikes)
```

#### **Parameters:**

PANDORA Toolbox: Prog.'s Maturaft v805 on 26007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

```
a_cip_trace: A cip_trace object.a_spikes: A spikes object.
```

#### **Returns:**

results: A structure associating test names with result values.

See also: cip\_trace (p. 54), spikes (p. 195), spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/30, Tom Sangrey

# A.2.9 Method cip\_trace/periodIniSpont

**Summary:** Returns the initial spontaneous activity period of cip\_trace, t.

Usage:

```
the_period = periodIniSpont(t)
```

### Parameters:

t: A trace object.

#### **Returns:**

the\_period: A period object.

**See also:** period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

### A.2.10 Method cip\_trace/spikes

**Summary:** Convert cip\_trace to spikes object for spike timing calculations.

**Usage:** 

```
obj = spikes(trace, plotit)
```

**Description:** Creates a spikes object by finding the spikes in the three separate periods, initial spontaneous activity period, CIP period, and final recovery period.

# **Parameters:**

trace: A trace object.

**plotit:** If non-zero, a plot is generated for showing spikes found (optional).

See also: spikes (p. 195), period (p. 137)

# A.2.11 Method cip\_trace/getCIPResults

Summary: Calculate test results about CIP protocol.

Usage:

```
results = getCIPResults(a_cip_trace, a_spikes)
```

#### **Parameters:**

```
a_cip_trace: A cip_trace object.
```

a\_spikes: A spikes object.

### **Returns:**

results: A structure associating test names with result values.

```
See also: cip_trace (p. 54), spikes (p. 195), spike_shape (p. 183)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/30

# A.2.12 Method cip\_trace/periodRecSpont1

**Summary:** Returns the first half of the recovery spontaneous activity period of cip\_trace, t.

### Usage:

```
the_period = periodRecSpont1(t)
```

# Parameters:

t: A trace object.

# **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

# A.2.13 Method cip\_trace/periodRecSpont2

**Summary:** Returns the second half of the recovery spontaneous activity period of cip\_trace, t.

### **Usage:**

```
the_period = periodRecSpont2(t)
```

### **Parameters:**

t: A trace object.

### **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

# A.2.14 Method cip\_trace/getProfileAllSpikes

**Summary:** Creates a cip\_trace\_allspikes\_profile object by collecting test results of a cip\_trace, analyzing each individual spike.

### Usage:

```
profile_obj = getProfileAllSpikes(a_cip_trace)
```

**Description:** Analyzes the spontaneous (periodIniSpont), pulse (periodPulse) and the recovery (periodRecSpont) periods separately and produces spike shape distribution results. Rate and CIP measurements are added to these.

### **Parameters:**

```
a_cip_trace: A cip_trace object.
```

# **Returns:**

profile\_obj: A cip\_trace\_allspikes\_profile object.

See also: cip\_trace (p. 54), cip\_trace\_allspikes\_profile (p. 68)

# A.2.15 Method cip\_trace/periodPulse

**Summary:** Returns the CIP period of cip\_trace, t.

Usage:

the\_period = periodPulse(t)

#### **Parameters:**

**t:** A trace object.

#### **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

# A.2.16 Method cip\_trace/calcRecSpontPotAvg

**Summary:** Calculates the average potential value of the recovery period of the cip\_trace, t.

# Usage:

avg\_val = calcRecSpontPotAvg(t)

# Parameters:

t: A cip\_trace object.

# **Returns:**

avg\_val: The avg value [dy].

See also: period (p. 137), trace (p. 255), trace/calcAvg (p. 259)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

# A.2.17 Method cip\_trace/periodPulseHalf1

Summary: Returns the first half of the CIP period of cip\_trace, t.

**Usage:** 

the\_period = periodPulseHalf1(t)

#### **Parameters:**

t: A trace object.

PANDORA Toolbox: Prog.'s MaDraft v805 or 2007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

#### **Returns:**

the\_period: A period object.

**See also:** period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

# A.2.18 Method cip\_trace/periodPulseIni50msRest1

**Summary:** Returns the first half of the rest after the first 50ms of the CIP period of cip\_trace, t.

# Usage:

the\_period = periodPulseIni50msRest1(t)

#### **Parameters:**

t: A trace object.

#### **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

# A.2.19 Method cip\_trace/periodPulseIni50msRest2

**Summary:** Returns the second half of the rest after the first 50ms of the CIP period of cip\_trace, t.

### **Usage:**

the\_period = periodPulseIni50msRest2(t)

#### **Parameters:**

t: A trace object.

#### **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

# A.2.20 Method cip\_trace/getPulseSpike

**Summary:** Convert a spike in the CIP period to a spike\_shape object.

### **Usage:**

```
obj = getPulseSpike(trace, spikes, spike_num, props)
```

**Description:** Creates a spike\_shape object from desired spike. Calls trace/getSpike method.

#### **Parameters:**

trace: A trace object.

spikes: (Optional) A spikes object obtained from trace,

calculated automatically if given as [].

spike\_num: The index of spike to extract.

**props:** A structure with any optional properties passed to getSpike.

# **Example:**

```
Get 2nd pulse spike and plot it:
» plotFigure(plotResults(getPulseSpike(t, [], 2)))
```

See also: spike\_shape (p. 183), trace/getSpike (p. 262)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/19

# A.2.21 Method cip\_trace/getRateResults

Summary: Calculate test results related to spike rate.

# Usage:

```
results = getRateResults(a_cip_trace, a_spikes)
```

#### **Parameters:**

```
a_cip_trace: A cip_trace object.
```

a\_spikes: A spikes object.

#### **Returns:**

results: A structure associating test names with result values.

```
See also: cip_trace (p. 54), spikes (p. 195), spike_shape (p. 183)
```

# A.2.22 Method cip\_trace/measureNames

**Summary:** Returns taxonomy of measurements collected by cip\_trace.

### **Usage:**

```
measures = measureNames(a_cip_trace)
```

**Description:** This is a static method, in the sense that it does not need the object passed as argument. Therefore it can be called directly by using the default constructor; e.g., measureNames(cip\_trace). The measure names are required for merging columns of a database generated by profiling these objects.

#### **Parameters:**

**a\_cip\_trace:** A cip\_trace object. It can be created by the the default constructor 'cip\_trace'.

#### **Returns:**

measures: A structure with cell arrays of types of measures, and measure names inside.

See also: getResults(p.??), getProfileAllSpikes(p.??), mergeMultipleCIPsInOne(p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

# A.2.23 Method cip\_trace/periodPulseIni100msRest1

### **Usage:**

```
the_period = periodPulseIni50msRest1(t)
```

# **Parameters:**

**t:** A trace object.

# **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

# A.2.24 Method cip\_trace/periodPulseIni100msRest2

#### Usage:

```
the_period = periodPulseIni50msRest2(t)
```

#### **Parameters:**

t: A trace object.

#### **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

### A.2.25 Method cip\_trace/periodRecSpontIniPeriod

# Usage:

```
the_period = periodRecSpont(t)
```

#### **Parameters:**

**t:** A trace object.

**iniPeriod:** the time following pulse offset that is kept, the rest of the time is ignored.

#### **Returns:**

the\_period: A period object.

**See also:** period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>,Tom Sangrey 2006/01/26

# A.2.26 Method cip\_trace/plot\_abstract

Summary: Plots a trace by calling plotData.

Usage:

```
a_plot = plot_abstract(t, title_str, props)
```

**Description:** If t is a vector of traces, returns a vector of plot objects.

# **Parameters:**

**t:** A trace object.

PANDORA Toolbox: Prog.'s MaDraft v805 or 62/007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/17

# A.2.27 Method cip\_trace/subsref

Summary: Defines generic indexing for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.2.28 Method cip\_trace/calcPulsePotAvg

**Summary:** Calculates the average potential value of the CIP period of the cip\_trace, t.

# **Usage:**

```
avg_val = calcPulsePotAvg(t)
```

#### **Parameters:**

t: A cip\_trace object.

### **Returns:**

```
avg_val: The avg value [dy].
```

See also: period (p. 137), trace (p. 255), trace/calcAvg (p. 259)

# A.2.29 Method cip\_trace/calcPulsePotSag

**Summary:** Calculates the minimal sag and sag amount of the CIP period of the cip\_trace, t.

# Usage:

```
[min_val, min_idx, sag_val] = calcPulsePotSag(t)
```

**Description:** The minimal sag is the minimal potential value of the first half of the CIP period. The sag amount is calculated by comparing this to the steady-state value at the end of the CIP period.

#### **Parameters:**

**t:** A cip\_trace object.

#### **Returns:**

min\_val: The min value [dy]. min\_idx: The index of the min value [dt]. sag\_val: The sag amount [dy].

See also: period (p. 137), trace (p. 255), trace/calcMin (p. 260)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

# A.2.30 Method cip\_trace/periodPulseIni100ms

Summary: Returns the first 100ms of the CIP period of cip\_trace, t.

#### **Usage:**

```
the_period = periodPulseIni100ms(t)
```

# **Parameters:**

t: A trace object.

#### **Returns:**

the\_period: A period object.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255)

# A.2.31 Method cip\_trace/periodRecSpont

**Summary:** Returns the recovery spontaneous activity period of cip\_trace, t.

Usage:

```
the_period = periodRecSpont(t)
```

#### **Parameters:**

t: A trace object.

### **Returns:**

the\_period: A period object.

**See also:** period (p. 137), cip\_trace (p. 54), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

### A.2.32 Method cip\_trace/getResults

**Summary:** Calculate test results given a\_spikes object.

Usage:

```
results = getResults(a_cip_trace, a_spikes)
```

#### **Parameters:**

a\_cip\_trace: A cip\_trace object.

a\_spikes: A spikes object.

# **Returns:**

results: A structure associating test names with result values.

See also: cip\_trace (p. 54), spikes (p. 195), spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.2.33 Method cip\_trace/getRecSpontSpike

**Summary:** Convert a spike in the CIP period to a spike\_shape object.

Usage:

```
obj = getRecSpontSpike(trace, spikes, spike_num, props)
```

**Description:** Creates a spike\_shape object from desired spike.

#### **Parameters:**

PANDORA Toolbox: Prog.'s MaDraft v805 or 627007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

trace: A trace object.

**spikes:** A spikes object on trace.

**spike\_num:** The index of spike to extract.

See also: spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/05/08

# A.3 Class cip\_trace\_allspikes\_profile

# A.3.1 Constructor cip\_trace\_allspikes\_profile/cip\_trace\_allspikes\_profile

**Summary:** Creates and collects test results of a cip\_trace.

**Usage:** 

obj = cip\_trace\_allspikes\_profile(a\_cip\_trace, a\_spikes, a\_spont\_spike\_shape,
results, id, props)

**Description:** This is a subclass of results\_profile. It is made to be used from subclass constructors.

#### **Parameters:**

a\_cip\_trace: A cip\_trace object.

a\_spikes: A spikes object.

spont\_spikes\_db, pulse\_spikes\_db, recov\_spikes\_db: tests\_dbs
with spontaneous, pulse and recovery period spike info.

results\_obj: A results\_profile object with test results.

id: Identification string.

**props:** A structure with any optional properties.

### Returns a structure object with the following fields:

trace, spikes, spont\_spikes\_db, pulse\_spikes\_db, recov\_spikes\_db, props

See also: cip\_trace (p. 54), spikes (p. 195), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/05/04

# A.3.2 Method cip\_trace\_allspikes\_profile/display

### A.3.3 Method cip\_trace\_allspikes\_profile/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.3.4 Method cip\_trace\_allspikes\_profile/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.3.5 Method cip\_trace\_allspikes\_profile/plotRowSpontSpikeAnal

**Summary:** Creates a row of plots that show spontaneous spikes, starting from the whole trace, zooming into the individual spike.

### **Usage:**

```
a_plot = plotRowSpontSpikeAnal(prof, title_str)
```

#### **Parameters:**

prof: A cip\_trace\_allspikes\_profile object.

title\_str: (Optional) String to append to plot title.

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), cip\_trace (p. 54), spike\_shape/plotCompareMethodsSimple (p. 189), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/05/23

# A.4 Class cip\_trace\_profile

# A.4.1 Constructor cip\_trace\_profile/cip\_trace\_profile

**Summary:** Creates and collects test results of a cip\_trace.

**Description:** The first usage is fully customizable to be used from subclass constructors. The second usage generates the spikes and spont\_spike\_shape objects, and collects some generic test results from them.

### **Parameters:**

data\_src: The trace column OR the filename.

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dt: Time resolution [s]

**dy:** y-axis resolution [ISI (V, A, etc.)]

pulse\_time\_start, pulse\_time\_width: Start and width of the pulse

[dt]

id: Identification string.props: See trace object.

# Returns a structure object with the following fields:

trace, spikes, spont\_spike\_shape, results, id, props.

See also: cip\_trace (p. 54), spikes (p. 195), spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

# A.4.2 Method cip\_trace\_profile/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.4.3 Method cip\_trace\_profile/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.4.4 Method cip\_trace\_profile/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.4.5 Method cip\_trace\_profile/subsref

**Summary:** Defines generic indexing for objects.

# A.4.6 Method cip\_trace\_profile/plot

**Summary:** Plots a cip\_trace\_profile object.

Usage:

h = plot(t)

Description: Plots contents of this object.

**Parameters:** 

t: A cip\_trace\_profile object.

**Returns:** 

h: Plot handle(s).

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/15

# A.5 Class cip\_traces\_dataset

# A.5.1 Constructor cip\_traces\_dataset/cip\_traces\_dataset

**Summary:** Dataset of cip\_traces objects, each with varying cip magnitudes.

**Usage:** 

obj = cip\_traces\_dataset(ts, cipmag, id, props)

**Description:** This is a subclass of params\_tests\_fileset.

**Parameters:** 

ts: A cell array of cip\_traces objects.

cipmag: A single cip magnitude to trace take from objects.

id: An identification string for the whole dataset.

**props:** A structure with any optional properties passed to cip\_trace\_profile.

# Returns a structure object with the following fields:

params\_tests\_dataset, cipmag, props (see above).

See also: cip\_traces (p.??), params\_tests\_fileset (p. 132), params\_tests\_db (p. 116)

# A.5.2 Method cip\_traces\_dataset/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.5.3 Method cip\_traces\_dataset/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.5.4 Method cip\_traces\_dataset/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.5.5 Method cip\_traces\_dataset/cip\_trace\_profile

Summary: Loads a raw cip\_trace\_profile given a index to this dataset.

Usage:

a\_cip\_trace\_profile = cip\_trace\_profile(dataset, index)

# Parameters:

dataset: A params\_tests\_dataset.

index: Index of file in dataset.

# **Returns:**

a\_cip\_trace\_profile: A cip\_trace\_profile object.

See also: cip\_trace\_profile (p. 69), params\_tests\_dataset (p. 111)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.5.6 Method cip\_traces\_dataset/subsref

Summary: Defines generic indexing for objects.

# A.5.7 Method cip\_traces\_dataset/paramNames

Summary: Returns the only parameter, 'pAcip,' for this fileset.

Usage:

```
param_names = paramNames(fileset)
```

**Description:** Looks at the filename of the first file to find the parameter names.

**Parameters:** 

```
fileset: A params_tests_fileset.
```

**Returns:** 

params\_names: Cell array with ordered parameter names.

```
See also: params_tests_fileset (p. 132), paramNames (p. ??), testNames (p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/06

# A.5.8 Method cip\_traces\_dataset/getItemParams

**Usage:** 

```
params_row = getParams(dataset, index)
```

# **Parameters:**

dataset: A params\_tests\_dataset.
index: Index of item in dataset.

## **Returns:**

params\_row: Parameter values in the same order of paramNames

```
See also: itemResultsRow (p.??), params_tests_dataset (p. 111), paramNames (p.??), testNames (p.??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/06

# A.5.9 Method cip\_traces\_dataset/loadItemProfile

**Summary:** Loads a profile object from a raw data item in the dataset.

# Usage:

```
a_profile = loadItemProfile(dataset, index)
```

**Description:** Subclasses should overload this function to load the specific profile object they desire. The profile class should define a getResults method which is used in the itemResultsRow method.

## **Parameters:**

```
dataset: A params_tests_dataset.
index: Index of item in dataset.
```

### **Returns:**

a\_profile: A profile object that implements the getResults method.

```
See also: itemResultsRow(p.??), params_tests_dataset(p.111), paramNames (p.??), testNames(p.??)
```

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.6 Class cip\_traceset

# A.6.1 Constructor cip\_traceset/cip\_traceset

**Summary:** A traceset with varying cip magnitudes from a single cip\_traces object.

## Usage:

```
obj = cip_traceset(ct, cip_mags, dy, props)
```

**Description:** This is a subclass of params\_tests\_fileset. This traceset can create a mini-database form a single cip\_traces object. The list contains cip\_mags. cip\_traceset\_dataset should be used to load multiple cip\_traceset objects.

### Parameters:

```
ct: A cip_traces object.
```

cip\_mags: An array of cip magnitudes to select from the object.

**dy:** y-axis resolution, [V] or [A] (default=1e-3).

**props:** A structure with any optional properties.

**offsetPotential:** Add this to physiology trace as compensation.

# Returns a structure object with the following fields:

```
params_tests_dataset, ct, props (see above).
```

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```
See also: cip_traces (p.??), params_tests_fileset (p. 132), params_tests_db (p. 116)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/30

# A.6.2 Method cip\_traceset/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.6.3 Method cip\_traceset/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.6.4 Method cip\_traceset/cip\_trace\_profile

**Summary:** Loads a raw cip\_trace\_profile given an index in this traceset.

Usage:

```
a_cip_trace_profile = cip_trace_profile(traceset, index)
```

### **Parameters:**

traceset: A cip\_traceset.
index: Index of item in traceset.

## **Returns:**

a\_cip\_trace\_profile: A cip\_trace\_profile object.

See also: cip\_trace\_profile (p. 69), params\_tests\_dataset (p. 111)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.6.5 Method cip\_traceset/paramNames

**Summary:** Returns the only parameter, 'pAcip,' for this traceset.

**Usage:** 

```
param_names = paramNames(traceset)
```

**Description:** Looks at the filename of the first file to find the parameter names.

**Parameters:** 

PANDORA Toolbox: Prog.'s MaDraft v805 on 75007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

traceset: A cip\_traceset.

## **Returns:**

params\_names: Cell array with ordered parameter names.

See also: params\_tests\_dataset (p. 111), paramNames (p. ??), testNames (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/06

# A.6.6 Method cip\_traceset/getItemParams

# **Usage:**

```
params_row = getParams(dataset, index)
```

## **Parameters:**

dataset: A params\_tests\_dataset.
index: Index of item in dataset.

**a\_profile:** A profile object for the item (optional).

## **Returns:**

params\_row: Parameter values in the same order of paramNames

See also: itemResultsRow (p.??), params\_tests\_dataset (p. 111), paramNames (p.??), testNames (p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/06

## A.6.7 Method cip\_traceset/loadItemProfile

**Summary:** Loads a profile object from a raw data item in the dataset.

## Usage:

```
a_profile = loadItemProfile(dataset, index)
```

**Description:** Subclasses should overload this function to load the specific profile object they desire. The profile class should define a getResults method which is used in the itemResultsRow method.

# **Parameters:**

dataset: A params\_tests\_dataset.
index: Index of item in dataset.

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## **Returns:**

a\_profile: A profile object that implements the getResults method.

See also: itemResultsRow(p.??), params\_tests\_dataset(p.111), paramNames (p.??), testNames(p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.7 Class cip\_traceset\_dataset

# A.7.1 Constructor cip\_traceset\_dataset/cip\_traceset\_dataset

**Summary:** Dataset of multiple cip magnitudes from cip\_traces objects .

# Usage:

```
obj = cip_traceset_dataset(cts, cip_mags, dy, id, props)
```

**Description:** This is a subclass of params\_tests\_dataset. Designed to extract a trace for each cip magnitude from the cip\_traceset objects contained. Uses cip\_traceset objects to extract multiple traces from each cip\_traces object.

#### **Parameters:**

cts: Array or cell array of cip\_traces objects.

cip\_mags: An array of cip magnitudes to select from each cip\_traces object.

**dy:** y-axis resolution, [V] or [A] (default = 1e-3).

id: An identification string.

**props:** A structure with any optional properties passed to cip\_traceset.

## Returns a structure object with the following fields:

```
params_tests_dataset, cip_mags
```

See also: physiol\_cip\_traceset (p. 144), params\_tests\_dataset (p. 111), params\_tests\_db (p. 116)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/01/28

# A.7.2 Method cip\_traceset\_dataset/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.7.3 Method cip\_traceset\_dataset/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.7.4 Method cip\_traceset\_dataset/loadItemProfile

**Summary:** Loads a cip\_trace\_profile object from a raw data file in the fileset.

**Usage:** 

a\_profile = loadItemProfile(fileset, neuron\_id, trace\_index)

#### **Parameters:**

**fileset:** A physiol\_cip\_traceset object.

neuron\_id : tells which item in fileset (corresponds to cells\_filename) to use
grab the cell information

trace index: Index of file in traceset.

## **Returns:**

a\_profile: A profile object that implements the getResults method.

See also: itemResultsRow (p.??), params\_tests\_fileset (p. 132), paramNames (p.??), testNames (p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14 and Tom Sangrey

# A.7.5 Method cip\_traceset\_dataset/readDBItems

**Summary:** Reads all items to generate a params\_tests\_db object.

**Usage:** 

```
[params, param_names, tests, test_names] = readDBItems(obj)
```

**Description:** This is a specific method to convert from cip\_traceset\_dataset to a params\_tests\_db, or a subclass. Output of this function can be directly fed to the constructor of a params\_tests\_db or a subclass.

## **Parameters:**

**obj:** A physiol\_cip\_traceset\_fileset

## **Returns:**

params, param\_names, tests, test\_names: See params\_tests\_db.

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```
See also: params_tests_db(p. 116), params_tests_fileset(p. 132), itemResultsRow testNames(p. ??), paramNames(p. ??), physiol_cip_traceset_fileset(p. 149)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/01/28

# A.8 Class cluster\_db

## A.8.1 Constructor cluster\_db/cluster\_db

**Summary:** A database of cluster centroids generated by a clustering algorithm from a rows of orig\_db.

## **Usage:**

```
a_cluster_db = cluster_db(data, col_names, orig_db, cluster_idx, id, props)
```

**Description:** This is a subclass of tests\_db. Use one of the clustering methods in tests\_db, such as kmeansCluster, to get an instance of this class.

## **Parameters:**

data: Database contents.

col\_names: The column names.

orig\_db: DB whose rows are clustered.

**cluster\_idx:** Array of cluster numbers that correspond to each row in orig\_db.

**id:** An identifying string.

**props:** A structure with any optional properties.

**sumDistances:** Total distance of elements within each cluster.

distanceMeasure: Measure used to find clusters (Default='correlation')

## Returns a structure object with the following fields:

tests\_db, orig\_db: original DB from which clusters were obtained, cluster\_idx: Array associating rows of orig\_db to each cluster here. props.

See also: tests\_db (p. 217), tests\_db/kmeansCluster (p. 228)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/08

\_\_\_\_\_

# A.8.2 Method cluster\_db/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.8.3 Method cluster\_db/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

## A.8.4 Method cluster\_db/plotHist

**Summary:** Creates a histogram plot showing the clustering memberships.

Usage:

```
a_plot = plotHist(a_cluster_db, title_str)
```

# Parameters:

```
a_cluster_db: A cluster_db object.
```

title\_str: (Optional) String to append to plot title.

## **Returns:**

a\_plot: A plot\_abstract object that can be plotted.

```
See also: plot_abstract (p. 153), plotFigure (p. ??), histogram_db (p. 94), histogram_db/plot_abstract (p. 96)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/08

## A.8.5 Method cluster\_db/plotQuality

**Summary:** Creates a plot\_abstract of the silhouette plot showing the clustering quality.

**Usage:** 

```
a_plot = plotQuality(a_cluster_db, title_str)
```

## Parameters:

```
a_cluster_db: A cluster_db object.
```

title\_str: (Optional) String to append to plot title.

# **Returns:**

a\_plot: A plot\_abstract object that can be plotted.

See also: plot\_abstract (p. 153), plotFigure (p. ??), silhouette (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/08

# A.8.6 Method cluster\_db/plot\_abstract

**Summary:** Creates a vertical plot\_stack of silhouette and membership histograms for the clusters.

## **Usage:**

```
a_plot = plot_abstract(a_cluster_db, title_str)
```

## **Parameters:**

```
a_cluster_db: A cluster_db object.
title_str: (Optional) String to append to plot title.
```

## **Returns:**

a\_plot: A plot\_abstract object that can be plotted.

See also: cluster\_db/plotQuality(p.80), cluster\_db/plotHist(p.80)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/08

# A.9 Class corrcoefs\_db

# A.9.1 Constructor corrcoefs\_db/corrcoefs\_db

**Summary:** A database of correlation coefficients generated from a column of another database.

## Usage:

```
a_coef_db = corrcoefs_db(col_name, coefs, coef_names, pages, id, props)
```

**Description:** This is a subclass of tests\_3d\_db. Allows generating a plot, etc.

# **Parameters:**

col name: The column with which the others are correlated.

coefs: Matrix where each column has another coefficient.

**coef\_names:** Cell array of column names corresponding to coefficients.

pages: Column vector of page indices pointing to the tests\_3d\_db.

id: An identifying string.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

```
tests_db.
```

```
See also: tests_db (p. 217), plot_simple (p. 163), tests_db/histogram (p. 225)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/06

# A.10 Class dataset\_db\_bundle

## A.10.1 Constructor dataset\_db\_bundle/dataset\_db\_bundle

**Summary:** The dataset and the DB created from it bundled together.

**Usage:** 

```
a_bundle = dataset_db_bundle(a_dataset, a_db, a_joined_db, props)
```

**Description:** This class is made to enable operations that require seamless connection between the high-level DB and the raw data. The raw DB is only required to bridge the gap between the high-level DB and the dataset. Therefore it only needs to contain columns necessary to make this connection. It is not required to include all raw DB columns, which is inefficient.

## **Parameters:**

**a\_dataset:** A params\_tests\_dataset object or a subclass.

**a** db: The raw tests db object (or a subclass) created from the dataset.

a\_joined\_db: The processed DB created from the raw DB.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

dataset, db, joined\_db, props.

```
See also: tests_db (p. 217), params_tests_dataset (p. 111)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/13

# A.10.2 Method dataset\_db\_bundle/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

## A.10.3 Method dataset\_db\_bundle/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.10.4 Method dataset\_db\_bundle/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.10.5 Method dataset\_db\_bundle/constrainedMeasuresPreset

**Summary:** Returns a dataset\_db\_bundle with constrained measures according to chosen preset.

# Usage:

[a\_bundle test\_names] = constrainedMeasuresPreset(a\_bundle, preset, props)

## **Parameters:**

**a\_bundle:** A dataset\_db\_bundle object.

**preset:** Choose preset measure list (default=1). **props:** A structure with any optional properties.

## **Returns:**

a bundle: Modified bundle.

See also: physiol\_bundle/constrainedMeasuresPreset (p. 140)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/13

## A.10.6 Method dataset\_db\_bundle/rankingReportTeX

**Summary:** Generates a report by comparing a\_bundle with the given match criteria, crit\_db from crit\_bundle.

## **Usage:**

```
tex_string = rankingReportTeX(a_bundle, crit_bundle, crit_db, props)
```

**Description:** Generates a LaTeX document with: - (optional) Raw traces compared with some best matches at different distances - Values of some top matching a\_db rows and match errors in a floating table. - colored-plot of measure errors for some top matches. - Parameter distributions of 50 best matches as a bar graph.

## **Parameters:**

**a\_bundle:** A dataset\_db\_bundle object that contains the DB to compare rows from.

**crit\_bundle:** A dataset\_db\_bundle object that contains the criterion dataset.

**crit\_db:** A tests\_db object holding the match criterion tests and STDs which can be created with matchingRow.

**props:** A structure with any optional properties.

**caption:** Identification of the criterion db (not needed/used?). **num\_matches:** Number of best matches to display (default=10).

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rotate: Rotation angle for best matches table (default=90).

## **Returns:**

tex\_string: LaTeX document string.

See also: displayRowsTeX (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/13

# A.10.7 Method dataset\_db\_bundle/matchingRow

**Summary:** Creates a criterion database for matching the tests of a row.

## **Usage:**

```
crit_db = matchingRow(a_bundle, row, props)
```

**Description:** Copies selected test values from row as the first row into the criterion db. Adds a second row for the STD of each column in the db.

## **Parameters:**

a\_bundle: A tests\_db object.

row: A row index to match.

**props:** A structure with any optional properties.

distDB: Take the standard deviation from this db instead.

## **Returns:**

crit\_db: A tests\_db with two rows for values and STDs.

# **Example:**

See also: rankMatching (p. ??), tests\_db (p. 217), tests2cols (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/21

## A.10.8 Method dataset\_db\_bundle/reportNeuron

**Summary:** Generates a report of neuron at given an\_index of a\_bundle.

## **Usage:**

```
a_doc_multi = reportNeuron(a_bundle, an_index, props)
```

**Description:** Generates a report document with preset layouts of annotated plots of the selected neuron. See reportLayout below for presets.

#### **Parameters:**

```
a_bundle: a dataset_db_bundle object which contains the neuronan index: The index to pass to ctFromRows method of a bundle.
```

**props:** A structure with any optional properties.

reportLayout: Allows choosing one of predefined report types (strings):

- 1: Only +/- 100 pA traces in one plot (default). 1a/b: Either one of the +/- 100 pA traces in one plot.
- 2: Only +/- 100 pA traces and spike shapes in one horiz. plot.
- 3: +100 pA raw trace and rate profile stacked vertically.
- **3b:** -100 pA raw trace and rate profile stacked vertically.
- **4:** Horiz stack of +/- 100 pA raw trace with rate profiles underneath.
- 5: 5-piece trace, spike shape, f-I curve, f-t curve quad-plot.

**numTraces:** Limit number of traces to show in plot (>=1).

traces: List of acceptable traces to load.

traceAxisLimits: If given, use these limits for trace plots.

rateAxisLimits: If given, use these limits for rate plots.

**flaxisLimits:** If given, use these limits for flcurve plots.

**fIstats:** Add a fI-stats plot in addition to the curve.

sshapeAxisLimits: If given, use these limits for spike shape plots.

**sshapeResults:** If 1, plot measures on the spike shape (default=1).

### Returns:

a\_doc\_multi: A doc\_multi object that can be printed as a PS or PDF file.

# **Example:**

# A.10.9 Method dataset\_db\_bundle/plotfICurve

**Summary:** Generates a f-I curve doc\_plot for neuron at given an\_index in a\_bundle.

# Usage:

```
a_plot = plotfICurve(a_bundle, trial_num, props)
```

### **Parameters:**

```
a_bundle: A dataset_db_bundle object.
an_index: An index with which to address the a_bundle.
props: A structure with any optional properties.
    shortCaption: This appears in the figure caption.
    plotMStats: If set, add the a_bundle stats plot.
    captionToStats: Use this as its legend label.
    quiet: if given, no title is produced
```

## **Returns:**

a\_plot: A plot\_superpose that contains a f-I curve plot.

## **Example:**

```
» a_p = plotfICurve(r, 1);
» plotFigure(a_p, 'The f-I curve of best matching model');

See also: plot_abstract (p. 153), plot_superpose (p. 167)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/16
```

(passed to plot\_superpose)

# A.10.10 Method dataset\_db\_bundle/subsref

**Summary:** Defines indexing for tests\_db objects for () and . operations.

# **Usage:**

```
obj = obj(rows, tests) obj = obj.attribute
```

**Description:** Returns attributes or selects the given test columns and rows and returns in a new tests\_db object.

### **Parameters:**

```
obj: A tests_db object.
rows: A logical or index vector of rows. If ':', all rows.
tests: Cell array of test names or column indices. If ':', all tests.
attribute: A tests_db class attribute.
```

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## **Returns:**

obj: The new tests\_db object.

See also: subsref (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17

# A.10.11 Method dataset\_db\_bundle/getNeuronRowIndex

**Summary:** Returns the neuron index from bundle.

## Usage:

```
a_row_index = getNeuronRowIndex(a_bundle, an_index, props)
```

**Description:** This is a polymorphic method. Therefor it is not defined for this class, but see subclasses of dataset\_db\_bundle for its more meaningful implementations.

#### **Parameters:**

a\_bundle: A dataset\_db\_bundle subclass object.

an\_index: An index number of neuron, or a DB row containing this.

**props:** A structure with any optional properties.

## **Returns:**

a\_row\_index: A row index of neuron in a\_bundle.joined\_db.

## **Example:**

```
» displayRows(mbundle.joined_db(getNeuronRowIndex(mbundle, 98364), :))
```

See also: dataset\_db\_bundle (p. 82)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/09

# A.10.12 Method dataset\_db\_bundle/subsasgn

Summary: Defines generic index-based assignment for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/06

# A.10.13 Method dataset\_db\_bundle/ctFromRows

**Summary:** Loads a cip\_trace object from a raw data file in the a\_bundle.

## **Usage:**

```
a_cip_trace = ctFromRows(a_bundle, a_db, cip_levels, props)
```

**Description:** This method is not implemented for the generic dataset\_db\_bundle class. See subclass implementations.

### **Parameters:**

- **a\_bundle:** A dataset\_db\_bundle object.
- a\_db: A DB created by the dataset in the a\_bundle to read the neuron index numbers from.

**cip levels:** A column vector of CIP-levels to be loaded.

**props:** A structure with any optional properties.

(passed to a\_bundle.dataset/cip\_trace)

## **Returns:**

a cip trace: One or more cip trace objects that hold the raw data.

See also: model\_ct\_bundle/ctFromRows (p. 103), physiol\_bundle/ctFromRows (p. 143)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/10/11

# A.11 Class doc\_generate

# A.11.1 Constructor doc\_generate/doc\_generate

Summary: Generic class to help generate printed or annotated documents with results.

# Usage:

```
a_doc = doc_generate(text_string, id, props)
```

**Description:** This constitutes the base class for other doc\_classes. For convenience, this class holds a text\_string to be printed when the document is generated with the printTeXFile option.

## **Parameters:**

text\_string: Contents of this document.

id: An identifying string.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

text, id, props.

See also: doc\_plot (p. 92), doc\_multi (p. 91)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.11.2 Method doc\_generate/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.11.3 Method doc\_generate/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.11.4 Method doc\_generate/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.11.5 Method doc\_generate/subsref

Summary: Defines generic indexing for objects.

# A.11.6 Method doc\_generate/printTeXFile

**Summary:** Creates a TeX file with the contents of this document.

**Usage:** 

printTeXFile(a\_doc, filename, props)

**Description:** Calls getTeXString to generate the contents. The filename is adjusted with a call to properFilename to generate an acceptable TeX filename. TeX-specific should only be added at this point or at getTeXString, because before we want the object to be a generic document container.

# **Parameters:**

a\_doc: A tests\_db object.

PANDORA Toolbox: Prog.'s MaDraft v805 or 2007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

**filename:** To write the TeX string.

**props:** A structure with any optional properties.

## **Returns:**

tex\_string: A string that contains TeX commands, which upon writing to a file, can be interpreted by the TeX engine to produce a document.

## **Example:**

```
» a_doc = doc_plot(a_plot, 'Results from cell.', 'Results.', struct, ");
» printTeXFile(a_doc, 'my_doc.tex')
then my_doc.tex can be used by including from a valid LaTeX document.
```

See also: doc\_generate (p. 88), doc\_plot (p. 92), string2File (p. ??), properFilename (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.11.7 Method doc\_generate/getTeXString

**Summary:** Returns the TeX representation for the document.

## **Usage:**

```
tex_string = getTeXString(a_doc, props)
```

**Description:** This is an abstract placeholder for this method. It specifies what this method should do in the subclasses that implement it. This method should create all the auxiliary files needed by the document. The generated tex\_string should be ready to be visualized.

## **Parameters:**

**a\_doc:** A tests\_db object.

**props:** A structure with any optional properties.

## **Returns:**

tex\_string: A string that contains TeX commands, which upon writing to a file, can be interpreted by the TeX engine to produce a document.

## **Example:**

See also: doc\_generate (p. 88), doc\_plot (p. 92)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.12 Class doc\_multi

## A.12.1 Constructor doc\_multi/doc\_multi

**Summary:** A document that is composed of multiple other doc\_generate objects.

Usage:

```
a_doc = doc_multi(docs, id, props)
```

## Parameters:

**docs:** A vector of doc\_generate objects.

id: An identifying string.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

docs, doc\_generate.

## **Example:**

```
» mydoc = doc_multi([doc_plot(a_plot1), doc_plot(a_plot2)], 'Two plots')
» printTeXFile(mydoc, 'two_plots.tex')
```

See also: doc\_generate(p. 88), getTeXString(p. ??), doc\_generate/printTeXFile(p. 89)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.12.2 Method doc\_multi/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.12.3 Method doc\_multi/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.12.4 Method doc\_multi/getTeXString

**Summary:** Returns the TeX representation for the document.

# Usage:

```
tex_string = getTeXString(a_doc, props)
```

**Description:** Concatenates TeX representations of doc\_generate, or subclass, objects it contains

### **Parameters:**

a\_doc: A tests\_db object.

**props:** A structure with any optional properties.

## **Returns:**

tex\_string: A string that contains TeX commands, which upon writing to a file, can be interpreted by the TeX engine to produce a document.

# **Example:**

See also: doc\_generate (p. 88), doc\_plot (p. 92)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.13 Class doc\_plot

# A.13.1 Constructor doc\_plot/doc\_plot

Summary: Generates a formatted plot for printing, annotated with captions.

## **Usage:**

```
a_doc = doc_plot(a_plot, caption, plot_filename, float_props, id, props)
```

**Description:** The generated file may take an extension according to chosen format.

# **Parameters:**

```
a_plot: A plot_abstract ready to be visualized.caption: Long caption to appear under the figure.
```

plot\_filename: Filename to be generated from the plot.

float\_props: Formatting instructions passed to TeXtable.

id: An identifying string.

PANDORA Toolbox: Prog.'s MaBuraft v805 or 92007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

**props:** A structure with any optional properties.

**orient:** Passed to the orient command before printing to figure file.

# Returns a structure object with the following fields:

plot, caption, plot\_filename, float\_props, doc\_generate.

# **Example:**

```
» a_doc = doc_plot(plotData(my_cip_trace), 'My CIP trace. Very interesting.',
...
'tracel', struct, 'first doc');
» printTeXFile(a_doc, 'my_doc.tex');
```

See also: doc\_generate (p. 88), TeXtable (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.13.2 Method doc\_plot/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.13.3 Method doc\_plot/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.13.4 Method doc\_plot/plot

**Summary:** Default plot method to preview the contained plot in a new figure.

# **Usage:**

```
figure_handle = plot(a_doc, props)
```

**Description:** Only generate the contained plot for previewing. Opens a new figure.

# **Parameters:**

a\_doc: A doc\_plot object.

**props:** A structure with any optional properties.

## **Returns:**

figure\_handle: Handle of newly opened figure.

PANDORA Toolbox: Prog.'s MaBuraft v805 or 92007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

# **Example:**

```
» figure_handle = plot(a_doc_plot)
```

See also: plot\_abstract/plotFigure (p. 155), doc\_generate (p. 88), doc\_plot (p. 92)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.13.5 Method doc\_plot/getTeXString

**Summary:** Returns the TeX representation for the plot document.

Usage:

```
tex_string = getTeXString(a_doc, props)
```

**Description:** Plots, prints EPS files and generates the necessary LaTeX code.

### **Parameters:**

a\_doc: A doc\_plot object.

**props:** A structure with any optional properties.

### **Returns:**

tex\_string: A string that contains TeX commands, which upon writing to a file, can be interpreted by the TeX engine to produce a document.

# **Example:**

See also: doc\_generate (p. 88), doc\_plot (p. 92)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.14 Class histogram\_db

## A.14.1 Constructor histogram\_db/histogram\_db

**Summary:** A database of histogram values generated for a column of another database.

## **Usage:**

```
a_hist_db = histogram_db(col_name, bins, hist_results, id, props)
```

**Description:** This is a subclass of tests\_db. Allows generating a histogram plot, etc.

PANDORA Toolbox: Prog.'s MaBuraft v805 or 924007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

## **Parameters:**

col\_name: The column name of the histogrammed value.

**bins:** The values for which the histogram values are calculated.

hist\_results: A column vector of histogram values.

id: An identifying string.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

tests\_db, props.

## **Example:**

```
» [hist_results, bins] = hist(my_data);
» a_hist_db = histogram_db('firing_rate', bins, hist_results, 'rate histogram db');
» plot(a_hist_db);

See also: tests_db (p. 217), plot_simple (p. 163), tests_db/histogram (p. 225)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/20

# A.14.2 Method histogram\_db/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

## A.14.3 Method histogram\_db/calcMode

Summary: Finds the mode of the distribution, that is, the bin with the highest value.

**Usage:** 

```
[mode_val, mode_mag] = calcMode(a_hist_db)
```

## **Parameters:**

**a\_hist\_db:** A histogram\_db object.

## **Returns:**

mode\_val: The center of the bin that has most members. mode\_mag: The value of the histogram bin.

See also: histogram\_db (p. 94)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/27

# A.14.4 Method histogram\_db/plot\_abstract

**Summary:** Generates a plottable description of this object.

Usage:

```
a_plot = plot_abstract(a_hist_db, title_str, props)
```

**Description:** Generates a plot\_simple object from this histogram.

## **Parameters:**

a\_hist\_db: A histogram\_db object.

props: Optional properties passed to plot\_abstract.

command: Plot command (Optional, default='bar')

logScale: If 1, use logarithmic y-scale.

quiet: If 1, don't include database name on title.

### **Returns:**

a plot: A object of plot abstract or one of its subclasses.

See also: plot\_abstract (p. 153), plot\_simple (p. 163)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

# A.14.5 Method histogram\_db/subsref

Summary: Defines generic indexing for objects.

## A.14.6 Method histogram\_db/plotPages

Summary: Generates a plot containing subplots from each page of histograms.

## Usage:

```
a_plot = plotPages(a_hist_db, command, an_orient)
```

**Description:** For each page of the histogram, a histogram is placed in a subplot.

## **Parameters:**

```
a_hist_db: A histogram_db object.
```

command: Plot command (Optional, default='bar')

an\_orient: Stack orientation. One of 'x', 'y', or 'z'.

## **Returns:**

a\_plot: A object of plot\_abstract or one of its subclasses.

PANDORA Toolbox: Prog.'s MaBuraft v805 or 92007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

See also: plotPages (p. ??), plot\_simple (p. 163)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/04

# A.14.7 Method histogram\_db/plotRowMatrix

Summary: Generates a subplot matrix of measure columns versus rows of databases.

## **Usage:**

```
a_plot = plotRowMatrix(hist_dbs, title_str, props)
```

**Description:** Each row in the hist\_dbs is assumed to come from a different DB. Columns represent histograms of different measurements. The plot is made such that histograms in each row have the same maximal count, and histograms in each column have the same axis limits.

### **Parameters:**

hist\_dbs: A matrix of histogram\_db objects.

title\_str: Title to go at the top.

props: A structure with any optional properties.

**rowLabels:** Cell array of y-axis labels for each row. (rest passed to histogram\_db/plot\_abstract)

## **Returns:**

a\_plot: A object of plot\_abstract or one of its subclasses.

See also: plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/11/22

A.14.8 Method histogram\_db/plotEqSpaced

**Summary:** Generates a histogram plot where the values are equally spaced on the x-axis. For use with non-linear parameter values.

## **Usage:**

```
a_plot = plotEqSpaced(a_hist_db, command, props)
```

**Description:** Generates a plot\_simple object from this histogram.

# **Parameters:**

```
a_hist_db: A histogram_db object.
command: Plot command (Optional, default='bar')
```

PANDORA Toolbox: Prog.'s MaBuraft v805 or 92007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

props: Optional properties passed to plot\_abstract.

#### **Returns:**

a\_plot: A object of plot\_abstract or one of its subclasses.

See also: plot\_abstract (p. 153), plot\_simple (p. 163)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

# A.15 Class model\_ct\_bundle

# A.15.1 Constructor model\_ct\_bundle/model\_ct\_bundle

**Summary:** The model cip\_trace dataset and the DB created from it bundled together.

# **Usage:**

```
a_bundle = model_ct_bundle(a_dataset, a_db, a_joined_db, props)
```

**Description:** This is a subclass of dataset\_db\_bundle, specialized for model datasets.

### **Parameters:**

- a\_dataset: A params\_cip\_trace\_fileset object.
- **a\_db:** The raw params\_tests\_db object created from the dataset. It only needs to have the pAcip, trial, and ItemIndex columns.
- a\_joined\_db: The one-model-per-line DB created from the raw DB.

**props:** A structure with any optional properties.

## Returns a structure object with the following fields:

dataset\_db\_bundle.

# **Example:**

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/13

# A.15.2 Method model\_ct\_bundle/getNeuronLabel

**Summary:** Constructs the neuron label from bundle.

## Usage:

```
a_label = getNeuronLabel(a_bundle, trial_num, props)
```

#### Parameters:

a\_bundle: A physiol\_cip\_traceset\_fileset object.trial\_num: The trial number of model neuron.props: A structure with any optional properties.

### **Returns:**

a\_label: A string label identifying selected neuron in bundle.

See also: dataset\_db\_bundle (p. 82)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/26

# A.15.3 Method model\_ct\_bundle/reportCompareModelToPhysiolNeuron

**Summary:** Generates a report by comparing given model neuron to given physiol neuron.

# Usage:

a\_doc\_multi = reportCompareModelToPhysiolNeuron(m\_bundle, trial\_num, p\_bundle, traceset\_index, props)

**Description:** Generates a report document with: - Figure displaying raw traces of the physiol neuron compared with the model neuron - Figure comparing f-I curves of the two neurons. - Figure comparing spont and pulse spike shapes of the two neurons.

### **Parameters:**

**m\_bundle**, **p\_bundle**: dataset\_db\_bundle objects of the model and physiology neurons.

trial\_num: Trial number of desired model neuron in m\_bundle.

traceset\_index: TracesetIndex of desired neuron in p\_bundle.

**props:** A structure with any optional properties.

horizRow: If defined, create a row-figure with all plots.

**numPhysTraces:** Number of physiology traces to show in plot (>=1).

### **Returns:**

a\_doc\_multi: A doc\_multi object that can be printed as a PS or PDF file.

PANDORA Toolbox: Prog.'s MaBuraft v805 or 92007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

# **Example:**

```
> printTeXFile(reportCompareModelToPhysiolNeuron(mbundle, 2222, pbundle, 34), 'a.tex')
```

```
See also: doc_multi(p.91), doc_generate(p.88), doc_generate/printTeXFile (p.89)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/24

# A.15.4 Method model\_ct\_bundle/plotCompareRanks

**Summary:** Generates a plots of given ranks from the ranked\_bundle.

## **Usage:**

```
plots = plotCompareRanks(m_bundle, p_bundle, a_ranked_db, ranks, props)
```

### **Parameters:**

m\_bundle: A model\_ct\_bundle object.

**p\_bundle:** A dataset\_db\_bundle object that originated the criterion.

**a\_ranked\_db:** A ranked\_db generated from ranking m\_bundle.

ranks: Vector of rank indices for which to generate the plots.

props: A structure with any optional properties.

## **Returns:**

plots: A structure that contains the joined\_db, and the plot vectors trace\_d100\_plots and trace h100 plots.

## **Example:**

```
» plots = plotCompareRanks(r, 1:10);
» plotFigure(plots.trace_d100_plots(1), 'The best matching +100 pA CIP trace');
```

## See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/16

# A.15.5 Method model\_ct\_bundle/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.15.6 Method model\_ct\_bundle/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.15.7 Method model\_ct\_bundle/getTrialNum

**Summary:** Extracts identifying neuron trial number from DB.

**Usage:** 

trial\_num = getTrialNum(a\_bundle, a\_db|trial\_num, props)

#### **Parameters:**

a\_bundle: A physiol\_cip\_traceset\_fileset object.

**a\_db:** DB rows representing deisred model neuron(s).

**trial\_num:** Trial numbers. If specified, this function does nothing but return them.

**props:** A structure with any optional properties.

### **Returns:**

trial\_num: The trial number(s) identifying selected neuron(s) in bundle.

See also: dataset\_db\_bundle (p. 82)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/26

## A.15.8 Method model\_ct\_bundle/collectPhysiolMatches

Summary: Compare model DB to given physiol criteria and return some top matches.

**Usage:** 

row\_index = collectPhysiolMatches(a\_mbundle, a\_crit\_bundle, props)

## **Parameters:**

a\_mbundle: A model\_ct\_bundle object.

**a\_crit\_bundle:** A physiol\_bundle object that holds the criterion neuron.

**props:** A structure with any optional properties.

**showTopmost:** Number of top matching models to return (default=50)

## **Returns:**

row index: Row indices of best matching models.

## See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/18

# A.15.9 Method model\_ct\_bundle/plotComparefICurve

**Summary:** Generates a f-I curve doc\_plot comparing m\_trial and to\_index.

## **Usage:**

```
a_plot = plotComparefICurve(m_bundle, m_trial, to_bundle, to_index, props)
```

**Description:** Note that this is not a general method. to\_bundle should have been able to accept any type of bundle. Most probably this method is redundant and deprecated.

## **Parameters:**

# **Returns:**

a\_plot: A plot\_superpose that contains a f-I curve plot.

## **Example:**

```
» a_p = plotComparefICurve(r, 1);
» plotFigure(a_p, 'The f-I curve of best matching model');

See also: plot_abstract (p. 153), plot_superpose (p. 167)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/16
```

# A.15.10 Method model\_ct\_bundle/getNeuronRowIndex

**Summary:** Returns the neuron index from bundle.

## **Usage:**

```
a_row_index = getNeuronRowIndex(a_bundle, trial_num, props)
```

### **Parameters:**

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

a\_bundle: A model\_ct\_bundle object.

trial\_num: The trial number of model neuron, or a DB row containing this.

**props:** A structure with any optional properties.

#### **Returns:**

a\_row\_index: A row index of neuron in a\_bundle.joined\_db.

See also: dataset\_db\_bundle (p. 82)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/09

# A.15.11 Method model\_ct\_bundle/ctFromRows

**Summary:** Loads a cip\_trace object from a raw data file in the a\_mbundle.

Usage:

```
a_cip_trace = ctFromRows(a_mbundle, a_db|trials, cip_levels, props)
```

**Description:** This is an overloaded method.

## **Parameters:**

**a\_mbundle:** A model\_ct\_bundle object.

**a\_db:** A DB created by the dataset in the a\_mbundle to read the trial numbers from.

trials: A column vector with trial numbers.

cip\_levels: A column vector of CIP-levels to be loaded.

**props:** A structure with any optional properties. (passed to a\_mbundle.dataset/cip\_trace)

## **Returns:**

a\_cip\_trace: One or more cip\_trace objects that hold the raw data.

See also: dataset\_db\_bundle/ctFromRows (p. 88)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/07/13

## A.15.12 Method model\_ct\_bundle/addToDB

**Summary:** Concatenate to existing DB in the bundle.

## Usage:

```
a_mbundle = addToDB(a_mbundle, a_raw_db, props)
```

**Description:** If joinedDb is not given in props, calls treatSimDB to get the joined\_db from this raw DB. Then concats to both db and joined\_db in bundle.

### **Parameters:**

```
a_mbundle: A model_ct_bundle object.
```

**a\_crit\_bundle:** A physiol\_bundle having a crit\_db as its joined\_db.

**props:** A structure with any optional properties.

joinedDb: The joined version of a\_raw\_db.

dataset: If given, this one is used to replace the fileset in the bundle.

## **Returns:**

a\_mbundle: a model\_ct\_bundle object containing the added DB.

## **Example:**

```
» mbundle = addToDB(mbundle, params_tests_db(mfileset, [19684:59956]))
```

See also: params\_tests\_fileset/addFiles(p. 133), multi\_fileset\_gpsim\_cns2005/addFileDir(p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/06

# A.15.13 Method model\_ct\_bundle/reportRankingToPhysiolNeuronsTeXFile

Summary: Compare model DB to given physiol criterion and create a report.

# Usage:

```
tex_filename = reportRankingToPhysiolNeuronsTeXFile(m_bundle, p_bundle,
a_crit_db, props)
```

**Description:** A LaTeX report is generated following the example in physiol\_bundle/matchingRow. The filename contains the neuron name, followed by the traceset index as an identifier of pharmacological applications, as in gpd0421c\_s34.

## Parameters:

```
m_bundle: A model_ct_bundle object.p_bundle: A physiol_bundle object.
```

**a\_crit\_db:** The criterion neuron chosen with a matchingRow method.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2407-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

props: A structure with any optional properties.

filenameSuffix: Append this identifier to the TeX filename.

(others passed to rankMatching)

#### Returns:

tex\_filename: Name of LaTeX file generated.

See also: tests\_db/rankMatching (p. 254), physiol\_cip\_traceset/cip\_trace (p. 149), physiol\_bundle/matchingRow (p. 140)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/18

# A.15.14 Method model\_ct\_bundle/rankMatching

Summary: Create a ranked\_db from given criterion db.

**Usage:** 

a\_ranked\_db = rankMatching(a\_mbundle, a\_crit\_db, props)

### **Parameters:**

a\_mbundle: A model\_ct\_bundle object.

**a\_crit\_db:** A crit\_db created by a matchingRow method.

**props:** A structure with any optional properties.

(passed to tests\_db/rankMatching)

## **Returns:**

a\_ranked\_db: a ranked\_db object containing the rankings.

See also: tests\_db/rankMatching (p. 254), ranked\_db (p. 170)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/18

# A.16 Class model\_ranked\_to\_physiol\_bundle

# A.16.1 Constructor model\_ranked\_to\_physiol\_bundle/model\_ranked\_to\_physiol\_bundle

**Summary:** A DB bundled with its dataset, ranked to a physiology DB bundle.

# **Usage:**

```
r_bundle = model_ranked_to_physiol_bundle(a_dataset, a_db, a_ranked_db, a_crit_bundle, props)
```

**Description:** This is a subclass of model\_ct\_bundle, specialized for model datasets.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2507-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

## **Parameters:**

- a\_dataset: A params\_cip\_trace\_fileset object.
- **a\_db:** The raw params\_tests\_db object created from the dataset. It only needs to have the pAcip, trial, and ItemIndex columns.
- **a\_ranked\_db:** The one-model-per-line DB created from the raw DB.
- **a\_crit\_bundle:** The bundle object associated with crit\_db that caused the ranking in a\_ranked\_db.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

```
crit_bundle, model_ct_bundle.
```

```
See also: model_ct_bundle (p. 98), ranked_db (p. 170), params_tests_dataset (p. 111)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/13

# A.16.2 Method model\_ranked\_to\_physiol\_bundle/plotCompareRanks

**Summary:** OBSOLETE - Generates a plots of given ranks from the ranked\_bundle.

## Usage:

```
plots = plotCompareRanks(r_bundle, crit_bundle, crit_db, props)
```

### **Parameters:**

r\_bundle: A ranked\_bundle object.

ranks: Vector of rank indices for which to generate the plots.

**props:** A structure with any optional properties.

### **Returns:**

plots: A structure that contains the joined\_db, and the plot vectors trace\_d100\_plots and trace\_h100\_plots.

# **Example:**

```
» plots = plotCompareRanks(r, 1:10);
» plotFigure(plots.trace_d100_plots(1), 'The best matching +100 pA CIP trace');
```

## See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/16

# A.16.3 Method model\_ranked\_to\_physiol\_bundle/plotfICurve

## **Usage:**

```
a_doc = docfICurve(r_bundle, crit_bundle, crit_db, props)
```

## **Parameters:**

**r\_bundle:** A ranked\_bundle object.

rank\_num: Rank index for which to generate the a\_doc.

**props:** A structure with any optional properties.

## **Returns:**

a\_doc: A doc\_plot that contains a f-I curve plot and associated captions.

# **Example:**

```
» a_d = docfICurve(r, 1);
» plot(a_d, 'The f-I curve of best matching model');

See also: doc_generate(p. 88), doc_plot(p. 92)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/16
```

## A.16.4 Method model\_ranked\_to\_physiol\_bundle/comparisonReport

**Summary:** OBSOLETE - Generates a report by comparing r\_bundle with the given match criteria, crit\_db from crit\_bundle.

## Usage:

```
a_doc_multi = comparisonReport(r_bundle, crit_bundle, crit_db, props)
```

**Description:** Generates a LaTeX document with: - (optional) Raw traces compared with some best matches at different distances - Values of some top matching a\_db rows and match errors in a floating table. - colored-plot of measure errors for some top matches. - Parameter distributions of 50 best matches as a bar graph.

## **Parameters:**

**r\_bundle:** A dataset\_db\_bundle object that contains the DB to compare rows from.

**crit\_bundle:** A dataset\_db\_bundle object that contains the criterion dataset.

**crit\_db:** A tests\_db object holding the match criterion tests and STDs which can be created with matchingRow.

**props:** A structure with any optional properties.

caption: Identification of the criterion db (not needed/used?).

```
num_matches: Number of best matches to display (default=10).
rotate: Rotation angle for best matches table (default=90).
```

## **Returns:**

tex\_string: LaTeX document string.

See also: displayRowsTeX (p. ??)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

# A.17 Class params\_cip\_trace\_fileset

# A.17.1 Constructor params\_cip\_trace\_fileset/params\_cip\_trace\_fileset

**Summary:** Description of a raw dataset consisting of cip\_trace files varying with parameter values.

# Usage:

```
obj = params_cip_trace_fileset(file_pattern, dt, dy, pulse_time_start,
pulse_time_width, id, props)
```

**Description:** This is a subclass of params\_tests\_fileset.

## **Parameters:**

```
file_pattern: File pattern mathing all files to be loaded.
```

dt: Time resolution [s]

dy: y-axis resolution [ISI (V, A, etc.)]

pulse\_time\_start, pulse\_time\_width: Start and width of the pulse
 [dt]

id: An identification string

**props:** A structure with any optional properties.

**profile\_class\_name:** Use this profile class (Default: 'cip\_trace\_profile'). (All other props are passed to cip\_trace objects)

# Returns a structure object with the following fields:

params\_tests\_fileset, pulse\_time\_start, pulse\_time\_width.

## **Example:**

```
» fileset = params_cip_trace_fileset('/home/abc/data/*.bin', 1e-4, 1e-3, 20001,
10000, 'sim dataset gpsc0501', struct('trace_time_start', 10001, 'type', 'sim',
'scale_y', 1e3))
```

See also: params\_tests\_fileset (p. 132), params\_tests\_db (p. 116)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.17.2 Method params\_cip\_trace\_fileset/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.17.3 Method params\_cip\_trace\_fileset/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.17.4 Method params\_cip\_trace\_fileset/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.17.5 Method params\_cip\_trace\_fileset/cip\_trace\_profile

**Summary:** Loads a raw cip\_trace\_profile given a file\_index to this fileset.

Usage:

```
a_cip_trace_profile = cip_trace_profile(fileset, file_index)
```

### Parameters:

fileset: A params\_tests\_fileset.
file index: Index of file in fileset.

# **Returns:**

a\_cip\_trace\_profile: A cip\_trace\_profile object.

See also: cip\_trace\_profile (p. 69), params\_tests\_fileset (p. 132)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.17.6 Method params\_cip\_trace\_fileset/ctFromRows

Summary: Loads a cip\_trace object from raw data files in the fileset.

**Usage:** 

```
a_cip_trace = ctFromRows(m_fileset, m_dball, a_db|itemIndices, cip_levels, props)
```

#### **Parameters:**

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```
m_fileset: A physiol_cip_traceset_fileset object.
```

m\_dball: A DB created by this fileset that contains the trial, pAcip, and ItemIndex cols.

**a\_db:** A DB that has one trial for each cip\_trace to be loaded.

itemIndices: A column vector with ItemIndex numbers.

cip\_levels: A column vector of CIP-levels to be loaded.

**props:** A structure with any optional properties.

**neuronLabel:** appropriate unique neuron label generated by the bundle.

(passed to params\_cip\_trace\_fileset/cip\_trace)

### **Returns:**

a\_cip\_trace: One or more cip\_trace objects that hold the raw data.

See also: loadItemProfile (p. ??), physiol\_cip\_traceset/cip\_trace (p. 149)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/07/13

### A.17.7 Method params\_cip\_trace\_fileset/loadItemProfile

Summary: Loads a cip\_trace\_profile object from a raw data file in the fileset.

### Usage:

```
[params_row, tests_row] = loadItemProfile(fileset, file_index)
```

### **Parameters:**

fileset: A params\_tests\_fileset.
file\_index: Index of file in fileset.

### Returns:

a\_profile: A profile object that implements the getResults method.

See also: itemResultsRow(p.??), params\_tests\_fileset(p.132), paramNames (p.??), testNames(p.??)

### A.17.8 Method params\_cip\_trace\_fileset/cip\_trace

**Summary:** Loads raw cip\_traces for each given file\_index in this fileset.

#### **Usage:**

```
a_cip_trace = cip_trace(fileset, file_index|a_db, props)
```

#### **Parameters:**

fileset: A params\_tests\_fileset.

file\_index: A single or array of indices of files in fileset.

**a\_db:** A DB created by this fileset to read the item indices from.

**props:** A structure with any optional properties.

**neuronLabel:** Used for annotation purposes.

#### **Returns:**

a\_cip\_trace: A cip\_trace object.

See also: cip\_trace (p. 54), params\_tests\_fileset (p. 132)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/13

# A.18 Class params\_tests\_dataset

### A.18.1 Constructor params\_tests\_dataset/params\_tests\_dataset

**Summary:** Contains a set of data objects or files of raw data varying with parameter values.

# **Usage:**

```
obj = params_tests_dataset(list, dt, dy, id, props)
```

**Description:** This is an abstract base class for keeping dataset information separate from the parameters-results database (params\_tests\_db). The list contents can be filenames or objects (such as cip\_traces) from which to get the raw data. The dataset should have all the necessary information to create a db when needed. This is an abstract class, thet it it cannot act on its own. Only fully implemented subclasses can actually hold datasets. See methods below.

#### **Parameters:**

**list:** Array of dataset items (filenames, objects, etc.).

dt: Time resolution [s]

dy: y-axis resolution [integral V, A, etc.]

id: An identification string.

**props:** A structure with any optional properties.

type: type of file (default = ")

### Returns a structure object with the following fields:

list, dt, dy, id, props (see above).

See also: params\_tests\_db (p. 116), params\_tests\_fileset (p. 132), cip\_traces\_dataset (p. 71)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/02

# A.18.2 Method params\_tests\_dataset/getItem

Summary: Returns the dataset item at given index.

Usage:

item = getItem(dataset, index)

#### Parameters:

dataset: A params\_tests\_dataset.
index: Index of item in dataset.

#### **Returns:**

item: Object, filename, etc.

See also: itemResultsRow (p.??), params\_tests\_dataset (p. 111), paramNames (p.??), testNames (p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/03

# A.18.3 Method params\_tests\_dataset/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.18.4 Method params\_tests\_dataset/get

Summary: Defines generic attribute retrieval for objects.

# A.18.5 Method params\_tests\_dataset/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

A.18.6 Method params\_tests\_dataset/params\_tests\_db

**Summary:** Generates a params\_tests\_db object from the dataset.

**Usage:** 

db\_obj = params\_tests\_db(obj, items, props)

**Description:** This is a converter method to convert from params\_tests\_dataset to params\_tests\_db. Uses readDBItems to read the files. A customized subclass should provide the correct paramNames, testNames, and itemResultsRow functions. Adds a ItemIndex column to the DB to keep track of raw data files after shuffling.

#### **Parameters:**

**obj:** A params\_tests\_dataset object.

**items:** (Optional) List of item indices to use to create the db.

**props:** Any optional params to pass to params\_tests\_db.

# **Returns:**

db\_obj: A params\_tests\_db object.

See also: readDBItems (p.??), params\_tests\_db (p. 116), params\_tests\_dataset (p. 111), itemResultsRow testNames (p. ??), paramNames (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/09

# A.18.7 Method params\_tests\_dataset/subsref

**Summary:** Defines generic indexing for objects.

# A.18.8 Method params\_tests\_dataset/subsasgn

Summary: Defines generic index-based assignment for objects.

# A.18.9 Method params\_tests\_dataset/getItemParams

**Summary:** Get the parameter values of a dataset item.

**Usage:** 

```
params_row = getItemParams(dataset, index, a_profile)
```

**Description:** This method can retrieve the item parameters by using either the dataset and the index to find the item or simply by using the item profile, a\_profile.

#### **Parameters:**

dataset: A params\_tests\_dataset.
index: Index of item in dataset.
a\_profile: An item profile.

#### **Returns:**

params\_row: Parameter values in the same order of paramNames

```
See also: itemResultsRow (p.??), params_tests_dataset (p. 111), paramNames (p.??), testNames (p.??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/10

### A.18.10 Method params\_tests\_dataset/itemResultsRow

**Summary:** Processes a raw data file from the dataset and return its parameter and test values.

### **Usage:**

```
[params_row, tests_row] = itemResultsRow(dataset, index)
```

**Description:** This method is designed to be reused from subclasses as long as the loadItemProfile method is properly overloaded. Adds an Index column to the DB to keep track of raw data items after shuffling.

### **Parameters:**

dataset: A params\_tests\_dataset.
index: Index of file in dataset.

#### **Returns:**

params\_row: Parameter values in the same order of paramNames tests\_row: Test values in the same order with testNames

```
See also: loadItemProfile (p.??), params_tests_dataset (p. 111), paramNames (p. ??), testNames (p. ??)
```

# A.18.11 Method params\_tests\_dataset/addItem

**Summary:** Returns the new dataset with the added item.

**Usage:** 

```
dataset = addItem(dataset, item)
```

**Description:** Note that, this is NOT the way to create a dataset. It is only intended for small additions to an existing dataset. This method is too slow for creating large datasets. The normal method for creating datasets is providing the full list of items to the class constructor.

#### **Parameters:**

dataset: A params\_tests\_dataset.
item: New item to add in dataset.

#### **Returns:**

dataset: With the added item.

See also: itemResultsRow(p.??), params\_tests\_dataset(p.111), paramNames (p.??), testNames(p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/01/25

### A.18.12 Method params\_tests\_dataset/testNames

Summary: Returns the ordered names of tests for this dataset.

**Usage:** 

```
test_names = testNames(dataset, item)
```

**Description:** Looks at the results of the first file to find the test names.

#### **Parameters:**

dataset: A params\_tests\_dataset.

#### **Returns:**

params\_names: Cell array with ordered parameter names. item: (Optional) If given, read names by loading item at this index.

See also: params\_tests\_dataset (p. 111), paramNames (p. ??), testNames (p. ??)

# A.18.13 Method params\_tests\_dataset/readDBItems

Summary: Reads all items to generate a params\_tests\_db object.

#### Usage:

```
[params, param_names, tests, test_names] = readDBItems(obj, items)
```

**Description:** This is a generic method to convert from params\_tests\_fileset to a params\_tests\_db, or a subclass. This method depends on the paramNames, testNames, and item-ResultsRow functions. Outputs of this function can be directly fed to the constructor of a params\_tests\_db or a subclass.

#### **Parameters:**

**obj:** A params\_tests\_fileset object.

items: (Optional) List of item indices to use to create the db.

#### **Returns:**

```
params, param_names, tests, test_names: See params_tests_db.
```

See also: params\_tests\_db (p. 116), params\_tests\_fileset (p. 132), itemResultsRow testNames (p. ??), paramNames (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/24

# A.19 Class params\_tests\_db

### A.19.1 Constructor params\_tests\_db/params\_tests\_db

**Summary:** A generic database of test results varying with parameter values, organized in a matrix format.

**Description:** This is a subclass of tests\_db. Defines all operations on this structure so that subclasses can use them.

### **Parameters:**

num\_params: Number of parameters.

a\_tests\_db: A tests\_db upon which to build the params\_tests\_db.

**props:** A structure with any optional properties.

### Returns a structure object with the following fields:

tests\_db num\_params: Number of variable parameters in simulations.

See also: tests\_db (p. 217), test\_variable\_db (N/I) (p. ??)

# A.19.2 Method params\_tests\_db/paramsTestsCoefsHists

**Summary:** Calculates histograms for all pairs of params and tests coefficients and returns in a cell array.

### **Usage:**

```
pt_coefs_hists = paramsTestsCoefsHists(a_db, p_coefs)
```

Description: Skips the 'ItemIndex' test.

#### **Parameters:**

a\_db: A tests\_db object.

**p\_coefs:** Cell array of tests coefficients for each parameter.

### **Returns:**

```
pt_coefs_hists: A cell array of corrcoefs_dbs for each param in a_db.
```

See also: params\_tests\_profile (p. 136)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/17

### A.19.3 Method params\_tests\_db/onlyRowsTests

**Summary:** Returns a tests\_db that only contains the desired tests and rows (and pages).

# Usage:

```
obj = onlyRowsTests(obj, rows, tests, pages)
```

**Description:** Selects the given dimensions and returns in a new tests\_db object.

### **Parameters:**

**obj:** A tests\_db object.

rows: A logical or index vector of rows. If ':', all rows.

tests: Cell array of test names or column indices. If ':', all tests.

pages: (Optional) A logical or index vector of pages. ':' for all pages.

### **Returns:**

obj: The new tests\_db object.

See also: subsref (p. ??), tests\_db (p. 217)

### A.19.4 Method params\_tests\_db/joinRows

**Summary:** Joins the rows of the given db with rows of with\_db with matching RowIndex values.

### Usage:

```
a_db = joinRows(db, tests, with_db, w_tests, index_col_name)
```

**Description:** Takes the desired columns in with\_db with rows having a row index and joins them next to dedired columns from the current db. Assumes each row index only appears once in with\_db. The created db preserves the ordering of with\_db.

#### **Parameters:**

```
db: A param_tests_db object.
```

tests: Test columns to take from db.

with\_db: A tests\_db object with a RowIndex column.

w\_tests: Test columns to take from with\_db.

index\_col\_name: (Optional) Name of row index column (default='RowIndex').

### **Returns:**

```
a_db: A params_tests_db object.
```

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/16

### A.19.5 Method params\_tests\_db/crossProd

**Summary:** Create a DB by taking the cross product of two database row sets.

# **Usage:**

```
cross_db = crossProd(a_db, b_db)
```

**Description:** Overloaded function to maintain correct number of parameters after cross product operation. See original in tests\_db/crossProd.

#### **Parameters:**

```
a_db, b_db: A tests_db object.
```

### **Returns:**

cross\_db: The tests\_db object with all combinations of rows.

See also: tests\_db/crossProd (p. 224)

# A.19.6 Method params\_tests\_db/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.19.7 Method params\_tests\_db/testsHists

**Summary:** Calculates histograms for all tests and returns them in a cell array.

**Usage:** 

t\_hists = testsHists(a\_db, num\_bins)

**Description:** Skips the 'ItemIndex' test.

**Parameters:** 

**a\_db:** One or more tests\_db objects in an array.

**num\_bins:** Number of histogram bins (Optional, default=100), or vector of histogram bin centers.

**Returns:** 

t\_hists: An array of histograms for each test in a\_db.

See also: params\_tests\_profile (p. 136)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/17

### A.19.8 Method params\_tests\_db/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.19.9 Method params\_tests\_db/set

Summary: Generic method for setting object attributes.

# A.19.10 Method params\_tests\_db/matchingRow

**Summary:** Creates a criterion database for matching the tests of a row.

#### Usage:

```
crit_db = matchingRow(a_db, row, props)
```

**Description:** Overloaded method for skipping parameter values. STD for param values will be NaNs.

#### **Parameters:**

a\_db: A tests\_db object.

row: A row index to match.

**props:** A structure with any optional properties.

distDB: Take the standard deviation from this db instead.

#### **Returns:**

crit\_db: A tests\_db with two rows for values and STDs.

```
See also: tests_db/matchingRow(p. 233), rankMatching(p. ??), tests_db (p. 217), tests2cols(p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/13

### A.19.11 Method params\_tests\_db/invarParam

**Summary:** Generates a 3D database of invariant values of a parameter and all test columns.

### **Usage:**

```
a_3D_db = invarParam(db, param)
```

**Description:** Finds all combinations when the rest of the parameters are fixed, and saves the variation of the selected parameter and all tests in a new database.

# **Parameters:**

**db:** A tests\_db object.

param: A parameter name/column number

### **Returns:**

```
a_3D_db: A tests_3D_db object of organized values.
```

```
See also: invarValues (p. ??), tests_3D_db (p. 212), corrCoefs (p. ??), tests_3D_db/plotPair (p. ??)
```

# A.19.12 Method params\_tests\_db/paramsHists

**Summary:** Calculates histograms for all parameters and returns in a cell array.

### Usage:

```
p_hists = paramsHists(a_db)
```

**Description:** Skips the 'ItemIndex' test. Useful for looking at subset databases and find out what parameter values are used most.

#### **Parameters:**

```
a_db: A tests_db object.
```

#### **Returns:**

p\_hists: An array of histograms for each parameter in a\_db.

See also: params\_tests\_profile (p. 136)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/20

# A.19.13 Method params\_tests\_db/makeGenesisParFile

**Summary:** Creates a Genesis parameter file with all the parameter values in a\_db.

### Usage:

```
makeGenesisParFile(a_db, filename, props)
```

**Description:** For each a\_db row, print the parameter names in a file formatted for Genesis.

#### **Parameters:**

```
a_db: A params_tests_db object.
```

**filename:** Genesis parameter file to be created.

**props:** A structure with any optional properties.

trialStart: If given, adds/replaces the trial parameter and counts forward.

#### **Returns:**

nothing.

# **Example:**

```
» blocked_rows_db = makeModifiedParamDB(ranked_for_gps0501a_db, 1, [1, 2], 10,
[-100 100]);

» makeGenesisParFile(blocked_rows_db, 'blocked_gps0501-03.par')
```

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See also: makeModifiedParamDB (p.??), scanParamAllRows (p.??), scaleParamsOneRow (p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/03/13

# A.19.14 Method params\_tests\_db/rankVsAllDB

**Summary:** Generates ranking DBs by comparing rows of a\_db with each row of to\_db.

### Usage:

```
tex_string = rankVsAllDB(a_db, to_db, a_dataset, to_dataset)
```

**Description:** Distance is each measure difference divided by the STD in to\_db, squared and summed. Returned DB contains only the selected to\_tests and the parameters from initial DB.

#### **Parameters:**

**a\_db:** A params\_tests\_db object to compare rows from.

to\_db: A tests\_db object to compare it with.

a\_dataset: Dataset for a\_db.

to\_dataset: Dataset for crit\_db.

### **Returns:**

ranked\_dbs: Array of created DBs with original rows and a distance measure, in ascending order. tex\_string: A LaTeX string for all tables created.

```
See also: rankVsDB(p.??), matchingRow(p.??), rankMatching(p.??), joinRows(p.??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/10

### A.19.15 Method params\_tests\_db/addParams

**Summary:** Inserts new parameter columns to tests db.

#### **Usage:**

```
obj = addParams(obj, param_names, param_columns)
```

**Description:** Adds new columns to the database and returns the new DB. This operation is expensive in the sense that the whole database matrix needs to be enlarged just to add a single new column. The method of allocating a matrix, filling it up, and then providing it to the tests\_db constructor is the preferred method of creating tests\_db objects. This method may be used for measures obtained by operating on raw measures.

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#### **Parameters:**

**obj:** A tests\_db object.

param\_names: A cell array of param names to be added.
param columns: Data matrix of columns to be added.

#### **Returns:**

obj: The tests\_db object that includes the new columns.

See also: allocateRows (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/10/11

# A.19.16 Method params\_tests\_db/mergeMultipleCIPsInOne

**Summary:** Merges multiple rows with different CIP data into one, generating a database of one row per neuron.

#### **Usage:**

```
a_db = mergeMultipleCIPsInOne(db, names_tests_cell, index_col_name)
```

**Description:** It calls invarParam to separate db into pages with different CIP level data. Then uses the names\_tests\_cell to choose tests from each page to be merged into the final database row. The tests will be suffixed with the field name so that they can be distinguished. RowIndex columns will be automatically included, and one of them can be chosen with index\_col\_name that has values for all cells. The suffixed for needs to be used to choose index\_col\_name, such as 'RowIndex\_H100pA', assuming 'H100pA' was the field name in names\_tests\_cell that corresponds to page -100 pA.

### **Parameters:**

**db:** A params\_tests\_db object.

names\_tests\_cell: A cell array alternating suffix names and test column
vectors.

The order of names correspond to each unique CIP level in db, with increasing order.

index\_col\_name: (Optional) Name of row index column
(default is 'RowIndex' suffixed with the first field name).

### **Returns:**

a\_db: A params\_tests\_db object of organized values.

### **Example:**

```
» control_phys_sdb =
mergeMultipleCIPsInOne(control_phys_db,
struct('_H100pA', [1:10], '_D100pA', [1:10 16:18]),
'RowIndex_H100pA')

See also: invarValues (p. ??), tests_3D_db (p. 212), corrCoefs (p. ??),
```

tests\_3D\_db/plotVarBox (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/01/13

# A.19.17 Method params\_tests\_db/subsref

Summary: Defines generic indexing for objects.

# A.19.18 Method params\_tests\_db/paramsParamsCoefs

**Summary:** Calculates a corrcoefs\_db for each param from correlations of variant params and invariant param coefs and collects them in a cell array.

#### Usage:

```
pp_coefs = paramsParamsCoefs(a_db, p_t3ds, p_coefs)
```

**Description:** Skips the 'ItemIndex' test.

### **Parameters:**

a\_db: A tests\_db object.

**p\_t3ds:** Cell array of invariant parameter databases.

**p\_coefs:** Cell array of tests coefficients for each parameter.

# **Returns:**

pp\_coefs: A cell array of corrcoefs\_dbs for each param combination in a\_db.

See also: params\_tests\_profile (p. 136)

# A.19.19 Method params\_tests\_db/displayRankingsTeX

**Summary:** Generates and displays a ranking DB by comparing rows of a\_db with the given match criteria.

### Usage:

```
tex_string = displayRankingsTeX(a_db, crit_db, props)
```

**Description:** Generates a LaTeX document with: - Values of 10 best matching a\_db rows in a floating table. - (optional) Raw traces compared with some best matches at different distances - Parameter distributions of 50 best matches as a bar graph.

#### **Parameters:**

**a\_db:** A params\_tests\_db object to compare rows from.

**crit\_db:** A tests\_db object holding the match criterion tests and STDs which can be created with matchingRow.

**props:** A structure with any optional properties.

**caption:** Identification of the criterion db (not needed/used?).

**a\_dataset:** Dataset for a\_db.

**a\_dball:** The non-joined DB for for a\_db.

crit\_dataset: Dataset for crit\_db.

crit\_dball: Dataset for crit\_db.

num\_matches: Number of best matches to display (default=10).
rotate: Rotation angle for best matches table (default=90).

Returns:

tex\_string: LaTeX document string.

See also: rankVsDB (p. ??), displayRowsTeX (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/20

### A.19.20 Method params tests db/getParamRowIndices

**Summary:** Returns indices of rows with matching parameter values from rows of this db.

# **Usage:**

```
row_indices = getParamRowIndices(a_db, rows, to_db)
```

### **Parameters:**

**a\_db:** A params\_tests\_db object.

rows: rows to find indices for.

to\_db: Where to find the matching rows.

#### **Returns:**

row\_indices: Array of row indices.

See also: makeModifiedParamDB (p.??), scanParamAllRows (p.??), scaleParamsOneRow (p.??), makeGenesisParFile (p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/01/14

### A.19.21 Method params\_tests\_db/plotParamsHists

**Summary:** Create a horizontal plot\_stack of parameter histograms.

### **Usage:**

```
a_ps = plotParamsHists(a_db, title_str, props)
```

**Description:** Skips the 'ItemIndex' test.

#### **Parameters:**

**a\_db:** A params\_tests\_db object.

title\_str: (Optional) A string to be concatanated to the title.

**props:** A structure with any optional properties.

quiet: Do not display the DB id on the plot title.

barAxisProps: passed to plotEqSpaced for each bar axis.

#### **Returns:**

a\_ps: A horizontal plot\_stack of plots

See also: plot\_stack (p. 164), paramsHists (p. ??), plotEqSpaced (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/07

### A.19.22 Method params\_tests\_db/rankVsDB

**Summary:** Generates a ranking DB by comparing rows of this db with the given test criteria.

#### Usage:

```
a_ranked_db = rankVsDB(a_db, crit_db)
```

**Description:** Distance is each measure difference divided by the STD in to\_db, squared and summed. Returned DB contains only the selected tests from crit\_db and the parameters from initial a\_db.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2607-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

#### **Parameters:**

**a\_db:** A params\_tests\_db object to compare rows from.

**crit\_db:** A tests\_db object holding the match criterion tests and STDs which can be created with matchingRow.

#### **Returns:**

a\_ranked\_db: The created DB with original rows and a distance measure, in ascending order.

See also: matchingRow (p. ??), rankMatching (p. ??), joinRows (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/20

# A.19.23 Method params\_tests\_db/delColumns

Summary: Deletes columns from tests\_db.

Usage:

```
index = delColumns(obj, tests)
```

**Description:** Overloaded function that maintains correct number of parameters. See original tests\_db/delColumns.

### **Parameters:**

obj: A tests\_db object.

tests: Numbers or names of tests (see tests2cols)

#### **Returns:**

obj: The tests\_db object that is missing the columns.

See also: tests\_db/delColumns (p. 245)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/10/11

# A.19.24 Method params\_tests\_db/paramsCoefs

Summary: Calculates a corrcoefs\_db for each param and collects them in a cell array.

**Usage:** 

```
p_coefs = paramsCoefs(a_db, p_t3ds)
```

**Description:** Skips the 'ItemIndex' test.

### **Parameters:**

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```
a_db: A tests_db object.
```

p\_t3ds: Cell array of invariant parameter databases.

#### **Returns:**

p coefs: A cell array of corrcoefs dbs for each param in a db.

```
See also: params_tests_profile (p. 136)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/17

# A.19.25 Method params\_tests\_db/getProfile

**Summary:** Create a profile object from a params\_tests\_db by collecting statistics.

# Usage:

```
a_pt_profile = getProfile(a_db, props)
```

**Description:** Calculates the following results items: idx: Name-index pairs for accessing results arrays. t\_hists: Cell array of histograms of each test. p\_t3ds: Cell array of invariant relations of each parameter with all tests. pt\_hists: Cell array of separate test value histograms for uniques value of each parameter. p\_stats: Cell array of test stats for each param. p\_coefs: Cell array of correlation coefficients for each parameter with all tests. pt\_coefs\_hists: Cell matrix of histograms of coefficients from correlations of each parameter with each test. pp\_coefs: Cell 3D matrix of mean coefficients from correlations of each parameter with correlation coefficients of each parameter with each test.

#### **Parameters:**

```
a_db: A params_tests_db object.
```

**props:** A structure with any optional properties.

#### Returns a params\_tests\_profile object.

```
See also: params_tests_profile (p. 136), results_profile (p. 176), params_tests_db (p. 116), params_tests_fileset (p. 132), tests_db (p. 217), tests_3D_db (p. 212), histogram_db (p. 94), stats_db (p. 206), corrcoefs_db (p. 81)
```

# A.19.26 Method params\_tests\_db/plotVarBoxMatrix

**Summary:** Create a stack of parameter-test variation plots organized in a matrix.

Usage:

```
a_plot_stack = plotVarBoxMatrix(a_db, p_t3ds)
```

**Description:** Skips the 'ItemIndex' test.

**Parameters:** 

a\_db: A tests\_db object.

p\_t3ds: Cell array of invariant parameter databases.

**Returns:** 

a\_plot\_stack: A plot\_stack with the plots organized in matrix form

See also: params\_tests\_profile (p. 136), plotVar (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/17

### A.19.27 Method params\_tests\_db/invarParams

Summary: Calculates invariant param dbs for all parameters and returns in an array.

Usage:

```
p_t3ds = invarParams(a_db)
```

**Description:** Skips the 'ItemIndex' test.

**Parameters:** 

a\_db: A tests\_db object.

**Returns:** 

p\_t3ds: An array of tests\_3D\_dbs for each param in a\_db.

See also: params\_tests\_profile (p. 136)

# A.19.28 Method params\_tests\_db/getDualCIPdb

**Summary:** Generates a database by merging selected tests of depolarizing and hyperpolarizing cip results.

#### **Usage:**

```
a_db = getDualCIPdb(db, depol_tests, hyper_tests, depol_suffix, hyper_suffix)
```

Description: depol\_tests need to have the RowIndex column in it.

#### **Parameters:**

db: A params\_tests\_db object.

#### **Returns:**

a\_db: A params\_tests\_db object of organized values.

### **Example:**

```
\mbox{\sc w} control_phys_sdb = getDualCIPdb(control_phys_db, depol_tests, hyper_tests, ", 'Hyp100pA') where depol_tests and hyper_tests are cell arrays of selected tests.
```

```
See also: invarValues (p. ??), tests_3D_db (p. 212), corrCoefs (p. ??), tests_3D_db/plotPair (p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/01/13

# A.19.29 Method params\_tests\_db/scanParamAllRows

**Summary:** Scans given parameter range for each row in DB.

### **Usage:**

```
a_params_db = scanParamAllRows(a_db, param, min_val, max_val, num_levels, props)
```

**Description:** Produces rows by replacing the desired parameter value, in all rows of DB, with num\_levels values between the given boundaries, min\_val and max\_val. This results in a DB with num\_levels times more rows than the original DB. Then, makeGenesisParFile can be used to generate a parameter file from this DB to drive new simulations.

### **Parameters:**

```
    a_db: A params_tests_db object whose first row is subject to modifications.
    param: The parameter to be varied (see tests2cols for param description).
    min_val, max_val: The low and high boundaries for the parameter value.
    num levels: Number of levels to produce, including the boundaries.
```

**props:** A structure with any optional properties.

renameTrial: If given, the 'trial' column is renamed to this name.

levelFunc: Use this function to get the parameter range with

feval(levelFunc, min\_val, max\_val, num\_levels). Example: 'logLevels'

### **Returns:**

a\_params\_db: A db only with params.

### **Example:**

```
Sets NaF to given range with 100 levels:
» naf_rows_db = scanParamAllRows(a_db(desired_rows, :), 'NaF', 0, 1000, 100);
```

See also: makeGenesisParFile (p.??), scaleParamsOneRow (p.??), ranked\_db/blockedDistances (p. 171), getParamRowIndices (p. ??), logLevels (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/16

# A.19.30 Method params\_tests\_db/scaleParamsOneRow

**Summary:** Scales chosen parameters in a row by multiplying with levels to create a new parameter db with as many rows as values in levels.

### Usage:

```
a_params_db = scaleParamsOneRow(a_db, params, levels)
```

**Description:** Produces rows by multiplying desired params, in the first row of DB, with each value in levels. Then, makeGenesisParFile can be used to generate a parameter file from this DB to drive new simulations.

#### **Parameters:**

**a\_db:** A params\_tests\_db object whose first row is subject to modifications.

params: Parameters to be varied (see tests2cols for param description).

**levels:** Column vector of parameter value multipliers (1=unity).

#### **Returns:**

a\_params\_db: A db only with params.

### **Example:**

```
Blocks NaF from 0» naf_rows_db = scanOneParam(a_db(desired_row, :), 'NaF',
0:0.1:1);
```

See also: ranked\_db/blockedDistances (p. 171), getParamRowIndices (p. ??), makeGenesisParFile (p. ??)

# A.20 Class params\_tests\_fileset

#### A.20.1 Constructor params\_tests\_fileset/params\_tests\_fileset

**Summary:** Description of a set of data files of raw data varying with parameter values. **Usage:** 

```
obj = params_tests_fileset(file_pattern, dt, dy, id, props)
```

**Description:** This is a subclass of params\_tests\_dataset. This class is used to generate params\_tests\_db objects and keep a connection to the raw data files. This class only keeps names of files and loads raw data files whenever it's requested. A database object can easily be generated using the convertion methods. Most methods defined here can be used as-is, however some should be overloaded in subclasses. The specific methods are loadItemProfile.

#### **Parameters:**

**file\_pattern:** File pattern, or cell array of patterns, matching all files to be loaded.

dt: Time resolution [s]

dy: y-axis resolution [ISI (V, A, etc.)]

id: An identification string

**props:** A structure with any optional properties.

**num\_params:** Number of parameters that appear in filenames.

param\_trial\_name: Use this name on the filename as the 'trial' parameter.

**param\_row\_filename:** If given, the 'trial' parameter will be used to address rows from this file and acquire parameters.

param\_desc\_filename: Contains the parameter range descriptions
 one per

each row. The parameter names are acquired from this file.

param\_names: Cell array of parameter names corresponding to the param\_row\_filename columns can be specified as an alternative to specifying param\_desc\_filename. These names are not for the parameters present in the data filename.

profile\_method\_name: It can be one of the profile-creating methods
 in this

class. E.g., 'trace\_profile', 'srp\_trace\_profile', etc. (See parent classes and cip\_trace object for more props)

### Returns a structure object with the following fields:

params\_tests\_dataset, path: The pathname to files.

See also: params\_tests\_db (p. 116), tests\_db (p. 217), test\_variable\_db (N/I) (p.  $\ref{prop:loop}$ )

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Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/09

### A.20.2 Method params\_tests\_fileset/addFiles

Summary: Adds to existing list of files in set.

Usage:

[a\_fileset, index\_list] = addFiles(a\_fileset, file\_pattern, props)

#### **Parameters:**

a\_fileset: A params\_tests\_fileset object.

**file\_pattern:** File pattern, or cell array of patterns, matching additional files.

**props:** A structure with any optional properties.

param\_row\_filename: Update parameters from here. The 'trial' parameter is used to address rows from this file and acquire parameters.

#### **Returns:**

a\_fileset: The augmented fileset object. index\_list: The vector of index numbers of the new files added. Can be used to selectively load the new files into a DB using params\_test\_db.

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/01

### A.20.3 Method params\_tests\_fileset/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.20.4 Method params\_tests\_fileset/get

Summary: Defines generic attribute retrieval for objects.

# A.20.5 Method params\_tests\_fileset/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

### A.20.6 Method params\_tests\_fileset/trace

**Summary:** Loads a raw trace given a file\_index to this fileset.

**Usage:** 

```
a_trace = trace(fileset, file_index)
```

#### Parameters:

fileset: A params\_tests\_fileset.
file index: Index of file in fileset.

#### **Returns:**

a\_trace: A trace object.

See also: trace (p. 255), params\_tests\_fileset (p. 132)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/13

### A.20.7 Method params\_tests\_fileset/paramNames

**Summary:** Returns the ordered names of parameters for this fileset.

### **Usage:**

```
param_names = paramNames(fileset, item)
```

**Description:** Looks at the filename of the first file to find the parameter names.

### **Parameters:**

fileset: A params\_tests\_fileset.

**item:** (Optional) If given, read param names by loading item at this index.

### **Returns:**

params\_names: Cell array with ordered parameter names.

See also: params\_tests\_fileset (p. 132), paramNames (p. ??), testNames (p. ??)

# A.20.8 Method params\_tests\_fileset/getItemParams

**Summary:** Get the parameter values of a dataset item.

**Usage:** 

```
params_row = getItemParams(dataset, index)
```

#### **Parameters:**

dataset: A params\_tests\_dataset.
index: Index of item in dataset.

#### **Returns:**

params\_row: Parameter values in the same order of paramNames

See also: itemResultsRow (p.??), params\_tests\_dataset (p. 111), paramNames (p.??), testNames (p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/03

# A.20.9 Method params\_tests\_fileset/trace\_profile

**Summary:** Loads a raw trace\_profile given a file\_index to this fileset.

Usage:

```
a_trace_profile = trace_profile(fileset, file_index)
```

# Parameters:

fileset: A params\_tests\_fileset.
file\_index: Index of file in fileset.

# **Returns:**

a\_trace\_profile: A trace\_profile object.

See also: trace\_profile (p. 264), params\_tests\_fileset (p. 132)

# A.20.10 Method params\_tests\_fileset/loadItemProfile

**Summary:** Loads a profile object from a raw data file in the fileset.

Usage:

```
a_profile = loadItemProfile(fileset, file_index)
```

**Description:** Subclasses should overload this function to load the specific profile object they desire. The profile class should define a getResults method which is used in the itemResultsRow method.

#### **Parameters:**

fileset: A params\_tests\_fileset.
file\_index: Index of file in fileset.

#### **Returns:**

a\_profile: A profile object that implements the getResults method.

```
See also: itemResultsRow(p.??), params_tests_fileset(p.132), paramNames(p.??), testNames(p.??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.21 Class params\_tests\_profile

# A.21.1 Constructor params\_tests\_profile/params\_tests\_profile

**Summary:** Holds the results profile from a params\_tests\_db.

Usage:

```
a_pt_profile = params_tests_profile(results, a_db, props)
```

### **Parameters:**

**a\_db:** A params\_tests\_db object.

results: A structure containing test results.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

results\_profile: Contains results of tests. db: The params\_tests\_db. props.

See also: results\_profile(p. 176), params\_tests\_db/params\_tests\_profile(p. ??)

# A.21.2 Method params\_tests\_profile/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.22 Class period

# A.22.1 Constructor period/period

**Summary:** Start and end times of a period in terms of the dt of the trace to which belongs.

### **Usage:**

```
obj = period(start_time, end_time)
```

#### **Parameters:**

(see below for the rest)

# Returns a structure object with the following fields:

start\_time, end\_time: Inclusive period [dt].

**See also:** trace (p. 255), spikes (p. 195), spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

# A.22.2 Method period/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.22.3 Method period/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.22.4 Method period/set

**Summary:** Generic method for setting object attributes.

# A.22.5 Method period/subsref

**Summary:** Defines generic indexing for objects.

# A.22.6 Method period/SpikeTimesinPeriod

#### Usage:

SpkTimes=Interval(times, period)

#### **Parameters:**

times: an array of spike times.

period: A period object

#### **Returns:**

the\_period: The cropped set of spike times that fall within a period.

See also: period (p. 137), cip\_trace (p. 54), trace (p. 255), spikes (p. 195)

Author: Tom Sangrey, 2006/01/26

# A.23 Class physiol\_bundle

### A.23.1 Constructor physiol\_bundle/physiol\_bundle

**Summary:** The physiology dataset and the DB created from it bundled together.

### Usage:

```
a_bundle = physiol_bundle(a_dataset, a_db, a_joined_db, props)
```

**Description:** This is a subclass of dataset\_db\_bundle, specialized for physiology datasets.

#### **Parameters:**

- a\_dataset: A physiol\_cip\_traceset\_fileset object.
- a\_db: The raw params\_tests\_db object created from the dataset.
  It only needs to have the pAcip, pAbias, TracesetIndex, and ItemIndex columns.
- **a\_joined\_db:** The one-treatment-per-line DB created from the raw DB.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

dataset\_db\_bundle, joined\_control\_db: DB of control neurons (no pharmacological applications).

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See also: dataset\_db\_bundle(p.82), tests\_db(p.217), params\_tests\_dataset (p.111)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/13

# A.23.2 Method physiol\_bundle/getNeuronLabel

Summary: Constructs the neuron label from dataset.

Usage:

a\_label = getNeuronLabel(a\_bundle, traceset\_index, props)

#### **Parameters:**

**a\_bundle:** A physiol\_cip\_traceset\_fileset object. **traceset\_index:** The traceset index of neuron. **props:** A structure with any optional properties.

#### **Returns:**

a\_label: A string label identifying selected neuron in bundle.

See also: dataset\_db\_bundle (p. 82)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/05

# A.23.3 Method physiol\_bundle/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

### A.23.4 Method physiol\_bundle/set

Summary: Generic method for setting object attributes.

# A.23.5 Method physiol\_bundle/constrainedMeasuresPreset

**Summary:** Returns a physiol\_bundle with constrained measures according to chosen preset.

### **Usage:**

```
[a_pbundle test_names] = constrainedMeasuresPreset(a_pbundle, preset, props)
```

#### **Parameters:**

```
a_pbundle: A physiol_cip_traceset_fileset object.
preset: Choose preset measure list (default=1).
props: A structure with any optional properties.
```

#### **Returns:**

a\_pbundle: One or more cip\_trace object that holds the raw data.

```
See also: loadItemProfile (p. ??), physiol_cip_traceset/cip_trace (p. 149)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/19

### A.23.6 Method physiol\_bundle/matchingRow

**Summary:** Creates a criterion database for matching the neuron at traceset index.

#### **Usage:**

```
a_crit_db = matchingRow(p_bundle, traceset_index, props)
```

**Description:** Copies selected test values from row as the first row into the criterion db. Adds a second row for the STD of each column in the db.

#### **Parameters:**

```
p_bundle: A physiol_bundle object.traceset_index: A TracesetIndex of the neuron and treatments to match.props: A structure with any optional properties.
```

# **Returns:**

a\_crit\_db: A tests\_db with two rows for values and STDs.

### **Example:**

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```
See also: rankMatching (p. ??), tests_db/matchingRow (p. 233)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/21
       A.23.7 Method physiol_bundle/plotfICurveStats
       Summary: Generates a f-I curve mean-std plot of physiology DB.
       Usage:
a_plot = plotfICurveStats(p_bundle, title_str, props)
       Parameters:
            p_bundle: A physiol_bundle object.
            title_str: (Optional) String to append to plot title.
            props: A structure with any optional properties.
                 quiet: if given, no title is produced
                     (passed to plot_superpose)
       Returns:
            a_plot: An f-I curve plot.
       Example:
» plotFigure(plotfICurveStats(pbundle));
       See also: dataset_db_bundle/plotfICurve(p. 86), plot_abstract(p. 153),
            plot_superpose (p. 167)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/16
       A.23.8 Method physiol_bundle/getNeuronRowIndex
       Summary: Returns the neuron index from bundle.
       Usage:
a_row_index = getNeuronRowIndex(a_bundle, traceset_index, props)
       Parameters:
            a_bundle: A physiol_bundle object.
            traceset_index: The TracesetIndex number of neuron, or a DB row con-
                 taining this.
            props: A structure with any optional properties.
```

#### **Returns:**

a\_row\_index: A row index of neuron in a\_bundle.joined\_db.

See also: dataset\_db\_bundle/getNeuronRowIndex (p. 87)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/09

# A.23.9 Method physiol\_bundle/bestMatchAllNeurons

Summary: Finds the best match among given database for each physiology neuron.

Usage:

```
all_ranks_db = bestMatchAllNeurons(p_bundle, joined_db, props)
```

**Description:** Returns a database of best matching entries from joined\_db for each entry in p\_bundle.joined\_control\_db.

#### **Parameters:**

```
p_bundle: A physiol_bundle object.
```

joined\_db: A database with neuron representations to rank against neurons.

**props:** A structure with any optional properties. (passed to rankMatching)

### **Returns:**

all\_ranks\_db: DB of best matching from joined\_db. Each row corresponds to p\_bundle.joined\_control\_db rows.

### **Example:**

```
» all_ranks_db = ...
bestMatchAllNeurons(constrainedMeasuresPreset(pbundle2, 6),
mbundle_maxcond.joined_db)
» plotXRows(all_ranks_db, 'Distance', 'maxcond DB distance per neuron',
'maxcond', ...
struct('LineStyle', '-', 'quiet', 1, 'PaperPosition', [0 0 4 3]))
```

See also: tests\_db/rankMatching(p. 254), tests\_db/matchingRow(p. 233)

# A.23.10 Method physiol\_bundle/ctFromRows

**Summary:** Loads a cip\_trace object from a raw data file in the a\_pbundle.

#### Usage:

```
a_cip_trace = ctFromRows(a_pbundle, a_db|traceset_idx, cip_levels, props)
```

#### **Parameters:**

```
a_pbundle: A physiol_cip_traceset_fileset object.
```

**a\_db:** A DB created by this fileset to read the traceset indices from.

traceset\_idx: A column vector with traceset indices.

cip\_levels: A column vector of CIP-levels to be loaded.

**props:** A structure with any optional properties.

traces: column vector of trace indices to load.

**showParamsList:** Cell array of params or treatments to include in the id field.

#### **Returns:**

a cip trace: One or more cip trace object that holds the raw data.

```
See also: loadItemProfile (p. ??), physiol_cip_traceset/cip_trace (p. 149)
```

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2005/07/13

### A.23.11 Method physiol\_bundle/matchingControlNeuron

**Summary:** Creates a criterion database for matching the neuron at traceset\_index.

#### Usage:

```
a_crit_bundle = matchingControlNeuron(a_bundle, neuron_id, props)
```

**Description:** Copies selected test values from row as the first row into the criterion db. Adds a second row for the STD of each column in the db.

### **Parameters:**

a\_bundle: A physiol\_bundle object.

**neuron\_id:** A NeuronId of the neuron to match.

**props:** A structure with any optional properties.

#### **Returns:**

a\_crit\_bundle: A tests\_db with two rows for values and STDs.

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# **Example:**

```
Matches gpd0421c from cip_traces_all_axoclamp.txt:
    a_crit_bundle = matchingControlNeuron(pbundle, 33)
    (see example in matchingRow)
```

See also: rankMatching (p. ??), tests\_db (p. 217), tests2cols (p. ??)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2005/12/21

# A.24 Class physiol\_cip\_traceset

# A.24.1 Constructor physiol\_cip\_traceset/physiol\_cip\_traceset

**Summary:** Dataset of cip traces from same PCDX file.

Usage:

obj = physiol\_cip\_traceset(trace\_str, data\_src, chaninfo, dt, dy, treatments, id,
props);

**Description:** This is a subclass of params\_tests\_dataset. Each trace varies in bias, pulse times and cip magnitude.

### **Parameters:**

trace\_str: Trace list in the format for loadtraces or just a Matlab vector.

data src: Absolute path of PCDX data source.

chaninfo: 4-element array containing vchan, ichan, vgain, igain

vchan, ichan: Current and voltage channels.

**vgain, igain:** External gain factors for voltage channel and current channel (vgain does NOT include the 10X amplification from the Axoclamp, so vgain = 1 would mean no additional amplification beyond the 10X.)

dt: Time resolution [s].

dy: Y-axis resolution [V] or [A].

**treatments:** Structure containing the names and concentrations of compounds.

id: Neuron name.

**props:** A structure with any optional properties.

nsHDF5: If 1, source is a NeuroSAGE HDF5 file.

**profile\_class\_name:** Use this profile class (Default: 'cip\_trace\_profile').

**cip\_list:** Vector of cip levels to which the current trace will be matched. (All other props are passed to cip\_trace objects)

PANDORA Toolbox: Prog.'s Mabraft v805 of 2407-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

# Returns a structure object with the following fields:

params\_tests\_dataset, data\_src, ichan, vchan, vgain, igain, treatments, id.

```
See also: cip_traces (p.??), params_tests_dataset (p.111), params_tests_db (p.116)
```

Author: Cengiz Gunay <cgunay@emory.edu> and Thomas Sangrey, 2005/01/17

## A.24.2 Method physiol\_cip\_traceset/setProp

Summary: Generic method for setting optional object properties.

**Usage:** 

```
obj = setProp(obj, prop1, val1, prop2, val2, ...)
```

**Description:** Modifies or adds property values. As many property name-value pairs can be specified.

### **Parameters:**

obj: Any object that has a props field.

attr: Property nameval: Property value.

### **Returns:**

obj: The new object with the updated properties.

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/22

# A.24.3 Method physiol\_cip\_traceset/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.24.4 Method physiol\_cip\_traceset/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.24.5 Method physiol\_cip\_traceset/cip\_trace\_profile

Summary: Loads a cip\_trace\_profile object from a raw data file in the traceset.

Usage:

```
a_profile = cip_trace_profile(traceset, trace_index)
```

#### **Parameters:**

traceset: A physiol\_cip\_traceset object.
trace\_index: Index of file in traceset.

#### **Returns:**

a\_profile: A profile object that implements the getResults method.

```
See also: itemResultsRow (p.??), params_tests_fileset (p. 132), paramNames (p.??), testNames (p.??)
```

Author: Cengiz Gunay <cgunay@emory.edu> and Thomas Sangrey, 2005/01/18

## A.24.6 Method physiol\_cip\_traceset/subsref

**Summary:** Defines generic indexing for objects.

## A.24.7 Method physiol\_cip\_traceset/CIPform

**Summary:** Extracts current bias and pulse information from the current channel.

Usage:

```
[ciptype, on, off, finish, bias, pulse] = ns_CIPform(traceset,trace_index)
```

# **Parameters:**

traceset: A physiol\_cip\_traceset object.
trace\_index: Index of item in traceset

See also: cip\_traces (p.??), params\_tests\_dataset (p.111), params\_tests\_db (p.116)

Author: Thomas Sangrey, 2005

## A.24.8 Method physiol\_cip\_traceset/paramNames

**Summary:** Returns the parameter names for this traceset.

Usage:

```
param_names = paramNames(traceset)
```

**Description:** Looks at the filename of the first file to find the parameter names.

**Parameters:** 

```
traceset: A params_tests_dataset.
```

**Returns:** 

param\_names: Cell array with ordered parameter names.

```
See also: params_tests_dataset (p. 111), paramNames (p. ??), testNames (p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/06

# A.24.9 Method physiol\_cip\_traceset/getItemParams

**Summary:** Get the parameter values of a dataset item.

Usage:

```
params_row = getItemParams(dataset, index, a_profile)
```

#### **Parameters:**

```
dataset: A params_tests_dataset.
index: Index of item in dataset.
a_profile: cip_trace_profile object
```

#### **Returns:**

params\_row: Parameter values in the same order of paramNames

```
See also: itemResultsRow (p.??), params_tests_dataset (p. 111), paramNames (p.??), testNames (p.??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/06

## A.24.10 Method physiol\_cip\_traceset/itemResultsRow

**Summary:** Processes a raw data file from the dataset and return its parameter and test values.

#### **Usage:**

```
[params_row, tests_row] = itemResultsRow(dataset, index)
```

**Description:** This method is designed to be reused from subclasses as long as the loadItemProfile method is properly overloaded. Adds an Index column to the DB to keep track of raw data items after shuffling.

#### **Parameters:**

dataset: A params\_tests\_dataset.
index: Index of file in dataset.

#### **Returns:**

params\_row: Parameter values in the same order of paramNames tests\_row: Test values in the same order with testNames

See also: loadItemProfile (p.??), params\_tests\_dataset (p. 111), paramNames (p. ??), testNames (p. ??)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/09/10

### A.24.11 Method physiol\_cip\_traceset/loadItemProfile

**Summary:** Loads a cip\_trace\_profile object from a raw data file in the traceset.

#### **Usage:**

```
a_profile = loadItemProfile(traceset, trace_index)
```

### Parameters:

traceset: A physiol\_cip\_traceset object.
trace\_index: Index of file in traceset.

### **Returns:**

a\_profile: A profile object that implements the getResults method.

See also: itemResultsRow (p.??), params\_tests\_fileset (p. 132), paramNames (p.??), testNames (p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

## A.24.12 Method physiol\_cip\_traceset/cip\_trace

**Summary:** Loads a cip\_trace object from a raw data file in the traceset.

#### **Usage:**

```
a_cip_trace = cip_trace(traceset, trace_index, props)
```

#### **Parameters:**

traceset: A physiol\_cip\_traceset object.
trace\_index: Index of file in traceset.

**props:** A structure with any optional properties.

**showParamsList:** Cell array of params to add to id field. **showName:** Show the name of the cell in the id field (default=1).

TracesetIndex: Indicates in the id field.

#### **Returns:**

a\_cip\_trace: A cip\_trace object that holds the raw data.

```
See also: itemResultsRow(p.??), params_tests_fileset(p.132), paramNames (p.??), testNames(p.??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/07/13

# A.25 Class physiol cip traceset fileset

### A.25.1 Constructor physiol\_cip\_traceset\_fileset/physiol\_cip\_traceset\_fileset

Summary: Physiological fileset of traceset objects (concatenated).

### **Usage:**

```
obj = physiol_cip_traceset_fileset(traceset_items, dt, dy, props)
```

Description: This is a subclass of params\_tests\_dataset. It contains a set of physiol\_cip\_traceset items that are tied to physical data sources. Each traceset can load a set of traces for an experimental recording. Most flexible usage is obtained when the input traceset\_items is given as a cell array of physiol\_cip\_traceset objects. These objects can each link to PCDX or NeuroSAGE HDF5 files independent of each other. A regular Matlab script can be used to create such a cell array. If a function is defined to return such an array, it can be passed as traceset\_items. Alternatively, the cell array can be constructed from an ASCII file as described below, such as for deprecated PCDX data files.

### **Parameters:**

traceset\_items: It can be a function handle, cell array or filename

string. Function should return a cell array of physiol\_cip\_traceset items. Finally this cell array can be provided directly. If it is an ASCII filename, then it should contain the following tab-delimited items: 1. Neuron ID (name to associate with the neuron). If left blank, use the filename with the 'all' extension removed. 2. The absolute path of the data file 3. The trace numbers to load, space-delimited (e.g. 1-21 24 26 27) 4. Vchan: voltage channel number 5. Ichan: current channel number 6. Vgain: external gain on voltage channel IN ADDITION to the 10X that automatically comes from the Axoclamp 2B. 7. Igain: external gain on current channel. 8. Pairs of condition names and molar concentrations in any order e.g.: TTX 1e-8 apamin 2e-7 picrotoxin 1e-4

### Returns a structure object with the following fields:

neuron\_idx: A structure that points from neuron names to NeuronId numbers. params\_tests\_dataset

```
See also: physiol_cip_traceset (p. 144), params_tests_dataset (p. 111), params_tests_db (p. 116)
```

Author: Cengiz Gunay <cgunay@emory.edu> and Thomas Sangrey, 2005/01/17

# A.25.2 Method physiol\_cip\_traceset\_fileset/setProp

**Summary:** Generic method for setting optional object properties.

**Usage:** 

```
obj = setProp(obj, prop1, val1, prop2, val2, ...)
```

**Description:** Modifies or adds property values. As many property name-value pairs can be specified.

## **Parameters:**

**obj:** Any object that has a props field.

attr: Property nameval: Property value.

## **Returns:**

obj: The new object with the updated properties.

### See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/22

## A.25.3 Method physiol\_cip\_traceset\_fileset/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

## A.25.4 Method physiol\_cip\_traceset\_fileset/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.25.5 Method physiol\_cip\_traceset\_fileset/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

### A.25.6 Method physiol\_cip\_traceset\_fileset/loadItemProfile

Summary: Loads a cip\_trace\_profile object from a raw data file in the fileset.

Usage:

a\_profile = loadItemProfile(fileset, traceset\_index, trace\_index)

### **Parameters:**

fileset: A physiol\_cip\_traceset object.

**traceset\_index**: Index of traceset item in this fileset (corresponds to row in cells\_filename) to use grab the cell information.

trace index: Index of item in the traceset.

### **Returns:**

a\_profile: A profile object that implements the getResults method.

See also: itemResultsRow(p.??), params\_tests\_fileset(p.132), paramNames(p.??), testNames(p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14 and Tom Sangrey

## A.25.7 Method physiol\_cip\_traceset\_fileset/cip\_trace

**Summary:** Loads a cip\_trace object from a raw data file in the fileset.

#### **Parameters:**

**fileset:** A physiol\_cip\_traceset\_fileset object.

traceset\_index: Index of traceset item in this fileset (corresponds

to row in cells\_filename) to find the cell information.

trace\_index: Index of item in the traceset.

**a\_db:** A DB created by this fileset to read the traceset and item indices from.

**props:** A structure with any optional properties, passed to physiol\_cip\_traceset/cip\_trace.

#### **Returns:**

a\_cip\_trace: One or more cip\_trace object that holds the raw data.

See also: loadItemProfile (p. ??), physiol\_cip\_traceset/cip\_trace (p. 149)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/07/13

### A.25.8 Method physiol\_cip\_traceset\_fileset/readDBItems

**Summary:** Reads all items to generate a params\_tests\_db object.

#### **Usage:**

```
[params, param_names, tests, test_names] = readDBItems(obj, items)
```

**Description:** This is a specific method to convert from physiol\_cip\_traceset\_fileset to a params\_tests\_db, or a subclass. Outputs of this function can be directly fed to the constructor of a params\_tests\_db or a subclass.

### **Parameters:**

obj: A physiol\_cip\_traceset\_fileset

items: (Optional) List of item indices to use to create the db.

### **Returns:**

```
params, param_names, tests, test_names: See params_tests_db.
```

See also: params\_tests\_db (p. 116), params\_tests\_fileset (p. 132), itemResultsRow (p. ??)

# A.26 Class plot\_abstract

### A.26.1 Constructor plot\_abstract/plot\_abstract

**Summary:** Abstract description of a single plot.

**Usage:** 

obj = plot\_abstract(data, axis\_labels, title, legend, command, props)

**Description:** Base class that holds the necessary data to draw a plot. This data can then be used to generate different plots. Subclasses define specific plots with additional data. Subclasses should conform to the standard that the series of commands found in plotFigure should produce a valid figure.

#### **Parameters:**

**data:** A cell array of data arrays (x, y, z, etc.) that can be fed to plot commands.

axis\_labels: Cell array of axis label strings.

title: Plot description string.

**legend:** Cell array of descriptions for each item plotted.

**command:** Plotting command to use (Optional, default='plot')

**props:** A structure with any optional properties.

axisLimits: Sets axis limits of non-NaN values in vector.

tightLimits: If 1, issues an "axis tight" command (default=0)

**border:** Relative size of border spacing around axis, between 0 - 1. (default=0)

If a scalar, equal border on all sides, give a four-element vector [left bottom right top] to define borders for each side.

fontSize: Set the fontsize.

grid: Display dashed grid in background.

**noXLabel:** No X-axis label. **noYLabel:** No Y-axis label.

noTitle: No title.

rotateXLabel: Rotates the X-axis label for smaller width. rotateYLabel: Rotates the Y-axis label for smaller width.

numXTicks: Number of ticks on X-axis.

**formatXTickLabels:** The sprintf format string for tick labels.

XTick, YTick: Point locations for axis ticks.

XTickLabel, YTickLabel: Axis tick labels.

ColorOrder: Set the ColorOrder of the axis.

**LineStyleOrder:** Set the LineStyleOrder of the axis.

**legendLocation:** Passed to legend(..., 'location', legendLocation).

legendOrientation: Passed to legend(..., 'orientation', legendLocation)

noLegends: If exists, no legends are displayed.

**axisProps:** Passed to set properties of the axis drawn. **plotProps:** Passed to set properties of the plot drawn.

**figureProps:** Passed to set properties of the figure drawn.

**PaperPosition:** Sets the figure property for printing at this size.

resizeControl: If 0, drawing after resize is disabled and prints at screen

size, if 1 (default), redraws figure after each resize event and prints at PaperPosition size.

### Returns a structure object with the following fields:

data, axis\_labels, title, legend, command, props

See also: plot\_abstract/plot(p. 158), plot\_abstract/plotFigure(p. 155)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

## A.26.2 Method plot\_abstract/setProp

Summary: Generic method for setting optional object properties.

Usage:

```
obj = setProp(obj, prop1, val1, prop2, val2, ...)
```

**Description:** Modifies or adds property values. As many property name-value pairs can be specified.

## **Parameters:**

**obj:** Any object that has a props field.

attr: Property nameval: Property value.

## **Returns:**

obj: The new object with the updated properties.

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/22

## A.26.3 Method plot\_abstract/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

## A.26.4 Method plot\_abstract/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

## A.26.5 Method plot\_abstract/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

### A.26.6 Method plot\_abstract/plotFigure

**Summary:** Draws this plot alone in a new figure window.

Usage:

handle = plotFigure(a\_plot)

#### **Parameters:**

a\_plot: A plot\_abstract object, or a subclass object.
title\_str: (Optional) String to append to plot title.

## **Returns:**

handle: Handle of new figure.

See also: plot\_abstract(p. 153), plot\_abstract/plot(p. 158), plot\_abstract/decorate (p. 159)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

## A.26.7 Method plot\_abstract/openAxis

**Summary:** Calculates the extents for the axis of this plot and opens it.

Usage:

```
[axis_handle, layout_axis] = openAxis(a_plot, layout_axis)
```

#### **Parameters:**

a\_plot: A plot\_abstract object, or a subclass object.

**layout\_axis:** The axis position to layout this plot (Optional).

If NaN, doesn't open a new axis.

#### **Returns:**

handles: Handles of graphical objects drawn.

See also: plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

### A.26.8 Method plot\_abstract/axis

**Summary:** Returns the estimated axis ranges of this plot according to its data.

**Usage:** 

```
ranges = axis(a_plot)
```

#### Parameters:

a\_plot: A plot\_abstract object, or a subclass object.

#### **Returns:**

ranges: The ranges as a vector in the same way 'axis' would return.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/13

## A.26.9 Method plot\_abstract/superposePlots

**Summary:** Superpose multiple plots with common command onto a single axis.

#### **Usage:**

```
a_plot = superposePlots(plots, axis_labels, title_str, command, props)
```

**Description:** The plot decoration will be taken from the last plot in the list, with the exception of legend labels.

#### **Parameters:**

```
plots: Array of plot_abstract or subclass objects.
```

axis\_labels: Cell array of axis label strings (optional, taken from plots).

title\_str: Plot description string (optional, taken from plots).

command: Plotting command to use (optional, taken from plots)

**props:** A structure with any optional properties.

**noLegends:** If exists, no legends are created.

#### **Returns:**

```
a_plot: A plot_abstract object.
```

```
See also: plot_abstract(p. 153), plot_abstract/plot(p. 158), plot_abstract/plotFigure (p. 155)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/23

## A.26.10 Method plot\_abstract/matrixPlots

Summary: Superpose multiple plots with common command onto a single axis.

## **Usage:**

```
a_plot = matrixPlots(plots, axis_labels, title_str, props)
```

### **Parameters:**

```
plots: Array of plot_abstract or subclass objects.
```

axis\_labels: Cell array of axis label strings (optional, taken from plots).

title\_str: Plot description string (optional, taken from plots).

**props:** A structure with any optional properties passed to the Y stack\_plot.

titlesPos: if specified, passed to the X stack\_plots.
rotateYLabel: if specified, passed to the X stack\_plots.
axisLimits: if specified, passed to the X stack\_plots.
goldratio: try to make the figure in this aspect ratio.

width, height: if specified, make the figure have this many plots in corresponding dimension.

#### **Returns:**

```
a_plot: A plot_abstract object.
```

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158), plot\_abstract/plotFigure (p. 155)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/07

# A.26.11 Method plot\_abstract/subsref

Summary: Defines generic indexing for objects.

### A.26.12 Method plot\_abstract/plot

**Summary:** Draws this plot in the current axis.

Usage:

handles = plot(a\_plot, layout\_axis)

#### **Parameters:**

a\_plot: A plot\_abstract object, or a subclass object.

layout\_axis: The axis position to layout this plot (Optional).

If NaN, doesn't open a new axis.

### **Returns:**

handles: Handles of graphical objects drawn.

See also: plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

## A.26.13 Method plot\_abstract/subsasgn

Summary: Defines generic index-based assignment for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/06

## A.26.14 Method plot\_abstract/decorate

**Summary:** Places decorations (titles, labels, ticks, etc.) on the plot.

### Usage:

```
handles = decorate(a_plot)
```

#### **Parameters:**

a\_plot: A plot\_abstract object, or a subclass object.

#### **Returns:**

handles: Handles of graphical objects drawn.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

## A.27 Class plot\_bars

## A.27.1 Constructor plot\_bars/plot\_bars

**Summary:** Bar plot with error lines in individual axes for each variable.

### **Usage:**

```
a_plot = plot_bars(mid_vals, lo_vals, hi_vals, n_vals, x_labels, y_labels, ...
title, axis_limits, props)
```

**Description:** Subclass of plot\_stack. The plot\_abstract/plot command can be used to plot this data. Rows of \*\_vals will create grouped bars, columns will create new axes.

### **Parameters:**

```
mid_vals: Middle points of error bars.
```

**lo\_vals:** Low points of error bars.

hi\_vals: High points of error bars.

**n\_vals:** Number of samples used for the statistic (Optional).

x\_labels, y\_labels: Axis labels for each bar group. Must match with data columns.

title: Plot description.

**axis\_limits:** If given, all plots contained will have these axis limits.

**props:** A structure with any optional properties.

**dispErrorbars:** If 1, display errorbars for lo\_vals and hi\_vals deviation from mid\_vals (default=1).

**dispNvals:** If 1, display n\_vals on top of each bar.

**groupValues:** Array of within-group numeric labels, instead of just a sequence of numbers.

truncateDecDigits: Truncate labels to this many decimal digits.
barAxisProps: props passed to plot\_abstract objects with bar commands

## Returns a structure object with the following fields:

plot\_abstract

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

### A.27.2 Method plot\_bars/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.28 Class plot\_errorbar

# A.28.1 Constructor plot\_errorbar/plot\_errorbar

Summary: Generic errorbar plot.

Usage:

a\_plot = plot\_errorbar(x\_vals, mid\_vals, lo\_vals, hi\_vals, line\_spec,
axis\_labels, title, legend, props)

**Description:** Subclass of plot\_abstract. The plot\_abstract/plot command can be used to plot this data. Needed to create this as a separate class to have the axis ranges method to measure the errorbars.

#### **Parameters:**

**x\_vals:** X coordinates of errorbars.

mid\_vals: Middle points of error bars.

lo vals: Low points of error bars.

hi\_vals: High points of error bars.

line\_spec: Plot line spec to be passed to errorbar

axis\_labels: Cell array for X, Y axis labels.

title: Plot description.

**legend:** For multiple errorbar plots (matrix form), description of each plot. **props:** A structure with any optional properties to be passed to plot\_abstract.

### Returns a structure object with the following fields:

plot\_abstract.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

### A.28.2 Method plot\_errorbar/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

### A.28.3 Method plot\_errorbar/axis

**Summary:** Returns the estimated axis ranges of this plot according to its data.

**Usage:** 

ranges = axis(a\_plot)

#### **Parameters:**

**a\_plot:** A plot\_abstract object, or a subclass object.

### **Returns:**

ranges: The ranges as a vector in the same way 'axis' would return.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/10/13

## A.29 Class plot\_errorbars

## A.29.1 Constructor plot\_errorbars/plot\_errorbars

**Summary:** Special plot for plotting distributions of variables in separate axes.

Usage:

a\_plot = plot\_errorbars(labels, mid\_vals, lo\_vals, hi\_vals, labels, title,
axis\_limits, props)

**Description:** Subclass of plot\_stack. The plot\_abstract/plot command can be used to plot this data.

#### **Parameters:**

**labels:** Labels of parameters to appear at bottom of each errorbar.

mid\_vals: Middle points of error bars.lo vals: Low points of error bars.

hi\_vals: High points of error bars.

title: Plot description.

**axis\_limits:** If given, all plots contained will have these axis limits.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

plot\_abstract, labels.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

# A.30 Class plot\_inset

# A.30.1 Constructor plot\_inset/plot\_inset

Summary: Superpose multiple plots with individual axis at arbitrary locations.

#### **Usage:**

```
a_plot = plot_inset(plots, axis_locations, title_str, props)
```

**Description:** Subclass of plot\_abstract. Contains other plot\_abstract objects or subclasses thereof to be layout in arbitaray format. Allows overlapping and therefore good for insets and special plots.

### **Parameters:**

plots: Cell array of plot\_abstract or subclass objects.

axis\_locations: Matrix of four-element vectors for each given plot.

title\_str: Title to go on top of the stack

props: A structure with any optional properties.

# Returns a structure object with the following fields:

plot\_abstract, plots, axis\_locations.

See also: plot\_abstract (p. 153), plot\_abstract/plotFigure (p. 155)

Author: Cengiz Gunay <cgunay@emory.edu>, 2007/06/05

# A.30.2 Method plot\_inset/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

## A.30.3 Method plot\_inset/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.30.4 Method plot\_inset/plot

**Summary:** Superposes contained plots in their own axes.

Usage:

handles = plot(a\_plot, layout\_axis)

# Parameters:

a\_plot: A plot\_superpose object.

layout\_axis: The axis position to layout this plot (Optional).

#### **Returns:**

handles: Handles of graphical objects drawn.

See also: plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2007/06/08

## A.31 Class plot\_simple

### A.31.1 Constructor plot\_simple/plot\_simple

**Summary:** Abstract description of a single plot.

**Usage:** 

a\_plot = plot\_simple(data\_x, data\_y, title, label\_x, label\_y, legend, command,
props)

**Description:** Subclass of plot\_abstract. The plot\_abstract/plot command can be used to plot this data.

### **Parameters:**

PANDORA Toolbox: Prog.'s MaBuraft v805 of 6007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

data\_x: X-axis values for the plot.

data\_y: Y-axis values for the plot.

title: Plot description.

**label\_x:** X-axis label string.

label\_y: Y-axis label string.

**legend:** Short description of data points.

command: Plotting command to use (Optional, default='plot')

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

plot\_abstract.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

## A.31.2 Method plot\_simple/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

## A.31.3 Method plot\_simple/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.32 Class plot\_stack

### A.32.1 Constructor plot\_stack/plot\_stack

Summary: A horizontal or vertical stack of plots.

Usage:

```
a_plot = plot_stack(plots, axis_limits, orientation, title_str, props)
```

**Description:** Subclass of plot\_abstract. Contains other plot\_abstract objects or subclasses thereof to be layout in stack format.

### **Parameters:**

plots: Cell array of plot\_abstract or subclass objects.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 6407-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

axis\_limits: If given, all plots contained will have these axis limits.
orientation: Stack orientation 'x' for horizontal, 'y' for vertical, etc.

title\_str: Title to go on top of the stack

**props:** A structure with any optional properties.

yLabelsPos: 'left' means only put y-axis label to leftmost plot.
yTicksPos: 'left' means only put y-axis ticks to leftmost plot.
xLabelsPos: 'bottom' means only put x-axis label to lowest plot.
xTicksPos: 'bottom' means only put x-axis ticks to lowest plot.

titlesPos: 'top' means only put title to top plot.

relaxedLimits: Add 10

relativeSizes: An array specifying relative size of each plot with one value

(Example: relativeSizes=[1 2] makes second plot twice wider than first.)

### Returns a structure object with the following fields:

plot\_abstract, plots, axis\_limits, orient.

See also: plot\_abstract (p. 153), plot\_abstract/plotFigure (p. 155)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/04

## A.32.2 Method plot\_stack/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

## A.32.3 Method plot\_stack/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

## A.32.4 Method plot\_stack/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.32.5 Method plot\_stack/superposePlots

**Summary:** Superpose multiple plot\_stack objects that contain exact same contents.

### Usage:

```
a_plot = superposePlots(plots, axis_labels, title_str, command, props)
```

**Description:** The plot decoration will be taken from the last plot in the list, with the exception of legend labels.

### **Parameters:**

plots: Array of plot\_stack objects.

axis\_labels: Cell array of axis label strings (optional, taken from plots).

title\_str: Plot description string (optional, taken from plots).

**command:** Plotting command to use (optional, taken from plots)

**props:** A structure with any optional properties.

noLegends: If exists, no legends are created.

#### **Returns:**

```
a_plot: A plot_stack object.
```

See also: plot\_abstract(p. 153), plot\_abstract/plot(p. 158), plot\_abstract/plotFigure (p. 155)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/14

### A.32.6 Method plot\_stack/plot

**Summary:** Draws this plot in the current axis or at the position in layout\_axis.

#### **Usage:**

```
handles = plot(a_plot, layout_axis)
```

#### **Parameters:**

**a\_plot:** A plot\_abstract object, or a subclass object.

layout\_axis: The axis position to layout this plot (Optional).

### **Returns:**

handles: Handles of graphical objects drawn.

See also: plot\_stack (p. 164), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/04

## A.32.7 Method plot\_stack/decorate

**Summary:** No additional decorations for stacked plots.

**Usage:** 

```
a_histogram_db = decorate(a_plot)
```

#### **Parameters:**

**a\_plot:** A plot\_abstract object, or a subclass object.

#### **Returns:**

handles: Handles of graphical objects drawn.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/04

# A.33 Class plot\_superpose

### A.33.1 Constructor plot\_superpose/plot\_superpose

**Summary:** Multiple plot\_abstract objects superposed on the same axis.

**Usage:** 

```
obj = plot_superpose(plots, axis_labels, title_str, props)
```

**Description:** Subclass of plot\_abstract. Contains multiple plot\_abstract objects to be plotted on the same axis. This is different than the plot\_abstract/superpose, where only using the same plot command is allowed. Here, each plot\_abstract can have its own special plotting command. Subclasses of plot\_abstract is also allowed here. The decorations comes from this object and not children plots. This behavior is different than plot\_stack, where each plot has its own decorations. If you want each plot to have its own axis (e.g. an inset, or plot with multiple axis labels) then you should use plot\_inset.

### **Parameters:**

plots: Cell array of plot\_abstract or subclass objects.

axis\_labels: Cell array of axis label strings.

title\_str: Plot description string.

**props:** A structure with any optional properties (passed to plot\_abstract).

### Returns a structure object with the following fields:

plot\_abstract, plots

See also: plot\_abstract/superpose (p.??), plot\_superpose/plot (p. 169)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/22

## A.33.2 Method plot\_superpose/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

## A.33.3 Method plot\_superpose/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.33.4 Method plot\_superpose/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

### A.33.5 Method plot\_superpose/axis

Summary: Returns the maximal axis ranges according to superposed subplots.

Usage:

ranges = axis(a\_plot)

## Parameters:

**a\_plot:** A plot\_abstract object, or a subclass object.

### **Returns:**

ranges: The ranges as a vector in the same way 'axis' would return.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/22

# A.33.6 Method plot\_superpose/superposePlots

**Summary:** Superpose multiple plot\_superpose objects by merging them into one.

**Usage:** 

```
a_plot = superposePlots(plots, axis_labels, title_str, command, props)
```

#### **Parameters:**

**plots:** Array of plot\_superpose objects.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 6807-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

```
axis_labels: Cell array of axis label strings (optional, taken from plots).
             title_str: Plot description string (optional, taken from plots).
             command: Plotting command to use (optional, taken from plots)
             props: A structure with any optional properties.
                  noLegends: If exists, no legends are created.
       Returns:
             a_plot: A plot_superpose object.
       See also: plot_abstract/superposePlots (p. 157), plot_stack/superposePlots
             (p. 166)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/14
       A.33.7 Method plot_superpose/plot
       Summary: Draws this plot in the current axis.
       Usage:
handles = plot(a_plot, layout_axis)
       Parameters:
             a_plot: A plot_superpose object.
             layout_axis: The axis position to layout this plot (Optional).
       Returns:
             handles: Handles of graphical objects drawn.
       See also: plot_abstract (p. 153)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/08
       A.33.8 Method plot_superpose/decorate
       Summary: Places decorations using the first plot of the superposed plots.
       Usage:
handles = decorate(a_plot)
```

### **Parameters:**

a\_plot: A plot\_abstract object, or a subclass object.

### **Returns:**

handles: Handles of graphical objects drawn.

See also: plot\_abstract (p. 153), plot\_abstract/plot (p. 158)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/11

## A.34 Class ranked db

## A.34.1 Constructor ranked\_db/ranked\_db

**Summary:** A database of distance values generated by ranking rows of orig\_db with the criterion in crit\_db.

#### **Usage:**

```
a_ranked_db = ranked_db(data, col_names, orig_db, crit_db, id, props)
```

**Description:** This is a subclass of tests\_db. It should contain a Distance column. A more general ranked db class may be needed later. Use the rankMatching method to get an instance of this class.

#### **Parameters:**

data: Database contents.

col\_names: The column names.

orig\_db: DB whose rows are ranked.

crit\_db: The criterion DB used for generating the ranking scores.

id: An identifying string.

**props:** A structure with any optional properties.

tolerateNaNs: If 0, rows with any NaN values are skipped (default=1).

# Returns a structure object with the following fields:

```
tests_db, orig_db, crit_db, props.
```

See also: tests\_db(p.217), tests\_db/rankMatching(p.254), tests\_db/matchingRow(p.233)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/21

### A.34.2 Method ranked\_db/blockedDistances

**Summary:** Creates a db of distances to blocked versions of top ranks.

### Usage:

[a\_db, ranked\_dbs] = blockedDistances(a\_ranked\_db, rows, blocked\_db, blocked\_param\_indices, block\_levels, crit\_db)

#### **Parameters:**

a\_ranked\_db: A ranked\_db object.

rows: Use the given row rankings.

**blocked\_db:** db with blocked versions of original ranks.

**blocked\_param\_indices:** Indices of parameters to be blocked.

**block\_levels:** Number of parameter levels for blocking.

crit\_db: Calculate distance from this criterion.

#### **Returns:**

a\_db: A tests\_db object with the matrix of distances. ranked\_dbs: A cell array of ranked\_dbs for each row.

### **Example:**

```
» dist_matx_db = blockedDistances(rankMatching(super_db,
matchingRow(rsuper_phys_db, 20)), 1:5, super_blocker_db, [1 2], 10,
matchingRow(rsuper_phys_db, 21))
```

See also: makeModifiedParamDB (p. ??), getParamRowIndices (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/01/14

### A.34.3 Method ranked\_db/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

### A.34.4 Method ranked\_db/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

### A.34.5 Method ranked\_db/plotDistMatrix

**Summary:** Create a color-coded matrix plot of with total errors from the ranked DB.

#### **Usage:**

```
a_plot = plotDistMatrix(db, rows, col_size, col_name, num_col_labels, row_name,
num_row_labels, title_str, props)
```

**Description:** The col\_size parameter is used to find the number of rows that make up the x-dimension of the color matrix plot.

#### **Parameters:**

**db:** A ranked\_db object.

rows: Indices of rows in db after joining (and sorting).

**col\_size:** Number of rows to take from DB to form the columns of matrix plot.

**col\_name**, **row\_name**: DB column to use for the figure column and row, respectively.

num\_col\_labels, num\_row\_labels: Number of labels to put on each
axis.

title\_str: If non-empty, replaces generic title with db name.

**props:** A structure with any optional properties.

**sortBy:** If specified, db is sorted after being joined with original using this column.

**colorbar:** Put a colorbar on the figure. (also passed to plot\_abstract)

### **Returns:**

a\_plot: A plot\_abstract object.

plotCompareDistMatx(p. ??)

## **Example:**

```
» plotFigure(plotDistMatrix(scored_blocked_sk_gps0503b_control_db, ':', 10, 'SK',
10, 'trial', 10, 'gps0503b (control), preset 6 - top 50 matches',
struct('sortBy', 'trial', 'colorbar', 1, 'PaperPosition', [0 0 5 3])));

See also: ranked_db(p. 170), plot_abstract(p. 153), getDistMatrix(p. ??),
```

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2005/12/12

## A.34.6 Method ranked\_db/plotCompareDistMatx

**Summary:** Compare differences and correlations of distance matrices from two ranked DBs.

### Usage:

```
a_plot = plotCompareDistMatx(db, rows, col_size, col_name, num_col_labels,
row_name, num_row_labels, title_str, props)
```

**Description:** Produces three plots: (1) distance difference matrix, (2) 2D cross-correlogram, and (3) repeated 1D cross-correlogram for each row.

#### **Parameters:**

**db**, **w\_db**: The ranked\_db objects to be compared.

rows: Indices of rows in db after joining (and sorting) for both DBs.

**col\_size:** Number of rows to take from DB to form the columns of matrix plot.

**col\_name**, **row\_name**: DB column to use fot the figure column and row, respectively.

num\_col\_labels, num\_row\_labels: Number of labels to put on each
axis.

title\_str: If non-empty, replaces generic title with db name.

**props:** A structure with any optional properties.

**sortBy:** If specified, db is sorted after being joined with original using this column.

**colorbar:** Put a colorbar on the figure. (also passed to plot\_abstract)

### **Returns:**

a\_plot: A plot\_abstract object.

See also: tests\_db (p. 217), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/12

### A.34.7 Method ranked\_db/plotRowErrors

**Summary:** Create plot of rankings with errors associated with each measure color-coded.

#### **Usage:**

```
a_plot = plotRowErrors(a_ranked_db, rows, props)
```

#### **Parameters:**

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```
a_ranked_db: A ranked_db object.
             rows: Indices of rows in a_ranked_db.
             title_str: (Optional) String to append to plot title.
             props: A structure with any optional properties.
                  sortMeasures: If specified, measure order is determined with increas-
                      overall distance.
                  RowName: Label to show on X-axis (default='Ranks')
                  rowSteps: Steps to jump in labeling rows on the x-axis.
                  superposeDistances: Superpose a white-colored distance line plot.
                      (rest passed to plot_abstract)
       Returns:
             a_plot: A plot_abstract object.
       See also: ranked_db (p. 170), tests_db/rankMatching (p. 254), plot_abstract
             (p. 153), plotImage (p. ??)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/12
       A.34.8 Method ranked_db/displayRows
       Summary: Displays rows of rankings together with errors associated with each mea-
             sure.
       Usage:
s = displayRows(db, rows)
       Parameters:
             db: A tests_db object.
             rows: Indices of rows in db.
       Returns:
             s: A structure of column name and value pairs.
       See also: tests_db (p. 217)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/15
       A.34.9 Method ranked db/subsref
       Summary: Defines generic indexing for objects.
```

## A.34.10 Method ranked\_db/joinOriginal

**Summary:** Joins the distance values to the original db rows with matching row indices.

### Usage:

```
a_db = joinOriginal(a_ranked_db, rows)
```

**Description:** Takes the parameter columns from orig\_db and all tests from crit\_db.

#### **Parameters:**

```
a_ranked_db: A ranked_db object.
rows: Join only the given rows.
```

#### **Returns:**

a\_db: A params\_tests\_db object (same type as a\_ranked\_db.orig\_db) containing the desired rows in ascending order of distance.

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/21

### A.34.11 Method ranked db/renameColumns

Summary: Rename an existing column or columns.

### Usage:

```
a_db = renameColumns(a_db, test_names, new_names)
```

**Description:** This method is an overloaded method for ranked\_db that keeps consistent the column names of the ranked, criterion and original DBs. The other DBs are not renamed for the Distance and RowIndex columns.

### **Parameters:**

```
a_db: A ranked_db object.
test_names: A cell array of existing test names.
new_names: New names to replace existing ones.
```

## **Returns:**

a\_db: The ranked\_db object that includes the new columns.

See also: tests\_db/renameColumns (p. 239)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/07

## A.34.12 Method ranked\_db/getDistMatrix

**Summary:** Create a matrix of total errors from the ranked DB.

**Usage:** 

```
distmatx = getDistMatrix(db, rows, col_size, props)
```

**Description:** The col\_size parameter is used to find the number of rows that make up the x-dimension of the matrix.

#### **Parameters:**

db: A tests\_db object.

rows: Indices of rows in db after joining (and sorting).

**col\_size:** Number of rows to take from DB to form the columns of matrix plot.

**props:** A structure with any optional properties.

**sortBy:** If specified, db is sorted after being joined with original using this column.

### **Returns:**

a\_plot: A plot\_abstract object.

See also: tests\_db (p. 217), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/12

## A.35 Class results\_profile

### A.35.1 Constructor results\_profile/results\_profile

**Summary:** Creates and collects result profiles for data objects.

**Usage:** 

```
obj = results_profile(results, id, props)
```

**Description:** This is the base class for all profile classes.

#### **Parameters:**

**results:** A structure containing test results.

id: Identification string.

**props:** A structure with any optional properties.

## Returns a structure object with the following fields:

results, id, props.

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See also: trace\_profile (p. 264), cip\_trace\_profile (p. 69)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

### A.35.2 Method results\_profile/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

## A.35.3 Method results\_profile/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

### A.35.4 Method results\_profile/subsref

Summary: Defines generic indexing for objects.

## A.35.5 Method results\_profile/plot

**Summary:** Generic method to plot a tests\_db or a subclass. Requires a plot\_abstract method to be defined for this object.

#### **Usage:**

```
h = plot(a_tests_db, title_str)
```

### **Parameters:**

a\_tests\_db: A histogram\_db object.

title\_str: (Optional) String to append to plot title.

### **Returns:**

h: The figure handle created.

See also: plot\_abstract (p. 153), plotFigure (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/06

## A.35.6 Method results\_profile/getResults

**Summary:** Return the results profile structure.

Usage:

```
results = getResults(p)
```

#### **Parameters:**

**p:** A result\_profile object.

### **Returns:**

results: A structure associating test names to values.

See also: results\_profile (p. 176)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.36 Class script\_array

# A.36.1 Constructor script\_array/script\_array

Summary: Generic class that provides the scripts for a repetitive array job.

**Usage:** 

```
obj = script_array(num_runs, id, props)
```

**Description:** This is the base class for all script\_array classes. Runs the runJob method as num\_runs many times.

#### **Parameters:**

num\_runs: The number of times the runJob script should be evoked.

id: Identification string.

**props:** A structure with any optional properties.

**runJobFunc:** A function name or handle to be used instead of default runJob.

#### Returns a structure object with the following fields:

num\_runs, id, props.

### **Example:**

```
» func1 = inline('x^2')
» runFirst(script_array(10, 'squares numbers up to 10'), struct('runJobFunc',
func1))
ans = [ 1] [ 4] [ 9] [ 16] [ 25] [ 36] [ 49] [ 64] [ 81] [100]

See also: runFirst (p. ??), runLast (p. ??), runJob (p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/01

## A.36.2 Method script\_array/runFirst

**Summary:** Method to be called at beginning of script\_array jobs.

Usage:

```
job_results = runFirst(a_script_array)
```

**Description:** This method initiates the script\_array jobs. It loops and calls runJob and finally calls runLast.

#### **Parameters:**

```
a_script_array: A script_array object.
```

### **Returns:**

job\_results: A cell array of results collected from each item of the vector jobs.

### **Example:**

See also: runLast (p. ??), runJob (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/01

# A.36.3 Method script\_array/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

### A.36.4 Method script\_array/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/06

## A.36.5 Method script\_array/runLast

**Summary:** Method to be called last after the script\_array jobs.

### Usage:

```
job_results = runLast(a_script_array, job_results)
```

**Description:** This method is provided as a placeholder and does nothing. It can filterout the results returned from the jobs run. Normally it is invoked internally by the runFirst method, after running and collecting results from the vector jobs with the runJob method.

#### **Parameters:**

```
a_script_array: A script_array object.
job results: The index within the vector job.
```

### **Returns:**

job\_results: Any output produced by the job.

## **Example:**

```
Call it directly:
» runLast(script_array(10, 'this one does nothing for 10 times'), );
See also: runJob(p.??), runFirst(p.??)
```

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2006/02/01

# A.36.6 Method script\_array/runJob

**Summary:** Method to be called for each of the script\_array jobs.

# **Usage:**

```
job_result = runJob(a_script_array, vector_index)
```

**Description:** This method is provided as a placeholder and does nothing. If the run\_job\_func property is defined, it will call that function.

### **Parameters:**

```
a_script_array: A script_array object.
vector_index: The index within the vector job.
```

### **Returns:**

job\_result: Any output produced by the job.

### **Example:**

```
See real example in script_array. Call the 5th job:
» runJob(script_array(10, 'this one does nothing for 10 times'), 5);

See also: runLast (p. ??), runFirst (p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/01

### A.36.7 Method script\_array/subsref

Summary: Defines generic indexing for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

### A.36.8 Method script\_array/subsasgn

Summary: Defines generic index-based assignment for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/06

# A.37 Class script\_array\_for\_cluster

## A.37.1 Constructor script\_array\_for\_cluster/script\_array\_for\_cluster

**Summary:** Generic class defining a repetitive vector job to be run on a Sun Grid Engine (SGE) computing cluster.

#### **Usage:**

a\_script\_cluster = script\_array\_for\_cluster(num\_runs, sge\_wrapper\_script, id,
props)

**Description:** This is a subclass of the script\_array class. The runFirst method spawns num\_runs copies of the runJob method in parallel on the cluster, followed by the invocation of the runLast method.

#### **Parameters:**

num\_runs: The number of times the runJob script should be evoked.

**sge\_wrapper\_script:** A script that can be submitted with qsub and can execute arbitrary

Matlab commands on the cluster nodes. It can have qsub options prepended to it such as '-p -100 -q all.q <abs\_path\_to>/sge\_matlab.sh'.

id: Identification string.

**props:** A structure with any optional properties.

**notifyByMail:** An SGE notification email is sent to this address after lastJob.

(others passed to script\_array)

# Returns a structure object with the following fields:

num\_runs, id, props.

See also: runFirst (p. ??), runLast (p. ??), runJob (p. ??), script\_array (p. 178)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2006/02/02

# A.37.2 Method script\_array\_for\_cluster/runFirst

**Summary:** Method to be called at beginning of script\_array\_for\_cluster jobs.

### Usage:

```
job_results = runFirst(a_script_cluster)
```

**Description:** This method initiates the script\_array\_for\_cluster jobs. It submits an SGE vector job for running each runJob and finally runLast. There is no way of collecting outputs from individual runJob calls.

#### **Parameters:**

```
a_script_cluster: A script_array_for_cluster object.
```

#### **Returns:**

job\_results: A cell array of results collected from each item of the vector jobs.

#### **Example:**

```
» runFirst(script_array_for_cluster(10, 'this one does nothing for 10 times'));
```

See also: script\_array\_for\_cluster (p. 181)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/01

# A.37.3 Method script\_array\_for\_cluster/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.37.4 Method script\_array\_for\_cluster/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/06

# A.38 Class script\_factory

### A.38.1 Constructor script\_factory/script\_factory

**Summary:** Generic class to automatically create a set of scripts.

Usage:

obj = script\_factory(num\_scripts, out\_name, id, props)

**Description:** This is the base class for all script\_factory classes.

#### **Parameters:**

num\_scripts: Number of scripts to create.

**out\_name:** The file name for the output scripts. A 'filename corresponds to the script number.

id: Identification string.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

num\_scripts, out\_name, id, props.

See also: script\_factory/writeScripts (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/11/28

#### A.38.2 Method script\_factory/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.39 Class spike\_shape

# A.39.1 Constructor spike\_shape/spike\_shape

**Summary:** An action potential shape trace.

**Usage:** 

obj = spike\_shape(data, dt, dy, id)

## **Parameters:**

data: A vector of data points containing the spike shape.

dt: Time resolution [s].

dy: y-axis resolution [ISI (V, A, etc.)]

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id: Identification string.

props: A structure with any optional properties.

**baseline:** Resting potential. **threshold:** Spike threshold.

init\_Vm\_method: Method to obtain spike initiation voltage.

1- maximum acceleration point 2- threshold crossing of acceleration (needs threshold) 3- threshold crossing of slope (needs threshold) 4-maximum acceleration in phase space (optionally specify maximal threshold as init\_threshold) 5- point of maximum curvature, when slope is between init\_lo\_thr and init\_hi\_thr 6- local maximum of second derivative in the phase space nearest slope crossing init\_threshold 7- threshold crossing of interpolated slope (needs threshold) 8- maximum curvature in phase-plane 9- Combined curvature and inflection method in time-domain.

init\_threshold: Spike initiation threshold (deriv or accel).
 (see above methods and implementation in calcInitVm)
init\_lo\_thr, init\_hi\_thr: Low and high thresholds for slope.

### Returns a structure object with the following fields:

trace, props.

See also: trace/spike\_shape (p. 263), trace/analyzeSpikesInPeriod (p. 264), trace (p. 255), spikes (p. 195), period (p. 137)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

## A.39.2 Method spike\_shape/calcInitVmSlopeThresholdSupsample

**Summary:** Estimates the AP threshold as the first slope threshold crossing by first supersampling the data using cubic spline interpolation.

## **Usage:**

[init\_idx, a\_plot] = calcInitVmSlopeThresholdSupsample(s, max\_idx, min\_idx, thr,
plotit)

## **Parameters:**

```
s: A spike_shape object.
max_idx: The index of the maximal point of the spike_shape [dt].
min_idx: The index of the minimal point of the spike_shape [dt].
thr: Threshold for time derivative of voltage.
plotit: If non-zero, plot a graph annotating the test results (optional).
```

### **Returns:**

init\_idx: AP threshold index in the spike\_shape [dt]. a\_plot: plot\_abstract, if requested.

See also: calcInitVm (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/03/23

# A.39.3 Method spike\_shape/plotCompareMethods

**Summary:** Creates a multi-plot comparing different action potential threshold finding methods.

#### **Usage:**

```
a_plot = plotCompareMethods(s, title_str)
```

#### Parameters:

**s:** A spike\_shape object.

title\_str: Title suffix (optional).

# **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: spike\_shape (p. 183), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/19

# A.39.4 Method spike\_shape/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.39.5 Method spike\_shape/calcMaxVm

**Summary:** Calculates the maximal value of the spike\_shape, s.

# Usage:

```
[max_val, max_idx] = calcMaxVm(s)
```

## **Parameters:**

**s:** A spike\_shape object.

#### **Returns:**

max\_val: The max value. max\_idx: Its index in the spike\_shape [dt].

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See also: period (p. 137), spike\_shape (p. 183), trace/calcMax (p. 260)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/08/02

### A.39.6 Method spike\_shape/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

## A.39.7 Method spike\_shape/calcInitVmV3hKpTinterp

**Summary:** Calculates candidates for action potential threshold using the first three time-domain derivatives.

### Usage:

[init\_idx, a\_plot] = calcInitVmV3hKpTinterp(s, max\_idx, min\_idx, lo\_thr, hi\_thr,
plotit)

**Description:** First uses interpolation to increase time points. Calculates h, the second derivative of phase-plane ( $d^2 v'/dv^2$ ), in terms of time-domain derivatives. Also calculates  $Kp = V''[1 + (V')^2]^{-(-3/2)}$ , the curvature. The maxima of these functions are used as candidates for AP thresholds.

#### **Parameters:**

**s:** A spike\_shape object.

max\_idx: The index of the maximal point of the spike\_shape [dt].

min\_idx: The index of the minimal point of the spike\_shape [dt].

lo\_thr, hi\_thr: Lower and higher thresholds for time derivative of voltage.

**plotit:** If non-zero, plot a graph annotating the test results (optional).

### **Returns:**

init\_idx: Indices of threshold candidates in the spike\_shape [dt]. a\_plot: plot\_abstract, if requested.

See also: calcInitVm (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/18

# A.39.8 Method spike\_shape/set

**Summary:** Generic method for setting object attributes.

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

### A.39.9 Method spike\_shape/calcInitVmSekerliV2

**Summary:** Calculates the action potential threshold using the maximum second derivative of the phase space of voltage-time slope versus voltage.

### **Usage:**

```
[init_idx, a_plot] = calcInitVmSekerliV2(s, max_idx, min_idx, plotit)
```

#### **Parameters:**

```
s: A spike_shape object.
```

max\_idx: The index of the maximal point of the spike\_shape [dt].

min\_idx: The index of the minimal point of the spike\_shape [dt].

**plotit:** If non-zero, plot a graph annotating the test results (optional).

#### **Returns:**

init\_idx: Its index in the spike\_shape [dt]. a\_plot: plot\_abstract, if requested.

See also: calcInitVm (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/18

# A.39.10 Method spike\_shape/calcMinVm

**Summary:** Calculates the minimal value of the spike\_shape, s.

#### **Usage:**

```
[min_val, min_idx, max_min_time] = calcMinVm(s, max_idx)
```

#### **Parameters:**

**s:** A spike\_shape object.

max\_idx: The index of the maximal point of the spike\_shape [dt].

### **Returns:**

min\_val: The min value [dy]. min\_idx: Its index in the spike\_shape [dt]. max\_min\_time: Time from max to min [dt].

See also: period (p. 137), spike\_shape (p. 183), trace/calcMin (p. 260)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/02

# A.39.11 Method spike\_shape/plotTPP

**Summary:** Plots the dV/dt vs. V phase-plane representation of the spike shape.

#### **Usage:**

```
a_plot = plotTPP(s)
```

**Description:** Uses the Taylor series estimation for finding the derivative dV/dt.

#### **Parameters:**

**s:** A spike\_shape object.

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: spike\_shape (p. 183), plot\_abstract (p. 153), diffT (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/16

# A.39.12 Method spike\_shape/calcInitVm

Summary: Calculates spike threshold related measures of the spike\_shape, s.

### **Usage:**

```
[init_val, init_idx, rise_time, amplitude, peak_mag, peak_idx, max_dlo, a_plot] =
calcInitVm(s, max_idx, min_idx)
```

### **Parameters:**

**s:** A spike\_shape object.

max\_idx: The index of the maximal point of the spike\_shape [dt].

min\_idx: The index of the minimal point of the spike\_shape [dt].

**plotit:** If non-zero, plot a graph annotating the test results (optional).

### **Returns:**

init\_val: The potential value [dy]. init\_idx: Its index in the spike\_shape [dt]. rise\_time: Time from initiation to maximum [dt]. amplitude: Magnitude from initiation to max [dy]. peak\_mag: Peak value [dy]. peak\_idx: Extrapolated spike peak index [dt]. max\_d1o: Maximal value of first voltage derivative [dy]. a\_plot: plot\_abstract, if requested.

See also: spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/02

# A.39.13 Method spike\_shape/calcInitVmSlopeThreshold

Summary: Calculates the AP threshold using the slope threhold crossing.

#### Usage:

```
[init_idx, a_plot] = calcInitVmSlopeThreshold(s, max_idx, min_idx, thr, plotit)
```

#### **Parameters:**

```
s: A spike_shape object.
```

max\_idx: The index of the maximal point of the spike\_shape [dt].

min\_idx: The index of the minimal point of the spike\_shape [dt].

**thr:** Threshold for time derivative of voltage.

**plotit:** If non-zero, plot a graph annotating the test results (optional).

### **Returns:**

init\_idx: AP threshold index in the spike\_shape [dt]. a\_plot: plot\_abstract, if requested.

See also: calcInitVm (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/24

# A.39.14 Method spike\_shape/plotCompareMethodsSimple

**Summary:** Creates a multi-plot comparing different action potential threshold finding methods.

#### **Usage:**

```
a_plot = plotCompareMethodsSimple(s, title_str)
```

### **Parameters:**

```
s: A spike_shape object.
```

title\_str: Title suffix (optional).

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: spike\_shape (p. 183), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/19

# A.39.15 Method spike\_shape/calcInitVmMaxCurvature

**Summary:** Calculates the action potential threshold using the maximum of the curvature equation.

### Usage:

```
[init_idx, a_plot] = calcInitVmMaxCurvature(s, max_idx, min_idx, plotit)
```

**Description:** Point of maximum curvature:  $Kp = V''[1 + (V')^2]^{-3/2}$  Taken from Sekerli, Del Negro, Lee and Butera. IEEE Trans. Biomed. Eng., 51(9): 1665-71, 2004.

### **Parameters:**

```
s: A spike_shape object.
max_idx: The index of the maximal point of the spike_shape [dt].
min_idx: The index of the minimal point of the spike_shape [dt].
plotit: If non-zero, plot a graph annotating the test results (optional).
```

#### **Returns:**

init\_idx: AP threshold index in the spike\_shape [dt]. a\_plot: plot\_abstract, if requested.

See also: calcInitVm (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/19

# A.39.16 Method spike\_shape/calcInitVmV2PPLocal

**Summary:** Calculates the action potential threshold by finding the local second derivative maximum in voltage-time slope versus voltage phase plane, nearest a slope threshold crossing.

#### **Usage:**

```
[init_idx, a_plot] = calcInitVmV2PPLocal(s, max_idx, min_idx, lo_thr, plotit)
```

# **Parameters:**

```
s: A spike_shape object.
max_idx: The index of the maximal point of the spike_shape [dt].
min_idx: The index of the minimal point of the spike_shape [dt].
lo_thr: Lower threshold for time voltage slope.
plotit: If non-zero, plot a graph annotating the test results (optional).
```

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### **Returns:**

init\_idx: Its index in the spike\_shape [dt]. a\_plot: plot\_abstract, if requested.

See also: calcInitVm (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/18

# A.39.17 Method spike\_shape/getResults

Summary: Runs all tests defined by this class and return them in a structure.

**Usage:** 

```
[results, a_plot] = getResults(s, plotit)
```

#### Parameters:

```
s: A spike_shape object.
```

**plotit:** If non-zero, plot a graph annotating the test results (optional).

#### **Returns:**

results: A structure associating test names to values in ms and mV. a\_plot: plot\_abstract, if requested.

See also: spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/02

# A.39.18 Method spike\_shape/calcWidthFall

**Summary:** Calculates the spike width and fall information of the spike\_shape, s.

**Usage:** 

```
[base_width, half_width, half_Vm, fall_time, min_idx, min_val, max_ahp,
ahp_decay_constant, dahp_mag, dahp_idx] = ... calcWidthFall(s, max_idx, max_val,
init_idx, init_val)
```

**Description:** max\_\* can be the peak\_\* from calcInitVm.

# **Parameters:**

```
s: A spike_shape object.
```

max\_idx: The index of the maximal point [dt].

max\_val: The value of the maximal point [dy].

init\_idx: The index of spike initiation point [dt].

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init\_val: The value of spike initiation point [dy].

#### Returns:

base\_width: Width of spike at base [dt] half\_width: Width of spike at half\_Vm [dt] half\_Vm: Half height of spike [dy] fall\_time: Time from peak to initialization level [dt]. min\_idx: The index of the minimal point of the spike\_shape [dt]. max\_ahp: Magnitude from initiation to minimum [dy]. ahp\_decay\_constant: Approximation to refractory decay after maxAHP [dt]. dahp\_mag: Magnitude of the double AHP peak dahp\_mag: Index of the double AHP peak

See also: spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/02

# A.39.19 Method spike\_shape/calcInitVmLtdMaxCurv

**Summary:** Calculates the action potential threshold using the maximum of the curvature equation only in the limited range given with two voltage slope thresholds.

### **Usage:**

[init\_idx, a\_plot] = calcInitVmLtdMaxCurv(s, max\_idx, min\_idx, lo\_thr, hi\_thr,
plotit)

**Description:** Point of maximum curvature:  $Kp = V''[1 + (V')^2]^{-3/2}$  Taken from Sekerli, Del Negro, Lee and Butera. IEEE Trans. Biomed. Eng., 51(9): 1665-71, 2004.

### **Parameters:**

**s:** A spike\_shape object.

max\_idx: The index of the maximal point of the spike\_shape [dt].

min\_idx: The index of the minimal point of the spike\_shape [dt].

lo\_thr, hi\_thr: Lower and higher thresholds for time derivative of voltage.

**plotit:** If non-zero, plot a graph annotating the test results (optional).

### **Returns:**

init\_idx: AP threshold index in the spike\_shape [dt]. a\_plot: plot\_abstract, if requested.

See also: calcInitVm (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/19

# A.39.20 Method spike\_shape/plotPP

**Summary:** Plots the dV/dt vs. V phase-plane representation of the spike shape.

### Usage:

```
a_plot = plotPP(s)
```

#### **Parameters:**

**s:** A spike\_shape object.

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: spike\_shape (p. 183), plot\_abstract (p. 153)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/11/16

# A.39.21 Method spike\_shape/plotResults

**Summary:** Plots the spike shape annotated with result characteristics.

### Usage:

```
a_plot = plotResults(s, title_str, props)
```

## **Parameters:**

**s:** A spike\_shape object.

### **Returns:**

a\_plot: A plot\_abstract object that can be visualized. title\_str: (Optional) String to append to plot title. props: A structure with any optional properties, passed to trace/plotData.

See also: spike\_shape (p. 183), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/17

### A.39.22 Method spike\_shape/calcInitVmMaxCurvPhasePlane

**Summary:** Calculates the voltage at the maximum curvature in the phase plane as action potential threshold.

### Usage:

```
[init_idx, max_dlo, a_plot, fail_cond] = calcInitVmMaxCurvPhasePlane(s, max_idx, min_idx, plotit)
```

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**Description:** First take the phase-plane v'-v from the beginning to  $\max(v')$ . Then regulate intervals by interpolation. Point of maximum curvature:  $Kp = V''[1 + (V')^2]^{-3/2}$  Taken from Sekerli, Del Negro, Lee and Butera. IEEE Trans. Biomed. Eng., 51(9): 1665-71, 2004.

#### **Parameters:**

```
s: A spike_shape object.
max_idx: The index of the maximal point of the spike_shape [dt].
min_idx: The index of the minimal point of the spike_shape [dt].
plotit: If non-zero, plot a graph annotating the test results (optional).
```

#### **Returns:**

init\_idx: AP threshold index in the spike\_shape [dt]. max\_d1o: Maximal value of first voltage derivative [dy]. a\_plot: plot\_abstract, if requested. fail\_cond: True if this algorithm fails to be trustable.

See also: calcInitVm (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/12

# A.40 Class spike\_shape\_profile

# A.40.1 Constructor spike\_shape\_profile/spike\_shape\_profile

Summary: Holds the results profile from a spike\_shape object.

### **Usage:**

```
a_ss_profile = spike_shape_profile(results, a_spike_shape, props)
```

#### **Parameters:**

```
results: A structure containing test results.

a_spike_shape: A spike_shape object.

props: A structure with any optional properties.
```

# Returns a structure object with the following fields:

results\_profile: Contains results of tests. spike\_shape: The spike\_shape object from which results were obtained. props.

See also: results\_profile (p. 176)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/08/17

# A.40.2 Method spike\_shape\_profile/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.40.3 Method spike\_shape\_profile/plot\_abstract

Summary: Plots the spike shape with measurements marked in red.

Usage:

```
a_plot = plot_abstract(s, props)
```

# Parameters:

**s:** A spike\_shape object.

**props:** A structure with any optional properties.

absolute\_peak\_time: Shift the peak to this point on the plot.

no\_plot\_spike: Do not plot the spike shape first.

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: spike\_shape (p. 183), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/08/17

# A.41 Class spikes

# A.41.1 Constructor spikes/spikes

**Summary:** Spike times from a trace.

**Usage:** 

```
obj = spikes(times, num_samples, dt, id)
```

#### **Parameters:**

times: The spike times [dt].

num\_samples: Number of samples in the original trace.

**dt:** Time resolution [s].

**id:** Identification string.

# Returns a structure object with the following fields:

times, num\_samples, dt, id.

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**See also:** trace/spikes (p. 258), trace (p. 255), spike\_shape (p. 183), period (p. 137)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

# A.41.2 Method spikes/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.41.3 Method spikes/SFA

**Summary:** Calculates the spike frequency accommodation (SFA) of the inter-spike-intervals (ISI).

# Usage:

```
sfa = SFA(s, a\_period)
```

**Description:** SFA is the ration of the last ISI to the first ISI in the period.

#### **Parameters:**

s: A spikes object.

a\_period: The period where spikes were found (optional)

### **Returns:**

sfa: Spike frequency accommodation.

**See also:** spikes (p. 195), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/13

# A.41.4 Method spikes/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.41.5 Method spikes/periodWhole

**Summary:** Returns the boundaries of the whole period of spikes, s.

Usage:

```
whole_period = periodWhole(s)
```

### **Parameters:**

s: A spikes object.

See also: period (p. 137), spikes (p. 195)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

# A.41.6 Method spikes/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.41.7 Method spikes/plotData

Summary: Plots a spikes object.

Usage:

```
a_plot = plotData(s, title_str)
```

**Description:** If s is a vector of spikes objects, returns a vector of plot objects.

### **Parameters:**

s: A spikes object.

# **Returns:**

a\_plot: A plot\_abstract object that can be visualized. title\_str: (Optional) String to append to plot title.

See also: trace (p. 255), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/10/21

# A.41.8 Method spikes/plotISIs

Summary: Plots a spikes object.

**Usage:** 

```
a_plot = plotISIs(s, title_str)
```

**Description:** If s is a vector of spikes objects, returns a vector of plot objects.

#### **Parameters:**

s: A spikes object.

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized. title\_str: (Optional) String to append to plot title.

See also: trace (p. 255), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/10/21

# A.41.9 Method spikes/spikeRateISI

**Summary:** Calculates the firing rate of the spikes found in the given period with an averaged inter-spike-interval approach.

# Usage:

```
freq = spikeRateISI(s, trace_index, times, period)
```

### **Parameters:**

s: A spikes object.

**period:** The period where spikes were found (optional)

### **Returns:**

freq: Firing rate [Hz]

See also: trace (p. 255), spikes (p. 195), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/03/09

# A.41.10 Method spikes/plotFreqVsTime

Summary: Plots a frequency-time graph from the spikes object.

### Usage:

```
a_plot = plotFreqVsTime(s, title_str, props)
```

**Description:** If s is a vector of spikes objects, returns a vector of plot objects.

### **Parameters:**

s: A spikes object.

title\_str: (Optional) String to append to plot title.

**props:** A structure with any optional properties.

type: If 'simple' plots 1/is for each spike time,

'manhattan' uses flat lines of 1/isi height between spike times (de-

fault). (others passed to plot\_abstract)

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized. title\_str: (Optional) String to append to plot title.

See also: trace (p. 255), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/05

# A.41.11 Method spikes/addSpikes

### **Usage:**

```
s = addSpike(s, times)
```

### **Parameters:**

s: A spikes object.

times: Times of spikes to add

### **Returns:**

s: The updated object.

See also: spikes (p. 195)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/08/16

# A.41.12 Method spikes/subsref

**Summary:** Defines generic indexing for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.41.13 Method spikes/spikeAmpSlope

**Summary:** Calculates the time constant and steady-state value of the spike amplitude for slow inactivating decays.

## Usage:

```
[a_tau, da_inf] = spikeAmpSlope(a_spikes, a_trace, a_period)
```

#### **Parameters:**

a\_spikes: A spikes object.

a\_trace: A trace object.

a\_period: The desired period (optional)

### **Returns:**

a\_tau: Approximate amplitude decay constant. da\_inf: Delta change in final spike peak value from initial.

See also: period (p. 137), spikes (p. 195), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/15

# A.41.14 Method spikes/intoPeriod

Summary: Shifts the spikes times to be within the given period.

## Usage:

```
obj = intoPeriod(s, a_period)
```

**Description:** Assuming this spikes object's length fits into the given period, it shifts all times to start from the beginning of the given period. This may be used to reconstruct the original spikes object from subperiods that were cut out previously, using the withinPeriod method.

### **Parameters:**

**s:** A spikes object.

a\_period: The desired period

```
Returns:
```

obj: A spikes object

See also: spikes (p. 195), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/31

# A.41.15 Method spikes/plot

Summary: Plots spikes.

Usage:

h = plot(t)

### **Parameters:**

t: A spikes object.

title\_str: (Optional) String to append to plot title.

### **Returns:**

h: Handle to figure object.

See also: spikes (p. 195), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.41.16 Method spikes/withinPeriod

**Summary:** Returns a spikes object valid only within the given period, subtracts the offset.

### Usage:

```
obj = withinPeriod(s, a_period)
```

# Parameters:

s: A spikes object.

a\_period: The desired period

### **Returns:**

obj: A spikes object

**See also:** spikes (p. 195), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/31

# A.41.17 Method spikes/ISICV

**Summary:** Calculates the coefficient of variation (CV) of the inter-spike-intervals (ISI).

# Usage:

```
cv = ISICV(s, a_period)
```

### **Parameters:**

s: A spikes object.

a\_period: The period where spikes were found (optional)

### **Returns:**

cv: Coefficient of variation.

See also: spikes (p. 195), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/13

# A.41.18 Method spikes/getResults

Summary: Runs all tests defined by this class and return them in a structure.

# Usage:

```
results = getResults(s)
```

# **Parameters:**

s: A spikes object.

# **Returns:**

results: A structure associating test names to values in ms and mV (or mA).

See also: spikes (p. 195)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/13

# A.41.19 Method spikes/vertcat

**Summary:** Vertical concatanation [a\_spikes;with\_spikes;...] operator.

### **Usage:**

```
a_spikes = vertcat(a_spikes, with_spikes, ...)
```

**Description:** Concatanates spike times of with\_spikes with that of a\_spikes. Overrides the built-in vertcat function that is called when [a\_spikes;with\_spikes] is executed.

### **Parameters:**

```
a_spikes, with_spikes, ...: Spikes objects.
```

#### **Returns:**

a\_spikes: A tests\_spikes that contains times of all given spikes objects.

```
See also: vertcat (p. ??), spikes (p. 195)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/08/16

# A.41.20 Method spikes/spikeRate

Summary: Calculates the average firing rate [Hz] of the given spike train.

# Usage:

```
freq = spikeRate(s, a_period)
```

### **Parameters:**

s: A spikes object.

a\_period: The period where spikes were found (optional)

## **Returns:**

```
freq: Firing rate [Hz]
```

See also: spikes (p. 195), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/03/09

# A.41.21 Method spikes/withinPeriodWOffset

**Summary:** Returns a spikes object valid only within the given period, keeps the offset.

### Usage:

```
obj = withinPeriodWOffset(s, a_period)
```

### **Parameters:**

s: A spikes object.

a\_period: The desired period

#### **Returns:**

obj: A spikes object

**See also:** spikes (p. 195), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/05/09

# A.41.22 Method spikes/getISIs

**Summary:** Calculates the firing rate of the spikes found in the given period with an averaged inter-spike-interval approach.

# Usage:

```
isi = getISIs(s, period)
```

# Parameters:

s: A spikes object.

period: The period where spikes were found (optional)

### **Returns:**

isi: Inter-spike-interval vector [dt]

See also: trace (p. 255), spikes (p. 195), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/03/09

# A.42 Class spikes\_db

## A.42.1 Constructor spikes\_db/spikes\_db

**Summary:** A database of spike shape results obtained from a period in a trace.

Usage:

```
a_spikes_db = spikes_db(data, col_names, a_trace, a_period, id, props)
```

**Description:** This is a subclass of tests\_db. Use trace/analyzeSpikesInPeriod to get an instance of this class.

#### **Parameters:**

data: Database contents.

col names: The column names.

a\_trace: The trace where the spikes were found.

**a\_period:** The period inside a\_trace where spikes were found.

id: An identifying string.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

```
tests_db, trace, period, props.
```

See also: tests\_db (p. 217), trace (p. 255), period (p. 137), trace/analyzeSpikesInPeriod (p. 264)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/08/17

# A.42.2 Method spikes\_db/plot\_abstract

**Summary:** Visualizes the spikes\_db by marking spike shapes measurements on the trace plot.

### **Usage:**

```
a_pm = plot_abstract(a_db, title_str, props)
```

# **Parameters:**

a\_db: A spikes\_db object.

title\_str: (Optional) A string to be concatanated to the title.

**props:** A structure with any optional properties passed to trace/plotData.

## **Returns:**

a\_pm: A trace plot.

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```
See also: plot_abstract/plot_abstract (p. 153), tests_db/plot_abstract (p. 237), plotFigure (p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/08/17

# A.43 Class stats\_db

# A.43.1 Constructor stats\_db/stats\_db

**Summary:** A database of rows corresponding to statistical distribution properties of tests. Multiple pages can be used to indicate another dimension.

## **Usage:**

```
a_stats_db = stats_db(test_results, col_names, row_names, page_names, id, props)
```

**Description:** This is a subclass of tests\_3D\_db. Allows generating a plot, etc.

#### **Parameters:**

```
test_results: The 3-d array of rows, columns, and pages.
```

col\_names: Test names in this db.

row\_names: Statistical test names for each row.

page names: Meaning of each separate page of data

(e.g., a different invariant parameter).

id: An identifying string.

**props:** A structure with any optional properties.

**axis\_limits:** Limits in the form of [xmin xmax ymin ymax] for errorbar axes.

yTicksPos: 'left' means only put y-axis ticks to leftmost plot. xTicksPos: 'bottom' means only put x-axis ticks to lowest plot.

## Returns a structure object with the following fields:

```
tests_3D_db.
```

```
See also: tests_3D_db (p. 212), plot_abstract (p. 153)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

# A.43.2 Method stats\_db/onlyRowsTests

**Summary:** Returns a tests\_db that only contains the desired tests and rows (and pages).

Usage:

```
obj = onlyRowsTests(obj, rows, tests, pages)
```

**Description:** Selects the given dimensions and returns in a new tests\_db object.

#### **Parameters:**

**obj:** A tests\_db object.

rows: A logical or index vector of rows. If ':', all rows.

tests: Cell array of test names or column indices. If ':', all tests.

pages: (Optional) A logical or index vector of pages. ':' for all pages.

#### **Returns:**

obj: The new tests\_db object.

See also: subsref (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17

# A.43.3 Method stats\_db/get

Summary: Defines generic attribute retrieval for objects.

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.43.4 Method stats\_db/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.43.5 Method stats\_db/plotVar

**Summary:** Generates a plot of the variation between two tests.

#### **Usage:**

```
a_plot = plotVar(a_stats_db, test1, test2, props)
```

**Description:** Creates a plot description where the mean values are used for solid lines and the std values of test2 is indicated with errorbars. It is assumed that each page of the stats\_db contains a value to be matched.

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### **Parameters:**

```
a_stats_db: A stats_db object.
```

**test1:** Test column for the x-axis, only mean values are used.

**test2:** Test column for the y-axis, std values are indicated with errorbars.

props: Optional properties.

plotType: 1, only errorbars (default); 2, errorbars extending from bars.
 (rest passed to plot\_abstract)

#### **Returns:**

a\_plot: A plot\_abstract object or one of its subclasses.

See also: plotVar (p. ??), plot\_simple (p. 163)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/13

# A.43.6 Method stats\_db/plotColorVar

Summary: Create a color-plot of parameter-test variations in a matrix.

## **Usage:**

```
a_plot = plotColorVar(p_stats, props)
```

**Description:** Skips the 'ItemIndex' test.

### **Parameters:**

**p\_stats:** Array of invariant parameter databases obtained from calling tests\_3D\_db/paramsTestsHistsStats.

title\_str: (Optional) String to append to plot title.

**props:** A structure with any optional properties, passed to plot\_stack.

plotMethod: 'plotVar' uses stats\_db/plotVar (default)
 'plot\_bars' uses stats\_db/plot\_bars

### **Returns:**

a\_plot: A plot\_abstract with the color plot

See also: paramsTestsHistsStats(p.??), params\_tests\_profile(p. 136), plotVar.(p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/17

# A.43.7 Method stats\_db/plotVarMatrix

**Summary:** Create a stack of parameter-test variation plots organized in a matrix.

### **Usage:**

```
a_plot_stack = plotVarMatrix(p_stats, props)
```

**Description:** Skips the 'ItemIndex' test.

#### **Parameters:**

**p\_stats:** Array of invariant parameter databases obtained from calling tests\_3D\_db/paramsTestsHistsStats.

**props:** A structure with any optional properties, passed to plot\_stack.

plotMethod: 'plotVar' uses stats\_db/plotVar (default)
 'plot\_bars' uses stats\_db/plot\_bars

rotateYLabel: Rotate row labels this much (default=60).

#### **Returns:**

a\_plot\_stack: A plot\_stack with the plots organized in matrix form

See also: paramsTestsHistsStats (p.??), params\_tests\_profile (p. 136), plotVar. (p.??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/17

### A.43.8 Method stats\_db/plot\_abstract

**Summary:** Generates an error bar graph for each db columns. Looks for 'min', 'max', and 'STD' labels in the row\_idx for drawing the errorbars.

### Usage:

```
a_plot = plot_abstract(a_stats_db, title_str, props)
```

**Description:** Generates a plot\_simple object from this histogram.

#### **Parameters:**

a\_stats\_db: A histogram\_db object.
title\_str: A title string on the plot

**props:** A structure with any optional properties.

### **Returns:**

a\_plot: A object of plot\_abstract or one of its subclasses.

See also: plot\_abstract (p. 153), plot\_simple (p. 163)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

### A.43.9 Method stats\_db/subsref

**Summary:** Defines generic indexing for objects.

# A.43.10 Method stats\_db/compareStats

**Summary:** Merges multiple stats\_dbs into pages of a single stats\_db for comparison.

Usage:

```
a_mult_stats_db = compareStats(a_stats_db)
```

**Description:** Generates a plot\_simple object from this histogram.

#### **Parameters:**

```
a_stats_db: A histogram_db object.
```

command: Plot command (Optional, default='bar')

#### **Returns:**

a\_mult\_stats\_db: A object of compareStats or one of its subclasses.

See also: plot\_abstract (p. 153), plot\_simple (p. 163)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

### A.43.11 Method stats\_db/plotYTests

**Summary:** Create an errorbar plot of database stats measures against given X-axis values.

### **Usage:**

```
a_p = plotYTests(a_stats_db, x_vals, tests, axis_labels, title_str, short_title,
command, props)
```

#### **Parameters:**

```
a_stats_db: A params_tests_db object.
```

**x\_vals:** A vector of X-axis values.

**tests:** A vector or cell array of columns to correspond to each value from  $x_{vals}$ .

title\_str: (Optional) A string to be concatanated to the title.

**short\_title:** (Optional) Few words that may appear in legends of multiplot.

command: (Optional) Command to do the plotting with (default: 'plot')

**props:** A structure with any optional properties.

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**LineStyle:** Plot line style to use. (default: 'd-') **quiet:** If 1, don't include database name on title.

#### **Returns:**

a\_p: A plot\_abstract.

#### **Example:**

```
» a_p = plotYTests(a_stats_db, [0 40 100 200], ...
'IniSpontSpikeRateISI_0pA', 'PulseIni100msSpikeRateISI_D40pA', ...
'PulseIni100msSpikeRateISI_D100pA', 'PulseIni100msSpikeRateISI_D200pA', ...
'current pulse [pA]', 'firing rate [Hz]', ', f-I curves', 'neuron 1');
» plotFigure(a_p);
```

See also: plotFigure (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/23

# A.43.12 Method stats\_db/plot\_bars

**Summary:** Creates a bar graph with errorbars for each db column.

#### **Usage:**

```
a_plot = plot_bars(a_stats_db, title_str, props)
```

**Description:** Looks for 'min', 'max', and 'STD' labels in the row\_idx for drawing the errorbars. Each page of the DB will produce grouped bars.

### **Parameters:**

```
a_stats_db: A stats_db object.
title_str: The plot title.
```

**props:** A structure with any optional properties, passed to plot\_bars/plot\_bars.

pageVariable: The column used for denoting page values.

### **Returns:**

a\_plot: A object of plot\_bars or one of its subclasses.

See also: plot\_abstract (p. 153), plot\_bars/plot\_bars (p. 159)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.44 Class tests\_3D\_db

#### A.44.1 Constructor tests\_3D\_db/tests\_3D\_db

**Summary:** A database multiple pages with rows of test columns. Each page may represent aspects of the data that are different, but not defined in this object.

### Usage:

```
a_3D_db = tests_3D_db(data, col_names, row_names, page_names, id, props)
```

**Description:** This is a subclass of tests\_db. Usually it contains a RowIndex column that points to an original db from which this data originated. The row indices can be used to reach the values associated with different pages of information contained in this object.

#### **Parameters:**

data: The 3-d vector of rows, columns, and pages.

col\_names: Colun names of the database.

id: An identifying string.

**props:** A structure with any optional properties.

invarName: Name of the invariant parameter for this db.

### Returns a structure object with the following fields:

tests\_db, page\_idx.

See also: tests\_db (p. 217), tests\_db/invarValues (p. 233)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/30

## A.44.2 Method tests\_3D\_db/joinPages

**Summary:** Joins the rows of the given db to the with\_db rows matching with the PageIndex column.

### Usage:

```
a_db = joinPages(db, tests, with_db, w_tests)
```

**Description:** Replicates the desired columns in the with\_db with rows having a page index and joins them next to desired columns from the current 3D\_db. Flattens the resulting 3D\_db to become a 2D db. Assumes each page index only appears once in with\_db.

## **Parameters:**

db: A tests\_3D\_db object.

with\_db: A tests\_db object with a PageIndex column.

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### **Returns:**

a\_db: A tests\_db object.

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/15

# A.44.3 Method tests\_3D\_db/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.44.4 Method tests\_3D\_db/diff2D

**Summary:** Creates a tests\_db by taking the derivative of the given test.

### Usage:

```
a_tests_db = diff2D(a_db, test, props)
```

**Description:** Applies the diff function to the chosen test, and collapses the middle dimension of the 3D DB to create a 2D DB.

### **Parameters:**

a\_db: A tests\_3D\_db object.

test: Test column.

props: Optional properties.

### **Returns:**

a\_tests\_db: A tests\_db that holds the requested differences of parameter values.

See also: boxplot (p. ??), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/05/22

# A.44.5 Method tests\_3D\_db/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

#### A.44.6 Method tests\_3D\_db/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

# A.44.7 Method tests\_3D\_db/plotParamPairImage

**Summary:** Generates an image plot of variation of a test with two parameters in the first page.

## **Usage:**

```
a_plot = plotParamPairImage(a_db, test, title_str, props)
```

**Description:** It is assumed that the 3D DB is created by invariant combinations of two parameters, which are the first two columns. Each page of the db must contain a same parameter values. This is the default character of tests\_3D\_db created by params\_tests\_db/invarParam. Parameter values will be enumerated and then an image plot is created.

#### **Parameters:**

```
a_db: A tests_3D_db object.
```

test: Test column to take the measure value.

title str: (Optional) String to append to plot title.

**props:** Optional properties to be passed to plot\_abstract.

truncateDecDigits: Truncate labels to this many decimal digits.

labelSteps: Skip this many labels between ticks to reduce to total

number.

**maxValue:** Maximal value to normalize colors and to annotate the colorbar.

## **Returns:**

a\_plot: A plot\_abstract object or one of its subclasses.

### Example:

### A.44.8 Method tests\_3D\_db/histograms

#### **Usage:**

```
a_histogram_db = histogram(db, col, num_bins)
```

**Description:** If one wants to get histograms of test values for each single value of the selected invariant parameter, then swapRowsPages should be done first on db.

### **Parameters:**

**db:** A tests\_3D\_db object.

col: Column to find the histogram.

num\_bins: Number of histogram bins (Optional, default=100)

### **Returns:**

a\_histogram\_db: A histogram\_db object containing the histogram.

See also: histogram\_db (p. 94), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/04

# A.44.9 Method tests\_3D\_db/swapRowsPages

**Summary:** Swaps the row dimension with the page dimension of the tests\_3D\_db.

# **Usage:**

```
a_3D_db = swapRowsPages(db)
```

**Description:** Assuming that this is a invariant parameter and tests relations db, this function swaps the pages with rows. Each resulting page correspond to a single value of the chosen parameter, with each row contianing a test result with different combinations of the rest of the parameters.

### **Parameters:**

**db:** A tests\_db object.

### **Returns:**

a\_3D\_db: A tests\_3D\_db object.

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/04

# A.44.10 Method tests\_3D\_db/paramsTestsHistsStats

Summary: Calculates histograms and statistics for DB.

**Usage:** 

```
[pt_hists, p_stats] = paramsTestsHistsStats(p_t3ds, props)
```

**Description:** Calculates histograms and statistics for all combinations of tests and params and returns them in a cell array. Skips the 'ItemIndex' test.

#### **Parameters:**

**p\_t3ds:** Array of invariant parameter databases obtained by calling the params\_tests\_db/invarParams method.

props: Optional properties.

**statsMethod:** method to call to get a stats\_db (default='statsMeanSE') **useDiff:** If 1, takes the derivative with diff on the 3D DBs (default=0).

#### **Returns:**

pt\_hists: An array of 3D histograms for each pair of param and test. p\_stats: An array of stats\_dbs for each param.

See also: invarParams (p. ??), params\_tests\_profile (p. 136)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/17

# A.44.11 Method tests\_3D\_db/mergePages

**Summary:** Merges tests from separate pages into a 2D params\_tests\_db.

Usage:

```
a_db = mergePages(db, page_tests, page_suffixes)
```

**Description:** Keeps uniqueness by adding suffixes to test names. If you're using invarParams, do swapRowsPages, then join with original db to get the parameter values.

#### **Parameters:**

db: A tests\_3D\_db object.

page\_tests: Cell array of list of tests to take from each page.

page\_suffixes: Cell array of suffixes to append to tests from each page.

#### **Returns:**

a\_db: A tests\_db object.

See also: tests\_db (p. 217), tests\_3D\_db (p. 212)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2005/01/13

# A.44.12 Method tests\_3D\_db/plotVarBox

**Summary:** Generates a boxplot of the variation between two tests.

#### Usage:

```
a_plot = plotVarBox(a_db, test1, test2, notch, sym, vert, whis, props)
```

**Description:** It is assumed that each page of the db contains a different parameter value.

#### **Parameters:**

```
a_db: A tests_3D_db object.
```

**test1:** Test column for the x-axis, only mean values are used.

test2: Test column for the y-axis, used for boxplot.

notch, sym, vert, whis: See boxplot, defaults = (1, '+', 1, 1.5).

props: Optional properties to be passed to plot\_abstract.

#### **Returns:**

a\_plot: A plot\_abstract object or one of its subclasses.

See also: boxplot (p. ??), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/10

## A.45 Class tests\_db

# A.45.1 Constructor tests\_db/tests\_db

Summary: A generic database of test results organized in a matrix format.

# Usage:

```
obj = tests_db(test_results, col_names, row_names, id, props)
```

**Description:** Defines all operations on this structure so that subclasses can use them.

#### **Parameters:**

```
test_results: A matrix that contains columns associated with tests and rows for separate observations.
```

col\_names: Cell array of column names of test\_results.

row\_names: Cell array of row names of test\_results.

id: An identifying string.

**props:** A structure with any optional properties.

# Returns a structure object with the following fields:

data: The data matrix. row\_idx, col\_idx: Structure associating row/column names to indices. id, props.

```
See also: params_tests_db (p. 116), params_db (p. ??), test_variable_db (N/I) (p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/01

# A.45.2 Method tests\_db/eq

**Summary:** Equality (==) operator. Returns logical indices of db rows that match with given row.

## **Usage:**

```
rows = eq(db, row)
```

#### Parameters:

db: A tests\_db object.

row: Row array to be compared with db rows.

#### **Returns:**

rows: A logical or index vector to be used in indexing db objects.

See also: eq (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17

# A.45.3 Method tests\_db/ge

**Summary:** Greater or equal to (>=) operator. Returns logical indices of db rows that are greater than or equal to given row.

# **Usage:**

```
rows = ge(db, row)
```

#### **Parameters:**

**db:** A tests\_db object.

row: Row array to be compared with db rows.

#### **Returns:**

rows: A logical or index vector to be used in indexing db objects.

**See also:** ge (p. ??), tests\_db (p. 217)

# A.45.4 Method tests\_db/gt

**Summary:** Greater than (>) operator. Returns logical indices of db rows that are greater than given row.

## **Usage:**

```
rows = gt(db, row)
```

## **Parameters:**

db: A tests\_db object.

row: Row array to be compared with db rows.

# **Returns:**

rows: A logical or index vector to be used in indexing db objects.

See also: gt (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17

# A.45.5 Method tests\_db/le

**Summary:** Less or equal (<=) operator. Returns logical indices of db rows that are less than or equal to given row.

# Usage:

```
rows = le(db, row)
```

## Parameters:

**db:** A tests\_db object.

row: Row array to be compared with db rows.

# **Returns:**

rows: A logical or index vector to be used in indexing db objects.

**See also:** le (p. ??), tests\_db (p. 217)

# A.45.6 Method tests\_db/lt

**Summary:** Less than (<) operator. Returns logical indices of db rows that are less than given row.

## **Usage:**

```
rows = lt(db, row)
```

## **Parameters:**

**db:** A tests\_db object.

row: Row array to be compared with db rows.

# **Returns:**

rows: A logical or index vector to be used in indexing db objects.

See also: lt (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17

# A.45.7 Method tests\_db/ne

Summary: Returns logical indices of db rows that doesn't match with given row.

# Usage:

```
rows = ne(db, row)
```

# **Parameters:**

db: A tests\_db object.

row: Row array to be compared with db rows.

# **Returns:**

rows: A logical or index vector to be used in indexing db objects.

See also: ne (p. ??), tests\_db (p. 217)

# A.45.8 Method tests\_db/onlyRowsTests

**Summary:** Returns a tests\_db that only contains the desired tests and rows (and pages).

# Usage:

```
obj = onlyRowsTests(obj, rows, tests, pages)
```

**Description:** Selects the given dimensions and returns in a new tests\_db object.

#### **Parameters:**

**obj:** A tests\_db object.

rows: A logical or index vector of rows. If ':', all rows.

tests: Cell array of test names or column indices. If ':', all tests.

pages: (Optional) A logical or index vector of pages. ':' for all pages.

#### **Returns:**

obj: The new tests\_db object.

See also: subsref (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17

# A.45.9 Method tests\_db/setProp

Summary: Generic method for setting optional object properties.

# Usage:

```
obj = setProp(obj, prop1, val1, prop2, val2, ...)
```

**Description:** Modifies or adds property values. As many property name-value pairs can be specified.

## **Parameters:**

**obj:** Any object that has a props field.

**attr:** Property name **val:** Property value.

### **Returns:**

obj: The new object with the updated properties.

### See also:

### A.45.10 Method tests\_db/isnanrows

**Summary:** Finds rows with any NaN values. Returns logical indices of db rows.

# Usage:

```
rows = isnanrows(db)
```

**Description:** Some operations need that no NaN values exist in the matrix. This method can be used to find and then remove NaN-contaminated rows from DB. Note that sometimes no rows can be found, and some columns should be discarded before this operation.

#### **Parameters:**

**db:** A tests\_db object.

#### **Returns:**

rows: A logical vector to be used in indexing db objects or passed through other logical operators.

See also: isnan (p.??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/08

### A.45.11 Method tests\_db/joinRows

**Summary:** Joins the rows of the given db with rows of with\_db with matching RowIndex values.

#### **Usage:**

```
a_db = joinRows(db, tests, with_db, w_tests, index_col_name)
```

**Description:** Takes the desired columns in with\_db with rows having a row index and joins them next to desired columns from the current db. Assumes each row index only appears once in with\_db. The created db preserves the ordering of with\_db.

### **Parameters:**

**db:** A param\_tests\_db object.

tests: Test columns to take from db.

with\_db: A tests\_db object with a RowIndex column.

w\_tests: Test columns to take from with\_db.

index\_col\_name: (Optional) Name of row index column (default='RowIndex').

#### **Returns:**

a\_db: A tests\_db object.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/16

#### A.45.12 Method tests\_db/setRows

**Summary:** Sets the rows of observations in tests\_db.

Usage:

index = setRows(obj, rows)

**Description:** Sets a new set of observations to the database and returns the new DB.

**Parameters:** 

**obj:** A tests\_db object.

rows: A matrix that contains rows for the DB.

**Returns:** 

obj: The tests\_db object with the new rows.

See also: allocateRows (p. ??), addRow (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/08

## A.45.13 Method tests\_db/plotTestsHistsMatrix

**Summary:** Create a matrix plot of test histograms.

**Usage:** 

a\_pm = plotTestsHistsMatrix(a\_db, title\_str, props)

**Description:** Skips the 'ItemIndex' test.

**Parameters:** 

a\_db: A params\_tests\_db object.

title\_str: (Optional) A string to be concatanated to the title.

**props:** A structure with any optional properties, passed to plot\_abstract.

orient: Orientation of the plot\_stack. 'x', 'y', or 'matrix' (default).

histBins: Number of histogram bins. quiet: Don't put the DB id on the title.

**axisLimits:** Only x-ranges are used from this expression.

### **Returns:**

a\_pm: A plot\_stack with the plots organized in matrix form

See also: params\_tests\_profile (p. 136), plotVar (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/17

# A.45.14 Method tests\_db/crossProd

**Summary:** Create a DB by taking the cross product of two database row sets.

Usage:

```
cross_db = crossProd(a_db, b_db)
```

**Description:** This is not a vector cross product operation. Each row of the two DBs are matched and added as a new row to a DB. The end is a DB with all combinations of rows from both DBs. The final DB contains columns of both DBs.

#### **Parameters:**

```
a_db, b_db: A tests_db object.
```

#### **Returns:**

cross\_db: The tests\_db object with all combinations of rows.

See also: allocateRows (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/10/11

# A.45.15 Method tests\_db/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.45.16 Method tests\_db/testsHists

Summary: Calculates histograms for all tests.

Usage:

```
t_hists = testsHists(a_db, num_bins)
```

### **Parameters:**

```
a_db: A tests_db object.
```

**num\_bins:** Number of histogram bins (Optional, default=100), or vector of histogram bin centers.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2407-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

### **Returns:**

t\_hists: An array of histograms for each test in a\_db.

See also: params\_tests\_profile (p. 136)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/27

# A.45.17 Method tests\_db/mtimes

Summary: Multiplies the DB with a scalar.

Usage:

```
a_db = mtimes(left_obj, right_obj)
```

### **Parameters:**

left\_obj, right\_obj: Operands of the multiplication. One must be of type tests\_db.

#### **Returns:**

a\_db: The resulting tests\_db.

See also: tests\_db/times (p. 236), mtimes (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/24

# A.45.18 Method tests\_db/histogram

**Summary:** Generates a histogram\_db object with rows corresponding to histogram entries.

## **Usage:**

```
a_histogram_db = histogram(db, col, num_bins)
```

### **Parameters:**

db: A tests\_db object.

**col:** Column to find the histogram.

**num\_bins:** Number of histogram bins (Optional, default=100), or vector of histogram bin centers.

#### **Returns:**

a\_histogram\_db: A histogram\_db object containing the histogram.

## **Example:**

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2507-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

```
» a_hist_db = histogram(my_db, 'spike_width');
» plot(a_hist_db);
       See also: histogram_db (p. 94), tests_db (p. 217)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17
       A.45.19 Method tests_db/cov
       Summary: Generates a database of the covariance of given DB.
       Usage:
a_cov_db = cov(db, props)
       Parameters:
             db: A tests_db object.
             props: A structure with any optional properties.
                 keepOrigDB: Keep db as origDB in the props.
                      (others passed to tests_db)
       Returns:
             a_cov_db: A tests_db which contains the covariance matrix.
       See also: cov (p. ??)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2007/05/25
       A.45.20 Method tests_db/end
       Summary: Overloaded primitive matlab function, returns maximal dimension size.
       Usage:
s = end(db, index, total)
       Parameters:
             db: A tests_db object.
       Returns:
             s: The size.
       See also: size (p. ??), tests_db (p. 217)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/06
```

# A.45.21 Method tests\_db/get

**Summary:** Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.45.22 Method tests\_db/sortrows

**Summary:** Returns a sorted\_db according to given columns.

Usage:

```
[sorted_db, idx] = sortrows(db, cols)
```

**Description:** For multi-page dbs, sorts only the first page and applies the ordering to all other pages (which is WRONG!)

#### **Parameters:**

db: A tests\_db object.

cols: Columns to use for sorting.

#### **Returns:**

sorted\_db: The sorted tests\_db. idx: The row index permutation vector, such that sorted\_db = db(idx, :).

See also: sortrows (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/11

# A.45.23 Method tests\_db/set

Summary: Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

## A.45.24 Method tests\_db/std

**Summary:** Returns the std of the data matrix of a\_db. Ignores NaN values.

Usage:

```
[a_db, n] = std(a_db, sflag, dim)
```

**Description:** Does a recursive operation over dimensions in order to remove NaN values. This takes considerable amount of time compared with a straightforward std operation.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

## **Parameters:**

a\_db: A tests\_db object.dim: Work down dimension.

#### **Returns:**

a\_db: The DB with std values. n: (Optional) Numbers of non-NaN rows included in calculating each column.

**See also:** std (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/06

## A.45.25 Method tests\_db/sum

**Summary:** Creates a tests\_db by summing all rows.

**Usage:** 

```
a_db = sum(a_db, props)
```

**Description:** Applies the sum function to whole DB. The resulting DB will have one row.

## **Parameters:**

a\_db: A tests\_db object.props: Optional properties.

#### **Returns:**

a\_db: The resulting tests\_db.

See also: sum (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/24

# A.45.26 Method tests\_db/kmeansCluster

**Summary:** Generates a database of cluster centers obtained from a k-means cluster analysis with the command kmeans.

## Usage:

```
a_cluster_db = kmeansCluster(db, num_clusters, props)
```

### **Parameters:**

**db:** A tests\_db object.

PANDORA Toolbox: Prog.'s MaDraft v805 of 2807-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

num\_clusters: Number of clusters to form.
props: A structure with any optional properties.

**DistanceMeasure:** Choose one appropriate for kmeans.

#### **Returns:**

a\_cluster\_db: A tests\_db where each row is a cluster center. a\_hist\_db: histogram\_db showing cluster membership from original db. idx: Cluster indices of each row or original db. sum\_distances: Quality of clustering indicated by total distance from centroid for each cluster.

See also: tests\_db (p. 217), histogram\_db (p. 94)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/06

### A.45.27 Method tests\_db/noNaNRows

Summary: Returns a DB by removing rows containing any NaN or Inf.

Usage:

```
a_db = noNaNRows(a_db)
```

# Parameters:

```
a_db: A tests_db object.
```

#### **Returns:**

a\_db: DB with missing rows.

See also: tests\_db/isnanrows (p. 222)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/09/21

# A.45.28 Method tests\_db/statsMeanStd

**Summary:** Generates a stats\_db object with two rows corresponding to the mean and std of the tests' distributions.

## **Usage:**

```
a_stats_db = statsMeanStd(db, tests, props)
```

### **Parameters:**

db: A tests\_db object.

tests: A selection of tests (see onlyRowsTests).

**props:** A structure with any optional properties for stats\_db.

PANDORA Toolbox: Prog.'s MaBuraft v805 of 2007-08-02 11:58:12 -0400 (Thu, 02 Aug 2007)

### **Returns:**

```
a_stats_db: A stats_db object.
```

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

# A.45.29 Method tests\_db/checkConsistentCols

**Summary:** Check if two DBs have exactly the same columns.

**Usage:** 

```
[col_names, with_col_names] = checkConsistentCols(db, with_db)
```

#### **Parameters:**

**db:** A tests\_db object.

with\_db: A tests\_db object whose column names are checked for consistency.

**props:** A structure with any optional properties.

**useCommon:** Tolerate mismatching column names and only return the common columns.

#### **Returns:**

col\_names, with\_col\_names: list of column names of each DB.

See also: vertcat (p. ??), tests\_db (p. 217)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2007/01/18

# A.45.30 Method tests\_db/rows2Struct

**Summary:** Convert given rows of database to a structure array.

Usage:

```
s = rows2Struct(db, rows, pages)
```

# **Parameters:**

db: A tests\_db object.

rows: Indices of rows in db.

pages: Pages of db.

### **Returns:**

s: A structure of column name and value pairs.

See also: tests\_db (p. 217)

# A.45.31 Method tests\_db/getColNames

Summary: Gets column names.

Usage:

```
col_names = getColNames(db, tests)
```

**Description:** Performs a light operation without touching the data matrix.

**Parameters:** 

**db:** A tests\_db object.

tests: Columns for which to get names (Optional, default = ':')

**Returns:** 

col\_names: A cell array of strings.

See also: getColNames (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/24

# A.45.32 Method tests\_db/plotrow

**Summary:** Creates a plot\_abstract describing the desired db row.

**Usage:** 

```
a_plot = plotrow(a_tests_db, row)
```

#### **Parameters:**

a\_tests\_db: A tests\_db object.

row: Row number to visualize.

**props:** A structure with any optional properties.

putLabels: Put special column name labels.

### **Returns:**

a\_plot: A plot\_abstract object that can be plotted.

See also: plot\_abstract (p. 153), plotFigure (p. ??)

# A.45.33 Method tests\_db/dbsize

**Summary:** Returns the size of the data matrix of db.

Usage:

```
s = dbsize(db)
```

#### **Parameters:**

**db:** A tests\_db object.

#### **Returns:**

s: The size values.

See also: size (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/06

# A.45.34 Method tests\_db/displayRowsTeX

**Summary:** Generates a LaTeX table that lists rows of this DB.

Usage:

```
tex_string = displayRowsTeX(a_db, caption, props)
```

**Description:** By default table is rotated 90 degrees and scaled to 90

#### **Parameters:**

a\_db: A tests\_db object.

caption: Table caption.

**props:** A structure with any optional properties, passed to TeXtable.

#### **Returns:**

tex\_string: LaTeX string for table float.

# **Example:**

```
» string2File(displayRowsTeX(a_db(1:10, 4:7), 'some values',
struct('rotate', 0)), 'table.tex')
```

See also: displayRows (p. ??), TeXtable (p. ??), cell2TeX (p. ??)

# A.45.35 Method tests\_db/matchingRow

Summary: Creates a criterion database for matching the tests of a row.

#### **Usage:**

```
crit_db = matchingRow(db, row, props)
```

**Description:** Copies selected test values from row as the first row into the criterion db. Adds a second row for the STD of each column in the db. Calculates the covariance for using the Mahalonobis distance in the ranking.

#### **Parameters:**

**db:** A tests\_db object.

row: A row index to match.

**props:** A structure with any optional properties.

distDB: Take the standard deviation and covariance of this db instead.

#### **Returns:**

crit\_db: A tests\_db with two rows for values and STDs.

#### **Example:**

```
» crit_db = matchingRow(phys_control_compare_db,
find(phys_control_compare_db(:, 'TracesetIndex') == 61))
```

See also: rankMatching (p. ??), tests\_db (p. 217), tests2cols (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/08

## A.45.36 Method tests\_db/invarValues

Summary: Generates a 3D database of invariant values of given columns.

### **Usage:**

```
a_tests_3D_db = invarValues(db, cols, main_cols)
```

**Description:** The invariant values of a column are its values when all other column values are fixed. The invariant values of desired columns forms a matrix of rows. This function finds all combinations of the rest of the columns and returns the invariant value matrices for each such combination in a page of a three-dimensional vector; i.e. a tests\_3D\_db. Each matrix page will contain an additional column for the original row index for the invariant values. This index can be used to find the test columns that were omitted. Note: the trial column will be ignored for finding invariant values.

### **Parameters:**

```
db: A tests_db object.
```

cols: Vector of column numbers to find invariant values.

main\_cols: Vector of columns that need to be unique in each page
 (Optional; used only if database is not symmetric, to ignore missing values
 of main\_cols)

#### **Returns:**

```
a_tests_3D_db: A tests_3D_db object of organized values.
```

```
See also: tests_3D_db(p. 212), tests_3D_db/corrCoefs(p. ??), tests_3D_db/plotPair(p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/30

# A.45.37 Method tests\_db/meanDuplicateRows

**Summary:** Row-reduces a db by finding sets of rows with same main\_cols values, and replacing each set with a single row containing main\_cols and the mean of rest\_cols.

#### **Usage:**

```
a_tests_db = meanDuplicateRows(db, main_cols, rest_cols)
```

**Description:** The database is sorted for the values of the columns of interest (main\_cols) and all rows with duplicate values of these columns are identified. The rest of the columns (rest\_cols) are averaged and reduced to a single row, and attached to the nominal values of main\_cols. Two additional parameter columns will be added to the database created. The NumDuplicates column is the the number of duplicates used in the mean operation, and RowIndex is the row number points to the first of a set of duplicate values.

### **Parameters:**

**db:** A tests\_db object.

main\_cols: Vector of columns in which to find duplicates.

rest\_cols: Vector of columns to be averaged for duplicate main\_cols.

# **Returns:**

a\_tests\_db: The db object of with the means on page 1 and standard deviations on page 2.

See also: tests\_db/mean (p. 240), tests\_db/std (p. 227), sortedUniqueValues (p. ??)

# A.45.38 Method tests\_db/minus

Summary: Subtracts two DBs or a scalar from one.

## **Usage:**

```
a_db = minus(left_obj, right_obj)
```

**Description:** If DBs have mismatching columns only the common columns will be kept. In any case, the resulting DB columns will be sorted in the order of the left-hand-side DB.

#### **Parameters:**

```
left_obj, right_obj: Operands of the subtraction. One must be of type
  tests_db
  and the other can be a scalar.
```

#### **Returns:**

a\_db: The resulting tests\_db.

See also: minus (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/24

# A.45.39 Method tests\_db/displayRows

Summary: Displays rows of data with associated column labels.

# **Usage:**

```
s = displayRows(db, rows, pages)
```

# Parameters:

db: A tests\_db object.

rows: Indices of rows in db.

pages: Pages of db.

#### **Returns:**

s: A cell array of trasposed database contents, prefixed with column names on each row. Meant to be displayed on the screen.

See also: tests\_db (p. 217)

# A.45.40 Method tests\_db/times

**Summary:** Multiplies the DB with a scalar.

#### **Usage:**

```
a_db = times(left_obj, right_obj)
```

#### **Parameters:**

left\_obj, right\_obj: Operands of the multiplication. One must be of type tests db.

#### **Returns:**

a\_db: The resulting tests\_db.

See also: times (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/24

# A.45.41 Method tests\_db/addLastRow

**Summary:** Inserts a row of observations at the end of tests\_db.

## Usage:

```
index = addLastRow(obj, row)
```

**Description:** Adds a new set of observations to the database and returns its row index. This operation is expensive because the whole database matrix needs to be duplicated and resized in order to add a single new row. The method of allocating a matrix, filling it up, and then providing it to the tests\_db constructor is the preferred method of creating tests\_db objects.

#### **Parameters:**

```
obj: A tests_db object.
```

row: A row vector that contains values for each DB column.

#### **Returns:**

obj: The tests\_db object that includes the new row.

See also: allocateRows (p. ??), addRow (p. ??), tests\_db (p. 217)

# A.45.42 Method tests\_db/diff

**Summary:** Creates a tests\_db by taking the derivative of all tests.

### **Usage:**

```
a_db = diff(a_db, props)
```

**Description:** Applies the diff function to whole DB. The resulting DB will have one less row.

#### **Parameters:**

```
a_db: A tests_db object.props: Optional properties.
```

#### **Returns:**

a\_db: The resulting tests\_db.

See also: diff (p. ??), tests\_3D\_db/getDiff2DDB (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/05/24

# A.45.43 Method tests\_db/plot\_abstract

**Summary:** Default visualization for a database.

# **Usage:**

```
a_pm = plot_abstract(a_db, title_str)
```

**Description:** Calls plotTestsHistsMatrix. Subclasses should override this method to provide their own visualization.

#### **Parameters:**

```
a_db: A params_tests_db object.
```

title\_str: (Optional) A string to be concatanated to the title.

**props:** A structure with any optional properties.

#### **Returns:**

a\_pm: A plot\_stack with the plots organized in matrix form

### **Example:**

```
> plot(my_db, ': first impression') will call this function and send the generated plot to the plotFigure function.
```

```
See also: plot_abstract/plot_abstract (p. 153), plotTestsHistsMatrix (p. ??), plotFigure (p. ??)
```

# A.45.44 Method tests\_db/addRow

**Summary:** Inserts a row of observations to tests\_db at the given row index.

#### **Usage:**

```
index = addRow(obj, row, index)
```

**Description:** Adds a new set of observations to the database and returns the new DB. This operation is expensive in the sense that the whole database matrix needs to be copied to be passed to this function just to add a single new row. The method of allocating a matrix, filling it up, and then providing it to the tests\_db constructor is the preferred method of creating tests db objects.

## **Parameters:**

**obj:** A tests\_db object.

row: A row vector that contains values for each DB column.

index: The row index.

#### **Returns:**

obj: The tests\_db object that includes the new row.

See also: addLastRow (p. ??), allocateRows (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/08

### A.45.45 Method tests\_db/subsref

**Summary:** Defines indexing for tests\_db objects for () and . operations.

#### **Usage:**

```
obj = obj(rows, tests) obj = obj.attribute
```

**Description:** Returns attributes or selects the given test columns and rows and returns in a new tests\_db object.

### **Parameters:**

**obj:** A tests\_db object.

rows: A logical or index vector of rows. If ':', all rows.

tests: Cell array of test names or column indices. If ':', all tests.

attribute: A tests\_db class attribute.

#### **Returns:**

obj: The new tests\_db object.

See also: subsref (p. ??), tests\_db (p. 217)

# A.45.46 Method tests\_db/shufflerows

**Summary:** Returns a db with rows of given test columns are shuffled.

**Usage:** 

```
s = shufflerows(db, tests, grouped)
```

**Description:** Can be used for shuffle prediction. Basically, shuffle rows of tests and rerun high order analyses.

#### **Parameters:**

```
db: A tests_db object.tests: Tests to shuffle.grouped: If 1 then shuffle tests all together, if 0 shuffle each test separately.
```

#### **Returns:**

```
a_db: The shuffled db.

See also: tests_db (p. 217)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/10

# A.45.47 Method tests\_db/renameColumns

**Summary:** Rename an existing column or columns.

Usage:

```
a_db = renameColumns(a_db, test_names, new_names)
```

**Description:** This is a cheap operation than modifies meta-data kept in object.

### **Parameters:**

```
a_db: A tests_db object.
```

test\_names: A cell array of existing test names. new\_names: New names to replace existing ones.

## **Returns:**

a\_db: The tests\_db object that includes the new columns.

## **Example:**

```
» new_db = renameColumns(a_db, 'PulseIni100msSpikeRateISI_D40pA', 'Firing_rate');
» new_db = renameColumns(a_db, 'a', 'b', 'c', 'd');

See also: allocateRows(p.??), tests_db(p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/09/30
```

# A.45.48 Method tests\_db/mean

**Summary:** Returns the mean of the data matrix of a\_db. Ignores NaN values.

# Usage:

```
[a_db, n] = mean(a_db, dim)
```

**Description:** Does a recursive operation over dimensions in order to remove NaN values. This takes more time, compared with a straightforward mean operation.

#### **Parameters:**

a\_db: A tests\_db object.dim: Work down dimension.

#### **Returns:**

a\_db: The DB with one row of mean values. n: (Optional) Numbers of non-NaN rows included in calculating each column.

See also: mean (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/06

# A.45.49 Method tests\_db/plot

**Summary:** Generic method to plot a tests\_db or a subclass. Requires a plot\_abstract method to be defined for this object.

# Usage:

```
h = plot(a_tests_db, title_str, props)
```

# Parameters:

a\_tests\_db: A histogram\_db object.

title\_str: (Optional) String to append to plot title.

**props:** A structure with any optional properties, passed to plot\_abstract.

### **Returns:**

h: The figure handle created.

See also: plot\_abstract (p. 153), plotFigure (p. ??)

# A.45.50 Method tests\_db/subsasgn

Summary: Defines generic index-based assignment for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/06

# A.45.51 Method tests\_db/plotXRows

**Summary:** Create a scatter plot with a test versus the row numbers on the X-axis.

Usage:

```
a_p = plotXRows(a_db, test_y, title_str, short_title, props)
```

#### **Parameters:**

a\_db: A params\_tests\_db object.

test\_y: Y variable.

title\_str: (Optional) A string to be concatanated to the title.

**short\_title:** (Optional) Few words that may appear in legends of multiplot.

**props:** A structure with any optional properties passed to plotScatter.

**RowName:** Label to show on X-axis (default='RowNumber')

# **Returns:**

a\_p: A plot\_abstract.

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2007/01/16

# A.45.52 Method tests\_db/princomp

**Summary:** Generates a database of the principal components of given DB.

**Usage:** 

```
a_pca_db = princomp(db, props)
```

#### **Parameters:**

**db:** A tests\_db object.

**props:** A structure with any optional properties.

normalized: If specified zscore is used before princomp.

### **Returns:**

a\_pca\_db: A tests\_db where each row is a principal component.

See also: princomp (p. ??), zscore (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/09/21

### A.45.53 Method tests db/allocateRows

**Summary:** Preallocates a NaN-filled num\_rows rows in tests\_db.

**Usage:** 

```
obj = allocateRows(obj, num_rows)
```

**Description:** Allocates the desired number of rows to speed up filling up the data matrix using assignRowsTests. Using addRow after this operation is still expensive. The method of allocating a matrix, filling it up, and then providing it to the tests\_db constructor is the preferred method of creating tests\_db objects.

#### **Parameters:**

```
obj: A tests_db object.
```

num\_rows: The predicted number of observations for this tests\_db.

### **Returns:**

obj: The new tests\_db object.

```
See also: assignRowsTests (p. ??), addRow (p. ??), setRows (p. ??), tests_db (p. 217)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/08

## A.45.54 Method tests\_db/compareRows

Summary: Returns comparison results of db and the given row in a column vector.

**Usage:** 

```
idx = compareRows(db, row)
```

**Description:** If the row argument has multiple rows, the comparison is done separately for each of its rows and the results are the logical AND of those. Note that, it uses summation of distance for magnitude comparison. That is, all columns have the same weight. If the db contains many columns and one row, then the columns are compared instead of rows.

### **Parameters:**

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```
db: A tests_db object.
```

row: Row array, matrix or database to be compared with db rows.

#### **Returns:**

idx: A inverted logical column vector of comparison results. (false if db == row, true otherwise)

```
See also: eq (p. ??), tests_db (p. 217)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17

# A.45.55 Method tests\_db/plotYTests

**Summary:** Create a plot given database measures against given X-axis values, for each row.

## Usage:

```
a_p = plotYTests(a_db, x_vals, tests, axis_labels, title_str, short_title,
command, props)
```

#### **Parameters:**

```
a_db: A params_tests_db object.
```

**x\_vals:** A vector of X-axis values.

**tests:** A vector or cell array of columns to correspond to each value from x vals.

title\_str: (Optional) A string to be concatanated to the title.

**short\_title:** (Optional) Few words that may appear in legends of multiplot.

command: (Optional) Command to do the plotting with (default: 'plot')

**props:** A structure with any optional properties.

**LineStyle:** Plot line style to use. (default: 'd-')

**ShowErrorbars:** If 1, errorbars are added to each point.

**StatsDB:** If given, use this stats\_db for the errorbar (default=statsMeanStd(a\_db)).

quiet: If 1, don't include database name on title.

# **Returns:**

a\_p: A plot\_abstract.

## **Example:**

```
» a_p = plotYTests(a_db_row, [0 40 100 200], ...
'IniSpontSpikeRateISI_0pA', 'PulseIni100msSpikeRateISI_D40pA', ...
'PulseIni100msSpikeRateISI_D100pA', 'PulseIni100msSpikeRateISI_D200pA', ...
'current pulse [pA]', 'firing rate [Hz]', ', f-I curves', 'neuron 1');
"> plotFigure(a_p);
```

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See also: plotFigure (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/23

## A.45.56 Method tests\_db/plotScatter

**Summary:** Create a scatter plot of the given two tests.

Usage:

```
a_p = plotScatter(a_db, test1, test2, title_str, short_title, props)
```

#### **Parameters:**

**a\_db:** A params\_tests\_db object.

test1, test2: X & Y variables.

title\_str: (Optional) A string to be concatanated to the title.

**short\_title:** (Optional) Few words that may appear in legends of multiplot.

**props:** A structure with any optional properties.

**LineStyle:** Plot line style to use. (default: 'x') **Regress:** Calculate and plot a linear regression. **quiet:** If 1, don't include database name on title.

# **Returns:**

a\_p: A plot\_abstract.

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/09/29

# A.45.57 Method tests\_db/addColumns

Summary: Inserts new columns to tests\_db.

**Description:** Adds new test columns to the database and returns the new DB. Usage 2 concatanates two DBs columnwise. This operation is expensive in the sense that the whole database matrix needs to be enlarged just to add a single new column. The method of allocating a matrix, filling it up, and then providing it to the tests\_db constructor is the preferred method of creating tests\_db objects. This method may be used for measures obtained by operating on raw measures.

# **Parameters:**

obj, b\_obj: A tests\_db object.

test\_names: A cell array of test names to be added.

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test\_columns: Data matrix of columns to be added.

### **Returns:**

obj: The tests\_db object that includes the new columns.

See also: allocateRows (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/09/30

## A.45.58 Method tests db/tests2idx

**Summary:** Find dimension indices from a test names/numbers specification.

Usage:

```
idx = tests2idx(db, dim_name, tests)
```

#### Parameters:

db: A tests\_db object.

dim\_name: String indicating 'row', 'col', or 'page'

**tests:** Either a single or array of column numbers, or a single test name or a cell array of test names. If ':', all tests.

#### **Returns:**

idx: Array of column indices.

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

# A.45.59 Method tests\_db/delColumns

**Summary:** Deletes columns from tests\_db.

Usage:

```
obj = delColumns(obj, tests)
```

**Description:** Deletes test columns from the database and returns the new DB. This operation is expensive in the sense that the whole database matrix needs to be copied just to delete a single column. The method of allocating a matrix, filling it up, and then providing it to the tests\_db constructor is the preferred method of creating tests\_db objects. This method may be used for measures obtained by operating on raw measures.

### **Parameters:**

```
obj: A tests_db object.
```

tests: Numbers or names of tests (see tests2cols)

#### **Returns:**

obj: The tests\_db object that is missing the columns.

See also: allocateRows (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/10/06

### A.45.60 Method tests db/factoran

**Summary:** Generates a database of factor loadings obtained from the factor analysis of db with factoran. Each row corresponds to a rotated factor and columns represent observed variables.

### **Usage:**

```
a_factors_db = factoran(db, num_factors, props)
```

**Description:** Uses the promax method to rotate common factors.

## **Parameters:**

db: A tests\_db object.

num factors: Number of common factors to look for.

**props:** A structure with any optional properties.

#### **Returns:**

a\_factors\_db: A corrcoefs\_db of the coefficients and page indices.

See also: tests\_db (p. 217), corrcoefs\_db (p. 81)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/08

#### A.45.61 Method tests db/statsAll

**Summary:** Makes a stats\_db with rows of mean, STD, SE, and CV of the tests' distributions in db.

#### **Usage:**

```
a_stats_db = statsAll(db, tests, props)
```

### **Parameters:**

**db:** A tests\_db object.

tests: A selection of tests (see onlyRowsTests).

**props:** A structure with any optional properties for stats\_db.

### **Returns:**

```
a_stats_db: A stats_db object.
```

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/08/24

### A.45.62 Method tests db/enumerateColumns

**Summary:** Replaces each value with an integer pointing to the index of enumerated unique values in a column.

### **Usage:**

```
a_db = enumerateColumns(a_db, tests, props)
```

**Description:** Finds unique values of each column, and replaces the original values with the enumerated indices of these unique values. Useful for normalizing all parameter values in a hypercube.

# **Parameters:**

```
a_db: A tests_db object.
```

**tests:** Array of tests to be enumerated.

props: Optional properties.

truncateDecDigits: Use only up to this many decimal digits after the point

when checking for uniqueness.

## **Returns:**

a db: The modified DB.

# **Example:**

```
» enumerated_db = enumerateColumns(a_db(:, 1:9));
```

See also: uniqueValues (p. ??)

# A.45.63 Method tests\_db/tests2cols

**Summary:** Find column numbers from a test names/numbers specification.

# Usage:

```
cols = tests2cols(db, tests)
```

#### **Parameters:**

**db:** A tests\_db object.

**tests:** Either a single or array of column numbers, or a single test name or a cell array of test names. If ':', all tests.

#### **Returns:**

cols: Array of column indices.

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

# A.45.64 Method tests\_db/plotrows

Summary: Creates a plot\_stack describing the db rows.

# Usage:

```
a_plot = plotrows(a_tests_db, axis_limits, orientation, props)
```

#### **Parameters:**

```
a_tests_db: A tests_db object.
```

axis\_limits: If given, all plots contained will have these axis limits.

orientation: Stack orientation 'x' for horizontal, 'y' for vertical, etc.

props: A structure with any optional properties passed to plot\_stack.

# **Returns:**

a\_plot: A plot\_stack object that can be plotted.

See also: plot\_abstract (p. 153), plotFigure (p. ??)

# A.45.65 Method tests\_db/statsMeanSE

**Summary:** Generates a stats\_db object with two rows corresponding to the mean and standard error (SE) of the tests' distributions.

# **Usage:**

```
a_stats_db = statsMeanSE(db, tests, props)
```

## **Parameters:**

db: A tests\_db object.

tests: A selection of tests (see onlyRowsTests).

**props:** A structure with any optional properties for stats\_db.

## **Returns:**

```
a_stats_db: A stats_db object.
```

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

# A.45.66 Method tests\_db/plot\_bars

**Summary:** Creates a bar graph comparing all DB rows in groups, with a separate axis for each column.

## **Usage:**

```
a_plot = plot_bars(a_tests_db, title_str, props)
```

# Parameters:

```
a_tests_db: A tests_db object.
```

title\_str: (Optional) The plot title.

**props:** A structure with any optional properties.

# **Returns:**

a\_plot: A object of plot\_bars or one of its subclasses.

See also: plot\_abstract (p. 153), plot\_simple (p. 163)

### A.45.67 Method tests\_db/corrCoefs

**Summary:** Generates a database of correlation coefficients by comparing col1 with other cols in the database. If db has multiple pages, then each page in db produces a row of coefficients and matching PageIndex.

## **Usage:**

```
a_coefs_db = corrCoefs(db, col1, cols, props)
```

**Description:** Assuming the db was created with invarValues, this function finds the invariant correlation coefficients between its columns. The invariant correlation coefficients are the correlation of one column value with another column value when some other column values are fixed. Since there are many occurences of the invariant coefficients, a histogram can then be created and returned from the created db. The other columns that are fixed are not in this db object, but can be reached using the row indices in the original db. The page number is saved in the created db, so that it can be used to find the page from which the coefficient came. Then row indices of the page points to original constant column values.

#### **Parameters:**

db: A tests\_db object.

**col1:** Column to compare.

**cols:** Columns to be compared with col1.

**props:** A structure with any optional properties.

**skipCoefs:** If coefficients of less confidence than should be skipped.

## **Returns:**

a\_coefs\_db: A corrcoefs\_db of the coefficients and page indices.

See also: tests\_db (p. 217), corrcoefs\_db (p. 81)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/30

# A.45.68 Method tests\_db/transpose

**Summary:** Transposes data matrix and swaps row and columns metadata as well.

# Usage:

```
a_db = transpose(a_db)
```

#### **Parameters:**

**a\_db:** A tests\_db.

### **Returns:**

a\_db: The transposed tests\_db.

See also: transpose (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2007/02/07

# A.45.69 Method tests\_db/vertcat

**Summary:** Vertical concatanation [db;with\_db;...] operator.

Usage:

```
a_db = vertcat(db, with_db, ...)
```

**Description:** Concatanates rows of with\_db to rows of db. Overrides the built-in vert-cat function that is called when [db;with\_db] is executed. If the first argument is a array of DBs, then this functionality is not needed; built-in vertcat is called.

## **Parameters:**

db: A tests\_db object.

with\_db: A tests\_db object whose rows are concatanated to db.

#### **Returns:**

a\_db: A tests\_db that contains rows of db and with\_db.

See also: vertcat (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/01/25

# A.45.70 Method tests\_db/assignRowsTests

**Summary:** Assign the values to the tests and rows (and pages) of the tests\_db.

Usage:

```
obj = assignRowsTests(obj, val, rows, tests, pages)
```

**Description:** Selects the given dimensions and returns in a new tests\_db object.

#### **Parameters:**

**obj:** A tests\_db object.

val: DB object or data matrix to be assigned to the addressed indices.

rows: A logical or index vector of rows. If ':', all rows.

tests: Cell array of test names or column indices. If ':', all tests.

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```
pages: (Optional) A logical or index vector of pages. ':' for all pages.
```

#### **Returns:**

obj: The new tests\_db object.

See also: subsref (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/02/08

# A.45.71 Method tests\_db/plotCovar

**Summary:** Generates an image plot of the covariance-type values in a\_db.

# Usage:

```
a_plot = plotCovar(a_db, title_str, props)
```

#### Parameters:

**a\_db:** A tests\_db object that resulted from a function like cov.

title\_str: (Optional) String to append to plot title.

**props:** Optional properties to be passed to plot\_abstract.

**inverse:** If 1, take inverse of the data matrix.

crosscoef: If 1, normalize matrix elements as crosscoef would a cov

matrix.

logScale: If 1, take logarithm of values before plotting.

## **Returns:**

a\_plot: A plot\_abstract object or one of its subclasses.

# **Example:**

```
» plotFigure(plotCovar(cov(get(constrainedMeasuresPreset(pbundle2, 6),
'joined_control_db'))));
```

```
See also: tests_db/cov(p. 226), plotImage(p. ??), tests_db/matchingRow. (p. ??)
```

### A.45.72 Method tests\_db/isinf

Summary: Returns logical row indices of Inf-valued columns.

**Usage:** 

```
rows = isinf(db, col)
```

#### **Parameters:**

```
db: A tests_db object.
```

**col:** Column to check (Optional, default = 1)

#### **Returns:**

rows: A logical column vector of rows.

```
See also: isinf (p. ??), tests_db (p. 217)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/08/16

# A.45.73 Method tests\_db/isnan

Summary: Returns logical row indices of NaN-valued columns.

### Usage:

```
rows = isnan(db, col)
```

### Parameters:

**db:** A tests\_db object.

**col:** Column to check (Optional, default = 1)

### **Returns:**

rows: A logical column vector of rows.

See also: isnan (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/06

### A.45.74 Method tests\_db/rankMatching

Summary: Create a ranking db of row distances of db to given criterion db.

Usage:

```
a_ranked_db = rankMatching(db, crit_db, props)
```

**Description:** crit\_db can be created with the matchingRow method. TestWeights modify the importance of each measure.

#### **Parameters:**

db: A tests db to rank.

crit\_db: A tests\_db object holding the match criterion tests and stds.

**props:** A structure with any optional properties.

**limitSTD:** limit any measure to this many STDs max.

tolerateNaNs: If 0, rows with any NaN values are skipped, if 1, NaN values are given a fixed 3xSTD penalty (default=1).

testWeights: Structure array associating tests and multiplicative weights.

**restoreWeights:** Reverse the testWeights application after calculating distances.

topRows: If given, only return this many of the top rows.

**useMahal:** Use the Mahalonobis distance from the covariance matrix in crit\_db.

#### **Returns:**

a\_ranked\_db: A ranked\_db object.

See also: matchingRow (p. ??), tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/08

### A.45.75 Method tests\_db/statsBounds

**Summary:** Generates a stats\_db object with three rows corresponding to the mean, min, and max of the tests' distributions.

#### Usage:

```
a_stats_db = statsBounds(a_db, tests, props)
```

**Description:** A page is generated for each page of data in db.

### **Parameters:**

a\_db: A tests\_db object.

tests: A selection of tests (see onlyRowsTests).

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**props:** A structure with any optional properties for stats\_db.

#### **Returns:**

a\_stats\_db: A stats\_db object.

See also: tests\_db (p. 217)

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/10/07

#### A.46 Class trace

### A.46.1 Constructor trace/trace

**Summary:** A generic trace from a cell. It can be voltage, current, etc.

### Usage:

```
obj = trace(data_src, dt, dy, id, props)
```

**Description:** Traces for specific experimental or simulation protocols can extend this class for adding new parameters. This object is designed to recognize most data file formats. See the data\_src parameter below.

#### **Parameters:**

data\_src: Trace data as a column vector OR name of a data file generated by

Genesis (.bin, .gbin, .genflac), PCDX (.all), or Matlab (.mat).

dt: Time resolution [s]

dy: y-axis resolution [ISI (V, A, etc.)]

id: Identification string

props: A structure with any optional properties.

**scale\_y:** Y-axis scale to be applied to loaded data.

offset\_y: Y-axis offset to be added to loaded and scaled data.

trace\_time\_start: Samples in the beginning to discard [dt]

baseline: Resting potential.

**channel:** Channel to read from file Genesis, PCDX, or Neuron file.

**file\_type:** Specify file type instead of guessing from extension:

'genesis': Raw binary files created with Genesis disk\_out method. 'genesis\_flac': Compressed Genesis binary files. 'neuron': Binary files created with Neuron's Vector.vwrite method. 'pcdx': .ALL data acquisition files from PCDX program. 'matlab': Matlab .MAT binary files with matrix data.

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traces: Traces to read from PCDX file.

```
spike_finder: Method of finding spikes
    (1 for findFilteredSpikes, 2 for findspikes).
init_Vm_method: Method of finding spike thresholds
    (see spike_shape/spike_shape).
init_threshold: Spike initiation threshold (deriv or accel).
    (see above methods and implementation in calcInitVm)
init_lo_thr, init_hi_thr: Low and high thresholds for slope.
threshold: Spike threshold.
quiet: If 1, reduces the amount of textual description in plots, etc.
```

#### Returns a structure object with the following fields:

data: The trace column matrix. dt, dy, id, props (see above)

```
See also: spikes (p. 195), spike_shape (p. 183), cip_trace (p. 54), period (p. 137)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

### A.46.2 Method trace/setProp

Summary: Generic method for setting optional object properties.

**Usage:** 

```
obj = setProp(obj, prop1, val1, prop2, val2, ...)
```

**Description:** Modifies or adds property values. As many property name-value pairs can be specified.

#### **Parameters:**

**obj:** Any object that has a props field.

attr: Property nameval: Property value.

#### **Returns:**

obj: The new object with the updated properties.

#### See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/22

#### A.46.3 Method trace/display

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

# A.46.4 Method trace/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

### A.46.5 Method trace/periodWhole

Summary: Returns the boundaries of the whole period of trace, t.

Usage:

whole\_period = periodWhole(t)

#### **Parameters:**

t: A trace object.

See also: period (p. 137), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

#### A.46.6 Method trace/set

**Summary:** Generic method for setting object attributes.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/08

#### A.46.7 Method trace/plotData

**Summary:** Plots a trace.

Usage:

```
a_plot = plotData(t, title_str, props)
```

**Description:** If t is a vector of traces, returns a vector of plot objects.

#### **Parameters:**

t: A trace object.

title\_str: (Optional) String to append to plot title.

**props:** A structure with any optional properties.

timeScale: 's' for seconds, or 'ms' for milliseconds.
(rest passed to plot\_abstract.)

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/17

### A.46.8 Method trace/spikes

Summary: Convert trace to spikes object for spike timing calculations.

**Usage:** 

```
obj = spikes(trace, a_period, plotit, minamp)
```

**Description:** Creates a spikes object.

### **Parameters:**

trace: A trace object.

**a\_period:** A period object denoting the part of trace of interest (optional, if empty vector, taken as wholePeriod).

**plotit:** If non-zero, a plot is generated for showing spikes found (optional).

**minamp:** minimum amplitude that must be reached if using findFilteredSpikes.

-> adjust as needed to discriminate spikes from EPSPs. (optional)

See also: spikes (p. 195)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

# A.46.9 Method trace/findFilteredSpikes

**Summary:** Runs a frequency filter over the data and then finds all peaks using find-spikes.

#### **Usage:**

```
[times, peaks, n] = findFilteredSpikes(t, a_period, plotit, minamp)
```

**Description:** Runs a 50-300 Hz band-pass filter over the data and then calls findspikes. The filter both removes low-frequency offset changes, such as cip period effects, and high-frequency noise that is detected as local peaks by findspikes. The spikes found are post-processed to make sure the rise and fall times are consistent. Note: The filter employed only works with data sampled at 10kHz.

#### **Parameters:**

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```
t: Trace object
             a_period: Period of interest.
             plotit: Plots the spikes found if 1.
                  minamp (optional): minimum amplitude above baseline that must be reached.
                  -> adjust as necessary to discriminate spikes from EPSPs.
             Returns:
             times: The times of spikes [dt].
             peaks: The peaks corresponding to the times of spikes.
             n: The number of spikes.
       See also: findspikes (p. ??), period (p. 137)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2004/03/08
       A.46.10 Method trace/calcAvg
       Summary: Calculates the average value of the given period of the trace, t.
       Usage:
avg_val = calcAvg(t, period)
       Parameters:
             t: A trace object.
             period: A period object (optional).
       See also: period (p. 137), trace (p. 255)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30
       A.46.11 Method trace/getDy
       Summary: Returns dy.
       Usage:
dy = getDy(t)
       Parameters:
             t: A trace object.
       Returns:
             dy: The dy value.
       See also: trace (p. 255)
       Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/31
```

# A.46.12 Method trace/calcMax

**Summary:** Calculates the maximal value of the given period of the trace, t.

### Usage:

```
[max_val, max_idx] = calcMax(t, period)
```

#### **Parameters:**

t: A trace object.

period: A period object (optional).

#### **Returns:**

max\_val: The max value. max\_idx: Its index in the trace.

See also: period (p. 137), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

### A.46.13 Method trace/calcMin

Summary: Calculates the minimal value of the given period of the trace, t.

### Usage:

```
[min_val, min_idx] = calcMin(t, a_period)
```

### **Parameters:**

t: A trace object.

a\_period: A period object (optional).

### **Returns:**

min\_val: The min value. min\_idx: Its index in the trace.

See also: period (p. 137), trace (p. 255)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/07/30

# A.46.14 Method trace/plot\_abstract

Summary: Plots a trace by calling plotData.

Usage:

```
a_plot = plot_abstract(t, title_str, props)
```

**Description:** If t is a vector of traces, returns a vector of plot objects.

#### **Parameters:**

t: A trace object.

title\_str: (Optional) String to append to plot title.

**props:** A structure with any optional properties.

timeScale: 's' for seconds, or 'ms' for milliseconds.
 (rest passed to plot\_abstract.)

#### **Returns:**

a\_plot: A plot\_abstract object that can be visualized.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/17

#### A.46.15 Method trace/subsref

Summary: Defines generic indexing for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

### A.46.16 Method trace/plot

Summary: Plots a trace.

Usage:

h = plot(t)

#### **Parameters:**

**t:** A trace object.

title\_str: (Optional) String to append to plot title.

**props:** A structure with any optional properties, passed to plot\_abstract.

#### **Returns:**

h: Handle to figure object.

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See also: trace (p. 255), plot\_abstract (p. 153)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

### A.46.17 Method trace/getSpike

**Summary:** Convert a spike in the trace to a spike\_shape object.

**Usage:** 

```
obj = getSpike(trace, spikes, spike_num, props)
```

**Description:** Creates a spike\_shape object from desired spike. It is more efficient if you already have the spikes object.

#### **Parameters:**

trace: A trace object.

spikes: (Optional) A spikes object obtained from trace,

calculated automatically if given as [].

spike\_num: The index of spike to extract.

**props:** A structure with any optional properties.

**spike\_id:** A prefix string added to the spike\_shape object's id.

### **Example:**

```
This will create an annotated plot of the third spike in my_trace: 
» plotFigure(plotResults(getSpike(my_trace, [], 3)))
```

See also: spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/19

#### A.46.18 Method trace/withinPeriod

Summary: Returns a trace object valid only within the given period.

Usage:

```
obj = withinPeriod(t, a_period)
```

### **Parameters:**

t: A trace object.

a\_period: The desired period

#### **Returns:**

obj: A trace object

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**See also:** trace (p. 255), period (p. 137)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/25

### A.46.19 Method trace/getResults

Summary: Runs all tests defined by this class and return them in a structure.

Usage:

results = getResults(t)

### Parameters:

t: A trace object.

#### **Returns:**

results: A structure associating test names to values in ms and mV (or mA).

See also: spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/13

### A.46.20 Method trace/spike\_shape

**Summary:** Convert averaged spikes in the trace to a spike\_shape object.

Usage:

obj = spike\_shape(trace, spikes, props)

**Description:** Creates a spike\_shape object.

**Parameters:** 

trace: A trace object.

spikes: A spikes object on trace.

See also: spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/04

### A.46.21 Method trace/analyzeSpikesInPeriod

**Summary:** Returns results and a db of spikes by collecting test results of a cip\_trace, analyzing each individual spike.

#### **Usage:**

[results period\_spikes a\_spikes\_db spikes\_stats\_db spikes\_hists\_dbs] =
analyzeSpikesInPeriod(a\_cip\_trace, a\_spikes, period, prefix\_str)

#### Parameters:

```
a_cip_trace: A cip_trace object.a_spikes: A spikes object from the a_cip_trace object.period: A period of object of a_cip_trace object of interest.prefix_str: Prefix string to be added to spike shape results.
```

#### **Returns:**

results: Results structure names prefixed with prefix\_str. period\_spikes: Corrected spikes object for this period. a\_spikes\_db: A mini spikes database of results from each spike in period. spikes\_stats\_db: Statistics from the mini spikes database. spikes\_hists\_dbs: Cell array of histograms from the mini spikes database.

```
See also: cip_trace (p. 54), spikes (p. 195), period (p. 137), spike_shape (p. 183), getProfileAllSpikes (p. ??)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/05/04

# A.47 Class trace\_profile

### A.47.1 Constructor trace\_profile/trace\_profile

**Summary:** Creates and collects test results of a trace.

**Description:** The first usage is fully customizable to be used from subclass constructors. The second usage generates the spikes and spike\_shape objects, and collects some generic test results from them. This usage is only provided as an example and is not used practically.

#### **Parameters:**

```
data_src: The trace column OR the .MAT filename.
dt: Time resolution [s]
dy: y-axis resolution [ISI (V, A, etc.)]
props: See trace object.
```

# Returns a structure object with the following fields:

trace, spikes, spike\_shape, results, id, props.

See also: trace (p. 255), spikes (p. 195), spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/13

# A.47.2 Method trace\_profile/get

Summary: Defines generic attribute retrieval for objects.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/14

# A.48 Utility functions

### A.48.1 Function functions/findspikes\_old

Author: <adelgado@biology.emory.edu>, 2003-03-31

#### A.48.2 Function functions/makeIdx

**Summary:** Prepare the idx structure from names.

Usage:

idx = makeIdx(names)

**Description:** Helper function.

**Parameters:** 

names: Cell array of names for a db dimension.

**Returns:** 

idx: Structure associating names to array indices.

See also: tests\_db (p. 217)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/17

### A.48.3 Function functions/diff2T\_h4

**Summary:** Estimate of second derivative using Taylor expansion (derived with same method as diffT).

#### **Usage:**

 $deriv2 = diff2T_h4(x, dy)$ 

**Description:** 
$$d^2 x x(k-2) - x(k-1) - x(k+1) + x(k+2) =$$

#### **Parameters:**

**x:** A vector of x = f(y).

dy: The resolution of the discrete points in the vector.

### **Returns:**

deriv2: Estimate of the derivative.

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/15

#### A.48.4 Function functions/diff2T

**Summary:** Estimate of second derivative using Taylor expansion.

### Usage:

deriv2 = diff2T(x, dy)

### **Parameters:**

**x:** A vector of x = f(y).

dy: The resolution of the discrete points in the vector.

### **Returns:**

deriv2: Estimate of the derivative.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/15

#### A.48.5 Function functions/diff3T

**Summary:** Estimate of third derivative using Taylor expansion.

### Usage:

$$deriv3 = diff3T(x, dy)$$

#### **Parameters:**

**x:** A vector of x = f(y).

dy: The resolution of the discrete points in the vector.

### **Returns:**

deriv3: Estimate of the derivative.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/15

### A.48.6 Function functions/diffT

**Summary:** Estimate of first derivative using Taylor expansion.

### Usage:

$$deriv = diffT(x, dy)$$

### **Parameters:**

**x:** A vector.

dy: The resolution of the discrete points in the vector.

#### **Returns:**

deriv: Estimate of the first derivative.

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/11/15

### A.48.7 Function functions/readNeuronVecBin

Author: Konstantin Miller <miller@cs.tu-berlin.de>, Aug 09, 2005.

### A.48.8 Function functions/logLevels

**Summary:** Returns a logarithmic-scaled series between min\_val and max\_val with num\_levels elements.

### **Usage:**

levels = logLevels(min\_val, max\_val, num\_levels)

#### **Parameters:**

min\_val, max\_val: The low and high boundaries for the output value.
num\_levels: Number of elements to produce, including the boundaries.

#### **Returns:**

levels: A column vector of logarithmic series between min\_val and max\_val.

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/18

#### A.48.9 Function functions/diff3T\_h4

**Summary:** Estimate of third derivative using Taylor expansion (derived with same method as diffT and diff2T\_h4).

# **Usage:**

 $deriv2 = diff3T_h4(x, dy)$ 

**Description:** 
$$d^3 x - x(k-2) + 2 * x(k-1) - 2 * x(k+1) + x(k+2) =$$

#### **Parameters:**

**x:** A vector of x = f(y).

dy: The resolution of the discrete points in the vector.

#### **Returns:**

deriv2: Estimate of the derivative.

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/18

#### A.48.10 Function functions/collectspikes

Author: <adelgado@biology.emory.edu>

#### A.48.11 Function functions/subTextLabel

**Summary:** Draws a text label on a plot.

**Usage:** 

handle = subTextLabel(x, y, text\_str, props)

#### **Parameters:**

**x**, **y**: 2D coordinates.

text\_str: String to be drawn on plot.

**props:** A structure with any optional properties.

Units: position units for the coordinates (see Units in axes properties).

#### **Returns:**

handle: Text object handle.

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/11

### A.48.12 Function functions/uniqueValues

**Summary:** Find unique rows in a matrix (or column vector). Version which makes use of sort and diff.

### Usage:

```
[rows, idx] = uniqueValues(data)
```

**Description:** Parameters: data: A matrix or column vector Returns: rows: A matrix or column vector of unique rows. idx: Indices of the unique rows in the original data matrix.

See also: sortedUniqueValues (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/24

#### A.48.13 Function functions/calcGraphNormPtsRatio

**Summary:** Return the ratios of normalized to point units for dimensions of axis.

Usage:

```
[ratio_x, ratio_y] = calcGraphNormPtsRatio(grfx_handle)
```

**Description:** Used for findind character sizes given the size of an axis. Normally if the plot is resized, the characters may take up too much space or may not fit anymore unless the spacing is corrected.

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#### **Parameters:**

grfx\_handle: A graphics handle to an existing axis or figure.

#### **Returns:**

ratio\_x, ratio\_y: Normalized to points ratio for axis width and height, respectively.

#### **Example:**

```
To find the normalized distance for a 10pt character:  >  dx = 10 *  calcGraphNormPtsRatio(my_figure_handle);
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/03/05

#### A.48.14 Function functions/findspikes

**Summary:** Performs spike discrimination on single tips exceeding or within a threshold and/or time window range.

#### **Usage:**

```
spikeTime = findspikes(traces, fs, threshold) [spikeTime spikePeak n] =
findspikes(traces, fs, threshold [,direction] [,win_range] [,'plot'])
```

#### **Parameters:**

**traces**: Multiple traces of signal. each trace in a column

fs: Sampling frequency, in KHz

threshold: Either a scalar, or [thres1 thres2] to define a range

direction: Optional.

A positive number to find positive-going spikes, and vice versa. The default value is +1 when threshold is a scalar, is sign(thres2-thres1) when threshold is a vector.

win\_range: Optional.

[win\_min win\_max] to define a time window range of the width at threshold. in ms. 'plot': Optional. Plot the result. Not to plot by default.

**spikeTime:** Returns a cell array, each cell is a vector of spike times in each trial

spikePeak: Returns the peak values of each spikes.

n: the total number of spikes

# A.48.15 Function functions/fillederrorbar

**Summary:** Plots an errorbar with the middle points filled with the pen color.

**Usage:** 

handles = fillederrorbar(...)

#### **Parameters:**

(see errorbar)

**Returns:** 

handles: Handles to graphics objects.

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/13

#### A.48.16 Function functions/findVectorInMatrix

Summary: Finds rows of data that match row.

**Usage:** 

idx = findVectorInMatrix(data, row)

**Description:** Matlab's eq (==) command unfortunately doesn't allow this directly.

**Parameters:** 

data: A matrix or column vector.

row: A row vector.

Returns:

idx: Indices of matching rows in the original data matrix.

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/09/1

### A.48.17 Function functions/string2File

Summary: Writes string verbatim into a file.

**Usage:** 

```
string2File(string, filename, props)
```

#### **Parameters:**

string: To be written into file.
filename: The file to be created.

**props:** A structure with any optional properties.

#### **Returns:**

See also: cell2TeX (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/10

### A.48.18 Function functions/chanTables2DB

**Summary:** Creates a DB with channel tables exported from Genesis.

### Usage:

```
a_chans_db = chanTables2DB(tables, id, props)
```

### **Parameters:**

tables: Structures returned from the dump files generated by dump\_chans.g.

id: String that identify the source of the tables structure.

**props:** A structure with any optional properties.

(rest passed to tests\_db.)

#### **Returns:**

a\_chans\_db: A tests\_db object containing channel tables.

See also: trace (p. 255), trace/plot (p. 261), plot\_abstract (p. 153), GP/common/dump\_chans.g (Genesis) (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2007/03/07

#### A.48.19 Function functions/maxima

Summary: Find all local maxima.

**Usage:** 

```
x_ix = maxima(x)
```

**Description:** Finds derivative sign-flipping points where the second derivative is less than zero

#### **Parameters:**

x: A vector.

#### **Returns:**

x\_idx: Indices of maxima.

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/04/18

### A.48.20 Function functions/sortedUniqueValues

**Summary:** Find unique rows in an already sorted matrix (or column vector). Uses the derivation method.

Usage:

```
[rows, idx] = sortedUniqueValues(data)
```

**Description:** Parameters: data: A ascending row-sorted matrix or column vector. Returns: rows: A matrix or column vector of unique rows. idx: Indices of the unique rows in the original data matrix.

See also: uniqueValues (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/27

# A.48.21 Function functions/TeXtable

Summary: Creates the LaTeX string for a floating table containing given contents.

**Usage:** 

```
tex_string = TeXtable(contents, caption, props)
```

### **Parameters:**

contents: Table contents in LaTeX.

caption: Table caption.

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**props:** A structure with any optional properties.

rotate: Degrees to rotate.
width: Resize to this width.
height: Resize to this height
center: Align to center.

**shortCaption:** Short version of caption to appear at list of tables.

**floatType:** LaTeX float to use (default='table').

### **Returns:**

tex\_string: LaTeX string for table float.

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/13

### A.48.22 Function functions/cell2TeX

**Summary:** Creates LaTeX string of a formatted table with the cell array's contents.

Usage:

```
tex_string = cell2TeX(a_cell, props)
```

#### **Parameters:**

**a\_cell:** A cell matrix to be tabularized.

**props:** A structure with any optional properties.

hasTitleRow: The first row contains titles.
hasTitleCol: The first column contains titles.

#### **Returns:**

tex\_string: LaTeX formatted table string.

See also:

**Author:** Cengiz Gunay <cgunay@emory.edu>, 2004/12/09

### A.48.23 Function functions/ns\_CIPlist

Author: Dawid Kurzyniec

# A.48.24 Function functions/parseGenesisFilename

**Summary:** Parses the GENESIS filename to get names and values of simulation parameters. Usage: names\_vals = parseGenesisFilename(filename)

Description: Parameters: filename: GENESIS filename (no need to exist)

**Returns:** 

names\_vals: A two-column cell array with names and values.

See also: cip\_profile (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/03/10

### A.48.25 Function functions/boxplotp

**Description:** BOXPLOTP(X,NOTCH,SYM,VERT,WHIS,PROPS) produces a box and whisker plot for each column of X. The box has lines at the lower quartile, median, and upper quartile values. The whiskers are lines extending from each end of the box to show the extent of the rest of the data. Outliers are data with values beyond the ends of the whiskers.

### A.48.26 Function functions/plotImage

Summary: Function that plots a color matrix on current figure.

**Usage:** 

```
h = plotImage(image_data, colormap_func, num_colors, props)
```

#### **Parameters:**

image\_data: 2D matrix with image data.

colormap\_func: Function name or handle to colormap (e.g., 'jet').

num\_colors: Parameter to be passed to the colormap\_func.

**props:** A structure with any optional properties.

colorbar: If defined, show colorbar on plot.

**truncateDecDigits:** Truncate labels to this many decimal digits. **maxValue:** Maximal value at num\_colors to annotate the colorbar.

#### **Returns:**

colors: RGB color matrix to be passed to colormap function.

See also: colormap (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/05

### A.48.27 Function functions/meanSpikeFreq

**Summary:** Returns the mean firing frequency in Hz according to mean \ inter-spike-interval of the given spike train and the time resolution dt.

#### **Usage:**

```
meanFreq = meanSpikeFreq( spike_train, dt, period )
```

**Description:** Parameters: spike\_train: Spike times returned by findspikes dt: Time step size [s] period: Duration of the total time period [dt]

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/03/08

# A.48.28 Function functions/interpValByIndex

**Summary:** Finds the interpolated value by using the real valued index from the data vector.

Usage:

```
val = interpValByIndex(idx, data)
```

**Description:** Parameters: idx: A real-valued index. data: A data vector.

**Returns:** 

val: the value taken from the nearest integer indices of data and interpolated.

#### **Example:**

```
» a= [1 2 3];
» interpValByIndex(1.5, a)
ans =
1.5000
```

See also: spike\_shape (p. 183)

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/08/02

# A.48.29 Function functions/growRange

Summary: Returns the maximal range from rows of axis limits.

**Usage:** 

```
range = growRange(ranges)
```

#### **Parameters:**

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ranges: A matrix where each row is return val of axis func.

### **Returns:**

range: The maximal range obtained that includes all given axes.

See also:

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/10/13

### A.48.30 Function functions/mergeStructs

**Summary:** Merges all the structures given as arguments and makes a single structure.

**Usage:** 

```
results = mergeStructs( struct1 [, struct2, ...] )
```

**Description:** The fields will in earlier arguments will have priority. So, while merging two structs, if there are duplicate fields, the fields in the first will be preserved.

#### **Parameters:**

struct(n): A structure.

#### **Returns:**

results: The merged structure.

# **Example:**

```
mergeStructs( struct('hello', 1), struct('bye', 2) );
=> struct('hello', 1, 'bye', 2)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/09/13

### A.48.31 Function functions/properTeXFilename

**Summary:** Replaces characters in string to make it a valid filename for inclusion in TeX documents.

### Usage:

```
filename = properTeXFilename( filename )
```

**Description:** It will replace characters like space, '/', '.', etc.

### **Parameters:**

**filename:** An input filename string (without extension!).

#### **Returns:**

filename: The corrected proper filename.

#### **Example:**

```
» fname = properTeXFilename('hello world/1')
ans = 'hello_world+1'
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2005/12/20

### A.48.32 Function functions/prefixStruct

**Summary:** Adds the given prefix to each of the field names in the structure.

### Usage:

```
new_struct = prefixStruct(a_struct, prefix_str)
```

#### Parameters:

**a\_struct:** A structure.

**prefix\_str:** A string to be prefixed to each field name.

### **Returns:**

new\_struct: The new structure.

### **Example:**

```
prefixStruct( struct('bye', 1), 'hello');
=> struct('hellobye', 1)
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2004/12/22

### A.48.33 Function functions/properTeXLabel

**Summary:** Replaces characters in string or cell array of strings to make it valid in TeX documents.

### Usage:

```
a_label = properTeXLabel( a_label )
```

**Description:** It will replace characters like space, '/', '.', etc.

#### **Parameters:**

a\_label: A label string.

#### **Returns:**

a\_label: The corrected proper a\_label.

#### **Example:**

```
» a_label = properTeXLabel('this_day')
ans = 'this\_day'
```

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/01/17

### A.48.34 Function functions/colormapBlueCrossRed

Summary: Blue to red crossing colormap, with a black-colored zero-crossing.

### **Usage:**

```
colors = colormapBlueCrossRed(num_half_colors)
```

**Description:** Colormap contains (2 \* num\_half\_colors + 1) colors, where (num\_half\_colors + 1) is the zero crossing.

#### **Parameters:**

num\_half\_colors: Number of colors to generate on one of the red or blue scales.

**props:** A structure with any optional properties.

### **Returns:**

colors: RGB color matrix to be passed to colormap function.

See also: colormap (p. ??)

Author: Cengiz Gunay <cgunay@emory.edu>, 2006/06/05

# A.48.35 Function functions/loadtraces

#### Parameters:

file: PCDX file.

**tracelist:** A string of trace description, such as '1-10'.

channel: Channel to read from.

quiet: (Optional) If 1, produce on print outs.

Author: <adelgado@biology.emory.edu>

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