

# SB pipe documentation

Release 1.5.0

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**CHAPTER** 

ONE

# **USER MANUAL**

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# Introduction

This package contains a collection of pipelines for dynamic modelling of biological systems. It aims to automate common processes and speed up productivity for tasks such as model simulation, single and double parameter scan, and parameter estimation.

# Requirements

In order to use SB pipe, the following software must be installed:

- Copasi 4.16 http://copasi.org/
- Python 2.7+ https://www.python.org/
- R 3.3.0+ https://cran.r-project.org/
- LaTeX 2013 (optional) https://latex-project.org/ftp.html

You should also make sure that the following packages are installed in your machine: python-pip, and (optionally) texlive-latex-base.

Before installing SB pipe Python and R dependencies the following environment variables must be added to your GNU/Linux \$HOME/.bashrc file:

```
# SB_PIPE
export SB_PIPE=/path/to/sb_pipe
export PATH=$PATH:${SB_PIPE}/sb_pipe

# Path to CopasiSE
export PATH=$PATH:/path/to/CopasiSE
```

The .bashrc file can then be reloaded from your shell using the command:

```
$ source $HOME/.bashrc
```

On Windows platforms, these environment variables are configured as any other Windows environment variable.

Now it is the time to install Python and R packages used by SB pipe. Two scripts are provided to perform these tasks automatically.

To install SB pipe Python dependencies, run:

```
cd ${SB_PIPE}/
./install_pydeps.py
```

To install SB pipe R dependencies, run:

```
cd ${SB_PIPE}/
$ R
# Inside R environment, answer 'y' to install packages locally
> source('install_rdeps.r')
```

If R package dependencies must be compiled, it is worth checking that the following additional packages are installed in your machine: build-essential, liblapack-dev, libblas-dev, libcairo-dev, libssl-dev, libcurl4-openssl-dev. After installing these packages, install\_rdeps.r must be executed again.

# Installation

Run the command inside the sb\_pipe folder:

```
python setup.py install
```

The correct installation of SB pipe and its dependencies can be checked by running the following commands inside the SB pipe folder:

```
cd tests
./test_suite.py
```

# How to use SB pipe

# **Preliminary configuration steps**

#### **Pipelines using Copasi**

Before using these pipelines, a Copasi model must be configured as follow using CopasiUI:

#### pipeline: simulate

- Tick the flag executable in the Time Course Task.
- Select a report template for the Time Course Task.
- Save the report in the same folder with the same name as the model but replacing the extension .cps with .csv.

# pipeline: single or double parameter scan

- Tick the flag executable in the Parameter Scan Task.
- Select a report template for the Parameter Scan Task.
- Save the report in the same folder with the same name as the model but replacing the extension .cps with .csv.

#### pipeline: param-estim

• Tick the flag executable in the Parameter Estimation Task.

- Select the report template for the Parameter Estimation Task.
- Save the report in the same folder with the same name as the model but replacing the extension .cps with .csv.

# Running SB pipe

SB pipe is executed via the command *run\_sb\_pipe.py*. The syntax for this command and its complete list of options can be retrieved by running *run\_sb\_pipe.py* -h.

As of Sep 2016 the output is as follows:

```
pdp@ariel:~/sb_pipe$ run_sb_pipe.py -h
Usage: run_sb_pipe.py [OPTION] [FILE]
Pipelines for systems modelling of biological networks.
List of mandatory options:
        -h, --help
                 Shows this help.
        -c, --create-project
                 Create a project structure using the argument as name.
        -s, --simulate
                 Simulate a model.
         -p, --single-param-scan
                Simulate a single parameter scan.
        -d, --double-param-scan
                Simulate a double parameter scan.
        -e, --param-estim
                 Generate a parameter fit sequence.
Exit status:
0 if OK,
1 if minor problems (e.g., a pipeline did not execute correctly),
2 if serious trouble (e.g., cannot access command-line argument).
Report bugs to sb_pipe@googlegroups.com
SB pipe home page: <a href="https://pdp10.github.io/sb_pipe">https://pdp10.github.io/sb_pipe</a>
For complete documentation, see \ensuremath{\mathtt{README.md}} .
```

The first step is to create a new project. This can be done with the command:

```
run_sb_pipe.py --create-project project_name
```

This generates the following structure:

Models must be stored in the Models/ folder. The folder Data/ is meant for collecting experimental data files and analyses in one place. Once the data files for Copasi (e.g. for parameter estimation) are generated, **it is advised** to move them into the Models/ folder so that the Copasi (.cps) file and its associated experimental data files are stored in the same folder. To run SB pipe, users need to create a configuration file for each pipeline they intend to run (see next section). These configuration files should be placed in the Working\_Folder/. This folder will eventually contain all the results generated by SB pipe.

For instance, the pipeline for parameter estimation configured with a certain configuration file can be executed by typing:

```
run_sb_pipe.py -e my_config_file.conf
```

# Pipeline configuration files

Pipelines are configured using files (here called configuration files). These files are INI files and are therefore structured as follows:

```
[pipeline_name]
option1=value1
option2=value2
...
```

In SB pipe each pipeline executes three tasks: data generation, data analysis, and report generation. Each task depends on the previous one. This choice allows user to analyse the same data without having to generate it every time, or to skip the report generation if not wanted. Assuming that the configuration files are placed in the Working\_Folder of a certain project, examples are given as follow:

### **Example 1:** configuration file for the pipeline *simulate*

```
[simulate]
# True if data should be generated, False otherwise
generate_data=True
# True if data should be analysed, False otherwise
analyse_data=True
# True if a report should be generated, False otherwise
generate_report=True
# The relative path to the project directory (from Working_Folder)
project_dir=..
# The Copasi model name
model=insulin_receptor_stoch.cps
# The cluster type. pp if the model is run locally,
# sge/lsf if run on cluster.
cluster=pp
# The number of CPU if pp is used, ignored otherwise
pp_cpus=7
# The number of simulations to perform.
\# n>=1 for stochastic simulations.
runs=40
# An experimental data set (or blank) to add to the
# simulated plots as additional layer
exp_dataset=insulin_receptor_dataset.csv
# True if the experimental data set should be plotted.
plot_exp_dataset=True
# The label for the x axis.
xaxis_label=Time [min]
# The label for the y axis.
yaxis_label=Level [a.u.]
```

# **Example 2:** configuration file for the pipeline *single\_param\_scan*

```
[single_param_scan]
# True if data should be generated, False otherwise
generate_data=True
# True if data should be analysed, False otherwise
analyse_data=True
# True if a report should be generated, False otherwise
generate_report=True
# The relative path to the project directory (from Working_Folder)
project_dir=..
# The Copasi model name
model=insulin_receptor_inhib_scan_IR_beta.cps
# The variable to scan (as set in Copasi Parameter Scan Task)
scanned_par=IR_beta
# The number of intervals in the simulation
simulate__intervals=100
# The number of simulations to perform for each scan
```

```
single_param_scan_simulations_number=1
# True if the variable is only reduced (knock down), False otherwise.
single_param_scan_knock_down_only=True
# True if the scanning represents percent levels.
single_param_scan_percent_levels=True
# The minimum level (as set in Copasi Parameter Scan Task)
min_level=0
# The maximum level (as set in Copasi Parameter Scan Task)
max_level=100
# The number of scans (as set in Copasi Parameter Scan Task)
levels_number=10
# True if plot lines are the same between scans
# (e.g. full lines, same colour)
homogeneous_lines=False
# The label for the x axis.
xaxis_label=Time [min]
# The label for the y axis.
yaxis_label=Level [a.u.]
```

### **Example 3:** configuration file for the pipeline *double\_param\_scan*

```
[double_param_scan]
# True if data should be generated, False otherwise
generate_data=True
# True if data should be analysed, False otherwise
analyse_data=True
# True if a report should be generated, False otherwise
generate_report=True
# The relative path to the project directory (from Working_Folder)
project_dir=..
# The Copasi model name
model=insulin_receptor_inhib_dbl_scan_InsulinPercent__IRbetaPercent.cps
# The 1st variable to scan (as set in Copasi Parameter Scan Task)
scanned par1=InsulinPercent
# The 2nd variable to scan (as set in Copasi Parameter Scan Task)
scanned_par2=IRbetaPercent
# The simulation length (as set in Copasi Time Course Task)
```

### **Example 4:** configuration file for the pipeline *param\_estim*

```
[param_estim]
# True if data should be generated, False otherwise
generate_data=True
# True if data should be analysed, False otherwise
analyse_data=True
# True if a report should be generated, False otherwise
generate_report=True
# True if a zipped tarball should be generated, False otherwise
generate_tarball=True
# The relative path to the project directory (from Working_Folder)
project_dir=..
# The Copasi model name
model=insulin_receptor_param_estim.cps
# The cluster type. pp if the model is run locally,
# sge/lsf if run on cluster.
cluster=pp
# The number of CPU if pp is used, ignored otherwise
# The parameter estimation round which is used to distinguish
# phases of parameter estimations when parameters cannot be
# estimated at the same time
round=1
```

```
# The number of parameter estimations
# (the length of the fit sequence)
runs=250
# The threshold percentage of the best fits to consider
best_fits_percent=75
# The number of available data points
data_point_num=33
# True if 2D all fits plots for 66% confidence levels
# should be plotted. This can be computationally expensive.
plot_2d_66cl_corr=True
# True if 2D all fits plots for 95% confidence levels
# should be plotted. This can be computationally expensive.
plot_2d_95cl_corr=True
# True if 2D all fits plots for 99% confidence levels
# should be plotted. This can be computationally expensive.
plot_2d_99cl_corr=True
# True if parameter values should be plotted in log space.
logspace=True
# True if plot axis labels should be plotted in scientific notation.
scientific_notation=True
```

Additional examples of configuration files can be found in:

```
${SB_PIPE}/tests/insulin_receptor/Working_Folder/
```

# Reporting bugs or requesting new features

SB pipe is a relatively young project and there is a chance that some error occurs. The following mailing list should be used for general questions:

```
sb_pipe AT googlegroups.com
```

All the topics discussed in this mailing list are also available at the website:

https://groups.google.com/forum/#!forum/sb\_pipe

To help us better identify and reproduce your problem, some technical information is needed. This detail data can be found in SB pipe log files which are stored in \${HOME}/.sb\_pipe/logs/. When using the mailing list above, it would be worth providing this extra information.

Issues and feature requests can also be notified using the github issue tracking system for SB pipe at the web page:

https://github.com/pdp10/sb\_pipe/issues.

# **DEVELOPER MANUAL**

Mailing list: sb\_pipe AT googlegroups.com

Forum: https://groups.google.com/forum/#!forum/sb\_pipe

# Introduction

This guide is meant for developers and contains guidelines for developing this project.

# **Development model**

This project follows the Feature-Branching model. Briefly, there are two main branches: master and develop. The former contains the history of stable releases, the latter contains the history of development. The master branch contains checkout points for production hotfixes or merge points for release-x.x.x branches. The develop branch is used for feature-bugfix integration and checkout point in development. Nobody should directly develop in here. The develop branch is versionless (just call it -dev).

#### **Conventions**

To manage the project in a more consistent way, here is a list of conventions to follow:

- Each new feature is developed in a separate branch forked from *develop*. This new branch is called *featureNUMBER*, where *NUMBER* is the number of the GitHub Issue discussing that feature. The first line of each commit message for this branch should contain the string *Issue #NUMBER* at the beginning. Doing so, the commit is automatically recorded by the Issue Tracking System for that specific Issue. Note that the sharp (#) symbol is required.
- The same for each new bugfix, but in this case the branch name is called bugfixNUMBER.
- The same for each new hotfix, but in this case the branch name is called hotfixNUMBER and is forked from *master*.

# Work flow

The procedure for checking out a new feature from the develop branch is:

```
$ git checkout -b feature10 develop
```

This creates the feature10 branch off develop. This feature10 is discussed in *Issue #10* in GitHub. When you are ready to commit your work, run:

```
$ git commit -am "Issue #10, summary of the changes. Detailed
description of the changes, if any."
$ git push origin feature10  # sometimes and at the end.
```

As of June 2016, the branches master and develop are protected and a status check using Travis-CI must be performed before merging or pushing into these branches. This automatically forces a merge without fast-forward. In order to merge any new feature, bugfix or simple edits into master or develop, a developer must checkout a new branch and, once committed and pushed, merge it to master or develop using a pull request. To merge feature10 to develop, the pull request output will look like this in GitHub Pull Requests:

```
base:develop compare:feature10 Able to merge. These branches can be automatically merged.
```

A small discussion about feature 10 should also be included to allow other users to understand the feature.

Finally delete the branch:

```
$ git branch -d feature10  # delete the branch feature10 (locally)
```

# **New releases**

When the develop branch includes all the desired feature for a release, it is time to checkout this branch in a new one called release-x.x.x. It is at this stage that a version is established. Only bugfixes or hotfixes are applied to this branch. When this testing/correction phase is completed, the master branch will merge with the release-x.x.x branch, using the commands above. To record the release add a tag:

```
git tag -a v1.3 -m "PROGRAM_NAME v1.3"
```

To transfer the tag to the remote server:

```
git push origin v1.3 # Note: it goes in a separate 'branch'
```

To see all the releases:

```
git show
```

# Package structure

This section presents the structure of the SB pipe package. The root of the project contains general management scripts for cleaning the package (clean\_pacakge.py), installing Python and R dependencies (install\_pydeps.py and install\_rdeps.r), and installing SB pipe (setup.py). Additionally, the logging configuration file (logging\_config.ini) is also at this level.

In order to automatically compile and run the test suite, Travis-CI is used and configured accordingly (.travis.yml).

The project is structured as follows:

These folders will be discussed in the next sections. In SB pipe, Python is the project main language. Instead, R is essentially used for computing statistics within the *data analysis tasks* (see section configuration file in User manual) and for generating plots. This choice allows users to run these scripts independently of SB pipe if needed using an R environment like Rstudio. This can be convenient if further data analysis are needed or plots need to be annotated or edited.

# docs

The folder *docs*/ contains the documentation for this project. The user and developer manuals are contained inside the subfolder *source*. In order to generate the complete documentation for SB pipe, the following packages must

be installed:

- python-sphinx
- pandoc
- · texlive-fonts-recommended
- texlive-latex-extra

By default the documentation is generated in html and LaTeX/PDF. Instruction for generating or cleaning SB pipe documentation are provided below.

To generate the source code documentation:

```
$ ./gen_doc.sh
```

To clean the documentation:

```
$ ./clean_doc.sh
```

If new folders containing new Python modules are added to the project, it is necessary to update the sys.path in *source/conf.py* to include these additional paths.

# sb\_pipe

This folder contains the main script for running SB pipe (run\_sb\_pipe.py). This script is an interface for the project.

# pipelines

The folder /sb\_pipe/pipelines/ contains the following pipelines within folders:

- create\_project: creates a new project
- *simulate*: simulates a model deterministically or stochastically using Copasi (this must be configured first), generate plots and report;
- single\_param\_scan: runs Copasi (this must be configured first), generate plots and report;
- double\_param\_scan: runs Copasi (this must be configured first), generate plots and report;
- param\_estim: generate a fits sequence using Copasi (this must be configured first), generate tables for statistics.

These pipelines are invoked directly via the script *sb\_pipe/run\_sb\_pipe.py*. Each pipeline extends the class *Pipeline*, which represents a generic and abstract pipeline. Each pipeline must implement the following methods of *Pipeline*:

```
def run(self, config_file)
def read_configuration(self, lines)
```

The method *run()* contains the procedure to execute for a specific configuration file. The method *read\_configuration()* is needed for reading the options required by the pipeline to execute. The class *Pipeline* contains already implements the INI parser and returns each pipeline the configuration file as a list of lines.

#### utils

The folder *sb\_pipe/utils/* contains the following structure:

- python: a collection of python utils.
- R: a collection of R utils (plots and statistics).

# tests

The folder *tests/* contains the script *run\_tests.py* to run a test suite. It should be used for testing the correct installation of SB pipe dependencies as well as reference for configuring a project before running any pipeline. Projects inside the folder tests/ have the SB pipe project structure:

- Data: (e.g. training / testing data sets for the model);
- *Model*: (e.g. Copasi models, datasets directly used by Copasi models);
- Working\_Folder: (e.g. pipelines configurations and parameter estimation results, time course, parameter scan etc)

Examples of configuration files (\*.conf) can be found in \${SB\_PIPE}/tests/insulin\_receptor/Working\_Folder/.

Travis-CI runs SB pipe tests using nosetests. Please see .travis.yml for detail.

# Miscellaneous of useful commands

#### Git

# Startup

```
$ git clone https://YOURUSERNAME@server/YOURUSERNAME/sb_pipe.git
# to clone the master
$ git checkout -b develop origin/develop
# to get the develop branch
$ for b in `git branch -r | grep -v -- '->'`; do git branch
--track ${b##origin/} $b; done  # to get all the other branches
$ git fetch --all  # to update all the branches with remote
```

#### **Update**

```
$ git pull [--rebase] origin BRANCH # ONLY use --rebase for private branches. Never use it for shared branches otherwise it breaks the history. --rebase moves your commits ahead. I think for shared branches, you should use `git fetch && git merge --no-ff`.

**[FOR NOW, DON'T USE REBASE BEFORE AGREED]**.
```

#### File system

```
$ git rm [--cache] filename
$ git add filename
```

#### Information

```
$ git status
$ git log [--stat]
$ git branch # list the branches
```

### Maintenance

```
$ git fsck  # check errors
$ git gc  # clean up
```

# Rename a branch locally and remotely

```
git branch -m old_branch new_branch  # Rename branch locally
git push origin :old_branch  # Delete the old branch
git push --set-upstream origin new_branch  # Push the new branch, set
local branch to track the new remote
```

### Reset

```
git reset --hard HEAD  # to undo all the local uncommitted changes
```

# Syncing a fork (assuming upstreams are set)

```
git fetch upstream
git checkout develop
git merge upstream/develop
```

# **SOURCE CODE**

# **Python modules**

# basic sync counter module

```
class basic_sync_counter.BasicSyncCounter
```

This is a monitor. It is a callback class for collecting information about finished processes. It is used by Parallel Python (pp).

add (pid, value)

The callback function

#### **Parameters**

- pid this is callbackargs passed to parallel python *submit()* method
- **value** the return value of the parallelised function. It is the callback value.

```
get_count()
```

Return the counter

**Returns** the number of running processes.

```
get value()
```

Return the internal status.

**Returns** True if the counter is empty.

# collect\_results module

```
collect_results.get_parameter_names_list (filein)
```

Return the list of parameter names from filein

Parameters filein – a Copasi parameter estimation report file

**Returns** the list of parameter names

Collect all the parameter estimates from the Copasi parameter estimation report. Results are stored in filename\_out.

- path\_in the path to the input files
- path\_out the path to the output files
- **filename\_out** the filename to store the final estimates

Collect the final parameter estimates from the Copasi parameter estimation report. Results are stored in filename out.

#### **Parameters**

- path\_in the path to the input files
- path\_out the path to the output files
- **filename** out the filename to store the final estimates

```
collect_results.retrieve_input_files (path)
```

Retrieve the input files in a path.

**Parameters** path – the path containing the input files to retrieve

**Returns** the list of input files

collect\_results.write\_all\_estimates (files, path\_out, filename\_out)

Write all the estimates to filename\_out

#### **Parameters**

- **files** the list of Copasi parameter estimation reports
- path\_out the path to store the file combining all the estimates
- filename\_out the file containing all the estimates

```
collect_results.write_final_estimates (files, path_out, filename_out)
```

Write the final estimates to filename\_out

#### **Parameters**

- **files** the list of Copasi parameter estimation reports
- path\_out the path to store the file combining the final (best) estimates (file-name\_out)
- **filename\_out** the file containing the final (best) estimates

```
collect_results.write_parameter_names (colNames, path_out, filename_out)
```

Write the list of parameter names to filename\_out

#### **Parameters**

- colNames the list of parameter names
- path\_out the path to store filename\_out
- **filename\_out** the output file to store the parameter names

# copasi parser module

```
class copasi_parser.CopasiParser
```

Retrieve information from a Copasi file.

```
retrieve_param_estim_values(file_in)
```

Parse a Copasi file and retrieve information on the parameters to estimate.

**Parameters** file\_in – the Copasi file including absolute path to parse

**Returns** a tuple containing the report file name, the parameter lower bounds, names, starting values, and upper bounds

# copasi\_utils module

```
copasi_utils.replace_str_copasi_sim_report (report)

Replace a group of annotation strings from a generated copasi report file
```

Parameters report – The report file with absolute path

# create project module

This module initialises the folder tree for a new project.

#### **Parameters**

- data\_folder the folder containing the data
- models\_folder the folder containing the models
- working\_folder the folder to store the results

run (project\_name)

Create a project directory tree.

Parameters project\_name - the name of the project
Returns 0

# double param scan module

This module provides the user with a complete pipeline of scripts for computing a double parameter scan using copasi.

**static analyse\_data** (*model*, *scanned\_par1*, *scanned\_par2*, *inputdir*, *outputdir*) The second pipeline step: data analysis.

#### **Parameters**

- model the model name
- scanned par1 the first scanned parameter
- scanned\_par2 the second scanned parameter
- inputdir the directory containing the simulated data sets to process
- **outputdir** the directory to store the performed analysis

static generate\_data (model, sim\_length, inputdir, outputdir)

The first pipeline step: data generation.

- model the model to process
- **sim\_length** the length of the simulation
- inputdir the directory containing the model
- **outputdir** the directory to store the results

**static generate\_report** (*model*, *scanned\_par1*, *scanned\_par2*, *outputdir*, *sim\_plots\_folder*) The third pipeline step: report generation.

#### **Parameters**

- model the model name
- scanned\_par1 the first scanned parameter
- scanned\_par2 the second scanned parameter
- outputdir the directory containing the report
- **sim\_plots\_folder** the folder containing the plots.

```
read configuration (lines)
```

run (config\_file)

# io\_util\_functions module

```
\verb"io_util_functions.files_with_pattern_recur" (\textit{folder}, \textit{pattern})
```

Return all files with a certain pattern in folder+subdirectories

#### **Parameters**

- folder the folder to search for
- pattern the string to search for

**Returns** the files containing the pattern.

```
io_util_functions.get_pattern_position(pattern, filename)
```

Return the line number (as string) of the first occurrence of a pattern in filename

# **Parameters**

- pattern the pattern of the string to find
- **filename** the file name containing the pattern to search

**Returns** the line number containing the pattern or "-1" if the pattern was not found

```
io_util_functions.refresh_directory(path, file_pattern)
```

Clean and create the folder if this does not exist.

# **Parameters**

- path the path containing the files to remove
- **file\_pattern** the string pattern of the files to remove

io\_util\_functions.replace\_string\_in\_file (filename\_out, old\_string, new\_string)
 Replace a string with another in filename\_out

#### **Parameters**

- filename\_out the output file
- old\_string the old string that should be replaced
- new\_string the new string replacing old\_string

io\_util\_functions.write\_matrix\_on\_file (path, filename\_out, data)

Write the matrix results stored in data to filename\_out

- path the path to filename\_out
- filename\_out the output file
- data the data to store in a file

# latex reports module

latex\_reports.get\_latex\_header(pdftitle='SB pipe report', title='SB pipe report', abstract='Generic report.')

Initialize a Latex header with a title and an abstract.

#### **Parameters**

- pdftitle the pdftitle for the LaTeX header
- title the title for the LaTeX header
- abstract the abstract for the LaTeX header

# Returns the LaTeX header

latex\_reports.latex\_report (outputdir, sim\_plots\_folder, model\_noext, filename\_prefix, cap-tion=False)

Generate a generic report.

#### **Parameters**

- outputdir the output directory
- **sim\_plots\_folder** the folder containing the simulated plots
- model noext the model name
- filename\_prefix the prefix for the LaTeX file
- caption True if figure captions (=figure file name) should be added

 $\label{latex_report_double_param_scan} \begin{tabular}{ll} latex_report_double_param_scan (outputdir, & sim_plots_folder, & filename\_prefix, & model_noext, & scanned\_par1, scanned\_par2) \\ \end{tabular}$ 

Generate a report for a double parameter scan task.

### **Parameters**

- $\bullet$   $\mbox{{\tt outputdir}}$  — the output directory
- **sim\_plots\_folder** the folder containing the simulated plots
- **filename\_prefix** the prefix for the LaTeX file
- model\_noext the model name
- scanned\_par1 the 1st scanned parameter
- scanned\_par1 the 2nd scanned parameter

latex\_reports.latex\_report\_param\_estim(outputdir, sim\_plots\_folder, model\_noext, file-name\_prefix)

Generate a report for a parameter estimation task.

#### **Parameters**

- outputdir the output directory
- **sim\_plots\_folder** the folder containing the simulated plots
- model\_noext the model name
- **filename\_prefix** the prefix for the LaTeX file

latex\_reports.latex\_report\_simulate(outputdir, sim\_plots\_folder, model\_noext, file-name\_prefix)

Generate a report for a time course task.

- outputdir the output directory
- sim plots folder the folder containing the simulated plots

- model noext the model name
- filename\_prefix the prefix for the LaTeX file

```
latex_reports.latex_report_single_param_scan(outputdir, sim_plots_folder, filename_prefix, scanned par) sim_plots_folder, model_noext,
```

Generate a report for a single parameter scan task.

#### **Parameters**

- **outputdir** the output directory
- sim\_plots\_folder the folder containing the simulated plots
- **filename\_prefix** the prefix for the LaTeX file
- model\_noext the model name
- scanned\_par the scanned parameter

```
latex_reports.pdf_report (outputdir, filename)
Generate a PDF report from LaTeX report using pdflatex.
```

#### **Parameters**

- **outputdir** the output directory
- filename the LaTeX file name

# parallel computation module

```
parallel_computation.parallel_computation(command, command_iter_substr, cluster_type, runs, output_dir, pp_cpus=1)

Generic funcion to run a command in parallel
```

#### **Parameters**

- command the command string to run in parallel
- **command\_iter\_substr** the substring of the iteration number. This will be replaced in a number automatically
- cluster\_type the cluster type among pp (multithreading), sge, or lsf
- runs the number of runs
- output\_dir the output directory
- pp\_cpus the number of cpus that pp should use at most

```
parallel_computation.run_command_instance(command)
Run a command using Python subprocess.
```

Parameters command - the string of the command to run

```
parallel_computation.run_command_pp (command, command_iter_substr, runs, server, sync-
Counter=<basic_sync_counter.BasicSyncCounter
instance>)
```

Run instances of a command in multithreading using parallel python (pp).

- command the command string to run in parallel
- **command\_iter\_substr** the substring of the iteration number. This will be replaced in a number automatically
- runs the number of runs
- server the server that pp should use

• syncCounter – the mutex object to count the jobs

parallel\_computation.run\_jobs\_lsf (command, command\_iter\_substr, outDir, errDir, runs)
Run jobs using a Load Sharing Facility (LSF) cluster.

#### **Parameters**

- command the full command to run as a job
- command\_iter\_substr the substring in command to be replaced with a number
- outDir the directory containing the standard output from bsub
- errDir the directory containing the standard error from bsub
- runs the number of runs to execute

parallel\_computation.run\_jobs\_pp (command, command\_iter\_substr, runs, pp\_cpus=1)
 Run jobs using parallel python (pp) locally.

#### **Parameters**

- command the full command to run as a job
- command\_iter\_substr the substring in command to be replaced with a number
- runs the number of runs to execute
- pp\_cpus The number of available cpus. If pp\_cpus <=0, all the available cores will be used

parallel\_computation.run\_jobs\_sge (command, command\_iter\_substr, outDir, errDir, runs)
Run jobs using a Sun Grid Engine (SGE) cluster.

#### **Parameters**

- command the full command to run as a job
- command\_iter\_substr the substring in command to be replaced with a number
- outDir the directory containing the standard output from qsub
- errDir the directory containing the standard error from qsub
- runs the number of runs to execute

# param estim module

This module provides the user with a complete pipeline of scripts for running a model parameter estimation using copasi

```
static analyse_data (model, inputdir, outputdir, fileout_final_estims, fileout_all_estims, fileout_approx_ple_stats, fileout_conf_levels, sim_plots_dir, best_fits_percent, data_point_num, plot_2d_66cl_corr=False, plot_2d_95cl_corr=False, plot_2d_99cl_corr=False, logspace=True, scientific_notation=True)

The second pipeline step: data analysis.
```

- model the model name
- inputdir the directory containing the simulation data
- outputdir the directory to store the results

- **fileout\_final\_estims** the name of the file containing final parameter sets with Chi^2
- **fileout\_all\_estims** the name of the file containing all the parameter sets with Chi^2
- **fileout\_approx\_ple\_stats** the file name of the PLE results
- fileout\_conf\_levels the file name of the confidence levels results
- **sim\_plots\_dir** the directory of the simulation plots
- best\_fits\_percent the percent to consider for the best fits
- data\_point\_num the number of data points
- plot\_2d\_66cl\_corr True if 2 dim plots for the parameter sets within 66% should be plotted
- plot\_2d\_95cl\_corr True if 2 dim plots for the parameter sets within 95% should be plotted
- plot\_2d\_99cl\_corr True if 2 dim plots for the parameter sets within 99% should be plotted
- logspace True if parameters should be plotted in log space
- scientific\_notation True if axis labels should be plotted in scientific notation

static generate\_data (model, inputdir, cluster\_type, pp\_cpus, nfits, outputdir, sim\_data\_dir, up-dated\_models\_dir)

The first pipeline step: data generation.

- Parameters
  - inputdir the directory containing the model
  - **cluster\_type** pp for parallel python, lsf for load sharing facility, sge for sun grid engine
  - **pp\_cpus** the number of cpu for parallel python
  - **nfits** the number of fits to perform

• model – the model to process

- **outputdir** the directory to store the results
- sim\_data\_dir the directory containing the simulation data sets
- updated\_models\_dir the directory containing the Copasi models with updated parameters for each estimation

 $\textbf{static generate\_report} \ (\textit{model}, \textit{outputdir}, \textit{sim\_plots\_folder})$ 

The third pipeline step: report generation.

#### **Parameters**

- model the model name
- **outputdir** the directory to store the report
- **sim\_plots\_folder** the folder containing the plots

read\_configuration(lines)

run (config\_file)

# pipeline module

#### **Parameters**

- data\_folder the folder containing the experimental (wet) data sets
- models\_folder the folder containing the models
- working\_folder the folder to store the results
- sim\_data\_folder the folder to store the simulation data
- sim\_plots\_folder the folder to store the graphic results

# config\_parser (config\_file, section)

Return the configuration for the parsed section in the config\_file

#### **Parameters**

- config\_file the configuration file to parse
- **section** the section in the configuration file to parse

**Returns** the configuration for the parsed section in the config\_file

#### get\_data\_folder()

Return the folder containing the experimental (wet) data sets.

Returns the experimental data sets folder.

### get\_models\_folder()

Return the folder containing the models.

Returns the models folder.

```
get_sim_data_folder()
```

Return the folder containing the in-silico generated data sets.

**Returns** the folder of the simulated data sets.

```
get_sim_plots_folder()
```

Return the folder containing the in-silico generated plots.

**Returns** the folder of the simulated plots.

```
get_working_folder()
```

Return the folder containing the results.

**Returns** the working folder.

### read\_common\_configuration(lines)

Parse the common parameters from the configuration file

**Returns** return a tuple containing the common parameters

```
read_configuration (lines)
```

Read the section lines from the configuration file. This method is abstract.

**Returns** a tuple containing the configuration

```
run (config_file)
```

Run the pipeline.

Parameters config\_file - a configuration file for this pipeline.

**Returns** 0 if the pipeline was executed correctly, 1 if the pipeline executed but some output was skipped, 2 if the pipeline did not execute correctly.

# random\_functions module

```
\verb"random_functions.get_rand_alphanum_str" (\textit{length})
```

Return a random alphanumeric string

Parameters length - the length of the string

**Returns** the generated string

```
\verb"random_functions.get_rand_num_str" (\textit{length})
```

Return a random numeric string

**Parameters** length – the length of the string

**Returns** the generated string

# randomise parameters module

```
class randomise_parameters.RandomiseParameters (path, filename_in)
```

This class generates multiple copies of a Copasi file configured for parameter estimation task, and randomises the starting values of the parameters to estimate.

#### **Parameters**

- path the path to filename\_in
- **filename\_in** the Copasi file to process.

```
generate_instances_from_template(num_files, idstr)
```

Generate num\_files files and randomise the starting values for the parameter to estimate.

#### **Parameters**

- num\_files the number of files (instances) to generate
- idstr an ID string to label the generated files (e.g. a timestamp)

```
get_copasi_obj()
```

Return the Copasi parser object

Returns the Copasi parser object

```
get_lower_bounds_list()
```

Return the list of parameter lower bounds

**Returns** the list of parameter lower bounds

```
get_param_names_list()
```

Return the list of parameter names

**Returns** the list of parameter names

```
get_path()
```

Return the path containing the template Copasi file

Returns the path to the Copasi file

```
{\tt get\_report\_filename\_template\_str}\:(\:)
```

Return the name of the template parameter estimation report

**Returns** the name of the report file name for parameter estimation

```
get_start_values_list()
```

Return the list of parameter starting values

**Returns** the list of parameter starting values

```
get_template_copasi_file()
```

Return the name of the template Copasi file

Returns the name of the Copasi file

```
get_upper_bounds_list()
```

Return the list of parameter upper bounds

**Returns** the list of parameter upper bounds

```
print_parameters_to_estimate()
```

Print the parameter names, lower/upper bounds, and starting value, as extracted from COPASI template file

# re utils module

```
re_utils.natural_sort_key(str)
```

The key to sort a list of strings alphanumerically (e.g. "file10" is correctly placed after "file2")

Parameters str - the string to sort alphanumerically in a list of strings

**Returns** the key to sort strings alphanumerically

# sensitivity module

This module provides the user with a complete pipeline of scripts for computing model sensitivity analysis using Copasi

```
static analyse_data(outputdir)
```

The second pipeline step: data analysis.

**Parameters** outputdir – the directory to store the performed analysis

static generate\_data (model, inputdir, outputdir)

The first pipeline step: data generation.

#### **Parameters**

- model the model to process
- inputdir the directory containing the model
- **outputdir** the directory to store the results

static generate\_report (model, outputdir, sim\_plots\_folder)

The third pipeline step: report generation.

#### **Parameters**

- model the model name
- **outputdir** the directory to store the report
- **sim\_plots\_folder** the directory containing the time courses results combined with experimental data

```
{\tt read\_configuration}~(\mathit{lines})
```

run (config\_file)

#### simulate module

This module provides the user with a complete pipeline of scripts for running a model simulation using copasi

static analyse\_data (model, inputdir, outputdir, sim\_plots\_dir, exp\_dataset, plot\_exp\_dataset, xaxis\_label, yaxis\_label)

The second pipeline step: data analysis.

### **Parameters**

- model the model name
- inputdir the directory containing the data to analyse
- outputdir the output directory containing the results
- **sim\_plots\_dir** the directory to save the plots
- exp\_dataset the full path of the experimental data set
- plot\_exp\_dataset True if the experimental data set should also be plotted
- **xaxis\_label** the label for the x axis (e.g. Time [min])
- yaxis\_label the label for the y axis (e.g. Level [a.u.])

**static generate\_data** (*model*, *inputdir*, *outputdir*, *cluster\_type='pp'*, *pp\_cpus=2*, *runs=1*) The first pipeline step: data generation.

#### **Parameters**

- model the model to process
- inputdir the directory containing the model
- outputdir the directory containing the output files
- cluster\_type pp for local Parallel Python, lsf for Load Sharing Facility, sge for Sun Grid Engine.
- pp\_cpus the number of CPU used by Parallel Python.
- runs the number of model simulation

static generate report (model, outputdir, sim plots folder)

The third pipeline step: report generation.

#### **Parameters**

- model the model name
- outputdir the output directory to store the report
- **sim\_plots\_folder** the folder containing the plots

```
{\tt read\_configuration}\ (\mathit{lines})
```

run (config\_file)

# single param scan module

This module provides the user with a complete pipeline of scripts for computing a single parameter scan using copasi.

static analyse\_data (model, scanned\_par, knock\_down\_only, outputdir, sim\_data\_folder, sim\_plots\_folder, simulations\_number, percent\_levels, min\_level, max\_level, levels\_number, homogeneous\_lines, xaxis\_label, yaxis\_label)

The second pipeline step: data analysis.

#### **Parameters**

- model the model name
- scanned\_par the scanned parameter
- knock\_down\_only True for knock down simulation, false if also scanning over expression.
- outputdir the directory containing the results
- **sim\_data\_folder** the folder containing the simulated data sets
- sim\_plots\_folder the folder containing the generated plots
- simulations\_number the number of simulations
- **percent\_levels** True if the levels are percents.
- min\_level the minimum level
- max\_level the maximum level
- levels number the number of levels
- homogeneous\_lines True if generated line style should be homogeneous
- **xaxis\_label** the name of the x axis (e.g. Time [min])
- yaxis\_label the name of the y axis (e.g. Level [a.u.])

#### **Parameters**

- model the model to process
- scanned\_par the scanned parameter
- sim number the number of simulations (for det sim: 1, for stoch sim: n>1)
- **simulate\_intervals** the time step of each simulation
- single\_param\_scan\_intervals the number of scans to perform
- inputdir the directory containing the model
- **outputdir** the directory to store the results

**static generate\_report** (*model*, *scanned\_par*, *outputdir*, *sim\_plots\_folder*)

The third pipeline step: report generation.

#### **Parameters**

• model – the model name

- $\bullet \ \, \textbf{scanned\_par} the \ scanned \ parameter$
- **outputdir** the directory containing the report
- **sim\_plots\_folder** the folder containing the plots

 ${\tt read\_configuration}\ (\mathit{lines})$ 

run (config\_file)

**CHAPTER** 

**FOUR** 

# **META INFORMATION**

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