

CBMPy User Guide

Release 0.8.8

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CBMPY: INSTALLATION GUIDE

1.1 Introduction

1.1.1 Support

PySCeS CBMPy is Open Source software released under the GNU GPL 3 licence (included with the source code) and is in constant development. All the latest downloads, documentation and development information is available on **SourceForge**: http://cbmpy.sourceforge.net

CBMPy is developed in the Department of Systems Bioinformatiocs at the Vrije Universiteit Amsterdam, as part of the BeBasic Metatoolkit project, by Brett Olivier (bgoli@users.sourceforge.net)

1.1.2 Python standard library modules

CBMPy is developed and tested on Python 2.7.x. but should also work on Python 3. The following Python Standard Library modules are used in CBMPy and should be available as part of any CPython distribution and not require additional installation:

```
'cPickle', 'cStringIO', 'cgi', 'copy', 'gc', 'itertools', 'locale', 'math',
'multiprocessing', 'os', 'pprint', 'random', 're', 'shutil', 'subprocess',
'time', 'urllib2', 'webbrowser', 'xml'
```

1.1.3 Required libraries (Python bindings)

Besides those mentioned above, the following packages are required for CBMPy's core functionality. Note that it is possible to install CBMPy using only *numpy* but that only very limited subset of functionality is then available. CBMPy is primarily developed on Microsoft Windows and Ubuntu Linux and where possible the package name is provided such that can be used with the software center or package manager sudo apt-get install package (please see the man pages for sudo and apt-get if you don't know what this). A comprehensive list of modules are listed at the end of this document. In the case of external C/C++ libraries the Python bindings should be installed (e.g. libSBML). Many of these are available in *batteries included* Python distributions and via the pip installer.

1.2 Overview

CBMPy has been designed to be used as a framework and can be used in different contexts. Here are the minimal requirements for each role. The fastest way to install the latest official version of CBMPy is using PyPI: https://pypi.python.org/pypi/cbmpy while development source, binaries, tools and utilities are available from SourceForge: https://sourceforge.net/projects/cbmpy/files

1.2.1 PyPI

Try one of:

```
pip install cbmpy
easy_install cbmpy
```

1.2.2 Minimal

Read, write and convert model files as well as view, create and edit model components and annotation.

- numpy
- libSBML (Python bindings)
- biopython

1.2.3 Normal

In addition to the minimal requirements a linear optimizer is required

- Optimization libraries (one or more of):
- CPLEX (LP, MILP): http:://www.ibm.com
- GLPK (LP): http://tfinley.net/software/pyglpk/
- matplotlib
- sympy
- xlwt
- xlrd

1.2.4 Full

This includes graphical interface support

- pyqt4
- wxPython
- suds
- scipy
- h5py
- networkx

1.2.5 User tools

These are highly recommended user tools but are not required for using CBMPy.

- iPython
- · iPython-notebook
- SCiTE

1.3 Installing on Ubuntu Linux

1.3.1 Python 2.7 (full install)

First we create a scientific Python workbench using the Ubuntu package manager:

```
sudo apt-get install python-dev python-numpy python-scipy python-matplotlib python-pip sudo apt-get install python-sympy python-suds python-xlrd python-xlwt python-h5py sudo apt-get install python-biopython python-wxgtk2.8 python-qt4 sudo apt-get install ipython ipython-notebook
```

1.3.2 libSBML

Installing libSBML is now easy using PiP and PyPI, first we need some dependencies:

```
sudo apt-get install libxml2 libxml2-dev
sudo apt-get install zlib1g zlib1g-dev
sudo apt-get install bzip2 libbz2-dev
```

Then we can install the latest and greatest with:

```
sudo pip install python-libsbml
```

1.3.3 glpk/python-glpk

Currently, CBMPy still requires GLPK version 4.47 to work with glpk-0.3 so it is important *not* to install the latest using the ubuntu package manager but rather use these instructions. First download the GLPK source and bindings:

```
https://sourceforge.net/projects/cbmpy/files/tools/glpk/glpk-4.47.tar.gz
https://sourceforge.net/projects/cbmpy/files/tools/glpk/glpk-0.3.tar.bz2
```

and then install the the following dependency:

```
sudo apt-get install libgmp-dev
```

Unpack the glpk-4.47 source and make it your current directory:

```
cd GLPK source (e.g. glpk-4.47)
```

Build and install GLPK:

```
./configure --with-gmp
make
make check
sudo make install
sudo ldconfig
```

Change to the python-glpk source (glpk-0.3) and make it the current directory:

```
cd to python-glpk source (glpk-0.3):
```

Then build and install it with the following commands:

```
make
sudo make install
```

Finally, install CBMPy either using PyPI:

```
pip install cbmpy
or
easy_install cbmpy
```

Or download a source file from either:

```
https://pypi.python.org/pypi/cbmpy
https://sourceforge.net/projects/cbmpy/files
```

and install manually:

```
python setup.py build sdist
sudo python setup.py install
```

1.4 Installing on Microsoft Windows

For the modeller that does not want to customize his installation and install all of the individual packages by hand there are some *batteries included* Python distributions which have many (if not all) of the required packages required.

CBMPy is developed using a 64 bit version of *Anaconda Python* 2.7 (http://continuum.io) has also been installed on 32 bit *Python*(*x*, *y*) and Enthought Python Distribution (EPD) http://www.enthought.com

1.4.1 Anaconda

Most of the required packages can be installed using the *conda* package manager for example:

```
conda install sympy pyqt4
```

or using pip (libSBML) or setuptools

1.4.2 Python(x,y)

Select the following packages from the *Python* branch of the Python(x,y) installation directory:

- WxPython
- Sympy
- NetworkX
- xlrd
- xlwt
- h5py
- · wxPython
- PyQT4

1.4.3 Installing CBMPy

There are two ways to install CBMPy either download the latest release as source bundle or binary from http://cbmpy. sourceforge.net and unzip or execute from a a temporary directory (recommended). Or, if you want the latest (greatest and potentially broken) version grab the latest revision from the the CBMPy Subversion repository:

```
svn co http://sourceforge.net/p/cbmpy/code/HEAD/tree/trunk/cbmpy cbmpy
```

In both cases you should should now have a directory that contains a file *setup.py* which can install by simply typing the following into a Windows shell (command line):

```
python setup.py build
python setup.py install
```

1.4.4 Installing libSBML with Python bindings

It is highly recommended to install libSBML which CBMPy uses to provide support for the Systems Biology Markup Language (SBML). First go to the libSBML download page http://sbml.org/Software/libSBML page follow the *Download libSBML -> Stable -> Windows -> 32bit* path and download libSBML (e.g. libSBML-5.10.0-win-x86.exe). The latest stable version can be found at http://sbml.org/Software/libSBML

```
http://sourceforge.net/projects/sbml/files/libsbml/5.10.0/stable/Windows/32-bit/libSBML-5.10. 0-win-x86.exe/download
```

Run the installer and make sure you select the Python Bindings during installation or install the appropriate Python bindings that match your Python(x,y) version directly e.g. (libSBML-5.10.0-win-py2.7-x86.exe)

1.4.5 Optmization (1): IBM cplex optimization studio (Academic)

If you have access to the the IBM CPLEX solver. It is a a good idea to use the latest available version. Again choose the appropriate 32 or 64 bit version and an installation path that suites your setup.

- Run cplex_studio126.win-x86-32.exe
- Select English language and accept licence
- Set "Program" install directory to C:\ILOG\CPLEX_Studio126
- Allow default associations to be set and PATH update

Once installation is complete we need to install the Python bindings

- · Open a terminal
- Execute cd c:\\ILOG\\CPLEX_Studio126\\cplex\\python\\x86_win32
- Execute python setup.py install

1.4.6 Optmization (2): GLPK

CBMPy 0.7.4 includes support for the free, Open Source GLPK solver. This allows access to CBMPy's LP functionality (MILP's requires CPLEX). A port of PyGLPK 0.3 is maintained by the OpenCOBRA project which is mirrored here:

https://sourceforge.net/projects/cbmpy/files/tools/glpk/

Select the binary or source distribution you require and either execute the binary:

• Execute glpk-0.3.win32-py2.7.exe

1.4.7 Testing your new installation

If everything has gone according to plan you can test your installation:

- · Open a terminal
- Execute ipython
- In ipython shell, execute import numpy, h5py, xlrd, xlwt

No import errors should occur.

- Execute import libsbml
- Execute libsbml.LIBSBML_VERSION_STRING

A successful test should return (for example):

```
In : libsbml.LIBSBML_VERSION_STRING
Out: '51000'
```

• Execute import cbmpy as cbm

This should return:

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Exit ipython with CTRL-D

If you installed CPLEX then try:

- · Open a terminal
- Execute ipython
- Execute import cplex
- Execute lp = cplex.Cplex()
- Execute lp.solve()

A succesful test should return:

```
In : lp.solve()
Tried aggregator 1 time.
No LP presolve or aggregator reductions.
Presolve time = 0.00 sec.
```

Exit ipython with CTRL-D

If you installed GLPK then try:

- · Open a terminal
- Execute ipython
- Execute import glpk
- Execute lp = glpk.LPX()

A successful test should return:

```
In : glpk.LPX() 
<glpk.LPX 0-by-0 at 0x036C24C8>
```

Exit ipython with CTRL-D

1.4.8 Install CBMPy (http://cbmpy.sourceforge.net)

Download the latest version of CBMPy

- Run **cbmpy-0.7.x.win32.exe** (or newer for 32 bit Windows)
- Run **cbmpy-0.7.x.amd64.exe** (or newer for 64 bit Windows)

Test installation:

- · Open a terminal
- Execute ipython

• Execute import cbmpy as cbm

This should return:

```
In [1]: import cbmpy as cbm
*****
Using GLPK
****
WX GUI tools available.
Qt4 GUI tools available
CBMPy environment
Revision: r346
         *********
* Welcome to CBMPy (0.7.4) - PySCeS Constraint Based Modelling
              http://cbmpy.sourceforge.net
* Copyright(C) Brett G. Olivier 2010 - 2015
* Dept. of Systems Bioinformatics
* Vrije Universiteit Amsterdam, Amsterdam, The Netherlands
* CBMPy is distributed under the GNU GPL v 3.0 licence, see
* LICENCE (supplied with this release) for details
*********
```

Exit ipython with CTRL-D

1.5 Linux: Ubuntu

On Linux many of the base dependencies are available as packages or from the Python Cheeseshop (http://pypi.python. org/pypi). For **libSBML**, **CPLEX** and/or **GLPK** please see the *Generic installation on Microsoft Windows (XP, 7, 2008)* for more details. For example using **Ubuntu** the base dependencies can be easily installed (depending on what functionality is required). If you don't know what these packages are please look them up before installing.

Required:

```
sudo apt-get install python-dev python-numpy
- libSBML for SBML support.
```

Please see http://sbml.org/Software/libSBML or try the following. Depending on your configurationyou need to install libxml2, bzip2 and their associated "dev" packages:

```
apt-get install libxml2 libxml2-dev
apt-get install zlib1g zlib1g-dev
apt-get install bzip2 libbz2-dev
easy_install pip
# for standard libSBML
```

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```
pip install python-libsbml
# for "experimental" libSBML (for FBC V2 and Groups support)
pip install python-libsbml-experimental
```

- Optimization (at least one of):
 - IBM CPLEX: http://www.ibm.com
 - PyGLPK: https://sourceforge.net/projects/cbmpy/files/tools/glpk/

Please note that due to changes in the GLPK API the current version of PyGLPK (0.3) **only supports GLPK up until version 4.47**. If your system has a newer version of GLPK then the current workaround is to uninstall the newer version and compile 4.47 from source (also available from the above directory). Dependencies are standard Linux build tools and GMP etc:

```
tar xzf glpk-4.47.tar.gz
cd glpk-4.47
./configure --with-gmp
make
make check
sudo make install
```

Graphical interfaces (highly recommended):

```
sudo apt-get install python-wxgtk2.8 python-qt4 python-matplotlib
```

Extended IO (highly recommended):

```
sudo apt-get install python-xlrd python-xlwt python-sympy
```

Web services and database:

```
sudo apt-get install python-suds python-pysqlite2
```

Advanced functionality:

```
sudo apt-get install python-scipy python-h5py python-networkx
```

User tools (highly recommended):

```
sudo apt-get install ipython ipython-notebook scite
```

1.6 Linux: Ubuntu 14.04

1.6.1 Python2

First we create a scientific Python workbench:

```
sudo apt-get install python-dev python-numpy python-scipy sudo apt-get install python-matplotlib python-pip sudo apt-get install python-sympy python-suds python-xlrd
```

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```
sudo apt-get install python-xlwt python-h5py
sudo apt-get install python-wxgtk2.8 python-qt4
sudo apt-get install ipython ipython-notebook
```

1.6.2 libSBML

Installing libSBML is now easy using Pip:

```
sudo apt-get install libxml2 libxml2-dev
sudo apt-get install zlib1g zlib1g-dev
sudo apt-get install bzip2 libbz2-dev
sudo pip install python-libsbml
```

1.6.3 glpk/python-glpk

GLPK needs to be version 4.47 to work with glpk-0.3:

```
sudo apt-get install libgmp-dev
```

cd GLPK source (e.g. glpk-4.47):

```
./configure --with-gmp
make
make check
sudo make install
sudo ldconfig
```

cd to python-glpk source (glpk-0.3):

```
make
sudo make install
```

1.6.4 CBMPy

Finally, install CBMPy:

```
python setup.py build sdist
sudo python setup.py install
```

1.6.5 Installing PyscesMarinerCBM

This will install PySCeS Mariner that adds SOAP web-services capability to CBMPy. First unpack pyscesmariner-0.7.7.zip and install the cherrypy webserver:

```
sudo apt-get install python-cherrypy
```

1.6.6 Install soaplib

cd <pysces_cbm_mariner>/misc:

```
tar -xf soaplib-0.8.1.tar.gz
cd soaplib-0.8.1
python setup.py build sdist
sudo python setup.py install
```

1.6.7 Install Mariner

cd <pysces_cbm_mariner> and set mariner configuration (not needed for Ubuntu, Windows or if the server does not read SBML):

cd to <pysces_cbm_mariner>:

```
python setup.py build sdist
sudo python setup.py install
```

1.6.8 Test installation

Open a new terminal window:

```
# cd <pysces_cbm_mariner>/demo
python cbm_server_demo.py
```

Open another terminal and run the client demo:

```
python cbm_client_demo.py
```

Kill the server by closing the terminal window.

1.6.9 Python3

Not all dependencies are available for Python3:

```
sudo apt-get install python3-dev python3-numpy python3-scipy
sudo apt-get install python3-matplotlib python3-pip
sudo apt-get install python3-xlrd python3-h5py

# need to find out what is going on with Python3 and xlwt suds
# easy_install3 sympy ???
# wxPython and PyQt4 not in Ubuntu P3 builds yet

sudo apt-get install ipython3 ipython3-notebook

sudo apt-get install libxml2 libxml2-dev
sudo apt-get install zliblg zliblg-dev
sudo apt-get install bzip2 libbz2-dev

sudo pip3 install python-libsbml

sudo apt-get install python-qt4 python-qt4-dev python-sip
sudo apt-get install python-sip-dev build-essential
```

1.7 Apple Macintosh: OS X

Installation is similar to Linux except packages are installed using distutils and pip. The first step is to install the Mac development tools xcode

Install Python packages:

```
sudo easy_install numpy ipython scipy matplotlib sudo easy_install xlrd xlwt sympy suds pyparsing pip
```

Use pip to install advanced Ipython and libsbml:

```
sudo pip install ipython[notebook]
ARCHFLAGS=-Wno-error=unused-command-line-argument-hard-error-in-future pip install
→python-libsbml
```

For solvers, either install your own copy of CPLEX or build PyGLPK which requires building both the GMP and GLPK libraries.

GMP (https://gmplib.org/):

```
download gmp
./configure --prefix=/usr/local
make
make check
sudo make install
```

GLPK (http://sourceforge.net/projects/cbmpy/files/tools/glpk):

```
download glpk-4.47.tar.gz
./configure --prefix=/usr/local --with-gmp
make
sudo make install
```

PyGLPK (http://sourceforge.net/projects/cbmpy/files/tools/glpk):

```
download python-glpk-0.3
python setup.py build
sudo python setup.py install
```

1.8 Installing PySCeS-CBM Mariner (Microsoft Windows and Linux)

The PySCeS Mariner module exposes the CBMPy functionality as SOAP web services (e.g. as a backend to FAME (http://F-A-M-E.org)). It is available for download from SourceForge:

PySCeS-CBM Mariner: http://sourceforge.net/projects/cbmpy/files/release/pysces_mariner/

1.8.1 Dependencies: CherryPy, libXML and SOAPlib

PySCeS-CBM Mariner requires (pure python) soaplib 0.8.1 (supplied with it) or downloadable from:

```
https//sourceforge.net/projects/cbmpy/files/tools/soaplib/
```

Soaplib itself has two dependencies which should be installed first:

- LXML (http://lxml.de)
 - Windows: install with easy_install lxml
 - Linux (Ubuntu) use sudo apt-get install python-lxml
- CherryPy (http://www.cherrypy.org)
 - Windows: install with easy_install cherrypy
 - Linux (Ubuntu) use sudo apt-get install python-cherrypy
- SOAPLIB 0.8.1:
 - Windows: Execute soaplib-0.8.1.win32.exe
 - Linux: Unpack the zip archive and run sudo python setup.py install

Test installation:

- · Open a terminal
- Execute "ipython"
- Execute "import cherrypy, lxml, soaplib" no errors or warnings should be generated
- Exit ipython with CTRL-D
- change directory to supplied soaplib tests e.g. "cd e:\cbmpy\tests\soaplib"
- Execute "python binary_test.py"
- Execute "python primitive_test.py"

All tests should pass.

1.8.2 PySCeS-CBM Mariner (http://cbmpy.sourceforge.net)

Download and install the latest version (0.7.4 or newer is required for CBMPy 0.7+):

- Windows: Execute pyscesmariner-0.7.7.zip
- Linux: unpack the archive and run sudo python setup.py install

To test installation, on Linux execute the commands in run_server.bat from the terminal directly.

- Open two terminals and in both
- Change directory to supplied PySCeS-CBM Mariner tests e.g. cd e:\\cbmpy\\tests\\pyscesmariner
- In terminal one Execute run_server.bat

Which should now display:

```
E:\\cbmpy\\tests\\pyscesmariner>python cbm_server_demo.py
Mariner using E:\\cbmpy\\tests\\pyscesmariner as a working directory
Mariner server name: 10.0.2.15
Mariner using port: 31313

Welcome to the PySCeS Constraint Based Modelling Toolkit (0.7.4)

<snipped>
Multiple Environment Module (0.6.2 [r1147])

PySCeSCBM/Mariner initialising ... this console is now blocked
```

In terminal two:

• Execute python cbm_client_demo.py

This should end without errors and display done. Congratulations you have successfully installed CBMPy and PySCeS-CBM Mariner!

CHAPTER

TWO

INTRODUCTION

PySCeS CBMPy is a new platform for constraint based modelling and analysis. It has been designed using principles developed in the PySCeS simulation software project: usability, flexibility and accessibility. Its architecture is both extensible and flexible using data structures that are intuitive to the biologist (metabolites, reactions, compartments) while transparently translating these into the underlying mathematical structures used in advanced analysis (LP's, MILP's).

PySCeS CBMPy implements popular analyses such as FBA, FVA, element/charge balancing, network analysis and model editing as well as advanced methods developed specifically for the ecosystem modelling: minimal distance methods, flux minimization and input selection.

To cater for a diverse range of modelling needs PySCeS CBMPy supports user interaction via:

- interactive console, scripting for advanced use or as a library for software development
- GUI, for quick access to a visual representation of the model, analysis methods and annotation tools
- SOAP based web services: using the Mariner framework much high level functionality is exposed for integration into web tools

For more information on the development and use of PySCeS CBMPy visit the website (http:cbmpy.sourceforge.net) for up to date information and feel free to contact the development team (bgoli@users.sourceforge.net).

CBMPY MODULE REFERENCE

3.1 CBMPy: CBCommon module

PySCeS Constraint Based Modelling (http://cbmpy.sourceforge.net) Copyright (C) 2009-2024 Brett G. Olivier, VU University Amsterdam, Amsterdam, The Netherlands

This program is free software: you can redistribute it and/or modify it under the terms of the GNU General Public License as published by the Free Software Foundation, either version 3 of the License, or (at your option) any later version.

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You should have received a copy of the GNU General Public License along with this program. If not, see http://www.gnu.org/licenses/

Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBCommon.py 710 2020-04-27 14:22:34Z bgoli \$)

class ComboGen

Generate sets of unique combinations

class MIRIAMannotation

The MIRIAMannotation class MIRIAM annotations: Biological Qualifiers

addIDorgURI(qual, uri)

Add a URI directly into a qualifier collection:

- qual a Biomodels biological qualifier e.g. "is" "isEncodedBy"
- uri the complete identifiers.org uri e.g. http://identifiers.org/chebi/CHEBI:58088

addMIRIAMannotation(qual, entity, mid)

Add a qualified MIRIAM annotation or entity:

- qual a Biomodels biological qualifier e.g. "is" "isEncodedBy"
- entity a MIRIAM resource entity e.g. "ChEBI"
- mid the entity id e.g. CHEBI:17158

checkEntity(entity)

Check an entity entry, this is a MIRIAM resource name: "chEBI". The test is case insensitive and will correct the case of wrongly capitalised entities automatically. If the entity is not recognised then a list of possible candidates based on the first letters of the input is displayed.

• entity a MIRIAM resource entity e.g. "ChEBI"

checkEntityPattern(entity)

For an entity key compile the pattern to a regex, if necessary.

• entity a MIRIAM resource entity

checkId(entity, mid)

Check that a entity id e.g. CHEBI:17158

• mid the entity id e.g. CHEBI:17158

deleteMIRIAMannotation(qual, entity, mid)

Deletes a qualified MIRIAM annotation or entity:

- qual a Biomodels biological qualifier e.g. "is" "isEncodedBy"
- entity a MIRIAM resource entity e.g. "ChEBI"
- mid the entity id e.g. CHEBI:17158

getAllMIRIAMUris()

Return a dictionary of qualifiers that contain ID.org URL'S

getAndViewUrisForQualifier(qual)

Retrieve all url's associated with qualifier and attempt to open them all in a new browser tab

• qual the qualifier e.g. "is" or "isEncoded"

getMIRIAMUrisForQualifier(qual)

Return all list of urls associated with qualifier:

• qual the qualifier e.g. "is" or "isEncoded"

viewURL(url)

This will try to open the URL in a new tab of the default webbrowser

• url the url

class StructMatrix(array, ridx, cidx, row=None, col=None)

This class is specifically designed to store structural matrix information give it an array and row/col index permutations it can generate its own row/col labels given the label src.

getColsByIdx(*args)

Return the columns referenced by index (1,3,5)

getColsByName(*args)

Return the columns referenced by label ('s','x','d')

getIndexes(axis='all')

Return the matrix indexes ([rows],[cols]) where axis='row'/'col'/'all'

getLabels(axis='all')

Return the matrix labels ([rows],[cols]) where axis='row'/'col'/'all'

getRowsByIdx(*args)

Return the rows referenced by index (1,3,5)

getRowsByName(*args)

Return the rows referenced by label ('s','x','d')

setCol(src)

Assuming that the col index array is a permutation (full/subset) of a source label array by supplying that src to setCol maps the row labels to cidx and creates self.col (col label list)

setRow(src)

Assuming that the row index array is a permutation (full/subset) of a source label array by supplying that source to setRow it maps the row labels to ridx and creates self.row (row label list)

class StructMatrixLP(array, ridx, cidx, row=None, col=None, rhs=None, operators=None)

Adds some stuff to StructMatrix that makes it LP friendly

getCopy(attr_str, deep=False)

Return a copy of the attribute with name attr_str. Uses the copy module copy.copy or copy.deepcopy

- attr_str a string of the attribute name: 'row', 'col'
- · deep [default=False] try to do a deepcopy. Use with caution see copy module docstring for details

checkChemFormula(cf, quiet=False)

Checks whether a string conforms to a Chemical Formula C3Br5 etc, returns True/False. Please see the SBML Level 3 specification and http://wikipedia.org/wiki/Hill system for more information.

- cf a string that contains a formula to check
- quiet [default=False] do not print error messages

checkId(s)

Checks the validity of the string to see if it conforms to a C variable. Returns true/false

• s a string

createAssociationDictFromNode(node, out, model, useweakref=True, cntr=0)

Converts a GPR string '((g1 and g2) or g3)' to a dictionary via a Python AST. In future I will get rid of all the string elements and work only with AST's.

- node a Python AST node (e.g. body)
- out a gpr dictionary
- model a CBMPy model instance
- useweakref [default=True] use a weakref as the gene object or alternatively the label

extractGeneIdsFromString(g, return_clean_gpr=False)

Extract and return a list of gene names from a gene association string formulation

- g a COBRA style gene association string
- return_clean_gpr [default=False] in addition to the list returns the "cleaned" GPR string

fixId(s, replace=None)

Checks a string (Sid) to see if it is a valid C style variable. first letter must be an underscore or letter, the rest should be alphanumeric or underscore.

- s the string to test
- replace [None] default is to leave out offensive character, otherwise replace with this one

${\bf func_getAssociationStrFromGprDict}(\mathit{gprd},\mathit{out},\mathit{parent=''})$

Get a old school GPR association string from a CBMPy gprDict, e.g. obtained from gpr.getTree()

- gprd the gprDictionary
- out the output string

• parent [default="] the string representing the current nodes parent relationship, used for recursion

getGPRasDictFromString(node, out)

Converts a GPR string '((g1 and g2) or g3)' to a gprDict which is returned

- *node* a Python AST note (e.g. *ast.parse(gprstring).body[0])*
- out a new dictionary that will be be created in place

parseGeneAssociation(gs)

Parse a COBRA style gene association into a nested list.

• gs a string containing a gene association

Disambiguate the chemical formula from either the Notes or the overloaded name

- s a species object
- getFromName [default=False] whether to try strip the chemical formula from the name (old COBRA style)
- overwriteChemFormula [default=False]
- overwriteCharge [default=False]

3.2 CBMPy: CBConfig module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBConfig.py 711 2020-04-27 14:22:34Z bgoli \$)

current_version()

Return the current CBMPy version as a string

current_version_tuple()

Return the current CBMPy version as a tuple (x, y, z)

3.3 CBMPy: CBCPLEX module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBCPLEX.py 710 2020-04-27 14:22:34Z bgoli \$)

cplx_FluxVariabilityAnalysis(fba, selected_reactions=None, pre_opt=True, tol=None, objF2constr=True, rhs_sense='lower', optPercentage=100.0, work_dir=None, quiet=True, debug=False, oldlpgen=False, markupmodel=True, default_on_fail=False, roundoff_span=10, method='o')

Perform a flux variability analysis on an fba model:

- fba an FBA model object
- selected reactions [default=None] means use all reactions otherwise use the reactions listed here
- pre_opt [default=True] attempt to presolve the FBA and report its results in the ouput, if this is disabled and objF2constr is True then the rid/value of the current active objective is used
- tol [default=None] do not floor/ceiling the objective function constraint, otherwise round of to tol
- rhs_sense [default='lower'] means objC >= objVal the inequality to use for the objective constraint can also be upper or equal
- *optPercentage* [default=100.0] means the percentage optimal value to use for the RHS of the objective constraint: optimal_value*(optPercentage/100.0)
- work_dir [default=None] the FVA working directory for temporary files default = cwd+fva
- debug [default=False] if True write out all the intermediate FVA LP's into work_dir
- quiet [default=False] if enabled, supress CPLEX output
- *objF2constr* [default=True] add the model objective function as a constraint using rhs_sense etc. If this is True with pre_opt=False then the id/value of the active objective is used to form the constraint
- markupmodel [default=True] add the values returned by the fva to the reaction.fva_min and reaction.fva_max
- default_on_fail [default=False] if pre_opt is enabled replace a failed minimum/maximum with the solution value
- roundoff_span [default=10] number of digits is round off (not individual min/max values)
- *method* [default='o'] choose the CPLEX method to use for solution, default is automatic. See CPLEX reference manual for details
 - 'o': auto
 - 'p': primal

```
- 'd': dual
```

- 'b': barrier (no crossover)

- 'h': barrier

- 's': sifting

- 'c': concurrent

Returns an array with columns: Reaction, Reduced Costs, Variability Min, Variability Max, abs(Max-Min), MinStatus, MaxStatus and a list containing the row names.

```
cplx_MinimizeNumActiveFluxes(fba, selected_reactions=None, pre_opt=True, tol=None, objF2constr=True, rhs_sense='lower', optPercentage=100.0, work_dir=None, quiet=False, debug=False, objective_coefficients=None, return_lp_obj=False, populate=None, oldlpgen=False)
```

Minimize the sum of active fluxes, updates the model with the values of the solution and returns the value of the MILP objective function (not the model objective function which remains unchanged). If population mode is activated output is as described below:

```
Min: sum(Bi)
    Bi = 0 -> Ci Ji = 0

Such that:
    NJi = 0 Jbio = opt

where:
    Binary Bi
```

Arguments:

- fba an FBA model object
- selected reactions [default=None] means use all reactions otherwise use the reactions listed here
- *pre_opt* [default=True] attempt to presolve the FBA and report its results in the ouput, if this is diabled and *objF2constr* is True then the vid/value of the current active objective is used
- tol [default=None] do not floor/ceiling the objective function constraint, otherwise round of to tol
- rhs_sense [default='lower'] means objC >= objVal the inequality to use for the objective constraint can also be upper or equal Note this does not necessarily mean the upper or lower bound, although practically it will. If in doubt use equal
- optPercentage [default=100.0] means the percentage optimal value to use for the RHS of the objective constraint: optimal_value * (optPercentage/100.0)
- work_dir [default=None] the MSAF working directory for temporary files default = cwd+fva
- debug [default=False] if True write out all the intermediate MSAF LP's into work_dir
- quiet [default=False] if enabled supress CPLEX output
- *objF2constr* [default=True] add the model objective function as a constraint using rhs_sense etc. If this is True with pre_opt=False then the id/value of the active objective is used to form the constraint
- *objective_coefficients* [default=None] a dictionary of (reaction_id: float) pairs that provide the are introduced as objective coefficients to the absolute flux value. Note that the default value of the coefficient (non-specified) is +1.
- return_lp_obj [default=False] off by default when enabled it returns the CPLEX LP object
- populate [default=None] enable search algorithm to find multiple (sub)optimal solutions. Set with a tuple of (RELGAP=0.0, POPULATE_LIMIT=20, TIME_LIMIT=300) suggested values only. RELGAP

[default=0.0] relative gap to optimal solution - *POPULATE_LIMIT* [default=20] terminate when so many solutions have been found - *TIME_LIMIT* [default=300] terminate search after so many seconds important with higher values of *POPULATION LIMIT*

• with_reduced_costs [default='uncsaled'] can be 'scaled', 'unscaled' or anything else which is None

With outputs:

- mincnt the objective function value OR
- mincnt, cpx the objective function and cplex model OR
- populate_data, mincnt a population data set OR
- populate_data, mincnt, cpx both the cps object and population data set

depending on selected flags.

```
cplx_MinimizeSumOfAbsFluxes(fba, selected_reactions=None, pre_opt=True, tol=None, objF2constr=True, rhs_sense='lower', optPercentage=100.0, work_dir=None, quiet=False, debug=False, objective_coefficients=None, return_lp_obj=False, oldlpgen=False, with reduced costs=None, method='o')
```

Minimize the sum of absolute fluxes $sum(abs(J1) + abs(J2) + abs(J3) \dots abs(Jn))$ by adding two constraints per flux and a variable representing the absolute value:

```
Min: Ci abs_Ji
    Ji - abs_Ji <= 0 Ji + abs_Ji >= 0

Such that:
    NJi = 0 Jopt = opt
```

returns the value of the flux minimization objective function (not the model objective function which remains unchanged from)

Arguments:

- fba an FBA model object
- selected reactions [default=None] means use all reactions otherwise use the reactions listed here
- *pre_opt* [default=True] attempt to presolve the FBA and report its results in the ouput, if this is disabled and *objF2constr* is True then the vid/value of the current active objective is used
- tol [default=None] do not floor/ceiling the objective function constraint, otherwise round of to tol
- rhs_sense [default='lower'] means objC >= objVal the inequality to use for the objective constraint can also be upper or equal
- *optPercentage* [default=100.0] means the percentage optimal value to use for the RHS of the objective constraint: optimal_value*(optPercentage/100.0)
- work dir [default=None] the MSAF working directory for temporary files default = cwd+fva
- debug [default=False] if True write out all the intermediate MSAF LP's into work_dir
- quiet [default=False] if enabled supress CPLEX output
- *objF2constr* [default=True] add the model objective function as a constraint using rhs_sense etc. If this is True with pre_opt=False then the id/value of the active objective is used to form the constraint
- objective_coefficients [default=None] a dictionary of (reaction_id: float) pairs that provide the are introduced as objective coefficients to the absolute flux value. Note that the default value of the coefficient (non-specified) is +1.
- return_lp_obj [default=False] off by default when enabled it returns the CPLEX LP object

- with_reduced_costs [default=None] if not None should be 'scaled' or 'unscaled'
- method [default='o'] choose the CPLEX method to use for solution, default is automatic. See CPLEX
 reference manual for details
 - 'o': auto
 - 'p': primal
 - 'd': dual
 - 'b': barrier (no crossover)
 - 'h': barrier
 - 's': sifting
 - 'c': concurrent

With outputs:

• *fba* an update instance of a CBModel. Note that the FBA model objective function value is the original value set as a constraint

Perform a flux variability analysis on a multistate LP

- lp a multistate LP
- selected reactions [default=None] means use all reactions otherwise use the reactions listed here
- pre_opt [default=True] attempt to presolve the FBA and report its results in the ouput
- tol [default=1e-10] do floor/ceiling the objective function constraint, otherwise floor/ceil to tol
- *rhs_sense* [default='lower'] means objC >= objVal the inequality to use for the objective constraint can also be *upper* or *equal*
- *optPercentage* [default=100.0] means the percentage optimal value to use for the RHS of the objective constraint: optimal_value*(optPercentage/100.0)
- work_dir [default=None] the FVA working directory for temporary files default = cwd+fva
- debug [default=False] if True write out all the intermediate FVA LP's into work_dir
- bypass [default=False] bypass everything and only run the min/max on lp

and returns an array with columns:

```
Reaction, Reduced Costs, Variability Min, Variability Max, abs(Max-Min), MinStatus, 
→MaxStatus
```

and a list containing the row names.

cplx_SolveMILP(c, auto mipgap=False)

Solve and MILP

• *auto_mipgap* auto decrease mipgap until mipgap == absmipgap

cplx_WriteFVAtoCSV(pid, fva, names, Dir=None, fbaObj=None)

Takes the resuls of a Flux Variability Analysis method and writes it to a nice csv file. Note this method has been refactored to *CBWrite.WriteFVAtoCSV()*.

• pid filename_base for the CSV output

- fva Flux Variability Analysis() OUTPUT_ARRAY
- names Flux Variability Analysis() OUTPUT NAMES
- Dir [default=None] if set the output directory for the csv files
- fbaObj [default=None] if supplied adds extra model information into the output tables

 $\label{lem:cplx_analyzeModel} \begin{subarray}{ll} $\tt cplx_analyzeModel($f$, $lpFname=None, $return_lp_obj=False, with_reduced_costs='unscaled', \\ with_sensitivity=False, $del_intermediate=False, build_n=True, $quiet=False, oldlpgen=False, \\ method='o') \end{subarray}$

Optimize a model and add the result of the optimization to the model object (e.g. *reaction.value*, *objectiveFunction.value*). The stoichiometric matrix is automatically generated. This is a common function available in all solver interfaces. By default returns the objective function value

- f an instantiated PySCeSCBM model object
- lpFname [default=None] the name of the intermediate LP file. If not specified no LP file is produced
- return_lp_obj [default=False] off by default when enabled it returns the CPLEX LP object
- with_reduced_costs [default='unscaled'] calculate and add reduced cost information to mode this can be: 'unscaled' or 'scaled' or anything else which is interpreted as 'None'. Scaled means s_rcost = (r.reduced_cost*rval)/obj_value
- with_sensitivity [default=False] add solution sensitivity information (not yet implemented)
- del_intermediate [default=False] redundant except if output file is produced and deleted (not useful)
- build_n [default=True] generate stoichiometry from the reaction network (reactions/reagents/species)
- quiet [default=False] suppress cplex output
- method [default='o'] choose the CPLEX method to use for solution, default is automatic. See CPLEX
 reference manual for details
 - 'o': auto
 - 'p': primal
 - 'd': dual
 - 'b': barrier (no crossover)
 - 'h': barrier
 - 's': sifting
 - 'c': concurrent

cplx_constructLPfromFBA(fba, fname=None)

Create a CPLEX LP in memory. - fba an FBA object - fname optional filename if defined writes out the constructed lp

cplx_fixConSense(operator)

Fixes the sense of inequality operators, returns corrected sense symbol

• operator the operator to check

This is a utility function that does a presolve for FVA, MSAF etc. Generates properly formatted empty objects if pre_opt == False

pre_opt a boolean

- fba a CBModel object
- objF2constr add objective function as constraint
- quiet [default=False] supress cplex output
- with_reduced_costs [default='unscaled'] can be 'scaled' or 'unscaled'
- method [default='o'] choose the CPLEX method to use for solution, default is automatic. See CPLEX
 reference manual for details
 - 'o': auto
 - 'p': primal
 - 'd': dual
 - 'b': barrier (no crossover)
 - 'h': barrier
 - 's': sifting
 - 'c': concurrent

Returns: pre_sol, pre_oid, pre_oval, OPTIMAL_PRESOLUTION, REDUCED_COSTS

cplx_func_SetObjectiveFunctionAsConstraint(cpx, rhs_sense, oval, tol, optPercentage)

Take the objective function and "optimum" value and add it as a constraint

- cpx a cplex object
- oval the objective value
- tol [default=None] do not floor/ceiling the objective function constraint, otherwise round of to tol
- *rhs_sense* [default='lower'] means objC >= objVal the inequality to use for the objective constraint can also be *upper* or *equal*
- *optPercentage* [default=100.0] means the percentage optimal value to use for the RHS of the objective constraint: optimal_value*(optPercentage/100.0)

cplx_getCPLEXModelFromLP(lptFile, Dir=None)

Load a LPT (CPLEX format) file and return a CPLX LP model

- *lptfile* an CPLEX LP format file
- Dir an optional directory

cplx_getDualValues(c)

Get the get the dual values of the solution

• c a CPLEX LP

Output is a dictionary of {name : value} pairs

${\tt cplx_getModelFromLP}(\mathit{lptFile}, \mathit{Dir} = \mathit{None})$

Load a LPT (CPLEX format) file and return a CPLX LP model

- lptfile an CPLEX LP format file
- Dir an optional directory

cplx_getModelFromObj(fba)

Return a CPLEX object from a FBA model object (via LP file)

cplx_getOptimalSolution(c)

From a CPLX model extract a tuple of solution, ObjFuncName and ObjFuncVal

cplx_getOptimalSolution2(c, names)

From a CPLX model extract a tuple of solution, ObjFuncName and ObjFuncVal

cplx_getReducedCosts(c, scaled=False)

Extract ReducedCosts from LP and return as a dictionary 'Rid': reduced cost

- c a cplex LP object
- scaled scale the reduced cost by the optimal flux value

cplx_getSensitivities(c)

Get the sensitivities of each constraint on the objective function with inpt

c a CPLEX LP

Output is a tuple of bound and objective sensitivities where the objective sensitivity is described in the CPLEX reference manual as:

```
... the objective sensitivity shows each variable, its reduced cost and the range_
    over
which its objective function coefficient can vary without forcing a change
in the optimal basis. The current value of each objective coefficient is
also displayed for reference.

- *objective coefficient sensitivity* {flux : (reduced_cost, lower_obj_sensitivity, _
    oceff_value, upper_obj_sensitivity)}
- *rhs sensitivity* {constraint : (low, value, high)}
- *bound sensitivity ranges* {flux : (lb_low, lb_high, ub_low, ub_high)}
```

cplx_getShadowPrices(c)

Returns a dictionary of shadow prices containing 'N row id': (lb, rhs, ub)

• c a cplex LP object

cplx_getSolutionStatus(c)

Returns one of:

- *LPS_OPT*: solution is optimal;
- LPS FEAS: solution is feasible;
- LPS_INFEAS: solution is infeasible;
- *LPS_NOFEAS*: problem has no feasible solution;
- LPS_UNBND: problem has unbounded solution;
- LPS_UNDEF: solution is undefined.
- LPS NONE: no solution

cplx_runInputScan(*fba*, *exDict*, *wDir*, *input_lb=-10.0*, *input_ub=0.0*, *writeHformat=False*, rationalLPout=False)

scans all inputs

cplx_setFBAsolutionToModel(fba, lp, with reduced costs='unscaled')

Sets the FBA solution from a CPLEX solution to an FBA object

• fba and fba object

- *lp* a CPLEX LP object
- with_reduced_costs [default='unscaled'] calculate and add reduced cost information to mode this can be: 'unscaled' or 'scaled' or anything else which is interpreted as None. Scaled is: s_rcost = (r.reduced_cost*rval)/obj_value

cplx_setMIPGapTolerance(c, tol)

Sets the the relative MIP gap tolerance

cplx_setObjective(*c*, *pid*, *expr=None*, *sense='maximize'*, *reset=True*)

Set a new objective function note that there is a major memory leak in *c.variables.get_names()* which is used when reset=True. If this is a problem use cplx_setObjective2 which takes *names* as an input:

- c a CPLEX LP object
- pid the r_id of the flux to be optimized
- expr a list of (coefficient, flux) pairs
- sense 'maximize'/'minimize'
- reset [default=True] reset all objective function coefficients to zero

cplx_setObjective2(c, pid, names, expr=None, sense='maximize', reset=True)

Set a new objective function. This is a workaround function to avoid the e is a major memory leak in *c.variables.get_names()* which is used in cplx_setObjective() when reset=True.

cplx_setOutputStreams(lp, mode='default')

Sets the noise level of the solver, mode can be one of:

- None silent i.e. no output
- 'file' set solver to silent and output logs to CPLX_RESULT_STREAM_FILE cplex_output.log
- 'iostream' set solver to silent and output logs to CPLX_RESULT_STREAM_IO csio
- 'default' or anything else noisy with full output closes STREAM IO and STREAM FILE (default)

cplx_setSolutionStatusToModel(m, lp)

Sets the lp solutions status to the CBMPy model

cplx_singleGeneScan(fba, r_off_low=0.0, r_off_upp=0.0, optrnd=8, altout=False)

Perform a single gene deletion scan

- fba a model object
- r off low the lower bound of a deactivated reaction
- r_off_upp the upper bound of a deactivated reaction
- optrnd [default=8] round off the optimal value
- *altout* [default=False] by default return a list of gene:opt pairs, alternatively (True) return an extended result set including gene groups, optima and effect map

cplx_singleReactionDeletionScan(fba, r_off_low=0.0, r_off_upp=0.0, optrnd=8)

Perform a single reaction deletion scan

- fba a model object
- r_off_low the lower bound of a deactivated reaction
- r_off_upp the upper bound of a deactivated reaction
- optrnd [default=8] round off the optimal value

cplx_writeLPsolution(fba_sol, objf_name, fname, Dir=None, separator=',')

This function writes the optimal solution, produced wth cplx_getOptimalSolution to file

- fba_sol a dictionary of Flux : value pairs
- objf name the objective function flux id
- fname the output filename
- Dir [default=None] use directory if not None
- separator [default=','] the column separator

cplx_writeLPtoLPTfile(c, filename, title=None, Dir=None)

Write out a CPLEX model as an LP format file

getReducedCosts(fba)

Get a dictionary of reduced costs for each reaction/flux

setReducedCosts(fba, reduced_costs)

For each reaction/flux, sets the attribute "reduced_cost" from a dictionary of reduced costs

- fba an fba object
- reduced_costs a dictionary of {reaction : value} pairs

3.4 CBMPy: CBDataStruct module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBDataStruct.py 710 2020-04-27 14:22:34Z bgoli \$)

3.5 CBMPy: CBGUI module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBGUI.py 710 2020-04-27 14:22:34Z bgoli \$)

loadCBGUI (*mod*, *version*=2)

Load an FBA model instance into the quick editor to view or change basic model properties

• mod a PySCeS CBMPy model instance

3.6 CBMPy: CBModel module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBModel.py 706 2020-03-23 21:31:49Z bgoli \$)

class Compartment(pid, name=None, size=1, dimensions=3, volume=None)

A compartment

containsReactions()

Lists the species contained in this compartment

containsSpecies()

Lists the species contained in this compartment

getDimensions()

Get the compartment dimensions

getSize()

Get the compartment size

setDimensions(dimensions)

Get the compartment dimensions

• dimensions set the new compartment dimensions

setId(fid)

Sets the object Id

• fid a valid c variable style id string

Reimplements @FBase.setId()

setSize(size)

Set the compartment size

• size the new compartment size

class ConstraintComponent(pid, coefficient, variable, ctype='linear')

A weighted flux that appears in an user defined constraint

class Fbase

Base class for CB Model objects

addMIRIAMannotation(qual, entity, mid)

Add a qualified MIRIAM annotation or entity:

- qual a Biomodels biological qualifier e.g. "is" "isEncodedBy"
- entity a MIRIAM resource entity e.g. "ChEBI"
- mid the entity id e.g. CHEBI:17158 or fully qualifies url (if only_qual_uri)

addMIRIAMuri(qual, uri)

Add a qualified MIRIAM annotation or entity:

- qual a Biomodels biological qualifier e.g. "is" "isEncodedBy"
- *uri* the fully qualified entity id e.g. http://identifiers.org/chebi/CHEBI:12345 (no validity checking is done)

clone()

Return a clone of this object. Cloning performs a deepcop on the object which will also clone any objects that exist as attributes of this object, in other words an independent copy of the original. If this is not the desired behaviour override this method when subclassing or implement your own.

deleteAnnotation(key)

Unsets (deltes) an objects annotation with key

• *key* the annotation key

deleteMIRIAMannotation(qual, entity, mid)

Deletes a qualified MIRIAM annotation or entity:

- qual a Biomodels biological qualifier e.g. "is" "isEncodedBy"
- entity a MIRIAM resource entity e.g. "ChEBI"
- mid the entity id e.g. CHEBI:17158

getAnnotation(key)

Return the object annotation associated with:

• key the annotation key

getAnnotations()

Return the object annotation dictionary

getCompartmentId()

Return the compartment id where this element is located

getId()

Return the object ID.

getMIRIAMannotations()

Returns a dictionary of all MIRIAM annotations associated with this object or None of there are none defined.

getMetaId()

Return the object metaId.

getModel()

Get the parent model object linked to in objecf, can return model or None for unlinked object

Overwritten by Model

getName()

Return the object name.

getNotes()

Return the object's notes

getPid()

Return the object ID.

getSBOterm()

Return the SBO term for this object.

hasAnnotation(key)

Returns a boolean representing the presence/absence of the key in the objext annotation

• key the annotation key

serialize(protocol=0)

Serialize object, returns a string by default

• protocol [default=0] serialize to a string or binary if required,

see pickle module documentation for details

Reimplemented in Model

serializeToDisk(filename, protocol=2)

Serialize to disk using pickle protocol:

- filename the name of the output file
- protocol [default=2] serialize to a string or binary if required,

see pickle module documentation for details

Reimplemented in Model

setAnnotation(key, value, ext=None)

Set an objects annotation as a key: value pair.

- key the annotation key
- value the annotation value
- ext a dictionary of extended properties in FBCv3 id, name, uri

setCompartmentId(compartment)

Set the compartment id where this element is located

setId(fid)

Sets the object Id

• fid a valid c variable style id string

Reimplemented by @Reaction, @Species, @Compartment, @Gene

setMetaId(mid=None)

Sets the object Id

• mid [default=None] a valid c variable style metaid string, if None it will be set as meta+id

setName(name)

Set the object name:

• name the name string

setNotes(notes)

Sets the object's notes:

• notes the note string, should preferably be (X)HTML for SBML

setPid(fid)

Sets the object Id

• fid a valid c variable style id string

setSBOterm(sbo)

Set the SBO term for this object.

• sbo the SBOterm with format: SBO:nnnnnnn"

class FluxBound(pid, reaction, operation, value)

A reaction fluxbound

getType()

Returns the type of FluxBound: 'lower', 'upper', 'equality' or None

getValue()

Returns the current value of the attribute (input/solution)

setReactionId(react)

Sets the reaction attribute of the FluxBound

setValue(value)

Sets the attribute "value"

class FluxBoundBase(pid, operator, value, parent=None)

A refactored and streamlined FluxBound base class that can be a generic bound, superclass to FluxBoundUpper and FluxBoundLower

getType()

Returns the type of FluxBound: 'lower', 'upper'

getValue()

Returns the current value of the attribute (input/solution)

setValue(value)

Sets the value attribute:

• value a float

property value

Returns the current value of the attribute (input/solution)

class FluxBoundLower(reaction, value=-inf)

class FluxBoundUpper(reaction, value=inf)

class FluxObjective(pid, reaction, coefficient=1, ctype='linear')

A weighted flux that appears in an objective function

NOTE: reaction is a string containing a reaction id

class FluxObjectiveQuadratic(pid, reaction, reaction2, coefficient=1, ctype='quadratic')

A weighted quadratic flux that appears in an objective function, this fluxobjective contains two reaction terms to define "quadratic" fluxobjectives of the type <coefficient>*<variable1>*<variable2> For example 2*R1*R2

NOTE: reaction is a string containing a reaction id

class Gene(pid, label=None, active=True)

Contains all the information about a gene (or gene+protein construct depending on your philosophy)

getLabel()

Returns the gene label

isActive()

Returns whether the gene is active or not

resetActivity()

Reset the gene to its default activity state

setActive()

Set the gene to be active

setId(fid)

Sets the object Id

• fid a valid c variable style id string

Reimplements Fbase method

setInactive()

Set the gene to be inactive

setLabel(label)

Sets the gene label, checks that the new label is unique

class GeneProteinAssociation(pid, protein, use_compiled=False)

This class associates genes to proteins.

addAssociation(assoc)

Add a gene/protein association expression

addGeneref(geneid)

Add a gene reference to the list of gene references

• geneid a valid model Gene id

createAssociationAndGeneRefsFromString(assoc, altlabels=None)

Evaluate the gene/protein association and add the genes necessary to evaluate it Note that this GPR should be added to a model with cmod.addGPRAssociation() before calling this method

- assoc the COBRA style gene protein association
- altlabels [default=None] a dictionary containing a label<->id mapping

createAssociationAndGeneRefsFromTree(gprtree, altlabels=None)

Evaluate the GPR tree and add the genes necessary to evaluate it Note that this GPR should be added to a model with cmod.addGPRAssociation() before calling this method

- gprtree the CBMPy GPR tree data structure
- altlabels [default=None] a dictionary containing a label<->id mapping

deleteGeneFromAssociation(gid)

Deletes a gene id from the gene association. WARNING this process is irreversible!!

• gid a valid gene identifier (not label)

deleteGeneref(gid)

Deletes a gene reference

• geneid a valid model Gene id

evalAssociation()

Returns an integer value representing the logical associations or None.

getActiveGenes()

Return a list of active gene objects

getAssociationStr(use_labels=False)

Return the gene association string, alternatively return string with labels

• use_lablels [default=False] return the gene association string with labels rather than geneId's (FBCv2 issue)

getGene(gid)

Return a gene object with id

getGeneIds()

Return a list of gene id's

getGeneLabels()

Return a list of gene labels associated with this GPRass

getGenes()

Return a list of gene objects associated with this GPRass

getProtein()

Return the protein associated with this set of genes

getTree()

Return the dictionary/tree representation of the GPR

getTreeCopy()

Return a copy of the dictionary/tree representation of the GPR

isProteinActive()

This returns a boolean which indicates the result of evaluating the gene association. If the result is positive then the protein is expressed and *True* is returned, otherwise if the expression evaluates to a value of 0 then the protein is not expressed and *False* is returned.

setAllGenesActive()

Activate all genes in association

setAllGenesInactive()

Deactivates all genes in association

setGeneActive(gid)

Set a gene to be inactive

setGeneInactive(gid)

Set a gene to be inactive

setProtein(protein)

Sets the protein associated with this set of genes

setTree(tree)

Add a GPR dictionary/tree representation to the GPR.

• tree a dictionary representation of a GPR.

class Group(pid)

Container for SBML groups

addMember(obj)

Add member CBMPy object(s) to the group

• *obj* either a single, tuple or list of CBMPy objects

addSharedMIRIAMannotation(qual, entity, mid)

Add a qualified MIRIAM annotation or entity to the list of members (all) rather than the group itself:

- qual a Biomodels biological qualifier e.g. "is" "isEncodedBy"
- entity a MIRIAM resource entity e.g. "ChEBI"
- mid the entity id e.g. CHEBI:17158 or fully qualifies url (if only_qual_uri)

assignAllSharedPropertiesToMembers(overwrite=False)

Assigns all group shared properties (notes, annotations, MIRIAM annotations, SBO) to the group members.

• overwrite [default=False] overwrite the target notes if they are defined

assignSharedAnnotationToMembers()

This function merges or updates the group member objects annotations with the group shared annotation.

assignSharedMIRIAMannotationToMembers()

This function merges or updates the group member objects MIRIAM annotations with the group shared MIRIAM annotation.

assignSharedNotesToMembers(overwrite=False)

Assigns the group shared notes to the group members.

• overwrite [default=False] overwrite the target notes if they are defined

assignSharedSBOtermsToMembers(overwrite=False)

Assigns the group shared member SBO term to the group members.

• overwrite [default=False] overwrite the target SBO term if it is defined

clone()

Return a clone of this object. Note the for Groups this is a shallow copy, in that the reference objects themselves are not cloned only the group (and attributes)

deleteMember(oid)

Deletes a group member with group id.

• oid group member id

getKind()

Return the group kind

getMember(mid)

Returns the group member object that corresponds to mid

• mid the id of the requested object

getMemberIDs(as_set=False)

Return the ids of the member objects.

• as_set return id's as a set rather than a list

getMembers(as_set=False)

Return the member objects of the group.

• as_set return objects as a set rather than a list

getSharedAnnotations()

Return a dictionary of the shared member annotations (rather than the group attribute).

getSharedMIRIAMannotations()

Return a dictionary of the shared member MIRIAM annotations (rather than the group attribute).

getSharedNotes()

Return the shared member notes (rather than the group attribute).

getSharedSBOterm()

Return the shared member SBO term (rather than the group attribute).

hasMember(mid)

Returns a boolean indicating whether a member is in the group.

• *mid* the id to check

serialize(protocol=0)

Serialize object, returns a string by default

• protocol [default=0] serialize to a string or binary if required,

see pickle module documentation for details

Reimplemented in Model

serializeToDisk(filename, protocol=2)

Serialize to disk using pickle protocol:

• filename the name of the output file

• protocol [default=2] serialize to a string or binary if required,

see pickle module documentation for details

Reimplemented in Model

setKind(kind)

Sets the kind or type of the group, this must be one of: 'collection', 'partonomy', 'classification'.

• kind the kind

setSharedAnnotation(key, value)

Sets the list of members (all) annotation as a key: value pair.

- key the annotation key
- value the annotation value

setSharedNotes(notes)

Sets the group of objects notes attribute (all):

• notes the note string, should preferably be (X)HTML for SBML

setSharedSBOterm(sbo)

Set the SBO term for the members of the group (all).

• sbo the SBOterm with format: "SBO:<7 digit integer>"

class GroupMemberAttributes

Contains the shared attributes of the group members (equivalent to SBML annotation on ListOfMembers)

class Model(pid)

Container for constraint based model, adds methods for manipulating:

- · objectives
- · constraints
- · reactions
- · species
- · compartments
- groups
- parameters
- N a structmatrix object

addCompartment(comp)

Add an instantiated Compartment object to the CBM model

• comp an instance of the Compartment class

addFluxBound(fluxbound, fbexists=None)

Add an instantiated FluxBound object to the FBA model

- fluxbound an instance of the FluxBound class
- *fbexists* [default=None] this is a list of strings which contains fluxbounds that have been added to the model, see sample code below.

The format of the string is 'reactionid boundtype'

"python fbexists = [] for fluxbound in list of fluxbounds:

 $model. addFluxBound (fluxbound, fbexists=fbexists) fbexists. append (``\{\}_{\{\}}". format (fluxbound.getReactionId(), fluxbound.getType()))$

...

Using the fbexists list drastically reduces the time it takes to add fluxbounds but circumvents any sort of existence checking and should *only* be used as shown above when constructing a model from scratch. Alternativel, you need to prepopulate fbexists with existing fluxbound component information:

```
`python fbexists = ["{}_{}".format(fluxbound.getReactionId(), fluxbound.getType()) for fluxbound in model.flux_bounds] `
```

addGPRAssociation(gpr, update_idx=True)

Add a GeneProteinAssociation instance to the model

• gpr an instantiated GeneProteinAssociation object

addGene(gene)

Add an instantiated Gene object to the FBA model

• gene an instance of the G class

addGroup(obj)

Add an instantiated group object to the model

• obj the Group instance

addMIRIAMannotation(qual, entity, mid)

Add a qualified MIRIAM annotation or entity:

- qual a Biomodels biological qualifier e.g. "is" "isEncodedBy"
- entity a MIRIAM resource entity e.g. "ChEBI"
- *mid* the entity id e.g. CHEBI:17158

addModelCreator(firstname, lastname, organisation=None, email=None)

Add a model creator to the list of model creators, only the first and fmaily names are mandatory:

- firstname
- lastname
- organisation [default=None]
- email [default=None]

addObjective(obj, active=False)

Add an instantiated Objective object to the FBA model

- obj an instance of the Objective class
- active [default=False] flag this objective as the active objective (fba.activeObjIdx)

addParameter(par)

Add an instantiated Parameter object to the model

• par an instance of the Parameter class

addReaction(reaction, create_default_bounds=False, silent=False)

Adds a reaction object to the model

• reaction an instance of the Reaction class

- create_default_bounds create default reaction bounds, irreversible 0 <= J <= INF, reversable -INF <= J <= INF
- silent [default=False] if enabled this disables the printing of information messages

addSpecies(species)

Add an instantiated Species object to the FBA model

• species an instance of the Species class

addUserConstraint(pid, fluxes=None, operator='>=', rhs=0.0)

Add a user defined constraint to FBA model, this is additional to the automatically determined Stoichiometric constraints.

- pid user constraint name/id, use None for auto-assign
- fluxes a list of (coefficient, reaction id) pairs where coefficient is a float
- *operator* is one of '=', '>=' or '<=' (< and > will be interpreted as >= or <=)
- rhs a float

addUserDefinedConstraint(udc)

Add a User Defined Constraint object to the FBA model

• obj an instance of the UserDefinedConstraint class

buildStoichMatrix(matrix_type='numpy', only_return=False)

Build the stoichiometric matrix N and additional constraint matrix CN (if required)

- matrix_type [default='numpy'] the type of matrix to use to generate constraints
 - numpy a NumPy matrix default
 - sympy a SymPy symbolic matrix, if available note the denominator limit can be set in CBModel.
 __CBCONFIG__['SYMPY_DENOM_LIMIT'] = 10**12
 - scipy_csr create using NumPy but store as SciPy csr_sparse
- *only_return* [default=False] **IMPORTANT** only returns the stoichiometric matrix and constraint matrix (if required), does not update the model

changeAllFluxBoundsWithValue(old, new)

Replaces all flux bounds with value "old" with a new value "new":

- · old value
- · new value

clone()

Return a clone of this object.

convertUserConstraintsToUserDefinedConstraints()

If a model is loaded with the old CBMPy specific constraint data structures json files and dictionaries, this function will upmark it to the new FBCv3 data structures

copyUserDefinedConstraintsToUserConstraints()

This is a workaround until I complete full UserDefinedConstraints support

createCompartment(cid, name=None, size=1, dimensions=3, volume=None)

Create a new compartment and add it to the model if the id does not exist

- cid compartment id
- name [None] compartment name
- size [1] compartment size
- dimensions [3] compartment size dimensions
- volume [None] compartment volume

Add genes to the model using the definitions stored in the annotation key. If this fails it tries some standard annotation keys: GENE ASSOCIATION, GENE_ASSOCIATION, gene_association, gene association.

- annotation_key the annotation dictionary key that holds the gene association for the protein/enzyme
- replace_existing [default=True] replace existing annotations, otherwise only new ones are added

$\label{lem:createGeneProteinAssociation} (\textit{protein, assoc, gid=None, name=None, gene_pattern=None, update_idx=True, altlabels=None)}$

Create and add a gene protein relationship to the model, note genes are mapped on protein objects which may or may not be reactions

- protein in this case the reaction
- assoc the COBRA style gene protein association
- gid the unique id
- name the optional name
- gene_pattern deprecated, not needed anymore
- update_idx update the model gene index, not used
- altlabels [default=None] alternative labels for genes, default uses geneIds

createGeneProteinAssociationFromTree(protein, gprtree, gid=None, name=None, gene_pattern=None, update_idx=True, altlabels=None)

Create and add a gene protein relationship to the model, note genes are mapped on protein objects which may or may not be reactions

- protein in this case the reaction
- gprtree the CBMPy GPR dictionary tree
- gid the unique id
- name the optional name
- gene_pattern deprecated, not needed anymore
- update_idx update the model gene index, not used
- altlabels [default=None] alternative labels for genes, default uses geneIds

createGroup(gid)

Create an empty group with

• gid the unique group id

createObjectiveFunction(*rid*, *coefficient=1*, *osense='maximize'*, *active=True*, *delete_current_obj=True*)

Create a single variable objective function:

- rid The
- **coefficient** [default=1]
- osense [default='maximize']
- active [default=True]
- **delete_current_obj** [default=True]

createParameter(pid, value, constant=True)

Instantiates a parameter

createReaction(*rid*, *name=None*, *reversible=True*, *create_default_bounds=True*, *silent=False*, *new=False*)

Create a new blank reaction and add it to the model:

- *id* the unique reaction ID
- name the reaction name
- reversible [default=True] the reaction reversibility. True is reversible, False is irreversible
- create_default_bounds create default reaction bounds, irreversible 0 <= J <= INF, reversable -INF <= J <= INF
- silent [default=False] if enabled this disables the printing of information messages

createReactionBounds(reaction, lb_value, ub_value)

Create a new lower bound for a reaction: value <= reaction

- reaction the reaction id
- lb_value the value of the lower bound
- ub_value the value of the upper bound

createReactionLowerBound(reaction, value)

Create a new lower bound for a reaction: value <= reaction

- reaction the reaction id
- value the value of the bound

Create a new blank reaction and add it to the model:

- id the unique reaction ID
- *name* the reaction name
- reversible [default=True] the reaction reversibility. True is reversible, False is irreversible
- create_default_bounds create default reaction bounds, irreversible 0 <= J <= INF, reversable -INF <= J <= INF
- silent [default=False] if enabled this disables the printing of information messages

createReactionReagent(reaction, metabolite, coefficient, silent=False)

Add a reagent to an existing reaction, both reaction and metabolites must exist

• reaction a reaction id

- metabolite a species/metabolite id
- coefficient the reagent coefficient

createReactionUpperBound(reaction, value)

Create a new upper bound for a reaction: reaction <= value

- reaction the reaction id
- value the value of the bound

createSingleGeneEffectMap()

This takes a model and analyses the logical gene expression patterns. This only needs to be done once, the result is a dictionary that has boolean effect patterns as keys and the (list of) genes that give rise to those patterns as values. This map is used by the single gene deletion method for further analysis.

Note this dictionary can also be stored and retrieved separately as long as the model structure is not changed i.e. the gene associations themselves or order of reactions (stored as the special entry 'keyJ').

Stored as self. __single_gene_effect_map___

Create a new species and add it to the model:

- id the unique species id
- boundary [default=False] whether the species is a variable (False) or is a boundary parameter (fixed)
- name [default="] the species name
- value [default=nan] the value not currently used
- compartment [default=None] the compartment the species is located in
- charge [default=None] the species charge
- chemFormula [default=None] the chemical formula

createUserDefinedConstraint(pid, lb, ub, components=None)

Create an FBCv3 UserDefinedConstraint

- pid unique id
- *lb* lower bound float/parameter
- ub upper bound float/parameter
- *componentents* optional, the user defined constraint componenents in the form of a list [(coefficient, variable, type, id), ...] and coefficient and variable can be parameters id is optional

deleteAllFluxBoundsWithValue(value)

Delete all flux bounds which have a specified value:

• value the value of the flux bound(s) to delete

deleteBoundsForReactionId(rid, lower=True, upper=True)

Delete bounds connected to reaction, rid

- rid a valid reaction id
- *upper* [default=True] delete the upper bound
- lower [default=True] delete the lower bound

deleteCompartment(sid, check_components=True)

Deletes a compartment object with id. Returns True if the compartment is deleted, False if not. In addition if components were checked a list of id's that reference the compartment are also returned.

- sid the compartment id
- check_components [default=True] if enabled check that no species or reactions makes

use of the compartment, fail if it does.

deleteGPRAssociation(gprid)

Delete a GPR association with id

• gprid the GPR association id

deleteGene(gid, also_delete_gpr=True)

Deletes the gene object with gid. Note if you want to delete a gene by label (locus tag etc) use the delete-GeneByLabel() function.

- gid the gene Id
- also_delete_gpr [default=True] automatically delete GPR's that contain no gene references

deleteGeneByLabel(label, also_delete_gpr=True)

Deletes the gene object with label (b2003 etc).

- label the gene with label to be deleted
- also_delete_gpr [default=True] automatically delete GPR's that contain no gene references

deleteGroup(gid)

Delete a group with

• gid the unique group id

deleteNonReactingSpecies(simulate=True)

Deletes all species that are not reagents (do not to take part in a reaction). Warning this deletion is permanent and greedy (not selective). Returns a list of (would be) deleted species

• simulate [default=True] only return a list of the speciesId's that would have been deleted if False

deleteObjective(objective_id)

Delete objective function:

objective_id the id of the objective function. If objective_id is given as 'active' then the active objective is deleted.

deleteReactionAndBounds(rid)

Delete all reaction and bounds connected to reaction

• rid a valid reaction id

deleteSpecies(sid, also_delete=None)

Deletes a species object with id

- sid the species id
- also_delete [default=None] only delete the species
- 'reagents' delete the species from the reactions it participates in as a **reagent** 'reactions' deletes the **reactions** that the species participates in

emptyUndelete()

Empties the undelete cache

exportFVAdata()

Export the fva data as an array and list of reaction id's

exportUserConstraints(filename)

Exports user constraints in json

findFluxesForConnectedSpecies(metab)

Returns a list of (reaction, flux value) pairs that this metabolite appears as a reagent of

• metab the metabolite name

getActiveObjective()

Returns the active objective object.

getActiveObjectiveReactionIds()

Returns the active objective flux objective reaction id's

getActiveObjectiveStoichiometry()

Returns a list of (coefficient, flux_objective) tuples

getAllFluxBounds()

DEPRECATED Returns a dictionary of all flux bounds [id:value]

getAllGeneActivities()

Returns a dictionary of genes (if defined) and whether they are active or not

getAllGeneProteinAssociations(use_labels=False)

Returns a dictionary of genes associated with each protein

• use_labels use V2 gene labels rather than ID's

getAllProteinActivities()

Returns a dictionary of reactions (if genes and GPR's are defined) and whether they are active or not

getAllProteinGeneAssociations(use_labels=False)

Returns a dictionary of the proteins associated with each gene

• use_labels use V2 gene labels rather than ID's

getBoundarySpeciesIds(rid=None)

Return all boundary species associated with reaction

• rid [default=None] by default return all boundary species in a model, alternatively a string containing a reaction id or list of reaction id's

getCompartment(cid)

Returns a compartment object with cid

• cid compartment ID

getCompartmentIds(substring=None)

Returns a list of compartment Ids, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getCompartmentObjects(substring=None)

Returns a list of compartment objects, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getDescription()

Returns the model description which was stored in the SBML <notes> field

getExchangeReactionIds()

Returns id's of reactions where the 'is_exchange' attribute set to True. This is by default reactions that contain a boundary species.

getExchangeReactions()

Returns reaction instances where the 'is_exchange' attribute set to True. This is by default reactions that contain a boundary species.

getFluxBoundByID(fid)

Returns a FluxBound/Parameter with id

• fid the fluxBound ID

getFluxBoundByReactionID(rid, bound)

Returns a FluxBound/Parameter instance

- rid the reaction ID
- bound the bound: 'upper', 'lower', 'equality'

getFluxBoundIds(substring=None)

Returns a list of fluxbound Ids, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getFluxBoundsByReactionID(rid)

Returns all FluxBound instances connected to a reactionId as a tuple of valid (lower, upper, None) or (None, None, equality) or alternatively invalid (lower, upper, equality).

• rid the reaction ID

under evaluation

getFluxesAssociatedWithCompartments(compartments)

Determines all reactions and flux values associated with a list of compartments. This function can be used to find all transport reactions between compartments, e.g. the cytosol and mitochondria. If the compartment IDs are 'cyt' and 'mit', respectively, you can call "your_model.getFluxesAssociatedWithCompartments(['cyt', 'mit'])" to get all fluxes between these compartments.

compartments: a list or set of compartment IDs.

To check the existing compartment IDs in your model call "your_model.getCompartmentIds()"

:returns a dictionary with reaction IDs as keys and corresponding flux values as values

getFluxesAssociatedWithSpecies(metab)

Returns a list of (reaction, flux value) pairs that this metabolite appears as a reagent in

• *metab* the metabolite name

${\tt getGPRIdAssociatedWithGeneId}({\it gid})$

Return the GPR(s) associated with the gene id:

• gid a gene id

getGPRIdAssociatedWithGeneLabel(label)

Return the GPR Id's associated with the gene label:

• label a gene label

getGPRIds(substring=None)

Returns a list of GPR Id's, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getGPRObjects(substring=None)

Returns a list of GPR objects, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getGPRassociation(gprid)

Returns a gene protein association object that has the identifier:

• gprid the gene protein identifier

getGPRforReaction(rid)

Return the GPR associated with the reaction id:

• rid a reaction id

getGPRforReactionAsDict(rid, useweakref=True)

Return the GPR associated with the reaction id as a nested dictionary structure:

• rid a reaction id

getGene(gid)

Returns a gene object that has the identifier:

• gid the gene identifier

getGeneByLabel(label)

Given a gene label return the corresponding Gene object

• label

getGeneIdFromLabel(label)

Given a gene label it returns the corresponding Gene id or None

• label

getGeneIds(substring=None)

Returns a list of gene Ids, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getGeneLabels(substring=None)

Returns a list of gene labels (locus tags), applies a substring search if substring is defined

• substring search for this pattern anywhere in the label

getGeneObjects(substring=None)

Returns a list of gene objects, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getGeneObjectsByLabel(substring=None)

Returns a list of gene objects, applies a substring search if substring is defined

• substring search for this pattern anywhere in the label

getGroup(gid)

Return a group with

• gid the unique group id

getGroupIds()

Get all group ids

getGroupMembership()

Returns group membership of items in groups. Returns {object_id: ['group_id1', 'group_id2']}

getGroupNames()

Get all group names

getIrreversibleReactionIds()

Return a list of irreversible reaction Id's

getModel()

Overrides the FBase inherited method, returns own instance.

getModelCreators()

Return model creator information

getObjFuncValue()

Returns the objective function value

getObject(pid)

• pid returns a model object with pid or None

getObjectiveIds(substring=None)

Returns a list of objective function Ids, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getOptimalValue()

Returns the optimal value of the objective function

getParameter(pid)

Returns a parameter object with pid

getReaction(rid)

Returns a reaction object with id

• rid reaction ID

getReactionActivity(rid)

If there is a GPR and genes associated with the reaction ID then return either active=True or inactive=False Note if there is no gene associated information then this will return active.

• rid a reaction id

getReactionBounds(rid)

Get the bounds of a reaction, returns a tuple of rid, lowerbound value, upperbound value and equality value (None means bound does not exist).

• rid the reaction ID

getReactionIds(substring=None)

Returns a list of reaction Ids, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getReactionIdsAssociatedWithSpecies(metab)

Returns a list of (reaction, flux value) pairs that this metabolite appears as a reagent in

• metab the metabolite name

getReactionLowerBound(rid)

Returns the lower bound of a reaction (it it exists) or None

• rid the reaction ID

getReactionNames(substring=None)

Returns a list of reaction names, applies a substring search if substring is defined

• substring search for this pattern anywhere in the name

getReactionObjects(substring=None)

Returns a list of reaction objects, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getReactionUpperBound(rid)

Returns the upper bound of a reaction (it it exists) or None

• rid the reaction ID

getReactionValues(only_exchange=False)

Returns a dictionary of ReactionID : ReactionValue pairs:

• only_exchange [default=False] only return the reactions labelled as exchange

getReversibleReactionIds()

Return a list of reversible reaction Id's

getSolutionVector(names=False)

Return a vector of solution values

• names [default=False] if True return a solution vector and list of names

getSpecies(sid)

Returns a species object with sid

• sid a specied ID

getSpeciesIds(substring=None)

Returns a list of species Ids, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

getSpeciesObjects(substring=None)

Returns a list of species objects, applies a substring search if substring is defined

• substring search for this pattern anywhere in the id

hasObject(pid)

• pid returns a boolean if there is a registered object with pid-

importUserConstraints(filename)

Exports user constraints in json

registerObjectInGlobalStore(obj)

object

renameObjectIds(prefix=None, suffix=None, target='all', ignore=None)

This method is designed for target="all" other use may result in inconsistent models. Update: "species" and "reactions" should also work as advertised, please check results.

- prefix [None] if supplied add as a prefix
- suffix [None] if supplied add as a suffix
- target ['all'] specify what class of objects to rename
- 'species'
- · 'reactions'
- · 'bounds'
- · 'objectives'
- 'all'
- ignore [default=None] a list of id's to ignore

resetAllGenes(update_reactions=False)

Resets all genes to their default activity state (normally on)

• *update_reactions* [default=False] update the associated reactions fluxbounds from the gene deletion bounds if they exist

resetAllInactiveGPRBounds()

Resets all reaction bounds modified by the ${\tt cmod.setAllInactiveGeneReactionBounds()}$ method to their previous values

serialize(protocol=0)

Serialize object, returns a string by default

• protocol [default=0] serialize to a string or binary if required,

see pickle module documentation for details

overloaded in CBModel

serializeToDisk(filename, protocol=2)

Serialize to disk using pickle protocol:

- filename the name of the output file
- protocol [default=2] serialize to a string or binary if required,

see pickle module documentation for details

overloaded in CBModel

setAllFluxBounds(bounds)

DEPRECATED! use setFluxBoundsFromDict()

Sets all the fluxbounds present in bounds

• bounds a dictionary of [fluxbound_id : value] pairs (not per reaction!!!)

setAllInactiveGPRBounds(lower=0.0, upper=0.0)

Set all reactions that are inactive (as determined by gene and gpr evaluation) to bounds:

- lower [default=0.0] the new lower bound
- upper [default=0.0] the new upper bound

setAllProteinActivities(activites, lower=0.0, upper=0.0)

Given a dictionary of activities [rid: boolean] pairs set all the corresponding reactions:

- activities a dictionary of [rid: boolean] pairs
- lower [default=0.0] the lower bound of the deactivated flux
- upper [default=0.0] the upper bound of the deactivated flux

setBoundValueByName(rid, value, bound)

Deprecated use setReactionBound

Set a reaction bound

- rid the reactions id
- value the new value
- bound this is either 'lower' or 'upper'

setCreatedDate(date=None)

Set the model created date tuple(year, month, day, hour, minute, second)

• *date* [default=None] default is now (automatic) otherwise (year, month, day, hour, minute, second) e.g. (2012, 09, 24, 13, 34, 00)

setDescription(html)

Sets the model description which translates into the SBML <notes> field.

• html any valid html or the empty string to clear "

setFluxBoundsFromDict(bounds)

DEPRECATED! This method will be modified to use reaction Idin CBMPy 0.9.0 Sets all the fluxbounds present in bounds

• bounds a dictionary of [fluxbound_id : value] pairs (not per reaction!!!)

setGeneActive(g_id, update_reactions=False)

Effectively restores a gene by setting it's active flag

- g_id a gene ID
- *update_reactions* [default=False] update the associated reactions fluxbounds from the gene deletion bounds if they exist

setGeneInactive(*g_id*, *update_reactions=False*, *lower=0.0*, *upper=0.0*)

Effectively deletes a gene by setting it's inactive flag while optionally updating the GPR associated reactions

- g_id a gene ID
- update_reactions [default=False] update the associated reactions fluxbounds
- lower [default=0.0] the deactivated reaction lower bound
- upper [default=0.0] the deactivated reaction upper bound

setModifiedDate(date=None)

Set the model modification date: tuple(year, month, day, hour, minute, second)

• *date* [default=None] default is now (automatic) otherwise (year, month, day, hour, minute, second) e.g. (2012, 09, 24, 13, 34, 00)

setObjectiveFlux(rid, coefficient=1, osense='maximize', delete_objflx=True)

Set single target reaction flux for the current active objective function.

- rid a string containing a reaction id
- coefficient [default=1] an objective flux coefficient
- osense the optimization sense must be maximize or minimize
- delete_objftx [default=True] delete all existing fluxObjectives in the active objective function

setPrefix(prefix, target)

This is alpha stuff, target can be:

- · 'species'
- · 'reactions'
- · 'constraints'
- · 'objectives'
- · 'all'

setReactionBound(rid, value, bound)

Set a reaction bound

- rid the reactions id
- value the new value
- bound this is either 'lower' or 'upper', or 'equal'

setReactionBounds(rid, lower, upper)

Set both the upper and lower bound of a reaction:

- rid the good old reaction id
- lower the lower bound value
- upper the upper bound value

setReactionLowerBound(rid, value)

Set a reactions lower bound (if it exists)

- rid the reactions id
- value the new value

$\verb"setReactionUpperBound" (rid, value)"$

Set a reactions upper bound (if it exists)

- rid the reaction id
- value the new value

setSuffix(suffix, target)

This is alpha stuff, target can be:

'species'

- · 'reactions'
- · 'constraints'
- · 'objectives'
- 'all'

sortReactionsById()

Sorts the reactions by Reaction.id uses the python string sort

sortSpeciesById()

Sorts the reaction list by Reaction.id uses the python string sort

splitEqualityFluxBounds()

Splits any equalit flux bounds into lower and upper bounds.

testGeneProteinAssociations()

This method will test the GeneProtein associations and return a list of protein, association pairs

unRegisterObjectInGlobalStore(sid)

• sid

updateNetwork(lower=0.0, upper=0.0, silent=False)

Update the reaction network based on gene activity. If reaction is deactivated then lower and upper bounds are used

- lower [default=0.0] deactivated lower bound
- upper [default=0.0] deactivated upper bound

class Objective(pid, operation)

An objective function

addFluxObjective(fobj, override=False)

Adds a FluxObjective instance to the Objective

- fobj the FluxObjective object
- override [default=False] override pushing the global id map, this should never be used

createFluxObjectives(fluxlist)

Create and add flux objective objects to this objective function.

• fluxlist a list of one or more ('coefficient', 'rid', 'type') triples

createQuadraticFluxObjectives(fluxlist)

Create and add quadratic flux objective objects to this objective function.

• fluxlist a list of one or more ('coefficient', 'rid', 'rid2', 'type') triples

deleteAllFluxObjectives()

Delete all flux objectives

getFluxObjective(foid)

Return the flux objective with id.

• foid the flux objective id returns either an object or a list if there are multiply defined flux objectives

getFluxObjectiveData()

Returns a list of ObjectiveFunction components as (coefficient, flux, type) pairs

getFluxObjectiveForReaction(rid)

Returns the FluxObjective associated with the suplied rid. If there is more than fluxObjective associated with a reaction (illegal) then a list of fluxObjectives is returned.

rid a reaction id

getFluxObjectiveIDs()

Returns a list of ObjectiveFlux ids, for the reaction id's use *getFluxObjectiveReactions()* or for coefficient, fluxobjective pairs use *getFluxObjectiveData()*

getFluxObjectiveReactions()

Returns a list of reactions that are used as flux_objectives

getFluxObjectives()

Returns the list of FluxObjective objects.

getLinearFluxObjectives()

Returns a list of linear variable flux objective objects

getOperation()

Returns the operation or sense of the objective

getQuadraticBivariateFluxObjectives()

Returns a list of bivariate quadratic variable flux objective objects

getQuadraticFluxObjectives()

Returns a list of quadratic variable flux objective objects

getValue()

Returns the current value of the attribute (input/solution)

setOperation(operation)

Sets the objective operation (sense)

• operation [default='maximize'] one of 'maximize', 'maximise', 'max', 'minimize', 'minimise', 'min'

setValue(value)

Sets the attribute "value"

class Parameter(pid, value, name=None, constant=True)

Holds parameter information

addAssociation(assoc)

Add an object ID to associate with this object

deleteAssociation(assoc)

Delete the object id associated with this object

getAssociations()

Return the Object ID's associated with this parameter

getValue()

Returns the current value of the attribute (input/solution)

setValue(value)

Sets the value attribute:

· value a float

property value

Returns the current value of the attribute (input/solution)

class Reaction(pid, name=None, reversible=True)

Holds reaction information

addReagent(reag)

Adds an instantiated Reagent object to the reaction

changeId(pid)

Changes the Id of the reaction and updates associated FluxBounds

createReagent(metabolite, coefficient)

Create a new reagent and add it to the reaction:

- metabolite the metabolite name
- · coefficient the

—negative coefficient is a substrate – positive coefficient is a product

Will fail if a species reference already exists

deactivateReaction(lower=0.0, upper=0.0, silent=True)

Deactivates a reaction by setting its bounds to lower and upper. Restore with reactivateReaction()

- lower [default=0.0] bound
- upper [default=0.0] bound

deleteReagentWithSpeciesRef(sid)

Delete a reagent (or reagents) that refers to the species id:

• sid a species/metabolite id

getEquation(reverse_symb='=', irreverse_symb='>', use_names=False)

Return a pretty printed string containing the reaction equation

- reverse_symb [default = '='] the symbol to use for reversible reactions
- *irreverse_symb* [default = '>'] the symbol to use for irreversible reactions
- use_names [defualt = False] use species names rather than id's

getFVAdata(roundnum=None, silent=True)

Returns the data generated by CBSolver.FluxVariabilityAnalysis() for this reaction as a tuple of (Flux, FVAmin, FVAmax, span) where span is abs(FVAmax - FVAmin). FVAmin or FVAmax is None this indicates no solution to that particular optimization (infeasible).

- roundnum [default=None] the integer number of roundoff decimals the default is no rounding
- *silent* [default=True] supress output to stdout

getGPRassociationString(use_labels=True)

Return the GPR string associated with this reaction (assuming it exists) or None.

• use_labels [default=True] return string with lab

getLowerBound()

Get the value of the reactions lower bound

getProductIds(use names=False)

Returns a list of the reaction products, species identifiers

• use_names [default = False] use species names rather than id's

getReagent(rid)

Return the one or more reagent objects which have *rid*:

• rid a reagent rid

getReagentObjIds()

Returns a list of the reagent id's. For the name of the reagents/metabolites use < reaction>.getSpeciesIds()

getReagentRefs()

Returns a list of the reagents/metabolites

getReagentWithSpeciesRef(sid)

Return the reagent object which refers to the *species* id. If there are multiple reagents that refer to the same species a list is returned.

• sid the species/metabolite id

getSpeciesIds()

Returns a list of the reagents/metabolites

getSpeciesObj()

Returns a list of the species objects that are reagents

getStoichiometry(use_names=False, altout=False)

Returns a list of (coefficient, species) pairs for this reaction

- use_names [default = False] use species names rather than id's
- *altout* [default = False] returns a dictionary [DEPRECATED]

getSubstrateIds(use_names=False)

Returns a list of the reaction substrates, species identifiers

• use_names [defualt = False] use species names rather than id's

getUpperBound()

Get the value of the reactions upper bound

getValue()

Returns the current value of the flux.

reactivateReaction(silent=True)

Activates a reaction deactivated with deactivateReaction

setId(fid)

Sets the object Id

• fid a valid c variable style id string

Reimplements @FBase.setId()

setLowerBound(value)

Set the value of the reactions lower bound

• value a floating point value

setStoichCoefficient(sid, value)

Sets the stoichiometric coefficient of a reagent that refers to a metabolite. Note *negative coefficients* are *substrates* while *positive* ones are *products*. At this point zero coefficients are not allowed

- sid the species/metabolite id
- *value* a floating point value != 0

setUpperBound(value)

Set the value of the reactions upper bound

• value a floating point value

setValue(value)

Sets the attribute *value* in this case the flux.

class ReactionNew(pid, name=None, lb=-inf, ub=inf, reversible=True)

Extended reaction class with new upper/lower bound structure

deactivateReaction(lower=0.0, upper=0.0, silent=True)

Deactivates a reaction by setting its bounds to lower and upper. Restore with reactivateReaction()

- lower [default=0.0] bound
- upper [default=0.0] bound

getLowerBound()

Get the value of the reactions lower bound

getUpperBound()

Get the value of the reactions upper bound

reactivateReaction(silent=True)

Activates a reaction deactivated with deactivateReaction

setId(fid)

Sets the object Id

• fid a valid c variable style id string

Reimplements @FBase.setId()

setLowerBound(value)

Set the value of the reactions lower bound

• value a floating point value

setUpperBound(value)

Set the value of the reactions upper bound

• value a floating point value

class Reagent(pid, species_ref, coef)

Has a reactive species id and stoichiometric coefficient:

- negative = substrate
- positive = product
- species_ref a reference to a species obj

getCoefficient()

Returns the reagent coefficient

getRole()

Returns the reagents role, "substrate", "product" or None

getSpecies()

Returns the metabolite/species that the reagent reference refers to

setCoefficient(coef)

Sets the reagent coefficient and role, negative coefficients are substrates and positive ones are products

• coeff the new coefficient

setSpecies(spe)

Sets the metabolite/species that the reagent reference refers to

Holds species/metabolite information

getCharge()

Returns the species charge

getChemFormula()

Returns the species chemical formula

getReagentOf()

Returns a list of reaction id's that this metabolite occurs in

getValue()

Returns the current value of the attribute (input/solution)

isReagentOf()

Returns a dynamically generated list of reactions that this species occurs as a reagent

setBoundary()

Sets the species so it is a boundary metabolite or fixed which does not occur in the stoichiometric matrix N

setCharge(charge)

Sets the species charge:

• charge a signed double but generally a signed int is used

setChemFormula(cf)

Sets the species chemical formula

• cf a chemical formula e.g. CH3NO2

setId(fid, allow_rename=False)

Sets the object Id

• fid a valid c variable style id string

Reimplements @FBase.setId()

setReagentOf(rid)

Adds the supplied reaction id to the reagent_of list (if it isn't one already)

• rid a valid reaction id

```
setValue(value)
```

Sets the attribute "value"

unsetBoundary()

Unsets the species boundary attribute so that the metabolite is free and therefore occurs in the stoichiometric matrix N

class UserDefinedConstraint(pid, lb, ub)

This is an FBCv3 class that defines a set of user defined constraints, it is similar to an objective constraint except allows parameters as coefficients and values in the constraint components

addConstraintComponent(cc)

• cc UserConstraintComponent

createConstraintComponent(pid, coefficient, variable, ctype)

getConstraintComponent(cid)

• cid get a constraint component that matches cid

getConstraintComponentData()

getConstraintComponentForVariable(rid)

rid a component id

getConstraintComponentIDs()

getConstraintComponentVariableTypes()

getConstraintComponentVariables()

getConstraintComponents()

3.7 CBMPy: CBModelTools module

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3.8 CBMPy: CBMultiCore module

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```
grouper(3, 'abcdefg', 'x') --> ('a', 'b', 'c'), ('d', 'e', 'f'), ('g', 'x', 'x')
```

runMultiCoreFVA(fba, selected_reactions=None, pre_opt=True, tol=None, objF2constr=True, rhs_sense='lower', optPercentage=100.0, work_dir=None, quiet=True, debug=False, oldlpgen=False, markupmodel=True, procs=2, override_bin=None)

Run a multicore FVA where:

- fba is an fba model instance
- procs [default=2] number of processing threads (optimum seems to be about the number of physical cores)
- python_override_bin allows customization of the Python bin used for the multicore process

3.9 CBMPy: CBMultiEnv module

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3.10 CBMPy: CBNetDB module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBNetDB.py 710 2020-04-27 14:22:34Z bgoli \$)

class DBTools

Tools to work with SQLite DB's (optimized, no SQL required).

checkEntryInColumn(table, col, rid)

Check if an entry exists in a table

- table the table name
- col the column name
- rid the row to search for

closeDB()

Close the DB connection and reset the DBTools instance (can be reconnected)

commitDB()

Commits all curent changes to DB, returns a boolean.

connectSQLiteDB(db_name, work_dir=None)

Connect to a sqlite database.

- db_name the name of the sqlite database
- work_dir the optional database path

createDBTable(table, sqlcols)

Create a database table if it does not exist:

- table the table name
- sqlcols a list containing the SQL definitions of the table columns: <id> <type> for example ['gene TEXT PRIMARY KEY', 'aa_seq TEXT', 'nuc_seq TEXT', 'aa_len INT', 'nuc_len INT']

Effectively writes CREATE TABLE "table" (<id> <type>, gene TEXT PRIMARY KEY, aa_seq TEXT, nuc_seq TEXT, aa_len INT, nuc_len INT) % table

dumpTableToCSV(table, filename)

Save a table as tab separated txt file

- table the table to export
- *filename* the filename of the table dump

dumpTableToTxt(table, filename)

Save a table as tab separated txt file

- table the table to export
- *filename* the filename of the table dump

executeSQL(sql)

Execute a SQL command:

• sql a string containing a SQL command

fetchAll(sql)

Raw SQL query e.g. 'SELECT id FROM gene WHERE gene="G""

```
getCell(table, col, rid, cell)
```

Get the table cell which correspond to rid in column. Returns the value or None

- table the database table
- · col the column id
- rid the row index id
- cell the column of the cell you want tp extract

getColumns(table, cols)

Fetch the contents of one or more columns of data in a table

- table the database table
- cols a list of one or more column id's

```
getRow(table, col, rid)
```

Get the table row(s) which correspond to rid in column. Returns the row(s) as a list, if the column is the primary key this is always a single entry.

- table the database table
- · col the column id
- rid the row index id

getTable(table, colOut=False)

Returns an entire database table

- table the table name
- *colOut* optionally return a tuple of (data,ColNames)

insertData(table, data, commit=True)

```
Insert data into a table: "INSERT INTO %s (?, ?, ?, ?, ?) VALUES (?, ?, ?, ?, ?)" % table,
```

```
(?, ?, ?, ?, ?)))
```

- table the DB table name
- data a dictionary of {id:value} pairs
- commit whether to commit the data insertions

updateData(table, col, rid, data, commit=True)

Update already defined data

- table the table name
- col the column name
- rid the row id to update
- data a dictionary of {id:value} pairs
- commit whether to commit the data updates

UPDATE COMPANY SET ADDRESS = 'Texas' WHERE ID = 6;

class KeGGSequenceTools(url, db_name, work_dir)

Using the KeGG connector this class provides tools to construct an organims specific sequence database

class KeGGTools(url)

Class that holds useful methods for querying KeGG via a SUDS provided soap client

$fetchSeqfromKeGG(k_gene)$

Given a gene name try and retrieve the gene and amino acid sequence

class MIRIAMTools

Tools dealing with MIRIAM annotations

class RESTClient

Class that provides the basis for application specific connectors to REST web services

Close()

Close the currently active connection

Connect(root)

Establish HTTP connection to

• root the site root "www.google.com"

Get(query)

Perform an http GET using:

- query e.g.
- reply_mode [default="] this is the reply mode

For example "/semanticSBML/annotate/search.xml?q=ATP"

GetLog()

Return the logged history

Log(txt)

Add txt to logfile history

• txt a string

class SemanticSBML

REST client for connecting to SemanticSBML services

parseXMLtoText(xml)

Parse the xml output by quickLookup() into a list of URL

• xml XML returns from SemanticSBML

quickLookup(txt)

Do a quick lookpup for txt using SemanticSBML (connectic if required) and return results. Returns a list of identifiers.org id's in descending priority (as return)

• *txt* the string to lookup

viewDataInWebrowser(maxres=10)

Attempt to view #maxres results returned by SemanticSBML in the default browser

• maxres default maximum number of results to display.

3.11 CBMPy: CBPlot module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBPlot.py 710 2020-04-27 14:22:34Z bgoli \$)

plotFluxVariability(fva_data, fva_names, fname, work_dir=None, title=None, ySlice=None, minHeight=None, maxHeight=None, roundec=None, autoclose=True, fluxval=True, type='png')

Plots and saves as an image the flux variability results as generated by CBSolver.FluxVariabilityAnalysis.

- fva_data FluxVariabilityAnalysis() FVA OUTPUT_ARRAY
- fva_names Flux Variability Analysis() FVA OUTPUT_NAMES
- fname filename_base for the CSV output
- work_dir [default=None] if set the output directory for the csv files
- title [default=None] the user defined title for the graph
- ySlice [default=None] this sets an absolute (fixed) limit on the Y-axis (+- ySlice)
- minHeight [default=None] the minimum length that defined a span
- maxHeight [default=None] the maximum length a span can obtain, bar will be limited to maxHeight and coloured yellow
- roundec [default=None] an integer indicating at which decimal to round off output. Default is no rounding.
- autoclose [default=True] autoclose plot after save
- fluxval [default=True] plot the flux value
- type [default='png'] the output format, depends on matplotlib backend e.g. 'png', 'pdf', 'eps'

3.12 CBMPy: CBRead module

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loadModel(sbmlfile)

Loads any SBML model in COBRA, FAME (SBML2FBA), SBML3FBCv1, SBML3FBCv2 format.

• sbmlfile an SBML model file

```
readCOBRASBML(fname, work_dir=None, return_sbml_model=False, delete_intermediate=False, fake_boundary_species_search=False, output_dir=None, skip_genes=False, scan_notes_gpr=True)
```

Read in a COBRA format SBML Level 2 file with FBA annotation where and return either a CBM model object or a (cbm_mod, sbml_mod) pair if return_sbml_model=True

- fname is the filename
- work dir is the working directory
- delete_intermediate [default=False] delete the intermediate SBML Level 3 FBC file
- fake_boundary_species_search [default=False] after looking for the boundary_condition of a species search for overloaded id's <id> b
- *output_dir* [default=None] the directory to output the intermediate SBML L3 files (if generated) default to input directory
- skip_genes [default=False] do not load GPR data
- scan_notes_gpr [default=True] if the model is loaded and no genes are detected the scan the <notes> field for GPR associationa

readExcel97Model(xlname, write_sbml=True, sbml_level=3, return_dictionaries=False)

Reads a model encoded as an Excel97 workbook and returns it as a CBMPy model object and SBML file. Note the workbook must be formatted exactly like those produced by cbm.writeModelToExcel97(). Note that reactions have to be defined in **both** the *reaction* and *network react* sheets to be included in the model.

- xlpath the filename of the Excel workbook
- return_model [default=True] construct and return the CBMPy model
- write sbml [default=True] write the SBML file to fname
- return_dictionaries [default=False] return the dictionaries constructed when reading the Excel file (in place of the model)
- sbml_level [default=3] write the SBML file as either SBML L2 FBA or SBML L3 FBC file.

readSBML2FBA(fname, work_dir=None, return_sbml_model=False, fake_boundary_species_search=False, scan_notes_gpr=True)

Read in an SBML Level 2 file with FBA annotation where:

- fname is the filename
- work_dir is the working directory if None then only fname is used
- return_sbml_model [default=False] return a a (cbm_mod, sbml_mod) pair
- fake_boundary_species_search [default=False] after looking for the boundary_condition of a species search for overloaded id's <id> b
- scan_notes_gpr [default=True] if the model is loaded and no genes are detected the scan the <notes> field for GPR associationa

readSBML3FBC(fname, work_dir=None, return_sbml_model=False, xoptions={'validate': False}, scan_notes_gpr=True)

Read in an SBML Level 3 file with FBC annotation where and return a CBM model object

- *fname* is the filename
- work_dir is the working directory
- return_sbml_model deprecated and ignored please update code
- xoptions special load options, enable with option=True except for nmatrix_type which has a type.
 - nogenes do not load/process genes
 - *noannot* do not load/process any annotations
 - validate validate model and display errors and warnings before loading
 - readcobra read the cobra annotation
 - read_model_string [default=False] read the model from a string (instead of a filename) containing an SBML document
 - nmatrix_type [default='normal'] define the type of stoichiometrich matrix to be built
 - * 'numpy' dense numpy array (best performance)
 - * 'scipy_csr' scipy sparse matrix (lower performance, low memory)
 - * 'sympy' a sympy rational matrix (low performance, high memory, cast to dense to analyse)
 - * None do not build matrix
- scan_notes_gpr [default=True] if the model is loaded and no genes are detected scan the <notes> field for GPR associationa

readSK_FVA(filename)

Read Stevens FVA results (opt.fva) file and return a list of dictionaries

readSK_vertex(fname, bigfile=True, fast_rational=False, nformat='%.14f', compression=None, hdf5file=None)
Reads in Stevens vertex analysis file:

- fname the input filename (.all file that results from Stevens pipeline)
- bigfile [default=True] this option is now always true and is left in for backwards compatability
- fast_rational [default=False] by default off and uses SymPy for rational—>float conversion, when on uses float decomposition with a slight (2th decimal) decrease in accuracy

- nformat [default='%.14f'] the number format used in output files
- compression [default=None] compression to be used in hdf5 files can be one of [None, 'lzf', 'gz?', 'szip']
- hdf5file [default=None] if None then generic filename '_vtx_.tmp.hdf5' is uses otherwise <hdf5file>.hdf5

and returns an hdf5 filename of the results with a single group named data which countains datasets

- · vertices
- rays
- lin

where all vectors are in terms of the column space of N.

readSK_vertexOld(fname, bigfile=False, fast_rational=False, nformat='%.14f', compresslevel=3)

Reads in Stevens vertex analysis file and returns, even more optimized for large datasets than the original.

- a list of vertex vectors
- a list of ray vectors
- the basis of the lineality space as a list of vectors

all vectors in terms of the column space of N

3.13 CBMPy: CBReadtxt module

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readCSV(model_file, bounds_file=None, biomass_flux=None, model_id='FBAModel', reaction_prefix='R_', has_header=False)

This function loads a CSV file and translates it into a Python object:

```
- *model_file* the name of the CSV file that contains the model
- *bounds_file* the name of the CSV file that contains the flux bounds
- *biomass_flux* the name of the reaction that is the objective function
- *reaction_prefix* [default='R _'] the prefix to add to input reaction ID's
- *has_header* [default=False] if there is a header row in the csv file
```

3.14 CBMPy: CBSolver module

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3.15 CBMPy: CBTools module

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addFluxAsActiveObjective(f, reaction_id, osense, coefficient=1)

Adds a flux as an active objective function

- reaction_id a string containing a reaction id
- osense objective sense must be maximize or minimize
- *coefficient* the objective funtion coefficient [default=1]

addGenesFromAnnotations(fba, annotation_key='GENE ASSOCIATION', gene_pattern=None)

THIS METHOD IS DERPRECATED PLEASE USE cmod.createGeneAssociationsFromAnnotations()

Add genes to the model using the definitions stored in the annotation key

- fba and fba object
- annotation_key the annotation dictionary key that holds the gene association for the protein/enzyme
- gene_pattern deprecated, not needed anymore

addSinkReaction(fbam, species, lb=0.0, ub=1000.0)

Adds a sink reactions that consumes a model *species* so that X ->

- fbam an fba model object
- species a valid species name
- *lb* lower flux bound [default = 0.0]
- *ub* upper flux bound [default = 1000.0]

addSourceReaction(fbam, species, lb=0.0, ub=1000.0)

Adds a source reactions that produces a model species so that -> X

- fbam an fba model object
- species a valid species name
- *lb* lower flux bound [default = 0.0]
- *ub* upper flux bound [default = 1000.0]

Note reversiblity is determined by the lower bound, default 0 = irreversible. If negative then reversible.

addStoichToFBAModel(fm)

Build stoichiometry: this method has been refactored into the model class - cmod.buildStoichMatrix()

checkExchangeReactions(fba, autocorrect=True)

Scan all reactions for exchange reactions (reactions containing a boundary species), return a list of inconsistent reactions or correct automatically.

- fba a CBMPy model
- autocorrect [default=True] correctly set the "is_exchange" attribute on a reaction

checkFluxBoundConsistency(fba)

Check flux bound consistency checks for multiply defined bounds, bounds without a reaction, inconsistent bounds with respect to each other and reaction reversibility. Returns a dictionary of bounds/reactions where errors occur.

checkIds(*fba*, *items='all'*)

Checks the id's of the specified model attributes to see if the name is legal and if there are duplicates. Returns a list of items with errors.

- fba a CBMPy model instance
- *items* [default='all'] 'all' means 'species,reactions,flux_bounds,objectives' of which one or more can be specified

checkReactionBalanceElemental(f, Rid=None, zero tol=1e-12)

Check if the reaction is balanced using the chemical formula

- \bullet f the FBA object
- *Rid* [default = None] the reaction to check, defaults to all
- zero_tol [default=1.0e-12] the floating point zero used for elemental balancing

This function is derived from the code found here: http://pyparsing.wikispaces.com/file/view/chemicalFormulas.py

createTempFileName()

Return a temporary filename

createZipArchive(zipname, files, move=False, compression='normal')

Create a zip archive which contains one or more files

- zipname the name of the zip archive to create (fully qualified)
- *files* either a valid filename or a list of filenames (fully qualified)
- move [default=False] attempt to delete input files after zip-archive creation
- compression [default='normal'] normal zip compression, set as None for no compression only store files (zlib not required)

deSerialize(s)

Deserializes a serialised object contained in a string

deSerializeFromDisk(filename)

Loads a serialised Python pickle from *filename* returns the Python object(s)

exportArray2CSV(arr, fname)

Export an array to fname.csv

- arr the an array like object
- fname the output filename
- sep [default=','] the column separator

exportArray2TXT(arr, fname)

Export an array to fname.txt

- arr the an array like object
- fname the output filename
- sep [default=','] the column separator

exportLabelledArray(arr, fname, names=None, sep=',', fmt='%f')

Write a 2D array type object to file

- arr the an array like object
- names [default=None] the list of row names
- fname the output filename
- sep [default=','] the column separator
- fmt [default='%s'] the output number format

exportLabelledArray2CSV(arr, fname, names=None)

Export an array with row names to fname.csv

- arr the an array like object
- fname the output filename
- names [default=None] the list of row names

exportLabelledArray2TXT(arr, fname, names=None)

Export an array with row names to fname.txt

- arr the an array like object
- names [default=None] the list of row names
- fname the output filename

exportLabelledArrayWithHeader(arr, fname, names=None, header=None, sep=',', fmt='%f')

Export an array with row names and header

- arr the an array like object
- names [default=None] the list of row names
- header [default=None] the list of column names
- *fname* the output filename
- sep [default=','] the column separator
- fmt [default='%s'] the output number format
- appendlist [default=False] if True append the array to fname otherwise create a new file

exportLabelledArrayWithHeader2CSV (arr, fname, names = None, header = None)

Export an array with row names and header to fname.csv

- arr the an array like object
- fname the output filename
- names [default=None] the list of row names
- header [default=None] the list of column names

exportLabelledArrayWithHeader2TXT(arr, fname, names=None, header=None)

Export an array with row names and header to fname.txt

- arr the an array like object
- names the list of row names
- header the list of column names
- fname the output filename

exportLabelledLinkedList(arr, fname, names=None, sep=',', fmt='%s', appendlist=False)

Write a 2D linked list [[...],[...],[...]] and optionally a list of row labels to file:

- arr the linked list
- fname the output filename
- names [default=None] the list of row names
- sep [default=','] the column separator
- fmt [default='%s'] the output number format
- appendlist [default=False] if True append the array to fname otherwise create a new file

findDeadEndMetabolites(fbam)

 $Finds \ dead-end \ (single \ reaction) \ metabolites \ rows \ in \ N \ with \ a \ single \ entry), \ returns \ a \ list \ of \ (metabolite, \ reaction) \ ids$

findDeadEndReactions(fbam)

Finds dead-end (single substrate/product) reactions (cols in N with a single entry), returns a list of (metabolite, reaction) ids

fixReversibility(fbam, auto_correct=False)

Set fluxbound lower bound from reactions reversibility information.

• fbam and FBAModel instance

• auto_correct (default=False) if True automatically sets lower bound to zero if required, otherwise prints a warning if false.

getBoundsDict(fbamod, substring=None)

Return a dictionary of reactions&bounds

getExchBoundsDict(fbamod)

Return a dictionary of all exchange reactions (as determined by the is_exchange attribute of Reaction)

• fbamod a CBMPy model

getModelGenesPerReaction(fba, gene_pattern=None, gene_annotation_key='GENE ASSOCIATION')

Parse a BiGG style gene annotation string using default gene_pattern='($(W^*w^*W^*)$)' or (<any non-alphanum><any alphanum>(any non-alphanum>)

Old eColi specific pattern '(bw*W)'

It is advisable to use the model methods directly rather than this function

loadObj(filename)

Loads a serialised Python pickle from *filename*.dat returns the Python object(s)

merge2Models(m1, m2, ignore=None, ignore_duplicate_ids=False)

Merge 2 models, this method does a raw merge of model 2 into model 1 without any model checking. Component id's in ignore are ignored in both models and the first objective of model 1 is arbitrarily set as active. Compartments are also merged and a new "OuterMerge" compartment is also created.

In all cases duplicate id's are tracked and ignored, essentially using the object id encountered first - usually that of model 1. Duplicate checking can be disabled by setting the *ignore_duplicate_ids* flag.

- *m1* model 1
- m2 model 2
- ignore [[]] do not merge these id's
- ignore_duplicate_ids [False] default behaviour that can be enabled

In development: merging genes and gpr's.

mergeGroups(m, groups, new_id, new_name=", auto_delete=False)

Merge a list of groups into a new group. Note, annotations are not merged!

- *m* the model containing the source groups
- groups a list of groups
- new_id the new, merged, group id
- new_name [default="] the new group name, the default behaviour is to merge the old names
- auto delete [default=False] delete the source groups

processBiGGannotationNote(fba, annotation_key='note')

Parse the HTML formatted reaction information stored in the BiGG notes field. This function is being deprecated and replaced by *CBTools.processSBMLAnnotationNotes()*

requires an annotation_key which contains a BiGG HTML fragment

${\tt processBiGGchemFormula}(\mathit{fba})$

Disambiguates the overloaded BiGG name NAME CHEMFORMULA into

- species.name NAME
- species.chemFormula CHEMFORMULA

processExchangeReactions(fba, key)

Extract exchange reactions from model using key and return:

- a dictionary of all exchange reactions without medium reactions
- a dictionary of *medium* exchange reactions (negative lower bound)

processSBMLAnnotationNotes(fba, annotation_key='note', level=3)

Parse the HTML formatted reaction information stored in the SBML notes field currently processes BiGG and PySCeSCBM style annotations it looks for the the annotation indexed with the *annotation key*

 annotation_key [default='note'] which contains a HTML/XHTML fragment in BiGG/PySCeSCBM format (ignored in L3)

removeFixedSpeciesReactions(f)

This function is a hack that removes reactions which only have boundary species as reactants and products. These are typically gene associations encoded in the Manchester style and there is probably a better way of working around this problem ...

• f an instantiated fba model object

roundOffWithSense(val, osense='max', tol=1e-08)

Round of a value in a way that takes into consideration the sense of the operation that generated it

- val the value
- osense [default='max'] the sense
- tol [default=1e-8] the tolerance of the roundoff factor

scanForReactionDuplicates(f, ignore_coefficients=False)

This method uses uses a brute force apprach to finding reactions with matching stoichiometry

scanForUnbalancedReactions(f, output='all')

Scan a model for unbalanced reactions, returns a tuple of dictionaries balanced and unbalanced:

- f an FBA model instance
- output [default='all'] can be one of ['all','charge','element']
- charge return all charge un balanced reactions
- element return all element un balanced reactions

setSpeciesPropertiesFromAnnotations (fbam, overwriteCharge=False, overwriteChemFormula=False)

This will attempt to set the model Species properties from the annotation. With the default options it will only replace missing data. With ChemicalFormula this is easy to detect however charge may have an "unknown value" of 0. Setting the optional values to true will replace any existing value with any valid annotation.

- overwriteChemFormula [default=False]
- overwriteCharge [default=False]

splitReversibleReactions(fba, selected_reactions=None)

Split a (set of) reactions into reversible reactions returns a copy of the original model

R1: A = B R1f: A -> B R1r: B -> A

- fba an instantiated CBMPy model object
- selected_reactions if a reversible reaction id is in here split it

splitSingleReversibleReaction(fba, rid, fwd_id=None, rev_id=None)

Split a single reversible reaction into two irreversible reactions, returns the original reversible reaction and bounds while deleting them from model.

R1: $A = B R1_fwd: A -> B R1_rev: B -> A$

- fba an instantiated CBMPy model object
- rid a valid reaction id
- fwd id [default=None] the new forward reaction id, defaults to rid fwd
- rev_id [default=None] the new forward reaction id, defaults to rid_rev

storeObj (*obj*, *filename*, *compress=False*)

Stores a Python *obj* as a serialised binary object in *filename*.dat

- *obj* a python object
- filename the base filename
- compress [False] use gzip compression not implemented

stringReplace(fbamod, old, new, target)

This is alpha stuff, target can be:

- · 'species'
- · 'reactions'
- · 'constraints'
- · 'objectives'
- 'all'

3.16 CBMPy: CBWrite module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBWrite.py 710 2020-04-27 14:22:34Z bgoli \$)

BuildHformatFluxBounds(fba, infinity_replace=None, use_rational=False)

Build and return a csio that contains the flux bounds in H format

- fba a PySCeS-CBM FBA object
- infinity replace [default=None] if defined this is the abs(value) of +-<infinity>

BuildLPConstraints(*fba*, *use rational=False*)

Build and return a csio that contains constraint constructed from the StoichiometeryLP object

- fba an fba model object which has a stoichiometry
- *use rational* write rational number output [default=False]

BuildLPConstraintsMath(*fba*, *use rational=False*)

Build and return a csio that contains the constaints in LP format Strict refers to dS/dt => 0 and dS/dt <= 0

BuildLPConstraintsRelaxed(fba)

Build and return a csio that contains the constaints in LP format Relaxed refers to $dS/dt \ge 0$

BuildLPConstraintsStrict(fba, use_rational=False)

Build and return a csio that contains the constaints in LP format Strict refers to dS/dt = 0

BuildLPFluxBounds(*fba*, *use_rational=False*)

Build and return a csio that contains the flux bounds in LP format

BuildLPUserConstraints(fba, use_rational=False)

Build and return a csio that contains constraint constructed from the StoichiometeryLP object

- fba an fba model object which has a stoichiometry
- *use_rational* write rational number output [default=False]

INFO: this method will be deprecated please update your scripts to use "writeFVAdata()"

WriteFVAtoCSV(id, fva, names, Dir=None, fbaObj=None)

INFO: this method will be deprecated please update your scripts to use "writeFVAtoCSV()"

INFO: this method will be deprecated please update your scripts to use "writeModelHFormatFBA2()"

INFO: this method will be deprecated please update your scripts to use "writeModelHFormatFBA2()"

WriteModelLP(fba, work_dir=None, fname=None, multisymb='', format='%s', use_rational=False, constraint mode=None, quiet=False)

INFO: this method will be deprecated please update your scripts to use "writeModelLP()"

INFO: this method will be deprecated please update your scripts to use "writeModelLPOld()"

WriteModelRaw(fba, work_dir=None)

INFO: this method will be deprecated please update your scripts to use "writeModelRaw()"

convertExcelToFloat(num)

Converts an Excel "number" to a float

• num a number

convertFloatToExcel(num, roundoff)

Converts a float to Excel compatible "number"

- num a number
- roundoff the number of roundoff digits for round()

exportModel(fba, fname=None, fmt='lp', work_dir=None, use_rational='both')

Export the FBA model in different formats:

- fba the FBA model
- fname [default=None] the exported filename if None then fba.getId() is used
- fmt [default='lp'] the export format can be one of: 'lp' (CPLEX), 'hformat' (Polyhedra), 'all' (both)
- *use_rational* [default='both'] if *all* or *hformat* is specified should hformat files be written using rational math or not. The default *both* is the legacy behaviour and writes both.

Note that 'hformat' ignores 'fname' and only uses fba.getId() this is a legacy behaviour

generateBGID(num, prefix)

Create a BGID generator, which is prefix><num> where perfix is two letters num is padded to 6 figures

- *num* the starting number
- prefix the two letter prefix

printFBASolution(fba, include_all=False)

Prints the FBA optimal solution to the screen.

- fba an FBA model object
- include_all include all variables

saveModel(model, filename, compress=False)

Saves the model to an SBML file using the lates SBML3 FBC version.

- model the CBMPy model
- filename the filename to write

writeCOBRASBML(fba, fname, directory=None)

Takes an FBA model object and writes it to file as a COBRA compatible:

- fba an fba model object
- fname the model will be written as XML to fname
- directory [default=None] if defined it is prepended to fname

Takes the resuls of a FluxVariabilityAnalysis method and writes it to a nice csv file. Note this method replaces the glpk/cplx_WriteFVAtoCSV methods. Data is output as a csv file with columns: FluxName, FVA_MIN, FVA_MAX, OPT_VAL, SPAN

- fvadata Flux Variability Analysis() FVA OUTPUT ARRAY
- names Flux Variability Analysis() FVA OUTPUT_NAMES
- fname filename_base for the CSV output
- work_dir [default=None] if set the output directory for the csv files

- roundec [default=None] an integer indicating at which decimal to round off output. Default is no rounding.
- scale min [default=False] normalise each flux such that that FVA MIN = 0.0
- appendfile [default=False] instead of opening a new file try and append the data
- info [default=None] a string added to the results as an extra column, useful with appendfile

writeFVAtoCSV(fvadata, names, fname, Dir=None, fbaObj=None)

Takes the resuls of a FluxVariabilityAnalysis method and writes it to a nice csv file. Note this method replaces the glpk/cplx_WriteFVAtoCSV methods.

- fvadata FluxVariabilityAnalysis() OUTPUT_ARRAY
- names FluxVariabilityAnalysis() OUTPUT_NAMES
- fname filename_base for the CSV output
- Dir [default=None] if set the output directory for the csv files
- fbaObj [default=None] if supplied adds extra model information into the output tables

 $\label{lem:writeMinDistanceLPwithCost} \textbf{(}\textit{fname}, \textit{fbas}, \textit{work_dir=None}, \textit{ignoreDistance=[]}, \textit{constraint_mode='strict'}\textbf{)}$

For backwards compatability only

Write an FBA-LP in polynomial H-Format file. This version has been replaced by *writeModelHFormatFBA2()* but is kept for backwards compatability.

- fba a PySCeS-CBM FBA object
- Work_dir [default=None] the output directory
- use_rational [default=false] use rational numbers in output (requires sympy)
- fullLP [default=True] include the default objective function as a maximization target
- format [default='%s'] the number format string
- infinity replace [default=None] if defined this is the abs(value) of +-<infinity>

Write an FBA-LP in polynomial H-Format file. This is an improved version of *WriteModelHFormatFBA()* which it replaces. Note that if a SymPy matrix is used as input then use_rational is automatically enabled.

- fba a PySCeS-CBM FBA object
- fname [default=None] the output filename, fba.getId() if not defined
- Work_dir [default=None] the output directory
- use_rational [default=false] use rational numbers in output (requires sympy)
- fullLP [default=True] include the default objective function as a maximization target
- format [default='%s'] the number format string
- infinity_replace [default=None] if defined this is the abs(value) of +-<infinity>

 $\textbf{writeModelInfoToFile} (\textit{fba}, \textit{fname}, \textit{Dir} = \textit{None}, \textit{separator} = \textit{'}, \textit{'}, \textit{only} _ \textit{exchange} = \textit{False}, \textit{met_type} = \textit{'all'})$

This function writes a CBModel to file

- fba an instance of an PySCeSCBM model
- fname the output filename

- Dir [default=None] use directory if not None
- separator [default=','] the column separator
- only_exchange [default=False] only output fluxes labelled as exchange reactions
- type [default='all'] only output certain type of species: 'all', 'boundary' or 'variable'

Writes an FBA object as an LP in CPLEX LP format

- fba an instantiated FBAmodel instance
- work_dir directory designated for output
- *fname* the file name [default=fba.getId()]
- *multisymb* the multiplication symbol (default: <space>)
- format the number format of the output
- *use_rational* output rational numbers [default=False]
- quiet [default=False] supress information messages

Writes a fba as an LP/LPT

- fba an instantiated FBAmodel instance
- work_dir directory designated for output
- *multisymb* the multiplication symbol (default: <space>)
- lpt the file format (default: True for lpt) or False for lp

writeModelRaw(fba, work_dir=None)

Writes a fba (actually just dumps it) to a text file.

- fba an instantiated FBAmodel instance
- work dir directory designated for output

Write a model in SBML and Excel format to a COMBINE archive using the following information:

- mod a model object
- fname the output base filename, archive will be <fname>.zip
- directory [default=None] created the combine archive 'directory'
- *sbmlname* [default='None'] If *sbmlname* is defined then SBML file is <sbmlname>.xml otherwise sbml will be <fname>.xml.
- with Excel [default=True] include a human readable Excel spreadsheet version of the model
- vc_given [default='CBMPy'] first name
- vc_family [default='Software'] family name
- vc_email [default='None'] email

- vc_org [default='None'] organisation
- add_cbmpy_annot [default=True] add CBMPy KeyValueData annotation. Replaces <notes>
- add_cobra_annot [default=True] add COBRA <notes> annotation

writeModelToExcel97(fba, filename, roundoff=6)

Exports the model as an Excel 97 spreadsheet

- fba a CBMPy model instance
- filename the filename of the workbook
- roundoff [default=6] the number of digits to round off to

writeOptimalSolution(fba, fname, Dir=None, separator=',', only_exchange=False)

This function writes the optimal solution to file

- fba an instance of an PySCeSCBM model
- fname the output filename
- Dir [default=None] use current directory if not None
- separator [default=','] the column separator
- only_exchange [default=False] only output fluxes labelled as exchange reactions

writeProteinCostToCSV(fba, fname)

Writes the protein costs 'CBM_PEPTIDE_COST' annotation to acsv file.

- fba an instantiated FBA object
- fname the exported file name

writeReactionInfoToFile(fba, fname, Dir=None, separator=',', only_exchange=False)

This function writes a CBModel to file

- fba an instance of an PySCeSCBM model
- fname the output filename
- Dir [default=None] use directory if not None
- *separator* [default=','] the column separator
- only_exchange [default=False] only output fluxes labelled as exchange reactions

writeSBML2FBA(fba, fname, directory=None, sbml_level_version=None)

Takes an FBA model object and writes it to file as SBML L2 with FBA annotations. Note if you want to write BiGG/FAME style annotations then you must use *sbml_level_version=(2,1)*

- fba an fba model object
- fname the model will be written as XML to fname
- sbml_level_version [default=None] a tuple containing the SBML level and version e.g. (2,1)

This is a utility wrapper for the function CBXML.sbml_writeSBML2FBA

Takes an FBA model object and writes it to file as SBML L3 FBC:

- fba an fba model object
- fname the model will be written as XML to fname

- directory [default=None] if defined it is prepended to fname
- *gpr_from_annot* [default=True] if enabled will attempt to add the gene protein associations from the annotations if no gene protein association objects exist
- add_groups [default=True] add SBML3 groups (if supported by libSBML)
- add_cbmpy_annot [default=True] add CBMPy KeyValueData annotation. Replaces <notes>
- add cobra annot [default=True] add COBRA <notes> annotation
- · xoptions extended options
 - fbc_version [default=1] write SBML3FBC using version 1 (2013) or version 2 (2015)
 - validate [default=False] validate the output SBML file
 - compress_bounds [default=False] try compress output flux bound parameters
 - zip_model [default=False] compress the model using PKZIP encoding
 - return_model_string [default=False] return the SBML XML file as a string

```
writeSBML3FBCV2(fba, fname, directory=None, gpr_from_annot=False, add_groups=True, add_cbmpy_annot=True, add_cobra_annot=False, validate=False, compress_bounds=False, zip_model=False, return_model_string=False)
```

Takes an FBA model object and writes it to file as SBML L3 FBCv2:

- fba an fba model object
- fname the model will be written as XML to fname
- directory [default=None] if defined it is prepended to fname
- *gpr_from_annot* [default=False] if enabled will attempt to add the gene protein associations from the annotations
- add_groups [default=True] add SBML3 groups (if supported by libSBML)
- add_cbmpy_annot [default=True] add CBMPy KeyValueData annotation. Replaces <notes>
- add_cobra_annot [default=False] add COBRA <notes> annotation
- validate [default=False] validate the output SBML file
- compress_bounds [default=True] try compress output flux bound parameters
- zip_model [default=False] compress the model using ZIP encoding
- return_model_string [default=False] return the SBML XML file as a string

```
writeSBML3FBCV3(fba, fname, directory=None, gpr_from_annot=False, add_groups=True, add_cbmpy_annot=True, add_cobra_annot=False, validate=False, compress_bounds=False, zip_model=False, return_model_string=False)
```

Takes an FBA model object and writes it to file as SBML L3 FBCv3:

- fba an fba model object
- fname the model will be written as XML to fname
- directory [default=None] if defined it is prepended to fname
- gpr_from_annot [default=False] if enabled will attempt to add the gene protein associations from the annotations
- add groups [default=True] add SBML3 groups (if supported by libSBML)
- add_cbmpy_annot [default=True] add CBMPy KeyValueData annotation. Replaces <notes>

- add_cobra_annot [default=False] add COBRA <notes> annotation
- validate [default=False] validate the output SBML file
- compress_bounds [default=True] try compress output flux bound parameters
- zip_model [default=False] compress the model using ZIP encoding
- return_model_string [default=False] return the SBML XML file as a string

writeSensitivitiesToCSV(sensitivities, fname)

Write out a sensitivity report using the objective sensitivities and bound sensitivity dictionaries created by e.g. cplx_getSensitivities().

- sensitivity tuple containing
- obj_sens dictionary of objective coefficient sensitivities (per flux)
- rhs_sens dictionary of constraint rhs sensitivities (per constraint)
- bound_sens dictionary of bound sensitivities (per flux)
- fname output filename e.g. fname.csv

writeSolutions(fname, sols=[], sep=',', extra_output=None, fba=None)

Write 2 or more solutions where a solution is a dictionary of flux:value pairs:

- fname the export filename
- sols a list of dictionaries containing flux:value pairs (e.g. output by cmod.getReactionValues())
- sep [default=','] the column separator
- extra_output [default=None] add detailed information to output e.g. reaction names by giving a CBModel object as an argument to extra_output.
- fba an fba model that canbe used for extra_output

writeSpeciesInfoToFile(fba, fname, Dir=None, separator=',', met_type='all')

This function writes a CBModel to file

- fba an instance of an PySCeSCBM model
- fname the output filename
- Dir [default=None] use directory if not None
- separator [default=','] the column separator
- met_type [default='all'] only output certain type of species: 'all','boundary' or 'variable'

Write an FBA-LP in polynomial H-Format file. This is an improved version of *WriteModelHFormatFBA()* which it replaces but is kept for backwards compatability.

- fba a PySCeS-CBM FBA object
- fname [default=None] the output filename, fba.getId() if not defined
- Work_dir [default=None] the output directory
- use_rational [default=false] use rational numbers in output (requires sympy)
- fullLP [default=True] include the default objective function as a maximization target

- format [default='%s'] the number format string
- infinity replace [default=None] if defined this is the abs(value) of +-<infinity>

3.17 CBMPy: CBWx module

PySCeS Constraint Based Modelling (http://cbmpy.sourceforge.net) Copyright (C) 2009-2024 Brett G. Olivier, VU University Amsterdam, Amsterdam, The Netherlands

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBWx.py 710 2020-04-27 14:22:34Z bgoli \$)

3.18 CBMPy: CBXML module

PySCeS Constraint Based Modelling (http://cbmpy.sourceforge.net) Copyright (C) 2009-2024 Brett G. Olivier, VU University Amsterdam, Amsterdam, The Netherlands

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: CBXML.py 710 2020-04-27 14:22:34Z bgoli \$)

class MLStripper

Class for stripping a string of HTML/XML used from: http://stackoverflow.com/questions/753052/strip-html-from-strings-in-python

```
SBML_NS = [('http://www.sbml.org/sbml/level3/version1/fbc/version3', 'L3V1FBC3'),
('http://www.sbml.org/sbml/level3/version1/fbc/version2', 'L3V1FBC2'),
('http://www.sbml.org/sbml/level3/version1/fbc/version1', 'L3V1FBC1'),
('http://www.sbml.org/sbml/level3/version2/fbc/version2', 'L3V2FBC2'),
('http://www.sbml.org/sbml/level3/version2/fbc/version1', 'L3V2FBC1'),
('http://www.sbml.org/sbml/level3/version2/core', 'L3V2core'),
('http://www.sbml.org/sbml/level3/version1/core', 'L3V1core'),
('http://www.sbml.org/sbml/level2/version4', 'L2'), ('http://www.sbml.org/sbml/level2',
'L2')]
```

print libsbml.BQB_ENCODES , 8 # "encodes", print libsbml.BQB_HAS_PART , 1 # "hasPart", print libsbml.BQB_HAS_PROPERTY , 10 # "hasProperty", print libsbml.BQB_HAS_VERSION , 4 # "hasVersion", print libsbml.BQB_IS , 0 # "isA", print libsbml.BQB_IS_DESCRIBED_BY , 6 # "isDescribedBy", print libsbml.BQB_IS_ENCODED_BY , 7 # "isEncodedBy", print libsbml.BQB_IS_HOMOLOG_TO , 5 # "isHomologTo", print libsbml.BQB_IS_PART_OF , 2 # "isPartOf", print libsbml.BQB_IS_PROPERTY_OF , 11 # "isPropertyOf", print libsbml.BQB_IS_VERSION_OF , 3 # "isVersionOf", print libsbml.BQB_OCCURS_IN , 9 # "occursIn", print libsbml.BQB_UNKNOWN , 12 # None

print libsbml.BQM_IS , 0 # None print libsbml.BQM_IS_DERIVED_FROM , 2 # None print libsbml.BQM_IS_DESCRIBED_BY , 1 # None print libsbml.BQM_UNKNOWN , 3 # None

sbml_convertCOBRASBMLtoFBC(fname, outname=None, work_dir=None, output_dir=None)

Read in a COBRA SBML Level 2 file and return the name of the created SBML Level 3 with FBC file that is created in the output directory

- fname is the filename
- outname the name of the output file. If not specified then <filename>.13fbc.xml is used as default
- work_dir [default=None] is the working directory
- output_dir [default=None] is the output directory (default is work_dir)

This method is based on code from libSBML (http://sbml.org) in the file "convertCobra.py" written by Frank T. Bergmann.

sbml_convertSBML3FBCToCOBRA(fname, outname=None, work_dir=None, output_dir=None)

Read in a SBML Level 3 file and return the name of the created COBRA file that is created in the output directory

- fname is the filename
- outname the name of the output file. If not specified then <filename>.cobra.xml is used as default
- work_dir [default=None] is the working directory
- output_dir [default=None] is the output directory (default is work_dir)

This method is based on code from libSBML (http://sbml.org) in the file "convertFbcToCobra.py" written by Frank T. Bergmann.

sbml_createAssociationFromAST(node, out)

Converts a GPR string '((g1 and g2) or g3)' to an association via a Python AST. In future I will get rid of all the string elements and work only with associations and AST's.

- *node* a Python AST note (e.g. body)
- out a new shiny FBC V2 GeneProductAssociation

sbml_createAssociationFromTreeV2(tree, out)

Converts a GPR tree to an association

- tree a GPR dict tree
- out a new shiny FBC V2 GeneProductAssociation

sbml_createModelL2(*fba*, *level=2*, *version=1*)

Create an SBML model and document:

- fba a PySCeSCBM model instance
- level always 2
- · version always 1

and returns:

model an SBML model

$\textbf{sbml_exportSBML2FBAModel} (document, filename, directory = None, return_doc = False, remove_note_body = False)$

Writes an SBML model object to file. Note this is an internal SBML method use *sbml_writeSBML2FBA()* to write an FBA model:

- model a libSBML model instance
- filename the output filename
- directory [default=None] by default use filename otherwise join, <dir><filename>
- return_doc [default=False] return the SBML document used to write the XML

sbml_fileFindVersion(f)

Try and find the SBML version and FBC support

• f the SBML file

sbml_fileValidate(f, level='normal')

Validate an SBML file and model

- f the SBML file
- level [default='normal'] the level of validation "normal" or "full"

sbml_getCVterms(sb, model=False)

Get the MIRIAM compliant CV terms and return a MIRIAMAnnotation or None

- sb a libSBML SBase derived object
- model is this a BQmodel term

sbml_getGPRasDictFBCv1(node, out)

Converts a GPR string '((g1 and g2) or g3)' to a gprDict which is returned

- node a Python AST note (e.g. ast.parse(gprstring).body[0])
- out a new dictionary that will be be created in place

sbml_getGPRasDictFBCv2(association, out, cntr)

Walk through an SBML L3FBCV2 gene protein association and return a dictionary/tree representation

sbml_getGeneRefs(association, out)

Walk through a gene association and extract GeneRefs inspired by Frank

sbml_getNotes(obj)

Returns the SBML objects notes

• obj an SBML object

sbml_readCOBRANote(s)

Parses a COBRA style note from a XML string

• s an XML string

sbml_readCOBRASBML(fname, work_dir=None, return_sbml_model=False, delete_intermediate=False, fake_boundary_species_search=False, output_dir=None, speciesAnnotationFix=True,

fake_boundary_species_search=False, output_dir=None, speciesAnnotationFix=True.skip_genes=False)

Read in a COBRA format SBML Level 2 file with FBA annotation where and return either a CBM model object or a (cbm_mod, sbml_mod) pair if return_sbml_model=True

- fname is the filename
- work dir is the working directory
- return_sbml_model [default=False] return a a (cbm_mod, sbml_mod) pair
- delete_intermediate [default=False] delete the intermediate SBML Level 3 FBC file
- fake_boundary_species_search [default=False] after looking for the boundary_condition of a species search for overloaded id's <id> b
- *output_dir* [default=None] the directory to output the intermediate SBML L3 files (if generated) default to input directory
- speciesAnnotationFix [default=True]
- skip_genes [default=False] convert GPR associations

sbml_readFBCv3KeyValuePairs(fbcp)

Reads FBCv3 KeyValue pair annotation and returns a dictionary of key:value pairs

• fbcp an FBC plugin

sbml_readKeyValueDataAnnotation(annotations)

Reads KeyValueData annotation (http://pysces.sourceforge.net/KeyValueData) and returns a dictionary of key:value pairs

sbml_readSBML2FBA(fname, work_dir=None, return_sbml_model=False, fake_boundary_species_search=False)

Read in an SBML Level 2 file with FBA annotation where and return either a CBM model object or a (cbm_mod, sbml_mod) pair if return_sbml_model=True

- fname is the filename
- work_dir is the working directory (only used if not None)
- return_sbml_model [default=False] return a a (cbm_mod, sbml_mod) pair
- fake_boundary_species_search [default=False] after looking for the boundary_condition of a species search for overloaded id's <id> b

sbml_readSBML3FBC(fname, work_dir=None, return_sbml_model=False, xoptions={})

Read in an SBML Level 3 file with FBC annotation where and return either a CBM model object or a (cbm_mod, sbml_mod) pair if return_sbml_model=True

- fname is the filename
- work dir is the working directory
- return sbml model [default=False] return a a (cbm mod, sbml mod) pair
- *xoptions* special load options enable with option = True
 - nogenes do not load/process genes
 - noannot do not load/process any annotations
 - validate validate model and display errors and warnings before loading
 - readcobra read the cobra annotation
 - read_model_string [default=False] read the model from a string (instead of a filename) containing an SBML document
 - nmatrix_type [default='normal'] define the type of stoichiometrich matrix to be built
 - model extension class extend CBModel class with new class (experimental, Python 3 only)

- model_metaclass add a custom metaclass to CBModel (experimental, Python 3 only)
 - * 'numpy' dense numpy array (best performance)
 - * 'scipy_csr' scipy sparse matrix (lower performance, low memory)
 - * 'sympy' a sympy rational matrix (low performance, high memory, cast to dense to analyse)
 - * None do not build matrix

sbml_setAnnotationsL3Fbc(cbmo, sbmlo)

Add CBMPy Fbase annotations to an SBML object, MIRIAM, SBO, Notes. Should be called last when creating SBML objects.

- cbmo the CBMPy object
- · sbmlo SBML object

Note: this function should be used for new code, old code still needs to be refactored.

sbml_setCVterms(sb, uridict, model=False)

Add MIRIAM compliant CV terms to a sbml object from a CBM object

- sb a libSBML SBase derived object
- uridict a dictionary of uri's as produced by getAllMIRIAMUris()
- model is this a BQmodel term [deprecated attribute, ignored and autodetected]

sbml_setCompartmentsL3(model, fba)

Sets the model compartments.

- model a libSBML model instance
- fba a PySCeSCBM model instance

$sbml_setDescription(model, fba)$

Sets the model description as a <note> containing txt in an HTML paragraph on the model object.

- model a libSBML model instance
- fba a PySCeSCBM model instance

sbml_setFBCv3KeyValuePairs(fbcp, kv_pairs)

Adds key value pairs to any FBCv3 SBase derived object

- fbcp an FBCv3 plugin
- kv_pairs a dictionary of CBMPy Key Value pairs (format will be upgraded over next few versions)

sbml_setGroupsL3(cs, fba)

add groups to the SBML model

- cs a CBMLtoSBML instance
- fba a CBMPy model instance

sbml_setNotes3(obj, s)

Formats the CBMPy notes as an SBML note and adds it to the SBMl object

- *obj* an SBML object
- s a string that should be added as a note

sbml_setParametersL3Fbc(*fbcmod*, *add_cbmpy_anno=True*, *fbc_version=2*)

Add non fluxbound related parameters to the model

- fbcmod a CBM2SBML instance
- add cbmpy anno [default=True] add CBMPy KeyValueData annotation.

sbml_setReactionsL2(model, fba, return_dict=False)

Add the FBA instance reactions to the SBML model

- model an SBML model instance
- fba a PySCeSCBM model instance
- return_dict [default=False] if True do not add reactions to SBML document instead return a dictionary description of the reactions

sbml_setReactionsL3Fbc(fbcmod, return_dict=False, add_cobra_anno=False, add_cbmpy_anno=True, fbc_version=2)

Add the FBA instance reactions to the SBML model

- fbcmod a CBM2SBML instance
- return_dict [default=False] if True do not add reactions to SBML document instead return a dictionary description of the reactions
- add_cbmpy_anno [default=True] add CBMPy KeyValueData annotation. Replaces <notes>
- add_cobra_anno [default=False] add COBRA <notes> annotation
- fbc version [default=2] writes either FBC v1 (2013) or v2 (2015) or v3 (2023)

sbml_setSpeciesL2(model, fba, return_dicts=False)

Add the species definitions to the SBML object:

- model [default="] a libSBML model instance or can be None if return_dicts == True
- fba a PySCeSCBM model instance
- return_dicts [default=False] only returns the compartment and species dictionaries without updated the SBML

returns:

• compartments a dictionary of compartments (except when give return_dicts argument)

sbml_setSpeciesL3(model, fba, return_dicts=False, add_cobra_anno=False, add_cbmpy_anno=True, substance_units=True, fbc_version=2)

Add the species definitions to the SBML object:

- *model* and SBML model instance or can be None if *return_dicts* == True
- fba a PySCeSCBM model instance
- return_dicts [default=False] only returns the compartment and species dictionaries without updating the SBML
- add_cbmpy_anno [default=True] add CBMPy KeyValueData annotation. Replaces <notes>
- add cobra anno [default=False] add COBRA <notes> annotation
- substance_units [default=True] defines the species in amounts rather than concentrations (necessary for default mmol/gdw.h)
- fbc version [default=2] the FBC version to use

returns:

• compartments a dictionary of compartments (except when given return_dicts argument)

sbml_setUnits(*model*, *units=None*, *give_default=False*, *L3=True*)

Adds units to the model:

- model a libSBML model instance
- units [default=None] a dictionary of units, if None default units are used
- give_default [default=False] if true method returns the default unit dictionary
- L3 [default=True] use the L3 defaults

sbml_setValidationOptions(D, level)

set the validation level of an SBML document

- D an SBML document
- level the level of consistency check can be either one of:
- · 'normal' basic id checking only
- · 'full' all checks enabled
- None disable all validation

sbml_validateDocument(D, fullmsg=False, docread=False)

Validates and SBML document returns three dictionaries, errors, warnings, other and a boolean indicating an invalid document:

- D and SBML document
- fullmsg [default=False] optionally display the full error message

sbml_writeAnnotationsAsCOBRANote(annotations)

Writes the annotations dictionary as a COBRA compatible SBML <note>

sbml_writeCOBRASBML(fba, fname, directory=None)

Takes an FBA model object and writes it to file as a COBRA compatible:

- fba an fba model object
- fname the model will be written as XML to fname
- directory [default=None] if defined it is prepended to fname

sbml_writeKeyValueDataAnnotation(annotations)

Writes the key:value annotations as a KeyValueData annotation (http://pysces.sourceforge.net/KeyValueData)

sbml_writeSBML2FBA(fba, fname, directory=None, sbml_level_version=None)

Takes an FBA model object and writes it to file as SBML L3 FBA:

- fba an fba model object
- fname the model will be written as XML to fname
- directory [default=None] if defined it is prepended to fname
- sbml_level_version [default=None] a tuple containing the SBML level and version e.g. (2,4) (ignored)

sbml_writeSBML3FBC(fba, fname, directory=None, sbml_level_version=(3, 1), autofix=True, return_fbc=False, gpr_from_annot=False, add_groups=False, add_cbmpy_annot=True, add_cobra_annot=False, xoptions={})

Takes an FBA model object and writes it to file as SBML L3 FBC:

- fba an fba model object
- fname the model will be written as XML to fname
- directory [default=None] if defined it is prepended to fname
- sbml_level_version [default=(3,1)] a tuple containing the SBML level and version e.g. (3,1)
- autofix convert <> to <=>=
- return_fbc return the FBC converter instance
- *gpr_from_annot* [default=False] if enabled will attempt to add the gene protein associations from the annotations if no gene protein association objects exist
- add_cbmpy_annot [default=True] add CBMPy KeyValueData annotation. Replaces <notes>
- add_cobra_annot [default=True] add COBRA <notes> annotation
- xoptions extended options
 - fbc_version [default=2] write SBML3FBC using version 1 (2013) or version 2 (2015) oe version (2023)
 - validate [default=False] validate the output SBML file
 - compress_bounds [default=False] try compress output flux bound parameters
 - zip_model [default=False] compress the model using ZIP encoding
 - return_model_string [default=False] return the SBML XML file as a string

setCBSBOterm(sbo, obj)

Given an SBOterm from libSBML, add it to a CBMPy object

- sbo the sbo term string
- · obj the CBMPy Fbase derived object

with_metaclass(meta, *bases)

Create a base class with a metaclass. Usage is: NewClass(with_metaclass(MetaClass, BaseClass*)

xml_addSBML2FBAFluxBound(document, rid, operator, value, fbid=None)

Adds an SBML3FBA flux bound to the document:

- document a minidom XML document created by xml_createSBML2FBADoc
- rid the reaction id
- operator one of ['greater', 'greaterEqual', 'less', 'lessEqual', 'equal', '>','>=','<','<=','=']
- value a float which will be cast to a string using str(value)
- fbid the flux bound id, autogenerated by default

xml_addSBML2FBAObjective(document, objective, active=True)

Adds an objective element to the documents listOfObjectives and sets the active attribute:

- document a minidom XML document created by xml_createSBML2FBADoc
- objective a minidom XML objective element created with xml_createSBML2FBAObjective
- active [default=True] a boolean flag specifiying whether this objective is active

xml_createListOfFluxObjectives(document, l)

Create a list of fluxObjectives to add to an Objective:

- document a minidom XML document created by xml_createSBML2FBADoc
- fluxobjs a list of (rid, coefficient) tuples

xml_createSBML2FBADoc()

Create a 'document' to store the SBML2FBA annotation, returns:

• DOC a minidom document

xml_createSBML2FBAObjective(document, oid, sense, fluxObjectives)

Create a list of fluxObjectives to add to an Objective:

- document a minidom XML document created by xml_createSBML2FBADoc
- oid the objective id
- sense a string containing the objective sense either: maximize or minimize
- fluxObjectives a list of (rid, coefficient) tuples

xml_getSBML2FBAannotation(fba, fname=None)

Takes an FBA model object and returns the SBML3FBA annotation as an XML string:

- *fba* an fba model object
- fname [default=None] if supplied the XML will be written to file fname

xml_stripTags(html)

Strip a string of HTML/XML, returns a string

• html the string containing html

xml_viewSBML2FBAXML(document, fname=None)

Print a minidom XML document to screen or file, arguments:

- · document a minidom XML document
- fname [default=None] by default print to screen or write to file fname

3.19 CBMPy: MultiCoreFVA module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: multicorefva.py 710 2020-04-27 14:22:34Z bgoli \$)

3.20 CBMPy: MultiCoreEnvFVA module

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Author: Brett G. Olivier PhD Contact developers: https://github.com/SystemsBioinformatics/cbmpy/issues Last edit: \$Author: bgoli \$ (\$Id: _multicoreenvfva.py 710 2020-04-27 14:22:34Z bgoli \$)

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