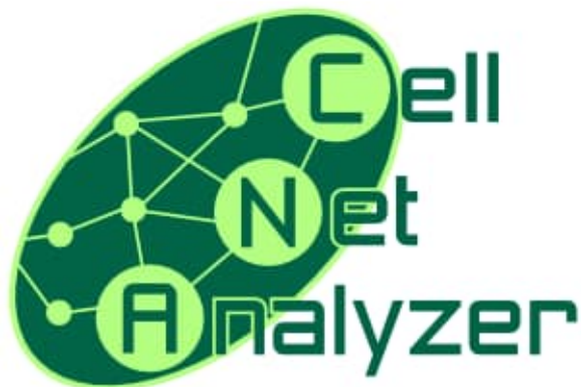


Manual

CellNetAnalyzer



Version 2025.2

For an online version see:

http://www.mpi-magdeburg.mpg.de/projects/cna/manual_cellnetanalyzer.pdf

This is the user manual of *CellNetAnalyzer* (CNA). CNA is a MATLAB toolbox for analyzing structure and function of biological networks on the basis of topological, stoichiometric, qualitative (logical) and semi-quantitative modeling approaches requiring no or only few parameters. In particular, metabolic networks (mass-flow) can be studied based on stoichiometric and constraint-based modeling approaches whereas signaling and regulatory networks (signal-flow networks) can be explored by qualitative and semi-quantitative modeling approaches (Boolean/logical models, interaction graphs, logic-based ODEs). CNA can be used via command-line based operations (supported by an application programming interface; API) or via a graphical user interface with embedded network visualizations.

The theoretical foundations of the methods supported by CNA are not described herein; for a detailed description please see the recommended references at the end of this manual.

For academic use, *CellNetAnalyzer* is freely available (from 2019 on with its source code). However, the user has to agree with our licensing conditions when downloading CNA. The academic licence can be found in LICENSE.txt in CNA's root directory and on the website <https://www.mpi-magdeburg.mpg.de/projects/cna/license.html>. Interested commercial users should send an email to: cellnetanalyzer@mpi-magdeburg.mpg.de

Requirements for using *CellNetAnalyzer* are:

- We suggest MATLAB version 8.2 (Release 2013b) or higher. Most functions also work with Octave 5.0 or higher.
- Some functions require an LP or (M)ILP solver. CNA supports the optimization toolbox of MATLAB (linprog) as well as [GLPKMEX](#) (which is part of the CNA distribution), CPLEX and Gurobi.

More information and a how-to tutorial on CNA can be found at:

<https://www2.mpi-magdeburg.mpg.de/projects/cna/cna.html>

CellNetAnalyzer has been developed at the Magdeburg Max Planck Institute by Steffen Klamt (since 2000) and Axel von Kamp (since 2007) as main developers and by contributions of several other colleagues listed in the CONTRIBUTORS.txt file in CNA's root directory.

CellNetAnalyzer is still (always ...) under construction. We appreciate any comments or suggestions for improvements and we are greatly interested in your feedback which you can give at this user survey: https://docs.google.com/forms/d/e/1FAIpQLSeD1LA6a1eMqv2-bZPCWNFUQu5DSbbljXsE_fO9z5qt7EIXWA/viewform

For questions and bug reports please contact us, either via the CNA user forum

<https://groups.google.com/forum/#!forum/cellnetanalyzer-user-forum> or via:

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Thank you for using *CellNetAnalyzer*!

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0. Quick start and general hints

0.1 Installation

After unzipping the downloaded file at an appropriate place, *CellNetAnalyzer* (in the following abbreviated by CNA) consists of a main directory (here indicated as “CellNetAnalyzer“), the “code” directory with the MATLAB functions and libraries, and several subdirectories, each related to one network project:

```
../CellNetAnalyzer - main directory of the CNA
../CellNetAnalyzer - directory containing the manual and several related papers
../CellNetAnalyzer/code - directory containing MATLAB functions and libraries
../CellNetAnalyzer/ECC2 - directory with specific files for the ECC2 network project (core
model of the central metabolism of E. coli)
../CellNetAnalyzer/SmallExample - specific files for the network project “SmallExample“
../CellNetAnalyzer/<.....> - other network projects
```

Network projects created by the user will normally be added as new subdirectories in CNA’s main directory by default.

0.2 Starting and closing CellNetAnalyzer

For starting CNA proceed as follows:

- Before starting CNA you should **update the paths in `configure_cplex_paths.m` and `configure_gurobi_paths.m`** (see the first lines in these files; these paths need to be set when you want to use the CPLEX / Gurobi solver) as well as in “**`javalibrarypath.txt`**” (required when the JAVA interface of CPLEX is used) according to your local system (see section 0.5)!
- Start MATLAB. **The best way** to start MATLAB when using CellNetAnalyzer is to start MATLAB directly from the main directory of CNA (e.g. via the command line in Linux) or to make CNA’s main directory the startup directory of MATLAB before starting MATLAB (e.g. under Windows). **Alternatively** you can navigate to the CellNetAnalyzer’s main directory or add it to the MATLAB path (right click -> add to path) once MATLAB is running. However, since the “`javalibrarypath.txt`“ file in CNA’s main directory is only read on MATLAB startup and required for functions that use the CPLEX-Java API, these functions would not work in this case [see also section 0.5!].
- Enter “**`startcna`**“. When calling `startcna` without an argument then the GUI will be initialized and the “Project Manager” comes up. Mass-flow (stoichiometric/metabolic) networks are indicated by “(M)” and signaling networks by “(S)”. Select a network project and click on „Start“. [Note for advanced users: If you want to load CNA without GUI (and only the API functions) you may start it with `startcna(1)`; see section 7.]
- The selected network will be loaded resulting (after a while) in the display of the associated network map(s) in separate windows. Each network map has different menu items (see Fig. 2), from which certain actions and calculations can be started (see section 3.3 and 4.3). Some frequently used actions (e.g. load/clear/reset scenario) can be accessed from CNA’s toolbar.

Note: CNA seeks to display the network maps in full resolution. If this is not possible (e.g. due to a low resolution of the monitor, recommended is 1600x1200), the network map might be resized to a smaller resolution often leading to a badly readable display. In this case try to resize the window of the network map or use the menu command “Set original network map size” (see section 3.3). If this does not succeed, you can either use CNA’s zoom facilities (section 3.3. and 4.3) or, for a first use of the CNA, you can try to start a smaller network (e.g. the “Small Network Example”). These issues are further discussed in section 2.

- After loading a network project, the default values are displayed in the text boxes on the network map(s). The sign “#” indicates an unknown (undefined) value. You can now perform calculations by choosing the respective menu functions. For example, when a mass-flow network has been loaded, your first action could be to determine a flux distribution resulting from some given values. For this purpose, write values into some text boxes (or use the default values) and choose via the menu “Analysis/Metabolic flux analysis/Flux analysis”. The calculable / balanceable rates that follow from the given values are displayed in the text boxes. Also take a look at the messages given by CNA in the command window during and after computations. After calculating a flux distribution (or anything else displayed in the text boxes) you can “Reset last values” or “Set default values” or “Clear all values” using the respective menu entries (or toolbar icons).
- A network session can be finished by closing any of the network maps. CNA will ask you whether you want to save your changes and whether the network variable is to be cleared from MATLAB’s workspace. Note that you may open several projects in parallel (even the same project can be loaded in several copies).

For a quick start you may also read the CNA tutorial which can be downloaded from the following web site: http://www.mpi-magdeburg.mpg.de/projects/cna/CNA_tutorial.pdf.

0.3. Hints for using CellNetAnalyzer with MATLAB under WINDOWS

CNA has been programmed with MATLAB under LINUX as operating system (KDE desktop). Thus, the display of maps, text boxes, control panels, message boxes and so on are checked on this platform. When using MATLAB under WINDOWS, the calculations work also fine but, unfortunately, the appearance of these graphical user interfaces differs sometimes pretty much from LINUX and they do then not look very well. We will check regularly how CNA appears under WINDOWS and try to avoid large discrepancies (but some might nevertheless occur).

Special hints for WINDOWS user are:

- It can happen that the cursor is not visible when clicking in the text boxes. Nevertheless, you will be able to enter values.
- After entering values in text boxes **you have to press the ENTER-key**, otherwise the new values you have entered might not be recognized by MATLAB.

CNA has been tested for different MATLAB versions. Unfortunately, sometimes it happens that the text boxes and the font size of the text appear differently. In such a case, you may try to adjust text box size and font size properly (by editing the project properties; see section 2).

0.4 MEX files

A few procedures of *CellNetAnalyzer* use (optionally) MEX files which can be recognized by their extensions “mexglx” (Linux 32-bit), mexa64 (Linux 64-bit), mexw32 (WINDOWS 32-bit), mexw64 (WINDOWS 64-bit) or “mexmaci” (Intel Mac). These files were written in C/C++ (see folder CNA/code/mexfiles) and contain an interface to MATLAB. The source files must be compiled with MATLAB with a suitable C-compiler. There are three groups of MEX-files used in CNA:

- calculation of elementary modes with CNA’s algorithm
- elementary-modes calculation with METATOOL
- linear optimization via GLPK/GLPKMEX.

The source code of MEX files for CNA’s elementary-modes-calculation procedure is located in directory “code/mexfiles” and the provided function “elmodes_compile.m” can be used to start the compilation process. Note that this can only be done if a compiler for MATLAB was properly configured (see MEX setup). After successful compilation, copy the created executable MEX files into the “code/lib” directory.

Note that the source code for METATOOL- and GLPK-MEX-files is not distributed with CNA. Instead we provide some available MEX-versions for 32-bit and 64-bit editions of MATLAB under WINDOWS and Linux. If you want to run these files on other systems please download the respective source code, compile them by yourself and put the resulting files in the code/lib directory.

In any case, if a compilation is not possible or if problems arise when using MEX files, the respective computations should be conducted with available alternative (MEX-independent) algorithms. Elementary-modes calculation by CNA’s algorithm version or by METATOOL can be circumvented by using the contained EFMTool (via Java; developed by Marco Terzer).

Links:

GLPK: <http://www.gnu.org/software/glpk/>

GLPKMEX: <http://glpkmex.sourceforge.net/>

EFMTool: <http://www.csb.ethz.ch/tools/efmtool>

METATOOL: <http://pinguin.biologie.uni-jena.de/bioinformatik/networks/>

IMPORTANT: some JAVA functions may sometimes deliver “strange” warnings (e.g., while using the *odefy* plugin or when *efmtool*’s rational arithmetic network compression routine is used the first time within a MATLAB session). Please ignore these messages (as long as the functions continue to work and do not stop with a true error message).

0.5 LP and MILP solvers

Some CNA functions require an LP or/and MILP solver. Regarding LP solvers, CNA supports the MATLAB optimization toolbox (function *linprog*), GLPK (via the MEX interface – see above; function *glpk*), the CPLEX MATLAB interface (function *cplexlp*) and Gurobi (function *gurobi*). Likewise, for MILP calculations, the MILP solver functionalities of MATLAB’s *intlinprog*, GLPK, CPLEX and Gurobi are supported.

Generally, for analyzing larger networks, we recommend to use CPLEX or Gurobi (if these two

solvers are not available then GLPK is preferred). The respective solvers must have been installed properly and then be coupled with CNA as described in the following:

1) MATLAB's `linprog` and `intlinprog`: check that the Optimization toolbox of MATLAB is available in your MATLAB installation. If this is the case, these solvers will directly work with CNA and nothing else needs to be done.

2) GLPK: Executable `GLPKMEX` files are included for some operating systems (Linux, Windows; see section 0.4) and should then directly work without further settings.

3.) Gurobi: Either purchase a Gurobi license or, if you are eligible (academic users), you may download Gurobi from:

<https://www.gurobi.com/academia/academic-program-and-licenses/>

Please set afterwards the respective paths for Gurobi and its license file in the `configure_gurobi_paths.m` file.

4) CPLEX: Either purchase a CPLEX license or, if you are eligible (academic user), you may download CPLEX from:

<https://community.ibm.com/community/user/datascience/blogs/xavier-nodet1/2020/07/09/cplex-free-for-students>

Please set afterwards the `cplex_root_path` in the `configure_cplex_paths` file.

For some API functions (CNAcomputeOptMDFpathway, CPLEX-based calculation of MIS, CNA(reg)MCSEnumerator and optionally for CNAMCSEnumerator2), also the Java CPLEX interfaces need to be installed and properly configured. For those API functions, the MATLAB JVM needs to have the CPLEX shared library on its library path which must be set up before (!) starting MATLAB. This is achieved by adding the path `CPLEX_ROOT/cplex/bin/OS/` (with OS being the designation of your system's binary executable format) to MATLAB's `javalibrarypath.txt` configuration file. The CNA main directory contains an example file for setting the shared library paths: `javalibrary_example.txt`. Adapt it to your paths, then rename it to `javalibrarypath.txt` and put it in the directory where you start MATLAB. Finally, you possibly also need to modify the `java.opts` file (in the CNA main directory) for specifying the heap space of your JAVA machine etc. To test whether your CPLEX installation and the JAVA interface work execute `setup_cplex_inner_class_access()` at the MATLAB prompt. Unless an error occurs the installation is working.

Note that Octave only works with the Java CPLEX interface and with the GLPK solver.

0.6 Compatibility with Octave

In version 2021.1, a larger revision of CNA was undertaken to improve compatibility with Octave (version 5.0 or higher for Octave is recommended) and most functionalities of CNA, including GUI-based analyses in the interactive network maps, now also work with Octave. However, some features do not work in Octave (e.g., autocomplete, see next section) and some menu items are accordingly disabled. Also, the GUI-based elements were optimized for MATLAB and might appear slightly differently in Octave. Sometimes you need to manually

increase the size of certain dialog boxes etc. for better readability or slightly move a window to get it rendered properly. We also observed that Octave has sometime problems to correctly display (network) graphics. In those cases, try to use another format (e.g. png instead of gif). Note also that Octave only works with the Java CPLEX interface and GLPK LP/MILP solver. If you want to use the Java CPLEX interface, Octave (64 bit version) needs access to a Java installation (64 bit version, install if necessary). The `configure_octave_java` script which is called by **startcna** will try to automatically find the Java installation but if that fails you can set the `JAVA_HOME` environment variable to point to the Java installation.

0.7 Notes on special features

Autocomplete functionality: several dialog boxes, where species or reaction IDs have to be entered by the user, are now equipped with autocomplete functionality. Enter the first letter of an ID and the first match will be displayed automatically. You may (1) chose the proposed ID (press the right cursor key), (2) switch to the next possible choice(s) by pressing `<Ctrl+Space>`, or (3) enter the next letter. This feature is not available under Octave.

0.8 New features

New features in CellNetAnalyzer 2025.2

- Resolving incompatibilities with MATLAB 2025a (and higher). Unfortunately, certain autocomplete functionalities (see section 0.7) will not be available anymore with MATLAB versions 2025a or higher (because some Java GUI objects can no longer be accessed).

New features in CellNetAnalyzer 2025.1

- Resolving some compatability issues and smaller performance improvements.

New features in CellNetAnalyzer 2023.1

- More flexibility in loading/combining scenarios.

New features in CellNetAnalyzer 2022.1

- Completely revised functions to treat redundant/infeasible systems in (classical) metabolic flux analysis and to check and resolve infeasibilities in FBA/FVA scenarios. A related API function was added.
- Several extended features in API functions.

New features in CellNetAnalyzer 2021.1

- Compatability with Octave (including GUI-based features) was largely improved.
- Support of Gurobi in all LP- and some MILP-based (CNAMCSEnumerator3) optimizations.
- MCSEnumerator3: new framework for MCS calculations. For example, this enables the calculation of bilevel (OptKnock-like) strain designs.
- MCSEnumerator2 and MCSEnumerator3 now support glpk, CPLEX and intlinprog and MCSEnumerator3 additionally Gurobi as MILP solver.

New features in CellNetAnalyzer 2020.3

- A CNA project of the (*E. coli* genome-scale) model iML1515 was added in the project

manager. This model also contains the gene rules (in the generic data fields of the reaction notes; see section 7.2.1) thus enabling, for example, computation of gene MCS.

- A dialog box for duality-based MCS computation was added to the GUI (accessible through the “Analysis/Minimal Cut Sets” menu item), which now enables convenient specification of an MCS problem which is then solved by duality-based algorithms (hence, elementary modes need not be calculated in a preprocessing step).
- CNAMCSEnumerator2 and CNAgeneMCSEnumerator2 now optionally use the MILP solver of the MATLAB Optimization Toolbox (intlinprog). Hence, while CPLEX is still preferred (the intlinprog variant is slower and does not support all features of the CPLEX solver) it is no longer mandatory for duality-based MCS computation.
- CNA functions are now callable from any working directory. It is no longer required to operate from the CellNetAnalyzer main path.
- If a metabolic network project is provided with GPR-rules (via the keyword “geneProductAssociation” in the generic data fields; see section 7.2.1 and the iML1515 model as example), the flux maps can be extended automatically with a stoichiometric representation of these GPR-rules by calling “Maps/Extend the network with GPR-rules” from the menu.
- Specified reactions can be placed on a separate and automatically generated map by calling “Maps – place reactions on separate map”.

New features in CellNetAnalyzer 2020.2

- The MILP-based (dual) algorithm for computing minimal cut sets (MCSEnumerator2) has been further extended and includes now new algorithmic variants that can be chosen via a set of options. As a result of this, the CPLEX MATLAB JAVA library is not mandatory anymore for this function.
- Improved options to display or to not display messages in some API functions.
- The best available LP solver is now automatically chosen in API functions such as CNAoptimizeFlux, CNAfluxVariability etc.

New features in CellNetAnalyzer 2020.1

- The MILP-based (dual) algorithm for computing minimal cut sets (MCS) has been significantly revised and extended. In particular, gene (instead of reaction) MCS can now be computed based on gene-protein-reaction rules; multiple desired and target regions can be defined and reaction/gene additions can be considered (in combination with classical reaction removal). The following new API functions have been added for this purpose: CNAMCSEnumerator2; CNAcharacterizeGeneMCS; CNAgeneMCSEnumerator2; CNAgenerateGPRrules; CNAintegrateGPRrules.

New features in CellNetAnalyzer 2019.3

- Structured data of mass-flow networks (such as geneProductAssociations for reactions or chemical formula for metabolites) can now separately be edited in the GUI.
- Extended “Check feasibility” functionality for mass-flow projects (in GUI only): if a given scenario is infeasible, the necessary (minimal) corrections are displayed.
- New API function (mass-flow networks) for computing elementary conservation relations.

New features in CellNetAnalyzer 2019.2

- For a given mass-flow project, basic network maps (listing reaction names and (optionally) reaction equations) to be used within the GUI environment of the project can now be generated in an automatic manner from the network model. This is especially

useful for larger (e.g. genome-scale) models (e.g. imported from an SBML file) where the manual creation of network maps would be a tedious task. Although the reaction names are only listed (no true network graphics), this feature is still useful as it pretty much eases the analysis of the model with the GUI.

- Many more model features contained in SBML model files (e.g., compartment, charge and chemical formula of a metabolite; name and gene product association of a reaction) are now stored in the generic data field of CNA when an SBML model is imported.

New features in CellNetAnalyzer 2019.1

- **CNA is now distributed with its complete source code and with a new (academic) license (see also LICENSE.txt, CONTRIBUTORS.txt, COPYRIGHT.txt, and README.txt in CNA's root directory).** Please read the license carefully; for your own purposes, you can do whatever you want with the code but when sharing modified CNA code you have to comply with some rules.
- Extended scenarios can now be saved in mass flow networks (via GUI + via API) comprising not only flux values but also flux bounds and the objective function.
- (Simple) SBML qual models can now be imported via GUI + via API function (requires MATLAB \geq 2014b and python). New libSBML version included.
- Improved flux variability analysis which is now automatically started in parallel mode in large networks. Extended CNAoptimizeFlux API function (e.g., for parsimonious FBA or with additional inequality constraints).
- New API function for ranking and characterizing minimal cut sets and other intervention strategies (CNAcharacterizeIS).
- New directory structure in CNA. All files now with Copyright info.
- Bugfix in CNAcompressMFNetwork (thanks to Mattia Gollub).
- New function to autocomplete input of reaction or species IDs in dialog boxes.
- New function to find location of a reaction's text box on large map(s).

New features in CellNetAnalyzer 2018.2

- New (GUI + API) function for parsimonious FBA.
- Improved display for flux variability analysis (GUI).
- New API function CNAwriteMFNText to write text (not only numbers) into text boxes.

New features in CellNetAnalyzer 2018.1

- New API functions to handle generic data for reactions and species/metabolites in a network project. Any parameter (e.g., ΔG_R^0 of a reaction) or textual information (e.g., KEGGID), which are not covered by CNA's reserved fields, can be stored under user-specified field names. The data are serialized in the reactions or species notes and can later be retrieved again (see section 7.2.1).
- Improved compatibility with COBRA toolbox (import/export of COBRA models). Note that also the COBRA toolbox itself provides now functions for interfacing COBRA with CNA and vice versa.
- New API functions CNAcomputeOptMDFpathway (to compute pathways with maximal thermodynamic driving force), CNAaddReactionMFN and CNAaddSpeciesMFN (to add reactions and species in mass-flow networks).

New features in CellNetAnalyzer 2017.4

- New API function CNAbasicNetworkProp.
- Improved import of SBML files (via libSBML).
- New options to set constraints when calculating elementary flux modes/vectors via

CNAcomputeEFM.

- New statistics function for elementary flux vectors: analysis of /selection via *rates*.
- Improved calculation of yield spaces + phase planes.

New features in CellNetAnalyzer 2017.3

- New API + GUI functions for calculating optimal yields and for calculating and drawing phase planes and yield spaces directly from the network (elementary modes or elementary vectors are not required).
- Better handling of and new analysis functions for calculated elementary flux vectors (EFVs): (1) phase plane of projected EFVs can be calculated and drawn; (2) improved yield space calculation based on EFVs; (3) whether the currently shown EFV is bounded or unbounded EFVs is displayed. All these functions are available via the “Elementary modes” panel.
- New function CNAgeneMCSEnumerator for calculating minimal gene cut sets based on a given gene-enzyme-reaction associations.

New features in CellNetAnalyzer 2017.2

- SBML models can now directly be imported via libSBML.
- Video tutorials for *CellNetAnalyzer* are now available on the CNA homepage and on the youtube channel of CNA.

New features in CellNetAnalyzer 2017.1

- CNA got a face lift – the menu structure has been completely reorganized.
- Easier handling of CNA paths (see section 0.2 and 0.5).
- New functions for mass-flow networks: flux bounds can now directly be used to calculate elementary flux vectors (= generalized elementary flux modes)
- New functions for mass-flow networks: SBML import directly from the GUI; display of objective function; calculation of net conversion of a given flux scenario.
- New API function to quickly calculate (single) minimal cut sets (CNAfindRegMCS).
- New functions for signal-flow networks: SBML export directly from the GUI.
- The network maps and their displayed values can now be saved as (bitmap) graphics file.
- The network can now directly be saved via the menu (not only via Network composer).
- New or adapted network projects for *E. coli* (including a slightly modified version of the iJO1366 genome-scale model (ECGS) and reduced versions of it (ECC2 and ECC2comp); see reference [28]).

New features in CellNetAnalyzer 2016.1

- We compiled a tutorial which is useful for a quick start of CNA. It can be downloaded from this web site: http://www.mpi-magdeburg.mpg.de/projects/cna/CNA_tutorial.pdf.
- Improved (accelerated) calculation of cut sets via CNAMCS(reg)Enumerator.
- CNAcompressMFNetwork: direct handling of conservation relations in rational format (the full network compression procedure can now be handled via rational calculation).
- CNAplot2DYieldSpace: a new function for calculating and plotting 2D yield spaces.
- CNAMFNetwork2sbmlSBML and CNAsbmlModel2MFNetwork.m: import and export of mass-flow (metabolic) networks now supports flux balance constraints including flux bounds and objective function (fbc package)
- CNA mass-flow projects can now also be exported as models for the COBRA toolbox.

New features in CellNetAnalyzer 2015.1

- New API function CNAcompressMFNetwork for (loss-free) compression of mass-flow

networks.

- New API function `CNAreduceMFNetwork` for reducing (pruning+compression) of mass-flow networks. See paper: [26].
- New API function `CNAremoveConsRel` for removing conservation relations.
- Several methods (including those mentioned above) now allow use of `efmtool`'s rational arithmetic routines (e.g. for network compression).

New features in CellNetAnalyzer 2014.1

- New API function `CNAregMCSEnumerator` which extends `CNAMCSEnumerator` by calculating smallest *regulatory* constrained Minimal Cut Sets (which are combinations of reaction knockouts and flux up- and down-regulations) in genome-scale networks (requires CPLEX) [Ref. 25].
- New API functions for importing / exporting COBRA models and for exporting signal-flow networks in SBML qual.
- New API function `CNAplotPhasePlane` for a 2D-projection of pairs of reaction fluxes.
- New API functions `CNAdeleteSpecies` and `CNAdeleteReaction` for convenient deletion of species or reaction(s) via command line.
- `CNAfluxVariability` can now be performed also for a subset of selected reactions.

New features in CellNetAnalyzer 2013.1

- CPLEX is now supported as LP solver for flux optimization, flux variability analysis and feasibility analysis (both in the GUI as well as in the API versions)
- Improved scrolling and panning of maps.
- Logical models can be exported in the new SBML qual format.
- New API function `CNAMCSEnumerator` for calculating (smallest) constrained Minimal Cut Sets without the requirement to compute the elementary modes first; can therefore also be applied to genome-scale networks (requires CPLEX) [Ref. 24].
- Bug fixes concerning minimal intervention sets in multi-value networks.
- Experimental implementation for the calculation of minimal intervention sets via an integer linear program (requires CPLEX).
- Updates for some Mex files to support also 64-bit MATLAB versions (special thanks to Ross Carlson and James Folsom for providing 64-bit files for MATLAB under Windows).

New features in CellNetAnalyzer 2012.1

- Version numbering format changed from 9.9 to 2012.1 (year + version number of the respective year)
- Mass-flow networks: the value for epsilon and the display format for fluxes can now be changed via the GUI.
- TRANSWESD can now also deal with maximal path lengths.

New features in CellNetAnalyzer 9.9

- For mass-flow-networks: the computation of elementary modes and convex bases (in flux cones) has been generalized to flux polyhedra (within the GUI and also as API function). This allows the specification of inhomogeneous constraints (e.g. fixing a rate to a particular value or restricting it within certain boundaries) and the subsequent calculation of elementary vectors, extreme points and rays characterizing/generating the solution space. This enables one, for example, to calculate the inner description of a polyhedron containing all optimal solutions for a given objective function.
- We have started to implement some computationally intensive algorithms in Java (which

are then used if a Java Virtual Machine is installed). First methods ported to Java include CNATranswedd and some path/cycle calculations. To properly set the Java paths please see section 0.5.

New features in CellNetAnalyzer 9.8

- For all networks: attributes of species, reactions and macromolecules can now directly be inspected and edited by right-clicking on the text boxes. It is not necessary anymore to search for a network element in the network composer and to open then the property dialog box.
- For mass-flow-networks: a two-dimensional yield-space plot can now be generated from the selected set of elementary modes (function is accessible from EM control panel).

New features in CellNetAnalyzer 9.7

- Major revision of the API functionality. Networks and scenarios can now be saved/loaded with or without GUI from command line.
- efmtool can now be used with rational numbers for computing elementary modes without numerical problems.

New features in CellNetAnalyzer 9.6

- Minimal cut sets (MCSs) calculation revised now allowing one to consider constraints: one may define sets of paths/cycles/modes that should not be hit (or where at least a minimum number of these paths/cycles/modes is not hit) by the cut sets to be computed. Relevant for MCSs calculations in signal-flow and mass-flow networks.
- Mass-flow networks: Flux Variability Analysis (GUI + API function). Useful for checking the feasible upper and lower boundaries of reaction rates for a given set of fixed fluxes.

New features in CellNetAnalyzer 9.5

- A project can now be started by double-clicking in the project manager.
- Mass-flow networks:
 - CASOP: a new computational Approach for identifying intervention strategies for strain optimization and Metabolic Engineering (as menu item + API function) [18].
 - Elementary modes can now also be selected with respect to their (product) yield.
- Signal-flow networks:
 - API function for TRANSWESD: a new method for transitive reduction useful for reverse engineering of regulatory networks (interaction graphs) [19].

New features in CellNetAnalyzer 9.4

- CellNetAnalyzer has now a logo!
- some bug fixes
- Mass-flow networks:
 - New API functions for loading/saving scenarios
- Signal-flow networks:
 - New *Odefy* version integrated (which reads species and reaction values before starting a simulation).
 - New API functions for loading/saving scenarios

New features in CellNetAnalyzer 9.3

New online manual: http://www.mpi-magdeburg.mpg.de/projects/cna/manual/toc_frame.htm
Mass-flow networks:

- EFMTTool ([15], developed by Marco Terzer, ETH Zürich), currently the most efficient method for computing elementary modes in large networks, can now optionally be used (on-the-fly) within CNA.
- Editing network elements directly from element selector.

Signal-flow networks:

- Computation of minimal intervention sets accelerated.
- Editing network elements directly from element selector.
- New network project: EGF/ErbB receptor signaling.

New features in CellNetAnalyzer 9.2

Signal-flow networks:

- Improved computation of minimal intervention sets with more options; the new API function (CNAcomputeMIS) also allows for considering multiple scenarios
- Element selector allows efficient searching and highlighting of network elements

Mass-flow networks:

- Element selector allows efficient searching and highlighting of network elements
- New API function CNAoptimizeFlux (for Flux Balance Analysis)

New features in CellNetAnalyzer 9.1

- API functionality improved and extended considerably (computation of logical steady states, dependency matrix, cut sets and others now possible via API)

Signal-flow networks:

- Improved calculation of dependency matrix and shortest signed paths
- Automatic comparison of (experimental) data with dependency matrix
- *Odefy*: Plugin developed by Fabian Theis' group (Jan Krumsiek, Dominik Wittmann) at the Helmholtz-Zentrum Munich for translating logical model into ODE models and exporting/simulating these ODE models (odefy also allows to simulate Boolean models with synchronous and asynchronous switching)

Mass-flow Networks

- Improved import/export of SBML models

New features in CellNetAnalyzer 9.0

CNA 9.0 has undergone a major revision: the internal variable structure of CNA has been changed in order to facilitate API functionalities. Also, several new functions have been added and other improvements (e.g. toolbar buttons) make daily work with CNA simply more convenient. The main changes are:

- Computation of paths and loops in signal-flow networks has been revised: more convenient and more efficient.
- Computation of convex basis (also of extreme pathways) now more efficient as it also uses Metatool.
- Convex basis and elementary modes are computed with Metatool's integer arithmetic (if the conversion of the stoichiometric matrix into integers is possible).
- Computation of strongly connected components (signal-flow networks).
- Improved computation of Minimal Cut Sets
- Statistical features of minimal intervention sets can be computed (signal-flow networks)
- Numerical values in text boxes can be displayed in "heatmap style" (text boxes become gradually green, red, and yellow; depending on their value).
- Conversion of (logical) hypograph into interaction graphs (signal-flow networks) can

be done with optional removal of duplicated arcs.

- Toolbar allowing quick access to CNA's zoom tools and for opening, closing, saving and resetting scenarios.
- Color selector can be used to define the text box / text colors of a network project.
- A network project can now be started without GUI. The user may use CNA's API functionality directly from the command line or do his own computations using the CNA project variable.
- Several network projects can now be loaded in parallel.
- A network project can be cloned.
- API functionality has been extended considerably.

New features in CellNetAnalyzer 8.0

- Bar chart showing in/out fluxes around a metabolite (mass-flow networks)
- Computation of (species) equivalence classes in (logical) signal-flow networks.
- Computation of Minimal Intervention Sets strongly accelerated.
- The internal variable structure in CNA has been revised cleaning up the workspace
- New algorithm for computing shortest signed (pos./neg.) paths and cycles in interaction graphs as required for building the dependency matrix (see section 4.3).

New features in CellNetAnalyzer 7.0

- CNA provides now a (small) API (Application Programming Interface) which enables the exchange of data and variables between CNA and other MATLAB functions and the integration of user-created functions in CNA's menu.
- For flux optimizations, as an alternative to MATLAB's *linprog* function from the optimization toolbox, the user may now optionally use the GLPK (GNU Linear Programming Kit) library via the GLPKMEX interface.
- Signal flow networks: species values can now be directly fixed by entering a value in the respective text box (instead of fixing signal flows pointing into that species).
- Improved algorithm for minimal intervention sets.

New features in CellNetAnalyzer 6.3

- zoom tools (scrollbars; zooming in and out) enable better handling of large maps with large resolution
- in signal-flow networks, an interaction can be defined to be non-monotone or monotone (in earlier versions, monotone behavior was implicitly assumed; read section 4). This feature is relevant for logical networks with multiple-valued discrete states.

New features in CellNetAnalyzer 6.2

- comments and notes can now be added to reactions and species making a documentation possible

New features in CellNetAnalyzer 6.1

- "interaction networks" are now referred to as signal-flow (or signaling) networks (S) whereas stoichiometric (metabolic) networks are now referred to as "mass-flow" networks (M)
- as a generalisation of logical minimal cut sets, logical minimal intervention sets can be computed in signal-flow networks: intervention goals can now be achieved not only by cutting (knocking-out) components but also by a (constitutive)

ACTIVATION (knock-in) of network species.

- an interaction can now be assigned an INCOMPLETE truth table. This is especially useful if it is not clear whether several species have to be combined with an AND or with an OR.
- in signaling networks, all input species / reactions can be set to zero in one step (“closing input/output gates”).

New features in CellNetAnalyzer / FluxAnalyzer 6.0

- Apart from stoichiometric networks, *CellNetAnalyzer* allows now also the construction and detailed analysis of signal-flow networks useful for analyzing the structure of signaling and regulatory networks. A number of tools is provided for studying graphical, hypergraphical and logical properties of signal-flow networks. Practically, *CellNetAnalyzer* facilitates now a functional topological analysis of any kind of cellular networks.
- for stoichiometric networks, most functions of *FluxAnalyzer* have been taken over in CNA, however, a few menu entries and functions have been revised.
- different sets of elementary modes (stored in several files) can now be merged.
- new MEX files (running also under MATLAB 7)

New features in FluxAnalyzer 5.3

- more efficient computation of elementary modes; a newly provided METATOOL module for computing the modes (which uses exclusively C-code and is therefore faster than the routines from *FluxAnalyzer*) can optionally be used
- some rates may be enforced to be involved in elementary modes to be computed; in some cases this can drastically reduce running time compared to the case where all modes are computed (however, sometimes it can even be slower)
- some more basic topological and graph-theoretical features are computed
- a new feature for minimal cut sets: searching for cut sets with lowest side-effects
- “variance weighted least squares method II” for flux analysis in redundant systems is no longer supported (hardly used in the literature; low relevance)

New features in FluxAnalyzer 5.2

- import/export SBML(Level 2)
- import/export from/to METATOOL format

New features in FluxAnalyzer 5.1

- using a new algorithm, the computation of elementary modes has been improved considerably, both with respect to time and memory demand.
- The menu function “Basic topological properties” has been revised and comprises some new features.
- Computation of (elementary) conservation relations: the user may choose to compute *all* or only *non-negative* elementary conservation relations.

New features in FluxAnalyzer 5.0

- Minimal cut sets (MCS; see [9]): calculation, display and detailed analysis. An MCS is a minimal set of reactions whose removal will make a certain function impossible; MCSs are useful for searching for targets for genetic modifications, for assessing network fragility, for finding a appropriate set of (flux) measurements making a certain reaction rate observable and for predicting inviable mutant phenotypes

New features in FluxAnalyzer 4.3

- calculation of graph-theoretical path lengths between metabolites (including the average path length and network diameter); can be done in a directed as well as undirected graph representation of the reaction network
- more intuitive arrangement of menu items
- separate calculation of (elementary) conservation relations
- flux analysis: calculation of variances of the estimated (calculable) rates in redundant systems
- new example networks

New features in FluxAnalyzer 4.2

- calculation of elementary (flux) modes /extreme pathways: the user has the option to consider isozymes only once
- elementary (flux) modes / extreme pathway matrices can be exported in ASCII format
- more convenient arrangement of text boxes on the network maps (see button “Move text boxes ...” in the network composer window)
- procedures for metabolic flux analysis do now automatically choose the procedures for redundant/non-redundant scenarios (menu-item: “Flux analysis ...”)
- text (“tooltip”) appears when the user moves and leaves the mouse pointer over a text box associated with a network element
- some new network properties can be calculated (e.g. detection of isozymes)
- manual construction of network projects is now described in the manual

New features in FluxAnalyzer 4.1

- computation of mutually excluding reaction pairs in a set of elementary modes
- “Select all” button allows for quick selection of the complete set of elementary modes
- improved graphical display of the stoichiometric matrix
- computation of elementary modes now also possible in MATLAB 6.5 (incompatibility removed)

New features in FluxAnalyzer 4.0

- calculation of “control effective fluxes” with respect to elementary flux modes (applied in [2])
- histogram of product yields in elementary flux modes
- arbitrary normalization of elementary modes
- improved display of control panels for low monitor resolution and for WINDOWS

New features in FluxAnalyzer 3.1

- bar chart for comparing the fluxes of a given scenario
- histogram displaying the frequency distribution of the connectivity number of metabolites
- histogram displaying the frequency distribution of the number of involved reactions in elementary modes
- new examples of network projects

New features in FluxAnalyzer 3.0

- metabolites can be defined as being external (not balanced) or internal (balanced)

- graphical display of the stoichiometric matrix including connectivity number of each metabolite
- the stoichiometric matrix can be exported as MATLAB or ASCII file
- enhanced algorithm for calculating elementary flux modes or of a convex basis; fast MATLAB-MEX-files can be used (needs MATLAB-Compiler)
- some small bugs have been removed
- start the FluxAnalyzer now by the command “startfa” (instead of “startmfa”); the menu item is now “FluxAnalyzer” (instead of MFA)

New features in FluxAnalyzer 2.4

- calculation of convex basis
- Minimum and Maximum rates may now also be $-\infty$ / $+\infty$
- checking the feasibility of a given network scenario (takes the given rates *and* the capacity constraints for the reaction rates into account) – see menu item “Check feasibility”
- layout of non-editable text boxes has been changed slightly

1. CellNetAnalyzer: Introduction and Overview

CNA is a MATLAB toolbox for analyzing structure and function of biological networks on the basis of topological, stoichiometric, qualitative (logical) and semi-quantitative modeling approaches requiring no or only few parameters. In particular, metabolic networks (mass-flow) can be studied based on stoichiometric and constraint-based modeling approaches whereas signaling and regulatory networks (signal-flow networks) can be explored by qualitative and semi-quantitative modeling approaches (Boolean/logical models, interaction graphs, logic-based ODEs). CNA can be used via command-line based operations (supported by an application programming interface; API) or via a graphical user interface with embedded network visualizations.

Basically, CNA is composed of two main parts (Fig. 1). The first part comprises network projects (some are distributed with CNA; others can be created by the user). A network project is either of type “mass-flow” (for modeling material flows as in metabolic or other stoichiometric reaction networks) or of type “signal-flow” (modeling information or signal flows as in signaling or regulatory networks). Each network project contains an abstract (formal) network representation and, optionally (for use with GUI), one or several network graphics visualizing the network under investigation. The abstract network model specifies network elements such as reactions or compounds. The network graphics must be provided by the user (for example, created with drawing programs or taken from other sources (see below)). In the GUI-based application, network model and network graphics are linked via user interfaces (text boxes) leading to *interactive network maps* (see Figure 2 for an example). The text boxes facilitate input and output (e.g. of reaction rates) directly within the network visualization. The interactive network maps are central for the GUI-based work with CNA but are not needed when CNA is used via its API.

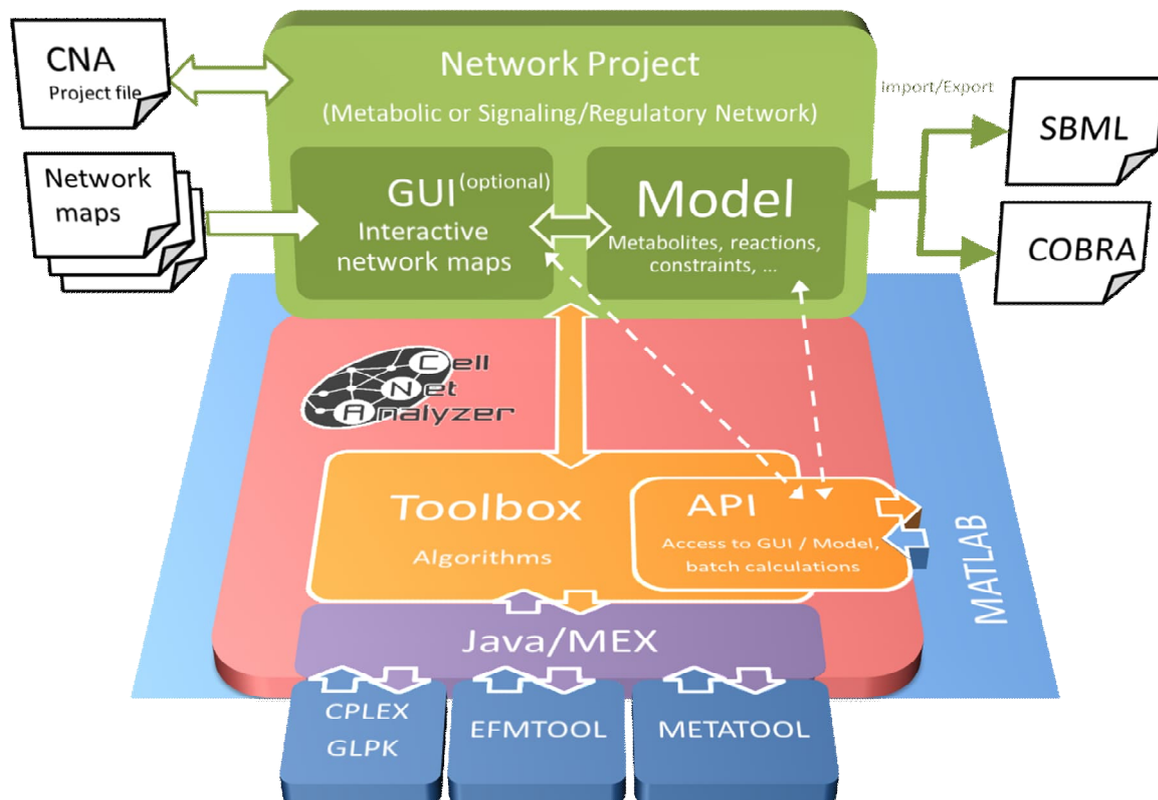


Fig. 1: Internal structure and architecture of CellNetAnalyzer

The second part of CNA comprises a large toolbox of functions and algorithms for analyzing a given network project. These functions can be conveniently started from pull-down menus in the network maps (when using the GUI; see chapters 3 and 4) or be started from command line (when using the API; see chapter 7). Some of the procedures are based on MATLAB's built-in functions, others are completely own implementations. A few algorithms use the MEX interface of MATLAB to call precompiled C/C++ functions (including those provided by Metatool) or use external JAVA functions (e.g., *efmtool*) to speed-up extensive computations. Calculations based on linear programming (LP) may use external solvers (GLPK, CPLEX, Gurobi).

Interactive network maps

When initializing a new network project in *CellNetAnalyzer* via the project manager (chapter 2), one or several graphic file must be specified for the network which will later serve as background visualization for the interactive network maps in the GUI-based model analysis (as surrogate you may also use a dummy graphic or create graphics with listed reaction names; see below). Importantly, CNA itself is not a pathway drawing engine and can therefore not be used to create the network maps from the network model (but see the exception with creating reaction lists below!). Algorithms for an automatic drawing of (metabolic or signaling) networks do exist but they are quite complex and do in almost all cases not lead to network visualizations as desired by the user. Therefore, CNA provides 4 different ways to link network visualizations (or other types of maps or network representations) to construct interactive network maps. The first two are:

- 1) The user may create network graphics in his favorite style with the help of external drawing programs (such as "CorelDraw", "inkscape", Xfig" etc.). This also includes specialized tools, e.g. OMIX for creating metabolic maps or ProMoT (also originally developed at the MPI in Magdeburg; www.mpi-magdeburg.mpg.de/projects/Promot) to build signal-flow networks where a logical network can be constructed in a visual manner and then be exported (together with the abstract model) to CNA. Note that there is no limitation in the style you draw the maps and they may contain arbitrary annotations or comments because, in CNA, the network graphics themselves will only be indirectly linked with the abstract network model.
- 2) Network graphics may also be obtained from other (e.g., internet) sources (e.g. KEGG, BioPath) with existing network visualizations.

You may have a look at the example projects provided with CNA to see what different kind of graphics are used in these projects. In general, for every network project, one may include as many graphic files as needed to represent the complete network, hence, the network visualization maybe split into several subnetworks. With one exception (see below), MATLAB (and CNA) can currently only read pixel-based graphics including tif, bmp, jpg, pcx; the jpg format can but should not be used). Therefore, the map size (in pixels) should ideally not be larger than the resolution of the monitor because the graphics can otherwise not be displayed in full resolution. However, the zoom tools of CNA facilitate also the use of large pixel-based network maps: zooming in and out and scrollbars provide a convenient way to navigate through large network maps. Zoom tools will be activated automatically when the resolution of the map is larger than the resolution of the monitor. When a network map has been resized, the menu command "Set original map size" resizes all network maps to the original resolution (or, if the resolution of the map is larger than that of the display, to the largest possible size).

A special opportunity for creating network maps are MATLAB figures (file extension “fig”) which may be drawn by using the “Plot editing tools” of MATLAB (which are rather limited, however). You may also try to use other tools such as SVG importers (see e.g. <https://de.mathworks.com/matlabcentral/fileexchange/66554-svg-import-export>). MATLAB figures are vector-oriented and can therefore be resized without loss of quality.

When loading a network project in CNA each registered network map serves as a background in a MATLAB figure, where text boxes - associated with elements of the abstract network model - can be placed by the user. The linkage of network graphics and user interfaces leads to *interactive network maps*. As an example, Fig. 2 shows the interactive network map belonging to a simple stoichiometric network project created by (and distributed with) CNA. The underlying network map was drawn by an external graphic program (xfig). Graphical user interfaces (text boxes) refer to reactions or biomass components linking the symbolically defined network (not shown) with the graphical scheme resulting in the interactive network map. The *CellNetAnalyzer* menu with functions for analyzing and editing the network appears in every interactive map from. For some frequently used functions (such as loading or saving a scenario), the CNA toolbar provides a short-cut to quickly start those actions.

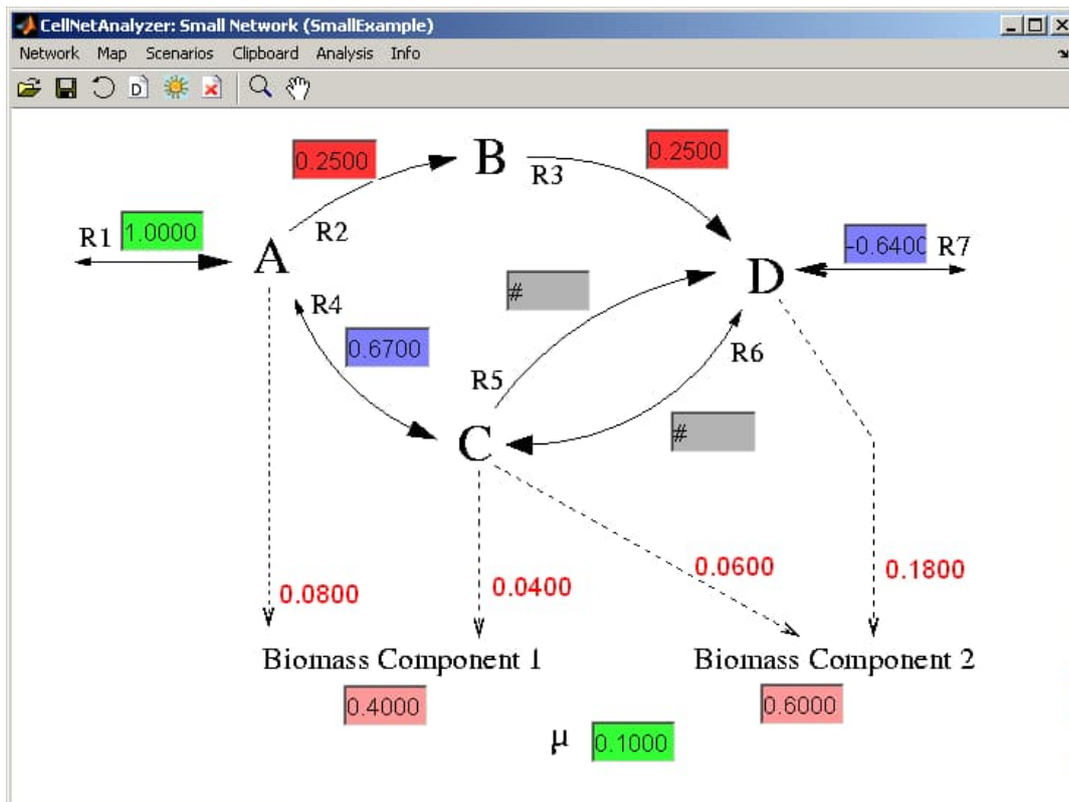


Fig. 2: Example of a stoichiometric (mass-flow) network project in CellNetAnalyzer.

For the case that the user does not have maps available for visualizing the network (e.g. when importing a large network from a database) or if he does not want to create one, there are still two additional possibilities to use CNA in GUI mode

- 3) For mass-flow projects, basic network maps (listing reaction names and (optionally) reaction equations) can now be generated in an automatic manner from the network model. This is especially useful for larger (e.g. genome-scale) models (e.g. imported from

an SBML file) where the creation of network maps would be a tedious task. Even though the reaction names are only listed (no true network graphics), this feature is very useful since it pretty much eases the analysis of the model with the GUI. (see also sections 3 and 6)

- 4) If even option (3) is not a way to go (e.g., in case of large imported signaling networks) then one may choose a “dummy” picture, for example CNA.pcx provided in the CNA main folder. Then, obviously, the user will not have a meaningful visualization of his network, however, he or she can nevertheless compose (or import) a network model and then analyze this network in the GUI with the provided menu functions. The text boxes can be placed anywhere or even outside of the dummy map (when a network is imported, all text boxes are put on the upper right corner of the map). Results, for example a computed set of elementary modes or signaling paths, can be exported and further analyzed via ASCII files.

Finally, apart from a GUI-based analysis of the model, the user may still load the network without a GUI (either using the ‘Start w/o GUI’ button in the project manager or just from command line using an API function; see chapter 7). One may then analyze the network with the provided API functions of CNA from command line (chapter 7).

Section 2 explains how a new network project can be created (and its properties be edited) by CNA’s Project Manager. The set-up of abstract network models and of interactive maps is described in sections 3.1, 3.2 (mass-flow networks) and 4.1, 4.2 (signal-flow networks), respectively. The toolboxes facilitating integrated network analysis are described in detail in sections 3.3 and 4.3, respectively.

2. Creating and managing network projects

As described in section 0, for using CNA one first starts MATLAB, ensures that CNA is either the current working directory or added to the MATLAB-path and then enters “startcna” to open CNA in GUI mode. The Project Manager (Fig. 3) appears showing all registered network projects. “(M)” indicates mass-flow (metabolic, stoichiometric) networks, “(S)” signal-flow (regulatory) networks. Existing network projects can be selected and then loaded, normally with GUI (button “Start”) but sometimes also without GUI (“Load w/o GUI”; see also chapter 7). In the only entry of the menu bar (‘Project’) you may alternatively “edit” general properties or “clone” (duplicate) or “remove” the selected project. Finally, you may initialize a new project. Initialization of a new project requires that the user defines some general properties of this project using the mask shown in Figure 4. First, the name of the network project (as it will appear later in the list of networks in the Project Manager) and the name (or relative path) of the project directory (where the project files will be stored) must be defined. It is recommended to declare only the name of the directory (e.g. ‘MyNetwork’), which then becomes a subdirectory of CNA’s root folder. It is also possible to define the network path relative to the CNA root folder (e.g. outside or inside of CNA: ‘../Networks/MyNetwork’ or ‘Networks/MyNetwork’). If the directory does not exist, CNA will create a new one.

Next, the user has to define the network type (“Mass-flow” or “Signal-flow”). Users not familiar with the fundamental differences between these two types of networks should first read sections 3 and 4 before assigning the respective network type. Note that the network type of a project cannot be changed once the project has been created.

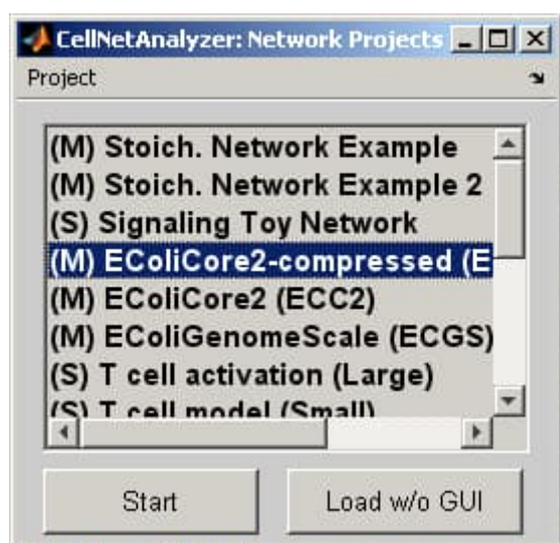


Fig. 3 (left): Project Manager

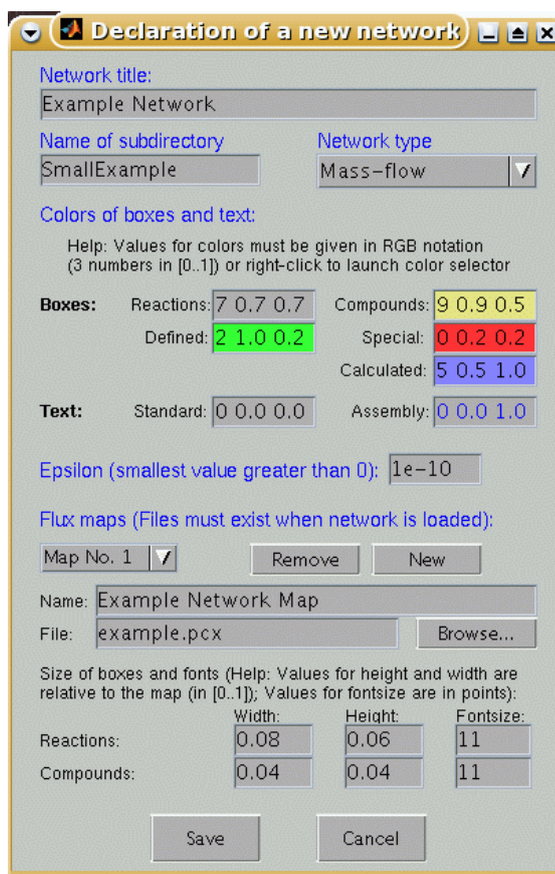


Fig. 4 (right): Declaring the properties of a new network project

Next, the background colors and text colors for the text boxes can be specified. Colors have to be given in RGB notation (three numbers between 0 and 1 defining the proportions of the colors red, green and blue). The text box color / text color resulting from the entered RGB values is directly exemplified in the respective box (see Figure 4). When initializing a new network, CNA inserts default color values which can then be changed by the user. A convenient way for selecting a color is to right-click on the respective text box which launches a color selector dialog.

The different colors have the following meaning (cf. colors of boxes and text in Fig. 2):

Text boxes:

- *Reactions* (default: gray): Standard background color for text boxes associated with elements of type “reaction/interaction” (see sections 3.1 and 4.1).
- *Compounds* (default: yellow): Standard background color for text boxes associated with compounds (“biomass components” in mass-flow networks and “species” in signal-flow networks; see sections 3.1, 3.2 and 4.1, 4.2)

After computations, box colors are often changed to indicate the context of each network element (cf. Fig. 2):

- *Calculated* (default: blue): Often used as text box color for network elements, for which a value has been calculated in a certain computation.
- *Defined* (default: green): Background color for boxes of those network elements whose value was defined before a computation.
- *Special* (default: red): Background color for marking elements which had a particular meaning in this computation (for example: balanceable rates in a redundant system; cf. section 3.2).

Text:

- *Standard* (default: black): Color of the text in all boxes, except:
- *Assembly* (default: blue, in Fig. 2 light red): Color of text used for displaying assembly rates. This is only of relevance in mass-flow networks.

Another project parameter to be defined by the user is *epsilon*, the smallest number larger than zero. *Epsilon* is used in various calculations for distinguishing “zero” from “non-zero”. We recommend $1e-10$.

Furthermore, as explained in detail in section 1, the associated networks maps of the current project (i.e. the graphics files) must be specified. Arbitrarily many network maps can be linked to a project; at least one must be declared for a GUI-based analysis. Regarding the drawing/generation of (different kind) of network maps or graphics, read section 1. By clicking on “New” in Fig. 4, a new interactive map can be declared, by “Remove” the currently selected map is removed from the network project (the graphics file itself is not deleted). Using the pull-down menu any of the maps associated with the project can be selected for editing its properties:

- *Name of the network map*: The map name appearing in the top of the window displaying the network map.
- *Name of the file*: Represents the graphic file storing the map (see section 1). Possible graphic formats are, for example, pcx, bmp, jpg, tif, png or a MATLAB figure (.fig). The button “Browse...” helps to search for the file, recommended is to have the file in the respective network directory (you then need only to specify the file name (without giving an explicit path)). The file will be loaded when the network project is started and each map serves as

background for an interactive network map in a separate window.

- *Size of boxes and fonts:* For each network map the *relative* width and height of the text boxes (separately for reactions and compounds) and the text box font size must be defined. These values are specific for each map. For example, the value 0.04 for height (or width) means that the box has a height (width) of 4% of the complete height (width) of the map. Maybe, you must try a little bit to find appropriate values.

Note again (see also section 1), for the case that the user does not have maps available for visualizing the network (e.g. when importing a large network from a database) or if he does not want to create one, there are still two additional possibilities to analyze a network in GUI mode:

- 1) *One may choose a “dummy” picture, for example CNA.pcx provided in the CNA main folder. Then, obviously, the user will not have a meaningful visualization of his network, however, he or she can nevertheless compose (or import) a network model (e.g., via SBML) and then analyze this network in the GUI with the provided menu functions. The text boxes can be placed anywhere or even outside of the dummy map (when a network is imported, all text boxes are put on the upper right corner of the map). Results, for example a computed set of elementary modes or signaling paths, can be exported and further analyzed via ASCII files.*
- 2) *An even more convenient opportunity exists for mass-flow projects. Here, basic network maps (listing reaction names and (optionally) reaction equations) can be generated in an automatic manner from the network model. This is especially useful for larger (e.g. genome-scale) models (e.g. imported from an SBML file) where the creation of network maps would be a tedious task. Even though the reaction names are only listed (no true network graphics), this feature is very useful since it pretty much eases the analysis of the model with the GUI. (see also sections 3 and 6). To use this option, choose initially the dummy graphics “CNA.pcx” from CNA’s root folder. Open then the project and import, for example, an SBML model. Choose then “Maps/Generate standard maps ...” from the menu and the maps will be generated from the network model and the text boxes automatically be placed in these generated maps.*

Furthermore, apart from a GUI-based analysis of the model, the user may still load the network without a GUI (either using the ‘Start w/o GUI’ button in the project manager or just from command line using an API function; see chapter 7). One may then analyze the network with the provided API functions of CNA from command line (chapter 7).

If all properties and parameters for the new network project have been defined the initialization of the network project is finalized by clicking on “Save”. CNA creates then all files (most of them are empty) needed for a network project. By clicking on “Cancel” you can quit without saving.

As mentioned above, having a network project selected in the Project Manager and clicking on “Start” loads the selected network. If the network project has just been initialized only the network maps will be displayed because a network model does not exist yet. In all maps, CNAs menu bar appears (see Figure 1) from which one can start various actions. In particular, one may open the Network Composer for manual compilation of the network structure (section 3.2 and 4.2) or import (e.g. an SBML) model. Note that you can also load several copies of the same network project (e.g. for comparing different scenarios). However, when saving a modified network structure in one copy, it will not yet be visible in the other copy in the memory. A network session can be finished by closing any of the network maps. CNA will ask you whether you want to save your changes and whether the network variable is to be cleared from MATLAB’s workspace. One may avoid clearing the variable in case one wants to apply API functions to the network project afterwards (without having the GUI loaded).

Using the “Edit” menu entry (under Project) in the Project Manager all properties described above can be changed for the selected network (same mask as in Fig. 4), except the network type. With “Remove” one may delete the selected network project from the network list (the files

of the network project are *not* deleted and can thus be re-installed later on).

With “Clone ...” in the Project Manager menu, it is possible to create a duplicate of an existing project in a separate directory. First the user is asked to specify the target directory for the project clone. If this directory does not exist it is created automatically. Then one can enter the name under which the clone will be registered in the project manager. If this is left empty, the project will be cloned but not registered. During cloning, the files describing the network and the network maps are copied. However, the network maps are copied only if they are located in the project directory of the source project. This allows sharing of network maps between projects. When cloning a project, possibly existing files in the target directory will not be overwritten and any attempt to do so will result in an error.

If one wants to register an existing network project (e.g. a copy of a network project received from a colleague), one should click on “New” in the project manager, insert the name of the project and the name of the (existing) project directory and then click on “Save”. You are then asked whether the “app_para.m – file” in the network directory should be overwritten. Click here on “No”. After this, the project is “visible” for *CellNetAnalyzer*.

NOTE: If a network map of an existing network project has been removed from the project, the user will possibly be requested to reassign the old map numbers to the new network maps (after loading the network project). This is necessary for positioning the text boxes in the correct network map.

3. Constructing and analyzing mass-flow (stoichiometric, metabolic) networks

3.1 How mass-flow networks are represented in CellNetAnalyzer

In CNA, a metabolic or mass-flow network is composed of elements of 4 distinct types (table 1):

Element Type	Metabolite (Species)	Reaction	Biomass Constituent	Assembly Route
	network node	(bio)chemical conversion	substantial macromolecule such as protein or RNA	efflux of a metabolite for synthesis of a biomass constituent
Text Box	none	reaction rate ([mmol/(gDW*h)])	relative biomass concentration ([g/gDW])	rate of metabolite consumption ([mmol/(gDW*h)])
Attributes	<ul style="list-style-type: none"> • <i>full name</i> • <i>identifier</i> • <i>external-flag</i> • <i>notes/comments/structured data</i> <p>(External metabolites are not considered to be in pseudo-steady state and therefore not balanced by eq. (1))</p>	<ul style="list-style-type: none"> • <i>identifier</i> • <i>reaction equation</i> • <i>default rate</i> • <i>rate minimum (lower flux bound)</i> • <i>rate maximum (upper flux bound)</i> • <i>coefficient in linear objective function</i> • <i>variance of measurements</i> • <i>text box parameters</i> • <i>notes/comments/structured data</i> 	<ul style="list-style-type: none"> • <i>full name</i> • <i>identifier</i> • <i>default concentration</i> • <i>cumulative synthesis equation</i> • <i>text box parameters</i> 	<ul style="list-style-type: none"> • <i>identifier</i> • <i>metabolite</i> • <i>biomass constituent</i> • <i>text box parameters</i>

Table 1: Element types (and their attributes) in mass-flow networks

- **Metabolites (species, storage):** Metabolites are the network nodes. Usually, in metabolic network models, metabolites represent small molecules like substrates, intermediary products, precursors and building blocks. The basic assumption of Metabolite Balancing techniques is that the concentration of these metabolites is in steady state (i.e. the sum of all fluxes producing or consuming a certain metabolite is zero):

$$\mathbf{N} \mathbf{r} = \mathbf{0} \quad (1)$$

(\mathbf{N} : stoichiometric matrix ($m \times q$) with m metabolites (rows) and q reactions (columns); \mathbf{r} : vector ($q \times 1$) of reaction rates, $\mathbf{0}$: null vector ($m \times 1$))

In CNA, a metabolite can also be considered to be “external”, i.e. it has not to fulfill eq. (1) and is therefore not included in the stoichiometric matrix if calculations are performed. Note that species are the only network elements that have no associated text box (input field) in interactive maps. (The reason is that almost all results are related to reactions).

In Fig. 2, for example, the capitals (A-D) are the (internal) metabolites and R1 and R7 represent uptake or excretion “reactions” of metabolites.

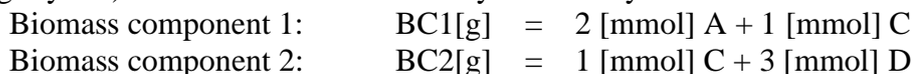
- **Reactions:** These elements correspond to biochemical reactions, each having a certain stoichiometry. Reactions are the (hyper-)edges in the network connecting the metabolites. The reaction *rates* of biochemical conversions (with unit [mmol/(g dryWeight * h)]) are the central focus of many computations relying on Metabolite Balancing. Each reaction is referenced by a text box which can be arranged by the user on its associated pathway in the network map. (Fig. 2: each of the 8 reactions (R1-R7, μ) has its own text box). Of course, an element of type *reaction* can also be a sequence of more than one biochemical reaction

(cumulative stoichiometry).

A special “reaction” is biomass synthesis, which in CNA is always referenced with the identifier *mue* (its rate is the growth rate; unit: 1/h). In contrast to all other reactions, its stoichiometry is not constant because this depends on the defined biomass composition (see *macromolecules*). Therefore, the stoichiometry of *mue* is determined before every computation on the basis of the currently defined biomass composition. In Fig. 2, the text box for *mue* (referencing the growth rate) is located in the network map next to the symbol μ .

Essentially, each (non-empty) mass-flow project comprises a certain number of reactions and metabolites. The following two network elements are optional. They facilitate the consideration of a flexible composition of the biomass and thus a flexible stoichiometry of the biomass synthesis reaction *mue*. In general, if biomass synthesis is considered in a mass-flow network, the user may either define a single biomass synthesis reaction with a fixed stoichiometry (in this case, do not name it *mue*). However, if he wants to investigate e.g. the consequences of changing the biomass composition on certain flux distributions, CNA provides a convenient way to that:

- **Biomass constituents (macromolecules):** Usually, this element type represents substantial components of the biomass like protein, DNA, RNA, lipids.... etc. In CNA, every biomass constituent (abbreviated as BC) has its own stoichiometry that defines which metabolites are consumed for its synthesis (where the user has to know, for example, the amino acid composition of proteins and so on). Additionally, every BC is referenced by its own text box as it was introduced for the reactions. Therein, the relative concentration of the biomass component can be declared by the user [g/(g dryWeight)]. As explained above, these values are used for calculating the stoichiometry of the biomass synthesis (reaction) *mue*. In Fig. 2, two biomass compositions were defined (BC1 and BC2; their associated text boxes are brightly red). The assumed stoichiometry for their synthesis is as follows:



There are some applications (as often given in the literature) where the user doesn't have or doesn't know the biomass composition but the overall demand of precursors for synthesizing the biomass. In such cases, the user can easily define only one BC (the biomass “itself”) and then define the overall efflux of precursors (energy, reduction equivalents, ...) into the biomass. Clearly, the relative concentration of the only BC must then be set to 1. Of course, alternatively he may also define a single reaction comprising the stoichiometry of precursors needed for building the complete biomass. Then, a BC needs not to be declared. However, in this case, assembly routes (below) can also not be defined and *mue* should not be used as reaction identifier.

- **Assembly routes (assembly rates):** Assembly route is an auxiliary element type and can be used for displaying the efflux of *one* metabolite into biosynthesis of *one* BC. For example, the amino acid serine is used for synthesis of lipids and for synthesis of proteins. For an explicit output of both fluxes in the interactive network map the element type *assembly route* can be assigned. The flux (i.e. the rate) along an assembly route from metabolite M into biomass component B is the product of the stoichiometric coefficient of M in the synthesis reaction of B with the concentration of B and the growth rate (the rate of the biosynthesis “reaction” *mue*). The product of the first two values is implicitly (cumulative with all effluxes of M into all BCs) contained in the biomass synthesis “reaction” *mue*. It is not mandatory to output all possible assembly routes (doing this for ATP would be very tedious, for example). In Fig. 2 one can see four assembly routes (dashed arrows) with red numbers

displaying the fluxes along these routes. Bear in mind: an assembly rate is not the same as a reaction rate in this context, because only the latter one is used for the balancing equations of the metabolites.

Each element type has a set of attributes which must be defined if a new element (an instance) of this type is declared (Table 1):

Attributes of *metabolites*:

- 1) **identifier** of this substance in CNA (e.g.: G6P) without blanks
- 2) **the full name** of the metabolite (e.g.: glucose-6-phosphate; no blanks allowed!)
- 3) **flag “external”**: if this flag is set, then the metabolite is considered to be external, i.e. the metabolite is considered to be in excess and therefore not balanced via eq. (1). This flag also allows to “remove” a metabolite temporarily from the network without removing it explicitly from the reaction equations
- 4) **notes/comments/structured data**: for documentation purposes. Can contain any strings including line breaks.

Note: With dedicated API functions (see section 7.2.1), the reaction and species notes can also be used to store generic (structured) data for reactions and species/metabolites if they are not covered by the default CNA fields. Those data are partially automatically extracted from SBML models (e.g., geneProductAssociation of a reaction) but they may also contain other (user-added) data (e.g., ΔG_R^0 of the reaction). The data are serialized in the reactions or species notes and can later be retrieved again.

Attributes of *reactions*:

- 1) **identifier** of this reaction without any blank
- 2) **symbolic reaction equation**: the metabolite identifiers can be used for compiling the reaction equation. The equal sign “=” denotes the reaction arrow. Identifiers, coefficients and equal sign must be separated by at least one blank. The stoichiometry can be defined in the usual way. An example (where NAD and ADP are not modeled explicitly):
$$G3P = PG + ATP + NADH$$

If a metabolite (e.g. Pyruvate (identifier Pyr)) is taken up or excreted the equation reads:
= Pyr or Pyr =
- 3) **default value**: this is the default value for the reaction rate (example: a reaction or pathway that is considered only in special cases could be assigned a default value of 0). When CNA is started all default rates are set in the associated text boxes. Of course, during a session, the rate values can be edited in the text boxes whenever the user wants to. Keep in mind that reaction rates have always the unit [mmol/(g dryWeight * h)], except the rate of *mue* ([1/h]). The default value “#” denotes that no default rate for this reaction is defined.
- 4) **minimum**: is the minimum of the reaction rate, e.g. 0 if the reaction is irreversible. This value is needed for consistency checks, optimization and calculation of elementary modes. If this value is unknown use a high (negative) value (e.g. -100) or even Infinity (“-Inf”). Note that the minimum value must be smaller or equal to 0.
- 5) **maximum**: is the maximum of the reaction rate. If unknown set a large value (e.g. 100) or even Infinity (value “Inf”). Note: The maximum value must be greater than zero. Thus, a reaction can not be defined in such a way that it can be driven only in reverse direction (e.g. minimum < 0 and maximum <= 0).

- 6) **Coefficient for objective function:** is the (default) coefficient of the linear objective function for optimization procedures (note: the objective function minimizes). Usually, most coefficients are zero, only some reactions have a coefficient unequal to zero. For maximizing the growth rate set this coefficient to -1 for *mue*.
- 7) **Variance:** variance of measurements of this reaction rate is given to be used in metabolic flux analyses. Whether the given value is a relative or absolute variance can be defined prior the analysis of redundant systems (section 3.3: „Procedures for redundant systems ...“).

Each reaction has an associated text box which must be arranged on one of the available network maps. Four parameters define (i) the position of the text box, (ii) the map number and (iii) whether the text box is editable or not. Size and color of the text box are basic properties of the complete network project and can be defined by editing the general project properties (see section 2).

- 8) **x-position** of the associated text box within the network map
- 9) **y-position** of the associated text box within the network map
- 10) **number of the network map** in which the text box has to be arranged
- 11) **text box style:** is “editable” or “non-editable”. The latter one could be useful if one rate depends only on a certain other rate and should therefore not be simultaneously editable by the user. However, calculated values will be displayed also for non-editable text. For instance, anabolic fluxes depend mostly only on biomass composition and growth rate. Because their rates cannot be measured directly it makes sense to define these fluxes as “non-editable”.

Usually, each reaction text box should be placed on the associated pathway (reaction arrow) in the network map. The position is related to the pixel position (in pixel-oriented graphics) or to the relative position (in MATLAB-graphics, *.fig). How the x/y-coordinates can be easily obtained is explained in section 3.2.

Note: If you don’t want to create/display a text box for a certain reaction you can define large negative coordinates for this text box. MATLAB places the text box then virtually outside the map.

- 12) **notes/comments/structured data:** for documentation purposes. Can contain any strings including line breaks.

Note: With dedicated API functions (see section 7.2.1), the reaction and species notes can also be used to store generic (structured) data for metabolites if these data are not covered by the default CNA fields. Those data are partially automatically extracted from SBML models (e.g., chemical formula and charge of the metabolite etc.) but they may also contain other (user-added) data (e.g., ΔG_f^0 of the metabolite). The data are serialized in the reactions or species notes and can later be retrieved again.

Note: The user should assign the identifier *mue* (as explained above) for the biomass synthesis “reaction” only if the biomass composition is explicitly modeled (via the “biomass constituent” element). A reaction equation for *mue* is not directly formulated, instead, the cumulative stoichiometry for biomass synthesis is calculated *dynamically and automatically* (prior any computation) by multiplying the current concentration of the biomass constituents and the synthesis equation for the biomass constituents. Therefore it is not necessary to define a reaction equation for *mue*.

As mentioned above, it is **not mandatory** to declare the reaction *mue*, one may also use another

reaction identifier (e.g. *biosyn*) and then assign a cumulative stoichiometry for biomass synthesis. However, in this case, the convenient definition of the biomass composition (and the resulting stoichiometry for biomass synthesis) by defining biomass constituents cannot be used.

Attributes of biomass constituents:

- 1) **identifier:** for an unique identification of this macromolecule (no blanks)
- 2) **full name** of the macromolecule (no blanks)
- 3) **default value** for the relative concentration in the biomass ([g/g dry weight]). This value is shown by default in the associated text box when the network application is started.
- 4) **synthesis equation:** contains the stoichiometries of the consumed metabolites for synthesis of 1 gram of this biomass component (unit: [mmol Metabolite / g biomass component]). With a small deviation (see section 3.2) the equation is formulated by using the metabolite identifiers in the same way as for reaction equations.

To specify the associated text box (containing the value of the relative concentration) for a biomass constituent the same four parameters as explained for the reactions are needed:

- 5) **x-position** of the associated text box in the network map
- 6) **y-position** of the associated text box in the network map
- 7) **number of the network map** in which the text box has to be arranged
- 8) **text box style** (“editable” or “non-editable”)

Attributes of assembly routes

- 1) **identifier of the metabolite** (consumed for synthesis of the biomass constituent (2))
- 2) **identifier for the biomass constituent** (in whose synthesis the metabolite (1) is consumed)
- 3) **x-position**, where the rate has to be displayed in the network map
- 4) **y-position** where the rate has to be displayed in the network map
- 5) **number of the network map** in which the rate has to be displayed

Metabolite (1) must be contained in the synthesis equation of the biomass component (2).

Each time a computation has been finished, CNA checks, whether the rate of *mue* (the growth rate) is known. If so, the efflux of metabolite (1) in biomass component (2) is displayed on position (3)/(4) in the network map (5). These text boxes always have the style “non-editable” because any user input would not make sense. As mentioned above, every (possible) assembly route is facultative.

So far we have learned how a metabolic or stoichiometric network is modeled in CNA. The external (chapter 5) and internal structure (chapter 7) of a mass flow network in CNA is explained at another place. The next paragraph explains how we can compose a network in CNA.

3.2 Composing and editing the structure of stoichiometric networks

Apart from importing a network via SBML (see SBML import in section 3.3), a convenient way to construct a network in CNA is the *Network composer* (Figure 5) which can be opened via the

menu “Network/Network composer”. The identifiers of the currently defined instances of the four element types are shown as alphabetically ordered lists. The properties of a selected element can be displayed and edited (“Edit/Info”) in a mask. For the declaration a new network element click on the “New” button of the desired element type. A mask comes up where all attributes of the new element can be defined as described in section 3.1. Figs. 6-9 show the masks of the element types; elements of the network shown in Fig. 1 were used as examples. The masks for existing reactions and biomass constituents can also be opened by right-clicking on the associated text boxes on the network map. It is not possible to open multiple editing masks at the same time. Therefore if you open a new editing mask any previously opened mask will be deleted and possible changes discarded.

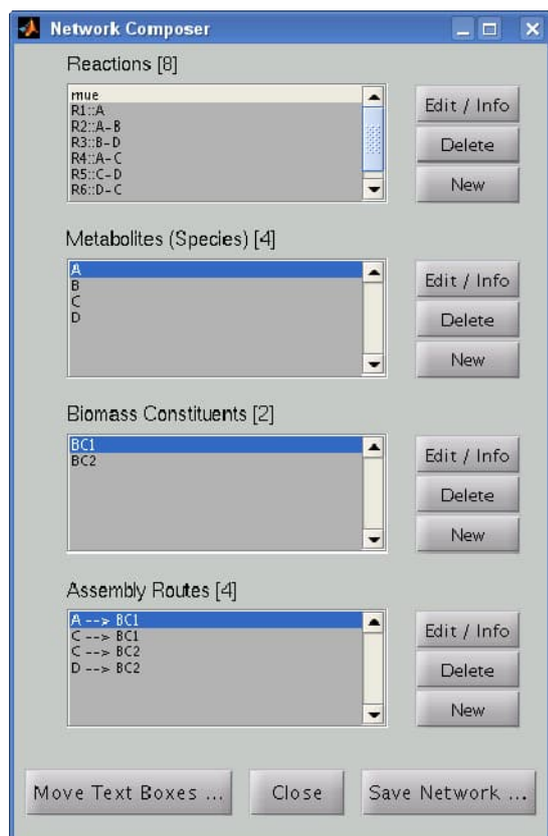


Fig. 5: Network Composer for editing the network structure (here: for example network Fig. 2)

Using the description of the element properties given in section 3.1 it should be no problem to fill out the masks. Some additional hints:

Metabolites: The participation of a certain metabolite in the reactions, synthesis of biomass constituents and assembly routes can be seen by clicking on the corresponding buttons in the metabolite mask (appearing after clicking on “Edit/Info”, Fig. 6). If structured data are contained in the “Notes” field, then they can be edited via the “Edit structured data ...” button.

Reactions: A reaction equation must be defined for each reaction except for the biomass synthesis *mue* (cf. section 3.1). Again, it is important that the rate maximum value is always greater than zero. Thus, a reaction should not be defined in such a way that it can be driven only in reverse direction (e.g. when minimum < 0 and maximum <= 0).

If structured data are contained in the “Notes” field, then they can be edited via the “Edit structured data ...” button.

The x/y-position of the associated text box for the reaction rate can be conveniently defined by clicking on the button “Get x/y-Pos” (Fig. 7) and positioning subsequently the mouse in the window of that network map where the text box is to be arranged. A crosshair appears which can be used to click on the place which is reserved for the text box. The x/y-coordinates (as pixel for pixel-based graphics or as relative coordinates for MATLAB-figures) are automatically inserted in the text boxes “X-Position” and “Y-Position”. After closing the mask (“OK”), the text box will be placed on the defined coordinates.

See below how you can change the coordinates for all text boxes conveniently in one step (button: “Move Text Boxes ...” in the Network Composer window).

Biomass constituents: A (cumulative) synthesis equation must be defined. The syntax is as follows: <coeff1> <MetID1> + <coeff2> MetID2 + ... (without equal sign). For determining the x/y-position of the associated text box (displaying the relative concentration) use the button “Get x/y-Pos” as described above for the reactions.

Assembly rates: For a selected biomass constituent (right pull-down menu in Fig. 9), CNA configures automatically the (left) metabolite pull-down menu, which lists all metabolites that are contained in the synthesis equation of this biomass constituent. Select one to determine a new assembly route. The text position for displaying the assembly rate is specified in the same way as for reaction text boxes.

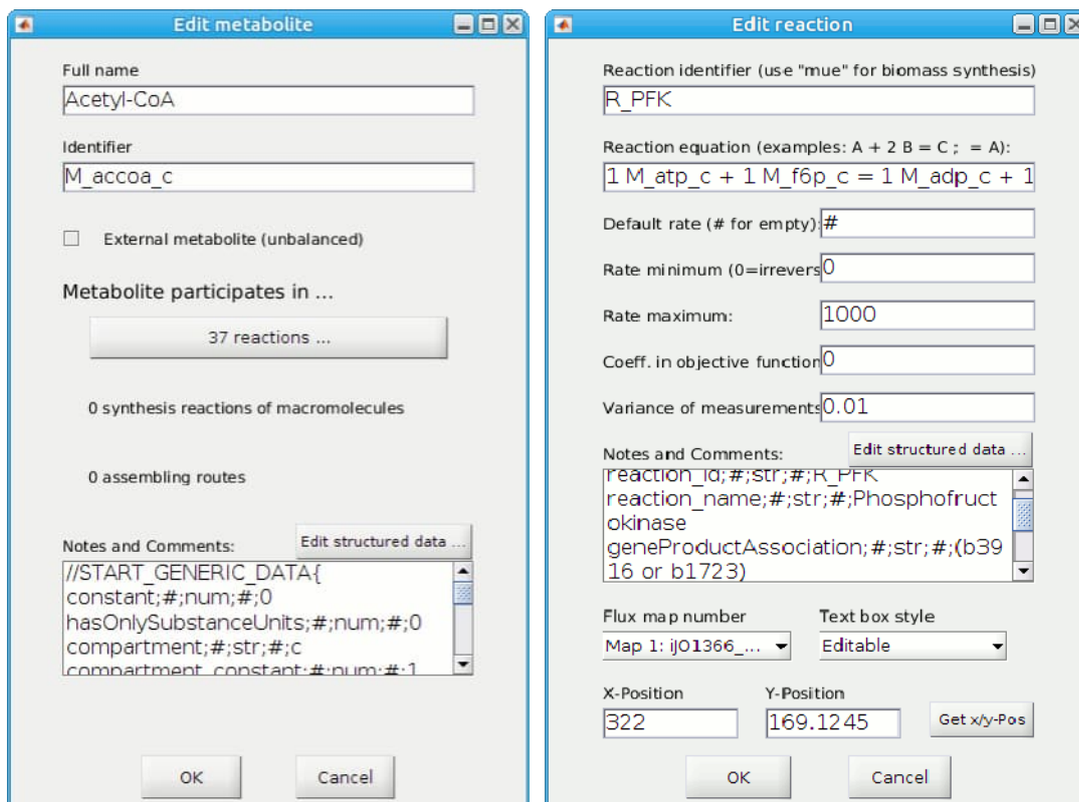


Fig. 6 (left): Mask for editing/declaring metabolites.

Fig. 7 (right): Mask for editing/declaring reactions.

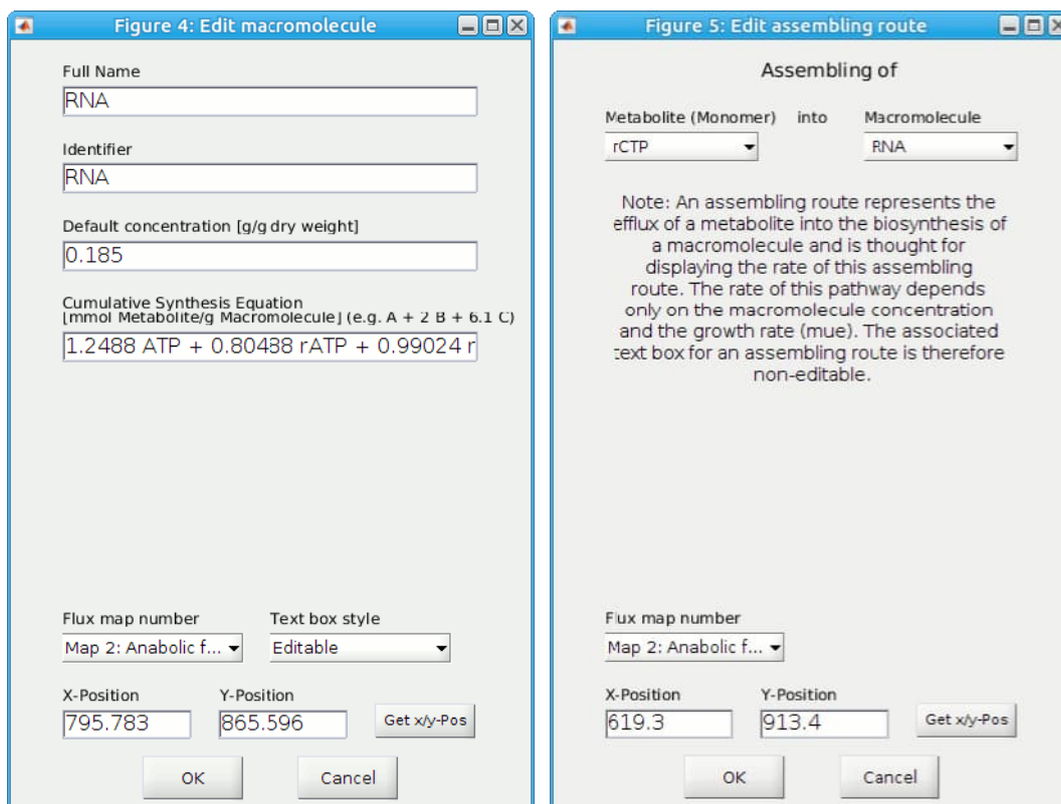


Fig. 8 (left): Mask for editing/declaring biomass constituents

Fig. 9 (right): Mask for editing/declaring assembly routes.

If all network elements have been defined or edited the network structure can be finally saved (“Save”). In case you press “Close”, the modifications are not lost but remain valid only for the current session with this network project. It is also possible to save the network later during the same session by opening the network composer again and then pressing “Save”.

Convenient positioning of all text boxes

The button “*Move Text Boxes ...*” in the network composer window allows convenient positioning and movement of text boxes directly on the network maps. Clicking on this button opens a small window with two buttons: “Accept changes” and “Discard changes”. The user will need this window when the rearrangement of the text boxes has been finished. Each text box (of an arbitrary map) can now be selected and then be moved (drag and drop) to the desired place on the map. It is important to use the LEFT mouse button since by the right button the selected text box will be copied. For selecting a text box of type reaction or biomass constituent one needs to click in the five pixel border of the box which becomes then highlighted. Selecting the text of assembly routes is a bit easier as you can click in the center of the text. When the user has finished the rearrangements he can accept or discard the changes by choosing the respective button in the above mentioned window. After this, the text boxes are “fixed” again and can not be moved anymore. Don’t forget to save the network.

When using this function the user should consider the following points:

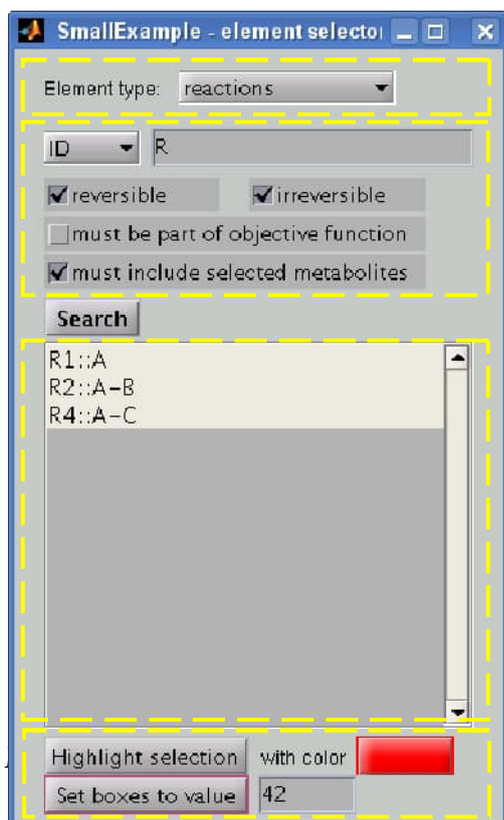
1. If a text box has been selected, it can also be resized. However, this will have no effect in CNA. After finishing the rearrangement (Accept/Discard changes), the text

boxes will be resized to the default size (that can be defined in the project properties, see section 2).

2. If a text box is moved from one network map into another one (or into any other MATLAB window) the text box will be COPIED (the original text box will remain in the original map). However, the new text box won't have any connection to *CellNet-Analyzer* and will thus not be used. If the user wants to move a text box from one network map into another he must change the network map number of the respective network element (see Figs. 7-9).
3. The toolbar of MATLAB should not be used as long as the text boxes are in "movable" state. The CNA menu is also in an inactive state as long as the rearrangement has not been finished. Finally, the user should also avoid changing the zoom factor (via CNA's zoom tools) while the text boxes are movable.

Searching for elements in the network (*element selector*)

Calling "Element selector" from the CNA menu launches a new window for searching elements in the network, highlighting their text boxes and setting values. Although the appearance of this window varies with the network type and the element type you are currently searching it consists of four main parts which are described in Fig. 10. The search string is taken to be a case-insensitive regular expression which follows MATLAB's `regexpi` syntax. This means that when you are searching for non-alphanumerical characters (most importantly `.*[>()`) they have to be quoted with a preceding backslash (e.g. use `\.` to search for a dot). The regular expression functionality is in particular useful for making an or-combination of two search strings using the pipe symbol (`string1|string2`). Leading and trailing white-space of the search string is ignored, but white-space within the search string is significant for the regular expression. If the search string is empty it is interpreted as matching all elements. For more advanced queries using regular expressions please refer to MATLAB's help.



Element type to search, influences the options below.

String search in the selected attribute of the element (can be, depending on the element type, the ID, name or notes); right-click for search history the other search options can be used to restrict the results

Search results; by default, all results are selected, but the selection can be modified by the user

- if IDs is searched they are displayed here
- if names or notes are searched they are displayed with the associated ID prepended to them

Subsets of the found search results can be (de-)selected.
Double-clicking on an element launches a window for editing the element properties (cf. Figs. 6-8).

Text boxes of selected elements from the search results can be highlighted or set to a specified value

irreversible and that include previously selected metabolites (in this case this is metabolite A).

The search results of a metabolites/species search can also be used in a subsequent search: When searching for reactions it is possible to restrict the results to those reactions in which all previously selected metabolites participate (the same can be done with species/interactions). For instance, in Fig. 10 this option is used to only select those reactions that involve metabolite A (which was selected in a previous search).

3.3 Toolbox (menu functions) for analyzing mass-flow networks

Each interactive network map displays *CellNetAnalyzer's* menu bar (see Figure 1) where you can open pull-down menus. If a menu function has been started it is recommended to pay attention to both the results displayed directly in the text boxes on the maps and to the output in the command window.

The menu bar for mass-flow networks has six entries (see also Fig. 1):

- **Network:** Editing/Saving/Importing the network model
- **Map:** Map generation tool, zooming tools and saving a map (with current values) as graphic
- **Scenario:** Saving/Setting/Loading flux scenarios
- **Clipboard:** Flux clipboard and arithmetic operations
- **Analysis:** toolbox of functions for the analysis of the network (see Fig. 11).
- **Info:** info about CellNetAnalyzer

In the following, all menu functions are described in detail (ordered by the menu groups):

3.3.1 Menu "Network": Editing/Saving/Importing the network model

Network / Network composer ...: This opens the main window for editing and saving network elements and network structure as it was described in detail in sections 3.1. and 3.2.

Network / Save ... : Saving the network structure. Note that saving the network structure (all network elements and its properties) always overwrites the old network version. If you want to maintain the older version, please use the "Clone" function (see section 2) from the network project manager before you save the new version.

Network / Export: the network model can be exported in four different formats:

Export stoichiometric matrix ...: Allows one to export the stoichiometric matrix (together with the identifiers of the reactions and metabolites) for further analysis outside of the *CellNetAnalyzer*. The user can choose whether the matrix is to be exported as ASCII or MATLAB file and whether the external metabolites should be included or not.

In ASCII format the matrix is saved as follows:

first row: <reaction names>

second row: reaction reversibilities (0 = reaction irreversible, 1 = reaction reversible)

third row <name of first metabolite> <first row of stoichiometric matrix>

fourth row: <name of second metabolite> <second row of stoichiometric matrix>

....

(m+2)-th row: <name of the m-th metabolite> <m-th row of stoichiometric matrix>

If the matrix is stored in MATLAB format then four variables are saved which can be reloaded in MATLAB later on:

metabolite_names: names of the m metabolites (char array)

reaction_names: names of the q reactions (char array)

reaction_reversibilities: reversibility of each reaction

stoichmat: stoichiometric matrix (m x q) where the m rows are the metabolites in *metabolite_names* and the q columns are the reactions in *reaction_names*

Export in SBML format: Exports the network (reaction + metabolite names, stoichiometric matrix, reaction reversibilities, flux bounds and objective function) in SBML (Level 3, version 1) format including the fbc (flux balance constraints) version 2. The user may choose name and location of the SBML file to be generated. Note that all internal metabolites (without “external flag”) are put into a compartment called “Internal species” and all external metabolites into compartment “External species”. Note that the reaction and metabolite ID’s are converted to valid SBML ID’s when saving. This means all illegal characters are replaced with a ‘_’ and all metabolites/reactions are prefixed with the letter ‘S’/‘R’ respectively followed by a consecutive number.

Export in COBRA toolbox format: Translates the CNA network project (reaction + metabolite names, stoichiometric matrix, reaction reversibilities, flux bounds and objective function) to a MATLAB structure representing a model that can be analyzed with the COBRA toolbox. The user may choose name and location of a MATLAB file in which the COBRA model will be saved.

Export in METATOOL format: Exports the network (basically the species and the reactions including their stoichiometry and reversibility) into a file which can be read by METATOOL. Note that in the output file reactions can occur which have no substrates or no products. This can happen when there are undeclared external metabolites in the network. In such cases, the METATOOL parser may show a warning about incorrect syntax, but this can be ignored because external metabolites don’t enter the stoichiometric matrix anyway and therefore don’t affect the calculations.

(See also chapter 7.4: Importing/Exporting from SBML and METATOOL).

Network / Import SBML model ... : Imports an SBML model (from an .xml file) via the libSBML/SBMLToolbox parser (the required files of the SBMLToolbox are included in the CNA distribution; see also <http://www.sbtoolbox.org/>). Generally, if you want to create a new CNA project from an SBML file proceed as follows:

- 1.) Create a new CNA project with the project manager (section 2). There, you can also specify which network map(s) you want to use. If you do not have such a map, just choose “CNA.pcx” from CNA’s root directory.
- 2.) Load the network project, which is empty at this stage.
- 3.) Start “Network/Import SBML model ...” and select the SBML file. You are then asked whether you want CNA to automatically generate basic maps where the reaction names (and optionally equations) are listed and the text boxes automatically positioned behind their names (see also menu item Maps/Generate basic maps ...).. After conversion, the network is saved and reloaded, either with the automatically generated networks maps or with the original map where the text boxes for the reactions are arranged (by default) in the upper right corner. In the

latter case you may then move the text boxes to their respective position on the network map (via Network composer; see section 3.2).

SBML models can also be imported via API functions (see section 7.4).

Network/Element Selector ...: Tool for searching network elements fulfilling a given set of properties. Explained in detail in section 3.2 (Fig. 10).

Network/Find reaction ...: Tool for searching the text box of a specific reaction. Enter the ID of the reaction (supported with autocomplete; see section 0.6) and after clicking on “Find” the respective map, where the text box of this reaction is placed, comes to the foreground and the box is blinking in red for a few seconds.

Network/Show all reactions equations: Displays all reaction equations in the command window (including the biomass synthesis equation based on the currently defined biomass composition). The reversibility is indicated for each reaction and the defined flux bounds (reacMin/reacMax) are also shown.

Network/Show names of network elements: In each text box, the name of the associated network element (reaction / biomass constituent / assembly route) is shown. This function is especially useful when moving text boxes (“Move Text Boxes ...”; see network composer in section 3.2).

The names of reactions and biomass constituents are also shown as tooltip strings when moving the mouse over the respective text box.

Network / Set epsilon and flux display format ...: The user may change the *epsilon* value (smallest number greater zero) and the display format, i.e. the number of digits displayed after the decimal point in the text boxes. The epsilon value is required for numerical calculations, e.g. for computation of elementary modes. Note that the epsilon will only be changed for the current session, not permanently. To change it also for later network session you must change it by editing general project properties; see chapter 2.

3.3.2 Menu “Map”

Map/Generate standard maps ...: Provides a convenient way to generate some basic interactive networks maps from the network structure. These basic maps list reaction names and (optionally) reaction equations. This is especially useful for larger (e.g. genome-scale) models that have been imported from an SBML file where the manual creation of network maps would be a tedious task. Even though only a list of all reaction names is displayed (no true network graphics), this feature is pretty useful as this representation as it eases the analysis of the model with the GUI. The maps will be generated from the network model and the text boxes automatically be positioned behind their corresponding reaction name. The project will be reloaded with the new maps. (Note that the original network file will not be deleted.)

Place reactions on separate map ...: Reactions can be specified whose text boxes will be relocated to a separate and automatically generated map. The reaction identifiers should be provided in the text box and separated by commas or spaces (e.g. “PFL, PDH”). Optionally, the

generated map shows also the reaction equation for each reaction. This feature is controlled by a checkbox. This feature is helpful, for example, when user-defined pathways were added to a network (without a suitable flux map) or if, in a very large network, the user wants some reactions to be concentrated at a separate map.

Extend the network with GPR-rules ...: If an MFN project holds information about the gene-protein-reaction-association for each reaction (stored in the serialized reaction data under the keyword “geneProductAssociation”; see section 7.2.1 and the iML1515 project as an example)), this feature can become accessible in the Maps menu.

The network can be extended with a stoichiometric representation of the GPR ruleset and maps and boxes for each GPR-pseudoreaction are generated automatically. This can be used for analyzing the effects of gene knockouts (or different expression levels) on the metabolism.

Map/Set original map size: Each network map is resized to its original resolution (pixel-based graphics) or the original size (MATLAB figures). The maps should then appear well-displayed. Ideally, the network graphics should have a resolution not larger than the monitor resolution.

Map/Zoom tools on/off: Displays the zoom tools in the network map in which this menu item has been selected (or the zoom tools disappear if they are currently visible). The zoom tools are always mounted in the upper right corner of the map. The scrollbars allow one to scroll the map left/right and up/down, respectively (note: sometimes, the scrollbars are not correctly displayed by MATLAB; you should then click somewhere in the map or just minimize and then maximize the window). The ‘+’ button enables to zoom in. Press the ‘+’ button and then select two points on the map using the cross-hair. The rectangle spanned by these two points will be the new region displayed in the window. The text boxes are also zoomed with the respective factor. By pressing the ‘-’ button one can zoom out (factor of 2). Finally, by pressing the ‘R’ (reset) button the entire map is displayed again. Note that the assignment of text box positions can also be done in the zoomed state.

Pan (select from toolbar): Can be used to activate the pan mode of the figure. When the pan mode is switched on you can click and drag the network map around inside the window. This will only have an effect if a part of the map is currently displayed (e.g. after zooming in). Due to limitations of the MATLAB GUI the textboxes are hidden while dragging the maps. Pan can be used in conjunction with zoom but then zoom must be activated first.

Map/Save map in graphics file ... : The map within which this menu item is selected can be saved as a graphics in a selectable format (e.g., EPS, bitmap graphics). The map will be saved together with the text boxes and the currently displayed values. You should not save the map as a MATLAB figure!

3.3.3 Menu “Scenarios”: Saving/Setting/Loading scenarios

The **Scenario** sub-menu provides several functions for saving, setting and loading scenarios. A scenario is the set of known numeric values of reactions (usually the reaction rates) and biomass constituents (usually the biomass proportion) as displayed in the respective text boxes.

Scenario/Clear all values: In each text box referencing a reaction or assembly rate the string “#” (meaning “undefined”) is displayed. The values for macromolecule concentrations remain unchanged (because they must have a value prior starting a calculation from the menu).

Scenario/Reset last scenario: The values defined prior the last calculation are displayed in the text boxes.

Scenario/Set default scenario: The default values for the reaction rates and the relative concentration of biomass components are displayed in the text boxes. If no default value exists for a certain network element the string “#” is displayed.

Scenario/Combine current with default scenario: Same as “Set default scenario” but with the difference that all flux values currently given in the boxes are maintained (and the default scenario fluxes are added to the scenario). If the default scenario contains a flux value for a reaction, for which a flux value was already defined in its box then the latter is maintained and not overwritten with the default value.

Scenario/Save scenario (fluxes only): Opens a dialog box for saving the currently shown text boxes values (reaction rates, biomass proportions) in a file. By default, the file is saved in the directory of the current network project and should have the file extension “val” (“values”).

Scenario/Save scenario (fluxes, flux bounds, objective) : Opens a dialog box for saving the currently shown text boxes values (reaction rates, biomass proportions) TOGETHER with the flux bounds and the objective function in a file. By default (but not mandatory), the file is saved in the directory of the current network project and should have the file extension “val” (“values”).

Scenario/Load scenario: Loads a scenario formerly saved by one of the two “Save scenario” functions (see above). A window opens for selecting a scenario file (by default with extension “val”). The loaded values are displayed in the text boxes and are highlighted in the same way as done by Scenario.../Highlight values (see below). Importantly, if the scenario contains also flux bounds and an objective function, these values will be overwritten in the model!

Note: problems may occur if names of network elements (reactions or biomass constituents) have been changed after saving the scenario. Network elements which have been added do not cause problems (they will get value “#”, i.e. “not defined”).

Scenario/Load scenario and combine with current scenario: Same as “Load scenario” but with the difference that all flux values currently given in the boxes are maintained (and the fluxes of the loaded scenario are added to the given (shown) fluxes). If the loaded scenario contains a flux value for a reaction, for which a flux value was already defined in its box then the latter is maintained and not overwritten with the loaded value.

Scenario/Highlight values (on/off): This function colors the reaction text boxes according to the numerical values displayed in them. Text boxes with a zero value are indicated by color “special” and those with non-zero values by color “computed”. Boxes with non-numerical values remain in standard color. This function is useful, for instance, to quickly identify zero fluxes in the network.

Scenario/Highlight values (heatmap): This function is similar to the previous one but uses the *heatmap* style to highlight the reaction text boxes. The user has to define a threshold value which defines the ‘zero’ or ‘neutral’ value. For all text boxes with a larger value, the text box color becomes green; the saturation of the green hue is determined by the ratio
(text box value)/(max_value_of_all text_boxes – threshold value).

Similarly, for all text boxes having a smaller value than the threshold value, the text box color

becomes red; the saturation of the red hue is determined by the ratio:

$(\text{text box value})/(\text{threshold value} - \text{min_value_of_all_text_boxes})$.

(Boxes with non-numerical values have standard color.) One may use this function, for instance, to display and highlight gene expression data of metabolic genes within a network context.

Scenario/Show current values in bar chart: Displays the values currently defined in the reaction text boxes in a bar chart. This is useful for comparing the fluxes of a given scenario and to identify the greatest absolute values. The function is also valuable for evaluating the results of arithmetic operations.

3.3.4 Menu “Clipboard”

Clipboard/Copy values to flux clipboard: Works like a usual clipboard: The currently defined values in the text boxes of reaction rates are copied to a clipboard (values of macromolecule (if those exist) are not copied!).

The following two functions can be applied to the values of the flux clipboard.

Clipboard/Paste values from flux clipboard: Writes the values from the flux clipboard into the text boxes.

Clipboard/Arithmetic operations ...: Useful for arithmetic linkage of two flux distributions, for example for computing the difference between two flux distributions. The four basic calculation operations are available for linking the values currently displayed in the text boxes with the values from the flux clipboard. Alternatively, arithmetic operations can also be performed between a number and a flux distribution (e.g.: normalization of a flux distribution on the substrate uptake rate). The result supersedes the old values in the text boxes.

3.3.5 Menu “Analysis”

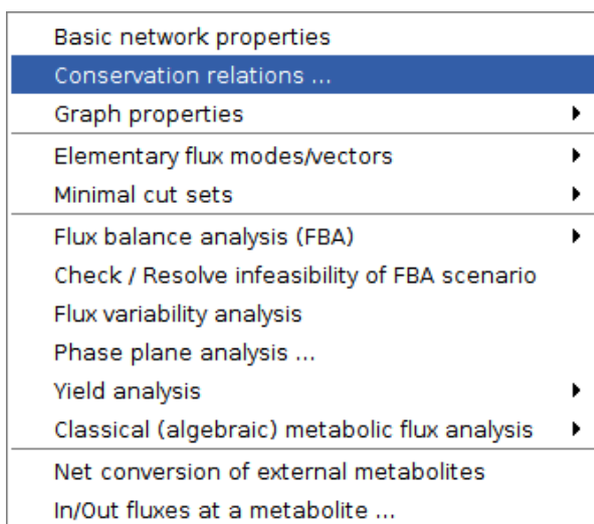


Fig. 11: The CNA sub-menu “Analysis” for mass-flow networks.

Analysis/Basic network properties: Calculates global topological properties of the network and displays them in the command window. This function is also useful to detect errors in the network structure after the network has been composed. Note that external metabolites are not considered when determining these properties.

- 1) Internal metabolites connected to no reaction and reactions without any participating internal metabolite are detected.
- 2) *Sinks and Sources:* Metabolites which can only be produced or only be consumed (dead-ends) are displayed.
- 3) *Simple uptake/excretion of one metabolite:* useful to detect transport (uptake/excretion) pseudo-reactions.
- 4) *Blocked (strictly detailed balanced) reactions:* The rates of such reactions are always zero in steady-state and are thus of no importance for many stoichiometric studies (for example, blocked reactions are never involved in any elementary mode). An example: the rate of a reaction in which a dead-end metabolite participates is always strictly detailed balanced (and must have the value zero to maintain the metabolite in steady state). Note that the identification of blocked reactions is done here by null-space analysis.
- 5) Reactions with *empty stoichiometry* (where none of the internal metabolites participates) are displayed.
- 6) *Enzyme subsets:* In each flux distribution obeying the steady state condition all reactions of one enzyme subset have always either zero flux or non-zero fluxes with fixed proportions.
- 7) All groups of *parallel reactions* (reactions having the same stoichiometry (often from isozymes) are displayed. Two reactions with reverse stoichiometry (reverse direction) are also considered to be isozymes if at least one of these reactions is reversible.
- 8) The linear objective function (required for the “Flux balance analysis”) –as currently defined– is displayed.
- 9) *Stoichiometric matrix:* Number of internal metabolites (rows) and reactions (columns) is given. The rank of the stoichiometric matrix is computed. If the rank is lower than the number of (internal) metabolites then conservation relations exist (linearly dependent rows) usually caused by „conserved moieties“. That means that the balance of (at least) one metabolite is implicitly contained in the balance of the other ones and can therefore be expressed as a linear combination of other balances. For instance, a simultaneous balancing of NAD and NADH usually leads to such a priori redundancy because the balance of NADH is the same as the balance of NAD multiplied with -1. Conservation relations can be eliminated by the removal of one (or more) metabolite(s) contained in the elementary conservation equations. The latter ones can be calculated and displayed by a separate menu function (mentioned below).

Next, *CellNetAnalyzer* computes a compressed version of the current network. This compressed network would have the same basic properties with respect to steady-state flux distributions (e.g. an equivalent set of elementary modes). In the compressed network, for example, all identified strictly detailed balanced reactions would be removed. The dimension of the compressed network tells you how complex your network really is. The number of reactions in the compressed network could be further reduced if parallel reactions (emerging, for example, by isozymes) are considered only once. Note that the network compression procedure is also employed before computing elementary modes.

Analysis/Conservation relations: This procedure calculates the *elementary* conservation relations (ECR) and displays them in the command window. Conservation relations occur in networks where the rank of the stoichiometric matrix is lower than the number of metabolites (see notes above: menu item “Basic topological ...”) and may be seen as redundant balance equations. Calculating the *elementary* conservation relations one obtains all non-decomposable conservation relations similarly to *elementary flux modes* (see reference [8]). The ECRs are

useful for detecting conserved moieties but also to find those metabolites, which are involved in conservation relations. The user is requested whether he wants to compute all ECRs or only the non-negative ECRs (which represent conserved moieties, see [8]), and he has also the option to use *CellNetAnalyzer*'s and/or METATOOL's MEX files for computing the ECR's (accelerates the computation). Regarding MEX files see also section 0.4 and under menu item "Elementary modes and pathway analysis ..."

CNA provides two functions for analyzing graph-theoretical properties of the stoichiometric network.

Analysis/Graph properties/Connectivity histogram: The distribution of the connectivities (connectivity = number of reactions in which a metabolite participates) is displayed in a histogram.

Analysis/Graph properties/Graph-theoretical path lengths ...: This enables the calculation of *graph-theoretical* (shortest) path lengths between metabolites (plus the average path length and the network diameter). Note that reaction networks are usually *hypergraphs*. Therefore, for applying graph-theoretical methods, this hypergraph must be transformed into a bipartite or substrate graph. Note also that the results from graph-theoretical paths analysis can differ considerably from pathway analysis using elementary modes (see below). For example, a graph-theoretical path from metabolite A to B can exist but, nevertheless, there might be no way to produce B from A. However, graph-theoretical path lengths are – compared with elementary modes – quickly computable and might be seen as a very rough characterization of routes in the network.

Prior to the computation, the user can choose several options in the appearing window:

- *Exclude external metabolites:* If this has been chosen, then connections via external metabolites are not considered, i.e. these nodes are considered as not existing in the network. (This also means that the path lengths between external and other metabolites are not calculated). This will usually increase the paths lengths.
- *Use directed graph:* If this option is selected the reversibility of the reactions is taken into account by using the rate minimum (see "Set constraints" and section 3.1), hence the directionality of the edges is then considered. In such a case, the shortest path length from A to B is, in general, not the same as that from B to A.
- *Display distance matrix:* Displays the shortest path lengths (distances) between each metabolite pair as a colored matrix. The larger the path lengths is the brighter the matrix cell. A blue color marks metabolite pairs, between which no path exists. In an undirected graph, the distance matrix is symmetric. In directed graphs, this is not necessarily the case. Then, the matrix element e_{AB} represents the shortest path length from A to B. In very large networks, the creation of this graphic representation of the matrix might fail in MATLAB. Use then:
- *Export distance matrix:* This allows for an export of the calculated shortest path lengths into an ASCII file. The user can then further study the path lengths matrix by its own algorithms and tools.
- *Exclude reactions whose rate is given:* All reactions having an arbitrary number (e.g. 0 or 1) in their associated text box are considered as being not existent during the calculation. This enables to hide certain reactions when calculating the path lengths.
- *Compute path lengths only for these metabolites:* This enables the user to select only some metabolites (using the corresponding identifiers) for which then the path lengths to all other metabolites are calculated. If, for example, metabolite 'A' is defined in this row, then only the path lengths from A to all others are computed. Several

metabolites may be defined here (separated by a blank). If no identifier is given, the shortest path lengths between all metabolite pairs are calculated.

After computation, depending on the chosen options, the distance matrix is shown and/or saved. Besides, the average path lengths, the network diameter (longest shortest path length) and the number of connected components in the graph are displayed in the command window.

Analysis/Elementary flux modes/vectors : This sub-menu opens the door for pathway analysis: elementary flux modes (elementary modes), elementary flux vectors (generalization of elementary modes in flux polyhedra) or a minimal generating set (convex basis) of the flux cone/polyhedron can be calculated and analyzed. Elementary modes are the support-minial vectors of a flux cone (flux cones arise by the steady-state and reversibility constraints) whereas additional inhomogeneous constraints (equality and inequalities; e.g. upper and lower flux bounds different from zero) give rise to flux polyhedra in which elementary flux vectors (short: elementary vectors) can be calculated. Elementary vectors generalize elementary modes and have similar properties and applications [23,29]. Both elementary modes and elementary vectors are a generating set of the flux cone or the flux polyhedron, respectively (i.e., any flux vector in the cone/polyhedron is a suitable non-negative linear combination of elementary modes/vectors). For this reason, they contain all extreme rays and (only for elementary vectors) all extreme points of the flux cone/flux polyhedron. However, elementary modes/vectors are, in general, not a *minimal* generating set (convex basis) and CNA therefore provides also a routine to compute a convex basis only.

Accordingly, two different scenarios can be considered: (i) In the *homogeneous* case (the standard case) elementary modes or a convex basis of the flux cone can be calculated. Note that whereas the elementary modes are unique (up to scaling), a convex basis is not if the cone is not pointed (has a lineality space). (ii) In the *inhomogeneous* case, inhomogeneous constraints (e.g., reaction rates fixed to a non-zero rate or/and upper and lower bounds (different from zero) for reaction rates) can additionally be considered giving rise to a flux polyhedron. There are two different ways to include inhomogeneous constraints in the calculations: they can either directly be taken from the specified min/max reaction rates OR they can (temporarily) be specified for (some) reaction rates within the text boxes (see the first two options under “Compute modes” below). With inhomogeneous specifications, the elementary vectors or a convex basis of the resulting flux polyhedron can be calculated. Note that then unbounded and bounded elementary vectors may exist (while elementary modes, as the whole flux cone, are always unbounded). Further, the zero point will not be delivered separately, even if it is an extreme point of the solution space. So far, most applications in the literature focus on elementary modes in the homogeneous setting and we will therefore mostly speak about (elementary) “modes”, although the same applications are, in principle also feasible for elementary vectors in flux polyhedra. Also, when analyzing elementary modes/vectors within CNA, the calculated vectors are often named as “modes” although it could also be elementary vectors.

For more information on elementary modes and elementary vectors see reference [23,29].

The sub-menu has three entries:

Load ...: Allows one to load modes saved in earlier sessions. Select a proper file (CNA saves modes in MATLAB files with extensions “mat”). Warning: modes should be newly calculated if they have been saved before the network structure has been modified. Otherwise, an error message could appear.

In case that a set of modes has already been loaded or computed, the user is asked whether the

set of modes to be loaded has to be combined with current set of modes (or whether the current set of modes has to be discarded). This enables to compute certain sets of modes independently and to merge them later.

Show ...: elementary modes which are still in the memory (loaded or calculated earlier in the current session) will be displayed again.

Compute ...: A dialog box comes up for specifying several options for computing elementary modes / elementary vectors / convex basis (=minimal generating set):

- *Consider constraints given in text boxes:* This option allows one to (a) *exclude* reactions or/and to (b) *enforce* the involvement of certain reactions or/and (c) to specify inhomogeneous constraints on reaction rates (see constraints (3)-(5) below). If this checkbox is activated, the following specifications in the reaction text boxes will be considered prior the computation:

(1) All reactions having a zero in their associated text box are considered to be off or inactive, hence, only those elementary modes (or vectors) will be computed that do not use this reaction (these reactions will have a 0 rate in all resulting elementary modes/vectors).

(2) In contrast, each reaction which has a numerical value unequal to zero in its corresponding text box is enforced to be involved in the modes to be computed. [Note that enforcement of reactions is not possible for computing convex bases (but exclusion is); see below.] Thus, this option can be used to compute a particular subset of all elementary modes of the network. The computation time may considerably be reduced by excluding reactions. Mostly, enforcing reactions does also have a positive effect on running time, but sometimes it may even take longer than computing the complete set of modes (without enforcing reactions) although this is a superset of those modes which involve all enforced reactions. **IMPORTANT:** Do not use EFMtool together with this option!!!!

(3) One may enforce equalities on reaction rates. For example, if a reaction rate should equal 3 then write “= 3” (with space between “=” and “3” !!) in the corresponding text box. Note the difference with constraints of type (2) – in the latter case, a non-zero value (without “=”) only demands that the reaction must be active in the modes to be computed, but no specific value is enforced.

(4) One may set lower boundaries for reaction rates, e.g. by writing “>= 5” (again with space between “>=” and “5”) in the text box.

(5) One may set upper boundaries for reaction rates, e.g. by writing “<= 7” (again with space between “<=” and “7”) in the text box.

Note that constraints (4) and (5) must explicitly be set in the text boxes, the standard min/max reaction rates (except for marking irreversible reactions will not be considered here (for doing so see option “*Use flux bounds to calculate elementary flux vectors*”). One may also combine constraints (4) and (5) (but no others) within one text box, e.g. “>= 1 <= 4” will constrain the respective reaction rate within the range [1,4]. Consistency should be ensured (e.g. “<= 1 >= 4” makes no sense and “<= -2” is not allowed if was rate minimum attribute of the reaction is larger than -2). Furthermore, one may set constraints (1)-(5) for arbitrary many reaction rates. One should not use constraints (3) for a value of “0”, as this can be better achieved by constraint (1).

As mentioned, any constraints of type (3), (4) or (5) render the problem to be inhomogeneous and the solution space to be a (flux) polyhedron. Note that, in the

inhomogeneous case, the returned vectors will be an unsorted mixture of bounded and unbounded generators which will be indicated when displaying the modes (see Fig. 12). These flags can also be found after the calculation in the variable “<projectname>.local.ray”.

If the option (checkbox) “*Consider constraints given in text boxes*” is not selected then all elementary modes/vectors (irrespective of constraints given in the text boxes) will be calculated.

One may also use the API function `CNAcomputeEFM` to calculate elementary modes or vectors (see chapter 7).

- *Use flux bounds to calculate elementary flux vectors:* If this box is checked, then the standard min/max reaction rates will be included as inhomogeneous constraints (hence, this will lead to calculation of elementary vectors). Flux bounds (e.g. lower flux bounds) of zeros and of “Inf” (infinity) will not be considered separately as they practically represent homogeneous constraints. Consequently, the number of inhomogeneous constraints and, hence, the complexity of the whole problem may increase considerably if many min/max reaction rates have values different from zero or Inf. Therefore, if this box is ticked, after starting the computation, CNA will allow you to set a threshold for the absolute (!) value of flux bounds above which flux bounds are considered as irrelevant. For example, in many metabolic models, upper bounds for reaction rates are set to an arbitrary large value (e.g., 1000) and the same for lower bounds of reversible reactions (-1000). To consider only absolute flux bounds below these values you may set, for example “500” as threshold. Then, in the example above, 1000 and -1000 will be treated as Inf and -Inf, respectively. If you really want to integrate all flux bounds as inhomogeneous constraints you may choose the default value “Inf”.

Note that the inhomogeneous constraints of this option can be combined with those from *Consider constraints given in text boxes* detailed above however, constraints set in the reaction text boxes have higher priority, hence, will overwrite min/max reaction rates.

- *Check reversibility:* If this option is selected (this is the standard case) the reversibility of the reactions is taken in account via the minimal reaction rates (see section 3.1 and 3.2). If this box is not checked, there will be no sign restrictions on the rates. This may drastically increase the number of modes but is useful for some applications in smaller networks.
- *Only a convex basis:* If this box is ticked, CNA computes only a “convex basis” (also: “minimal set of generating vectors”) of the flux cone / flux polyhedron which (as implemented herein) forms always a subset of the elementary modes/vectors (in a few cases, elementary modes/vectors may coincide with the convex basis). In particular, the convex basis contains all extreme rays and extreme points (in flux polyhedra). Note that the convex basis is not unique if the cone/polyhedron is not pointed (has non-trivial lineality space, which is the space of reversible flux vectors [29]). A convex basis suffices to represent each feasible flux distribution in the network by suitable linear combinations of the convex basis vectors. However, for many applications one needs the full set of elementary modes/vectors. In large networks with many reversible reactions the number of elementary modes/vectors can be several-fold greater than the number of vectors in the convex basis. Note that the

“convex basis” option can also be used to compute Extreme Pathways – provided that the network has been configured properly (see also ref. [4]).

The convex basis cannot be calculated with EFMtool!

- *Consider isozymes only once*: Selecting this option means that from each isozyme group (group of reactions having the same stoichiometry) only one representative is considered for calculating the modes. The other reactions of the isozyme group will then not occur in any of the determined modes but their “equivalent modes” can easily be identified by the modes where the representative reaction occurs (one only has to substitute the representative by another reaction from the isozyme group). Two reactions with reverse stoichiometry (reverse direction) are also considered to be isozymes if at least one of these reactions is reversible. In this case, the reversible reaction is retained. For example, in Fig.1, R5 and R6 would be one isozyme group, R6 would be retained as representative and R5 would be dropped. In this example, the only “true” information lost is the mode (2-cycle) built up by R5 and R6. This option is especially useful in larger (reconstructed) networks. Therein, many isozymes may occur, which then increase drastically the number of modes. Many of these modes would then carry redundant (equivalent) information due to the parallel isozyme reactions.
- *Calculation method*: Currently four different calculation methods are supported:
 1. *METATOOL + CNA MEX files* (default): When this method is selected, *CellNetAnalyzer* uses METATOOLs algorithm for computing elementary modes which proved to be faster than *CellNetAnalyzer*’s routines (one reason is that the main computation is done *completely* externally). The required MEX-file (for WINDOWS: DLL-file) is distributed together with CNA (cf. sect. 0.4). In fact, CNA and METATOOL MEX files are used together with this option.
 2. *EFM Tool*: Currently the most popular method algorithm [15]. However, calculation of the convex basis is not supported and it may fail when too many reactions are enforced (c.f. bugs section on www.csb.ethz.ch/tools/efmtool). Therefore, an error message appears if *efmtool* is selected together with these options. When choosing EFM Tool, an additional checkbox “*Use rational numbers*” appears. If this is selected EFM Tool uses arbitrary length rational numbers during calculation. This yields exact results (at least on the EFM Tool side) but increases memory usage and computation time. For additional information see README.txt in *CellNetAnalyzer/code/ext/efmtool/*.
 3. *CNA MEX files*: CNA’s original algorithm for computing elementary modes also uses MEX files accelerating the computation of elementary modes. If the METATOOL module does not work on your system (for reasons described above), you may try to use CNA’s MEX version (cf. sect. 0.4). If no executable CNA MEX file is found the pure MATLAB version will be used which, however, is slow in large networks.
 4. *CNA functions*: Provided as fall-back option in case the MEX-files do not work.

In general, which version (EFMTool, METATOOL MEX, CNA MEX or CNA MATLAB) is actually used is displayed in the command window after starting the computation.

When the computation has been started the progress of the calculation can be seen in the command window. The algorithm starts with preprocessing steps mapping the original network

to a smaller one with the same final number of modes (e.g. by combining all reactions from an enzyme subset into one overall reaction). Calculating elementary modes/vectors or a convex basis needs information on the reversibility of each reaction which is taken from the rate minimum (if rate minimum < 0 then the reaction is considered to be reversible, otherwise not). It is important that the rate maximum value is always greater than zero. Thus, a reaction should not be defined in such a way that it can be driven only in reverse direction (minimum < 0 and maximum ≤ 0). If METATOOL is used, METATOOL will try to convert the stoichiometric matrix into an integer representation. If this is successful (if you have not too many real numbers in your matrix with many decimal places) METATOOL will employ integer arithmetic (and thus produce exact intermediate results). EFMtool can also be used with rational arithmetic (see above).

The overall calculation time depends on the network structure, on the selected options described above and on your hardware. It can take a few seconds or several days (or even longer ...). Besides, it can occur that the algorithm stops with an error message: “Out of memory”. In this case you have not enough RAM or your network is simply too large to be treated with elementary-modes analysis.

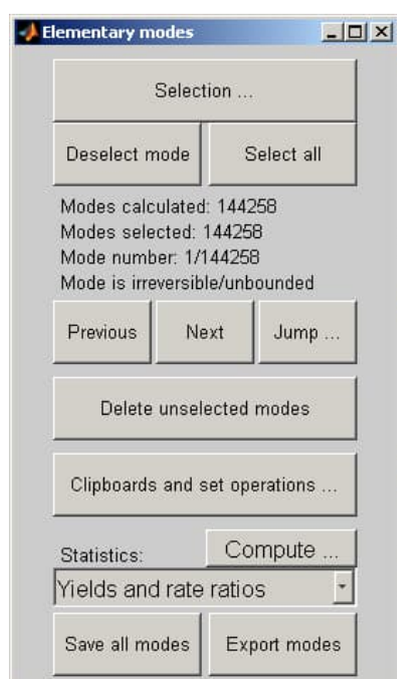


Fig. 12: Control panel for displaying/analyzing computed elementary modes

If the calculation has been finished (or if a set of modes has been loaded) the first mode is shown in the network map and a control panel comes up (Fig. 12; please note again that, in the following, the term “(elementary) mode” may also represent an elementary vector). In the map, all participating reactions of a mode (having a rate unequal to zero in this mode) get the color “calculated” and reactions having a zero get the color “standard” (see section 2). A reaction which does not occur in any mode (for instance when it was excluded by using the option “Exclude reactions” as described above) is designated by the color of type “defined”. Each elementary mode is normalized in such a way that the smallest rate has an absolute value of 1, except if biomass synthesis occurs (μ_{ue} unequal to zero). In the latter case the flux mode is normalized to a growth rate of 0.1. The number of the current mode as well as its reversibility is displayed in the “Elementary modes” control panel (Fig. 12). Again, it depends on the chosen (possibly inhomogeneous) constraints and options (convex basis) whether the shown mode is an

(unbounded) elementary mode or a bounded/unbounded elementary vector. This will also be indicated for the currently selected mode (see Fig. 12). Note that the variable “<projectname>.local.ray” indicates which of the elementary vectors are bounded (0) or unbounded (1).

The “Elementary modes” control panel (Fig. 12) provides the following functions that are useful for changing the currently displayed flux mode, for selecting a certain subset of all modes, for calculation of statistical properties and for saving the computed set of modes:

- *Selection*: This tool allows the selection of a certain subset of all flux modes. First, a panel opens where the following 8 specifications for selecting a subset can be made:
 - first row: reactions which *must not* be contained in any mode (i.e. those reactions must have a rate of zero in the selected modes)
 - second row: reactions which *must* be contained in all modes (i.e. those reactions must have a rate unequal to zero in the selected modes)
 - third row: reactions which may only proceed in backward direction (i.e. those reactions *must* have a rate being negative *or zero* in the selected modes)
 - fourth row: reactions which may only proceed in forward direction (i.e. those reactions *must* have a rate being positive *or zero* in the selected modes)
 - metabolites which must not participate in any mode
 - metabolites which must participate in all modes
 - minimal number of participating reactions
 - maximal number of participating reactions

For each row no, one or several reaction or metabolite name(s) can be defined, respectively (input of reaction / species IDs is supported with autocomplete; see section 0.6). If nothing is specified the complete set of modes is selected (see also button “*Select all*” below).

After pressing the button OK the user is asked whether the selection should be performed on the current selection (if a subset is chosen currently) or on the complete set of elementary flux modes. In the first case the user specifies a subset of the current subset thereby refining the selection to incorporate the new constraints.

For example, if in the first row *mue* is entered the user will get all flux modes without growth. Another example: First row: uptake of oxygen; second row: *mue* - all flux modes which allow anaerobic growth will be selected.

Except button “*Save all modes*”, all functions described in the following operate only on the current selection of modes.

Important: additional options to select elementary modes/rates (namely with respect to their product yields or absolute rates) is provided when selecting “*Yields and rate ratios ...*” or “*Rates ...*” (see below).

- *Deselect mode*: Deselects the currently displayed mode. Thus, a certain flux mode can be deselected separately without using the *Selection* tool.
- *Select all*: Selects the complete set of elementary modes.
- *Next mode / Previous mode*: Switches to the display of the next or previous mode of the set of currently selected modes.
- *Jump ...*: Here, a certain mode can be reached by its number. This mode becomes the current-

ly displayed mode. (Note: the mode number must be contained in the set of currently selected modes).

- *Delete unselected modes*: Removes all unselected modes irreversibly from the memory (in contrast to the *Selection* tool). This tool could be useful if some modes are not of interest and can be dropped.
- *Clipboard and set operations*: Useful for obtaining subsets of elementary modes which can not be generated by the *Selection* tool alone. The user can copy the current selection to clipboard 1 or 2 as well as restore a subset from clipboard 1 or 2 to the current selection. Furthermore an additional window can be opened by “*Set operations*” where set operations (union, difference, intersection) for combining the subsets of the current selection, clipboard 1 and clipboard 2 are provided. After performing a set operation the obtained subset becomes the current selection. Example (cf. Fig. 1): Flux modes for growth on substrate A and D have been calculated. For selecting all flux modes, where substrate A or D is taken up (reaction $R1::A$ or $R7::D$ have rate unequal to zero) select first all modes, where $R1::A$ is unequal to zero (*Selection* tool) and copy this selection (*Clipboard* tool) to clipboard 1. Then select all modes, where $R7::D$ is unequal to zero (*Selection* tool) and combine then (*Clipboard* tool) the current selection with clipboard 1 by set operation “union”. The selection tool would only allow one to get a subset of modes in which $R1::A$ and $R7::D$ participate but not those where only one of these both is unequal to zero.
- *Statistics*: A number of useful statistical features of *the current selection* of modes may be determined by selecting the respective feature from the pull-down menu (Fig. 12) and clicking on button “Compute”:
 - *Yields and rate ratios*: This function operates on the **current selection of modes** (which can be elementary modes or elementary vectors) and, for a user-specified yield function (or rate ratio) Y , it (1) searches for the mode(s) with maximum yield, it (2) shows the yield distribution within the set of selected modes, and (3) it can optionally be used to select modes above/below a specific minimum/maximum yield. The yield Y has to be defined by the user as a linear-fractional (yield) function of the type $Y = \mathbf{c}^T \mathbf{e} / \mathbf{d}^T \mathbf{e}$, where \mathbf{e} is the mode vector and \mathbf{c} and \mathbf{d} are vectors indicating weighting factors for the reaction rates in \mathbf{e} . In typical applications, \mathbf{c} contains a single non-zero entry (usually “1”) for the product (P) excretion reaction and \mathbf{d} contains a single non-zero entry (usually “1”) for the substrate (S) uptake reaction, hence, the yield of P/S would be maximized. However, more general yield functions (with multiple “substrate” and “product” reactions) or rate ratios can be considered as well by choosing appropriate factors for \mathbf{c} and \mathbf{d} . The user will be asked to specify the reaction identifiers (input of reaction IDs is supported with autocomplete; see section 0.6) with non-zero values in \mathbf{c} (“product reaction”) and \mathbf{d} (“substrate reactions”), respectively, together with the respective weighting factor (only weights different from “1” need to be defined; negative weights can also be used, e.g., if a substrate uptake reaction is defined in the direction of excretion thus having negative fluxes for substrate uptake). For example, specifying “R1 2” for the product reactions (nominator) and “R2 R3 2” for the substrate reactions (denominator) would specify the yield function $Y=2*R1/(R2 + 2* R3)$.
Only those modes of the current selection will be considered where the denominator of ($\mathbf{d}^T \mathbf{e}$) is positive and where the nominator ($\mathbf{c}^T \mathbf{e}$) is non-negative. The mode with the maximal yield becomes the current mode and is displayed. If there are two or more modes with the same maximal yield then the first of these is displayed and the numbers

of the others are shown in the command window. Besides calculating the maximal yield, a figure comes up displaying the yield distribution over all selected modes (again: only modes with meaningful substrate uptake / product excretion will be considered).

In addition to the calculations above, modes can be optionally selected whose yield is greater than a definable minimum yield or/and smaller than a defineable maximum yield (these boundaries can be entered in the same window where the yield function must be specified). According to these specifications, all modes of the *current* selection which are in the valid yield range will be selected (modes with negative nominator ($\mathbf{c}^T\mathbf{e}$) or/and zero or negative denominator ($\mathbf{d}^T\mathbf{e}$) become unselected!) The control panel for displaying the modes will be updated when the selection has been made. If neither a minimum nor a maximum yield is specified (empty lines) then the current selection of modes remains unaltered (and the first optimal mode will be selected as described above). Note that if a minimal or maximal yield has been specified for selection, then the figure with the distribution of yields will not be displayed.

- *2D Yield space plot*: From the **current set of selected modes** this function generates a two-dimensional yield space plot displaying the dependency between two yields (yield functions or rate ratios) Y1 (x-axis) and Y2 (y-axis). Yield space plots are useful, for example, to study the trade-off between growth (biomass) yield (Y1) and certain product yields (Y2). A yield space plot also indicates the feasibility of coupled growth and product synthesis [27]. The two yield functions $Y1 = \mathbf{c}1^T\mathbf{r} / \mathbf{d}1^T\mathbf{r}$ and $Y2 = \mathbf{c}2^T\mathbf{r} / \mathbf{d}2^T\mathbf{r}$ must be defined in the same way as explained above for the previous function “*(Product) Yields and rate ratios*”. The yield space plot is easy to interpret if the set of selected modes either contains only unbounded or only bounded elementary vectors/modes [30]. Yield space plots of mixtures of bounded and unbounded elementary vectors must sometimes be interpreted with care because the yield space might be open at some boundaries [30] (a corresponding warning will come up in this case). Bounded (*) and unbounded (o) modes/vectors are indicated by different symbols in the plot.
- *Rates ...*: This function is analogous to “*Yield and rate ratios ...*” but refers here to the *absolute value of a single reaction rate* in all selected modes/vectors. Since unbounded elementary vectors (which include the case of elementary modes) can be scaled to infinity this function makes only sense if the selected reaction rate is exclusively contained in bounded elementary vectors (calculated from an inhomogeneous system [30]). Otherwise a warning is displayed. The elementary vector with the maximal rate becomes the current mode and is displayed. If there are two or more vectors with the same maximal yield then the first of these is displayed and the numbers of the others are shown in the command window. Besides calculating the maximal value of the specified rate, a figure comes up displaying the rate distribution over all selected vectors.
- *2D Phase plane plot*: As for “*Rates ...*”, this function should only be used in conjunction with (bounded) elementary vectors calculated from an inhomogeneous system. It calculates the projection of the solution space (spanned by the **currently selected** set of elementary vectors) onto two selected reaction rates and draws the resulting phase plane [30]. This is frequently used, for example, to study the dependency between growth rate and synthesis rate of a product of interest (then also called production envelope) but is also useful for many other applications. The user has to specify the identifiers of the selected reactions (input of reaction IDs is supported with autocomplete; see section 0.6). If more than two reactions are defined (say the number is x), then all $x*(x-1)/2$ phase planes of all pairs of reactions will be drawn as subplots in one figure. Note that the two

reaction rates must be 0 in all (selected) *unbounded* modes / elementary vectors since otherwise the phase plane would be unbounded as well and can then not be drawn.

- *Control-effective fluxes*: The concept of control-effective fluxes (CEFs) was introduced and explained in [2]. It tells you something about the “importance” of each reaction in the current set of modes and involves a weighting procedure for each mode with respect to (user-)defined objectives. It has even been applied to predict transcript ratios [2]. Clicking on this button opens a dialog box where you have to define (the last parameter is optional):
 - *Objective reactions*: Here one defines the reaction IDs (input of reaction IDs is supported with autocomplete; see section 0.6) which are supposed to be objectives of cellular optimization (e.g. growth rate ‘*mue*’ or ATP production).
 - *Substrate uptake reactions*: Here you can define (arbitrary many) reaction names which represent uptake of ‘sources’ or ‘input’, hence, usually substrate uptake reaction(s).
 - *Reference substrate uptake reactions*: This is similar to the previous parameter but selects a reference ‘source’ or ‘reference input’ in order to calculate the ratios of the CEF.

The CEFs for each reaction (or CEF ratios when “reference substrate uptake reactions” is defined) are calculated and displayed in the text boxes.

(Hint: a CEF ratio could alternatively be computed by calculating the CEFs for Substrate uptake reaction(s) A, copying the values to the flux clipboard, then calculating the CEFs for Substrate uptake reaction(s) B and dividing the CEF values of A by the CEF values of B by using the menu function ‘Arithmetic operations ...’).

- *Reaction participation*: Determines the absolute (displayed in the command window) and relative (in %; displayed in the network maps) frequency of the appearance of each reaction (rate unequal to zero) in the selected modes.
- *Histogram of pathway length*: Displays a histogram representing the frequency distribution of the number of participating reactions in the elementary modes. Quite useful to assess whether there are many/few small/large pathways.
- *Structural couplings of reactions*: Determines enzyme subsets [= groups of pathways which operate in the current selection of flux modes always together (if one reaction of an enzyme subset has a rate unequal to zero in a certain flux mode then all other member of this enzyme subset have a rate unequal to zero, too)] and mutually excluding reaction pairs [= two reactions which never occur together (with rate unequal to zero) in any mode]. The results are displayed in the command window.
- *Minimal cut sets*: Starts minimal cut set calculation, display and analysis. Pressing this button is a short-cut because it is equivalent to starting the menu item “**Analysis/Minimal cut sets (via EFM/EFV) ...**” from the pull-down menu (for a description see below).
- *Normalization...*: This feature allows the normalization of the (currently selected) elementary modes to a certain reaction or number. Accordingly, a dialog box opens allowing one to enter
 - a reaction name: All modes involving this reaction are normalized to the rate of this reaction. Modes in which this reaction is not involved remain unchanged. If more than one reaction is defined the modes are normalized to the sum of the defined reactions (input of reaction IDs is supported with autocomplete; see section 0.6).
 - AND/OR a number: All rates in each elementary mode are divided by this number. Note, if reaction(s) have been defined in the first row of the dialog box, then

only modes involving this (or at least one of these) reaction(s) are normalized as described above and additionally divided by the given number.

An example: If you want to normalize all growth-related modes (with rate of 'mue' > 0) to a growth rate of 0.5 then write in the first row of the "Normalization ..." dialog box 'mue' and in the second row the value 2.

- *Save all modes*: All modes (not only the selected subset) including the current values of the relative concentrations of biomass components are saved in a file whose name has to be provided. The extension of this file must be "mat" (MATLAB-format).
- *Export modes*: All SELECTED modes can be exported to an ASCII file: either as list of modes (using the reaction names) or as a matrix (rows: elementary modes, columns: reactions). In the latter case, the reaction names are given in the first row.

The **Analysis/Minimal cut sets** sub-menu provides several functions related to computation and analysis of minimal cut sets useful, for example, for computational strain (or rmetabolic network) design.

Analysis/Minimal cut sets/Compute (directly via duality): Computation of minimal cut sets (MCSs) with the duality-based MCS algorithm and posterior display and analysis. This metabolic design function finds a minimum set of reaction or gene deletions (and optionally additions) that shape the solution space of feasible flux vectors for a given metabolic network according to user-defined specifications. It can be used, for example, to design microbial production strains or to identify gene or reaction knockouts that are lethal for a given strain (synthetic lethals). The user must define the design specifications in the form of inequality and optionally also bi-level optimality constraints (e.g. required for computing OptKnock-like strain designs). Undesired flux states (all to be deleted) are described by one or more *target regions* (T), i.e. user defined subspaces of the wildtype flux space that should be rendered infeasible by knockouts. Undesired flux spaces may, for instance, be those that lead to the production of an undesired by-product (Q) observed in an exchange reaction r_Q . In this case, a target region can be defined as $r_Q \geq 0.01$ to make the MCS algorithm find knockouts that block flux vectors with r_Q greater than zero (≥ 0.01). It is also possible to define target regions that cover flux vectors with low and zero product yields (e.g. $r_{EtOH} / r_{Glc} \leq 1$) and thus *enforce production above* a certain yield. In the example of synthetic lethals, the undesired behavior is the growth rate r_{BM} and the target constraint reads $r_{BM} \geq 0.01$.

Furthermore, the user may optionally define *desired regions* (D) to prevent flux spaces from complete removal. In the context of production host design, such a constraint can be a lower growth rate threshold, e.g. $r_{BM} \geq 0.1$ to prevent designed strains from being unable to grow. Both target and desired regions should intersect with the feasible (wildtype) solution space. If their constraints are incompatible with the wildtype, MCS computation is either obsolete or not possible and the computation will return an error. For more information on computing MCS via duality see paper [32].

It is furthermore possible to use optimality constraints to define target or desired region. These can be used to define OptKnock- or RobustKnock-line strain design goals. Optimality constraints *cannot* be used to optimize production directly. Like the inequality constraints before, they are used to specify flux states that should be maintained or removed. For an OptKnock-like design for instance, one demands (through a desired region) to maintain flux states in which simultaneously (1) production is larger than some threshold (e.g. $r_{EtOH} \geq 1$), (2) growth is maximal (optimality constraint: $r_{BM} = r_{BM_max}$) and (3) growth is larger than some

threshold ($r_{BM} \geq 0.1$).

General comments: a guide to successful MCS computations

For the success of your first own MCS computation, three cornerstones are vital: (1) your model, (2) your MCS setup, and (3) the optimization (MILP) parameters. **First**, make sure your model is suited for MCS computation. How does your model cope with knockouts? Make sure that your initial model is feasible and that there are at least some reactions that can be knocked out without rendering your model infeasible, because this is the engineering potential MCS is going to exploit. Many too specific flux bounds can also hamper MCS computation, because they might force many reactions to be active that may then not be possible to knockout. Furthermore, consider only relevant exchange reactions to be active and “close” exchanges that are likely to be irrelevant (e.g., only one substrate exchange possible at a time, and only keep open the exchange of truly relevant fermentation products, e.g., ethanol, formate, acetate, succinate and lactate for *E. coli*; all other carbon exchanges should be closed by setting their fluxes to zero). **Second**, make sure your target and desired region(s) are defined correctly. There are hints for the definition of both regions, given above in the function introduction, and also below in the description of the respective fields of the MCS dialog. Most importantly, make sure to not include the zero flux vector in your target region (e.g. a minimum (non-zero) ATP maintenance demand excludes the zero flux vector in many network models by default). MCS will try to eliminate the target region through suitable reaction/gene knockouts. This will be impossible for the zero flux vector where all reactions are inactive anyway. So, for instance, if your target region reads $r_{product} \leq 0.1$ and MCS doesn't find any solution, try adding a second constraint to the target region, e.g. $r_{substrate_uptake} \geq 0.1$, to explicitly exclude the zero flux vector. In your first MCS computation for production strain design, try to select only small subspaces in your target region (e.g. $r_P \leq 0.1$ instead of $r_P \leq 0.4$), and rather large subspaces for your desired region (e.g. $r_{BM} \geq 0.1$ instead of $r_{BM} \geq 0.4$), chances are higher that MCS for your problem setup exist. You will also want to use your wildtype strain as a reference for MCS computation. Take into account maximal growth and production rate in your threshold definitions. **Third**, MILP solver parameters may have a big impact on the success of your MCS computation. Best practice is to first set parameters that make it as easy as possible to solve the MCS problem. This means you should set Max size to a high number, e.g. 30, set Max solutions to (only) 1 or 10 and set a moderate time limit, e.g. 30 minutes (1800 s). Use the “any MCS (fast)” setting and select a solver, ideally CPLEX or Gurobi, if these are not available, you may also use GLPK or intlinprog from the MATLAB optimization toolbox. If your first test run does not succeed, consider (1) revising your model (too many inhomogenous constraints, a large number of open exchange reactions that allow the uptake of various substrates or the exchange of random metabolites) or (2) revising the MCS setup, i.e. target and desired region (zero-flux-vector, target or desired region too greedy).

Once MCS computation succeeds you can confine your MCS setup; set higher production or yield demands, decrease Max size to the number of interventions you're willing to implement experimentally, increase Max solutions, remove the time limit or do a full enumeration of MCS by cardinality. You can also speed up MCS computation and reduce the number of redundant solutions by declaring some reactions (or genes) not knockable (or not addable) or set specific costs.

Example MCS setups are provided with the iML1515 *E. coli* model. Load first scenario “flux_bounds_core_metabolism.val”, open then the MCS dialog box (“Analysis/MCS (directly via duality) ...”) and load then the MCS setup “MCS_ ethanol_yield_0.8.mcs” and start the computation.

In the following we give an overview on the different fields of the dialog for duality-based MCS computation (Fig. 13).

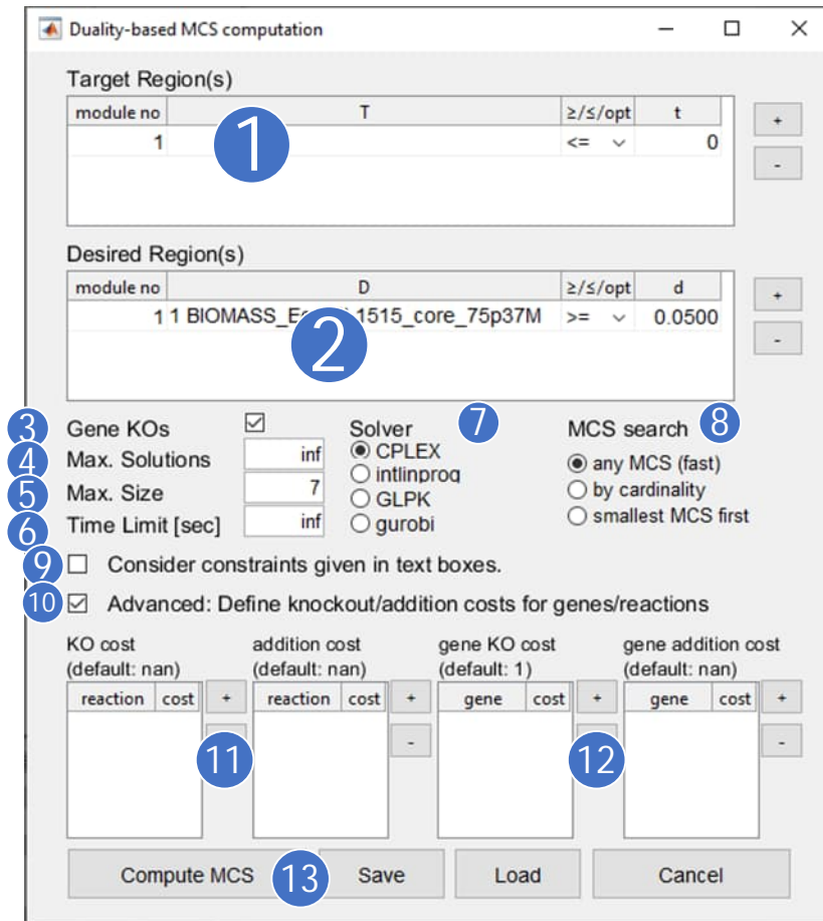


Fig. 13: Dialog for duality-based MCS computation.

MCS problem setup:

1. Target Region(s):

Each row of a target region corresponds to a (target) constraint (flux vectors of the target region fulfill all target constraints for this target region). Target constraints must be stated as linear inequalities or as optimality constraints. When stated as inequalities, the left-hand side term may contain sums, differences, or ratios of rates, the right-hand side is a single (float-point) value. Each rate can furthermore be associated with individual factors. Examples for valid target constraints are:

- a. $r_{\text{product}} \leq 1$
- b. $r_{\text{byproduct}} \geq 1$
- c. $r_{\text{product}} / r_{\text{substrate}} \leq 0.4$
- d. $r_1 - 5 r_3 \geq 0$

Examples for *invalid* target constraints are:

- e. $r_1 \leq r_2$
- f. $r_{\text{product}} * r_{\text{biomass}} \leq 1$

When an optimality constraint is used to specify the target (or desired) region, enter the identifier of the reaction that is to be minimal (or maximal) and select “min” or “max” where you otherwise choose the inequality sign. Leave the right-hand side (“t”) empty. Note that this constraint is used to describe the optimality in the *final* (MCS-mutant) strain. For

instance, “r_biomass max” refers to growth maximal flux states in the mutant. If additionally, inequalities like $r_{\text{product}} \leq 1$ (and $r_{\text{biomass}} \geq 1$) are defined in the same target module, the MCS algorithm will attempt to remove flux states that are at the same time growth maximal and have low production rates.

Constraints can be added or removed with the + and – buttons to the right of the table. The first column (region no) is used to group constraints if multiple target regions are to be defined. When two constraints are grouped in one target region, their combined (convex) subspace is eliminated by MCS, i.e. a target region $r_1 \geq 1$ and $r_2 \geq 1$ demands the deletion of all flux states where both, r_1 and r_2 are active together (above the threshold of 1 each). Flux states where the reactions are active individually are still permitted. When the two constraints are grouped in two different target regions (only supported with CPLEX), each subspace is eliminated so that neither of r_1 or r_2 can be active (above 1), neither together nor separately.

2. Desired Region(s):

The desired region(s) follow the same pattern as the target region(s). Desired constraints are used to define subspaces of the (wildtype) flux space that should still be, at least partially, feasible after applying the computed MCS. In the context of a bioproduction host design, a classical desired region is a lower threshold for the growth rate. The desired region $r_{\text{biomass}} \geq 0.1$ demands that, after applying the MCS, the model must still be feasible at growth rates above 0.1. This *does not mean* that *all* flux states from the original desired region must be maintained, *nor* that *all* remaining flux states after implementing the MCS must fulfil this constraint. It instead demands that $r_{\text{biomass}} \geq 0.1$ must still be possible. If your model has an objective function (commonly growth), it will be automatically put into the first row of your desired region (but this can be changed, of course).

3. Gene knockouts:

This checkbox is only available when the network contains gene-protein-reaction associations (GPR-rules). Those are stored in the reaction notes as structured data under the keyword “geneProductAssociation” (see CNAgetGenericReactionData_as_array and section 7.2.1). The “iML1515” *E. coli* is an example model that provides information about the GPR-rules.

Default: off. When checked, MCS are computed based on genes instead of reaction knockouts and additions. The computation of gene-MCS usually requires more computational resources but also generates more relevant intervention strategies. After computation, a representation of the gene MCS can be viewed on the flux maps through the MCS explorer panel (see below). However, the displayed MCS only present the effect of the computed gene MCS on the level of the metabolic reaction network and the actual gene-MCS are stored in the MATLAB variable gMCS (which can be inspected in the MATLAB workspace). Often, different combinations of gene knockouts lead to identical sets of knockouts in the reaction network. Duplicate reaction cut sets are not shown and all gene-MCS that lead to a specific reaction cut set are listed together in a column of the gMCS variable.

MILP Parameters:

4. Max solutions:

Number of (different) solutions after which the MCS algorithm stops. Default: Inf. For small networks, up to around 50 reactions, a complete enumeration is usually possible. In larger networks, max solutions should be set carefully in consideration of “Max cost” and the network size. Note that the final number of MCS delivered can be larger since many of the found (compressed) MCS often correspond to multiple equivalent solutions.

5. Max size:

Maximal number of deletions or additions in each MCS. Default: 7. For MCS computation in small networks this value can be set to inf, in larger networks it should be set to a lower value, e.g. 7. Best practice is to first set Max size to the number of interventions you would realistically be able to implement experimentally.

6. Time Limit:

Time after which MCS computation is stopped. Default: Inf. This time limit does not include pre- and postprocessing time, only pure MILP time.

7. Solver:

Default: CPLEX. In most cases, this is the best choice. To make this option available, set up CNA with CPLEX correctly (see section 0.5). If CPLEX is not available, Gurobi may be used. To set up Gurobi, follow the instructions in section 0.5. Finally also GLPK (part of CNA) or intlinprog, the MILP solver from the MATLAB Optimization toolbox, can be used. Both solvers are slower, rely on a more error-prone MCS-MILP formulation (that uses the Big-M method) and might fail in large networks where CPLEX or Gurobi may still find solutions. However, they will be useful (e.g. for demonstration purposes) in smaller or core networks.

8. MCS search method:

In which manner should MCS be computed, (1) any MCS (fast), (2) by cardinality or (3) smallest MCS first? Default: any MCS (fast). Options 1 and 3 compute only one MCS at a time, store the solution and repeat the search. Option 1 searches any valid solution to the MCS problem setup (a cut sets(and then minimizes the number of interventions (to obtain a minimal cut set) before repeating the previous steps (but this does not necessarily find the MCS with minimal cardinality), while (3) identifies the globally smallest MCS in every iteration step. In larger networks, (1) and (3) should be used with a fixed value for Max Solutions. Option 2 uses the CPLEX populate feature (hence it is not available with intlinprog) and is the best choice for enumerating all MCS up to a Max size (the latter should then not be too high)

Other parameters:

9. Constraints in text boxes:

If checked, values from the reactions text boxes on the reaction maps are taken into account for the MCS computation. Default: off. Usually only the network structure and the upper and lower bounds for each reaction are used.

10. Advanced: knockout and addition costs:

When checked, tables to define individual addition or deletion costs for every reaction become available. When “gene KOs” is also checked, individual cost factors can be set for each gene, too. Reactions that are not targetable are marked with “nan”, any other value represents an intervention cost. If you selected MCS search mode 2, all cost factors must be integers (or nan). Under point 11 and 12, it is explained how the intervention costs can be specified. By default (without specifying intervention costs), the following settings will be used for the intervention costs:

In reaction MCS: All reactions may be knocked out (cost 1) by the MCS algorithm to meet the demands from the MCS setup. No reactions can be added (cost nan). If you want to use reaction additions, first add your custom reaction to the model, then call the MCS dialog, select the “Advanced” checkbox, press “+” next to the “reaction additions” table, enter the reaction identifier and set its cost to 1 (or any other value). The reaction will now be treated as if it was not part of the wild type. It will only be added as part of an MCS when it is absolutely necessary to meet the demands of the MCS setup, otherwise it will be left out.

In gene MCS: With standard settings, all genes can potentially be knocked out (cost 1). No reactions can be knocked out or added (cost nan) and also no genes can be added (cost nan). If you want to change this behavior, use the “+” or “-“ buttons to define your deviant

preferences. As explained above, gene MCS require the field “geneProductAssociation” in the structured reaction data (see CNAsetGenericReactionData). If you want to use a gene addition candidate, add the catalyzed reaction to your model and enter the corresponding gene or gene rule using the keyword of “geneProductAssociation”. When you then call the MCS dialog and press the “+” button next to the table of gene additions you’ll be able to choose the gene together with a cost factor of your choice. Again, the gene will only be added as part of an MCS when it is absolutely necessary to meet the demands of the MCS setup, otherwise it will be left out.

The definitions of reaction knockout or addition costs in the context of gene MCS is occasionally of value. Exchange reactions, like oxygen or co-substrate supplies can be equipped with intervention costs to enable their addition or removal aligned with the genetic interventions.

11. Reaction knockout and addition costs:

Add “+” or remove “-“ rows to define individual cost factors for the knockout or addition of each reaction. If you want to prevent a reaction from being knocked out, enter “nan” in the cost field. If (non-nan) addition and knockout costs are defined for the same reaction, addition costs overwrite the knockout costs.

12. Gene knockout and addition costs:

Add “+” or remove “-“ rows to define individual cost factors for the knockout or addition of each gene. If you want to prevent a gene from being knocked out, enter “nan” in the cost field. If (non-nan) addition and knockout costs are defined for the same reaction, addition costs overwrite the knockout costs.

13. Compute, Save, Load, Cancel buttons:

The Save and Load buttons can be used to save/load MCS problem setups (not the MCS itself!) in/from files with the suffix “.mcs”. Cancel stops the MCS problem specification and closes the dialog box. With Compute the computation of MCS is started. After successful MCS computation, a panel comes up that can be used to view and analyze the MCS as described in the following.

Exploring computed Minimal Cut Sets via the MCS panel:

[Functions and actions described in the following are partially similar to those for the control panel of elementary modes.] When the calculation of MCS has been finished (or if a file with stored MCSs has been loaded) the first MCS is shown in the network map and a control panel comes up. All reactions that are knocked-out by the current MCS are indicated by color “special” (and value “0”). Reactions without intervention in the MCS get the “standard” color (see section 2). For the special case where MCS have been computed with “knockins” (see above), “knockin reactions” that need to be added for the current MCS are designated by the box color of type “defined”, otherwise they get the color “computed” and a zero value in their box indicating that they are not needed.

The number of the current MCS as well as the overall number of MCSs is shown in the cut sets control panel which is analogous to that in Fig. 12. Similar as for elementary modes, the following functions in the cut sets control panel are useful for stepping through the set of MCSs, for selecting a certain subset of MCSs, for assessing statistical features or for saving/exporting the MCSs:

- *Selection*: This tool facilitates the definition of a selection of a certain subset of all MCSs. For this purpose a panel opens where the following six specifications can be made:
 - reactions which *must not* be contained in the MCSs to be selected
 - reactions which *must* be contained in the MCSs
 - metabolites which must not participate in any reaction involved in the MCSs

- metabolites which must participate in all reactions involved in the MCSs
- minimal size of the MCSs (min. number of contained reactions)
- maximal size of the MCSs (max. number of contained reactions)

For each row no, one or several reaction or metabolite name(s) can be defined, respectively (separated by a blank; input of reaction / species IDs is supported with autocomplete; see section 0.6)). If nothing is specified the complete set of MCSs is selected (equivalent to “*Select all*”). After entering these specifications the user is requested whether the selection should be performed on the current selection (when a selection is currently already active) or on the complete set of MCSs. In the first case the user specifies a subset of the current subset of MCSs thereby refining the selection to incorporate the new constraints.

All functions described as follows operate only on the current selection of MCSs, except “*Save cut sets*” and “*Expand cut sets*”.

- *Deselect set*: Deselects the currently displayed MCS. Thus, a certain MCS can be deselected separately without using the *Selection* tool.
- *Select all*: Selects the complete set of MCSs.
- *Next / Previous*: Switches the display to the next or previous MCS of the currently selected MCS.
- *Jump ...*: Here, the number of an MCS to be displayed can be entered. This MCS becomes the currently displayed MCS. (Note: the MCS number must be contained in the set of currently selected MCSs).
- *Delete unselected cut sets*: Removes all unselected cut sets from memory (in contrast to the *Selection* tool). Useful if some cut sets are not of interest and can be dropped for further analysis.
- *Statistics*: A number of useful statistical features of the current selection of MCSs can be computed by selecting the respective feature from the pull-down menu (similar as in Fig. 12) and clicking on button “*Compute*”:
 - *Select non-affected modes*: Clicking on this button all those elementary modes (currently in memory) that are not affected by the current MCS (i.e. all modes remaining operable after removing the MCS from the network) will be selected. This enables the user to check which network functions are not affected by the current MCS. Of course, this function makes only sense if the current sets of MCSs and elementary modes are corresponding, i.e. if the set of MCSs has been calculated on the basis of the current set of modes.
 - *Average cut set size*: Determines and displays for each reaction the average size of all selected MCSs in which it is a member of. This gives you a clue how robust the network (or a function) with respect to a failure of that reaction is. The average MCS size of all selected MCSs and the average *average MCSs size of each reaction* as well as the averaged fragility coefficient (see [9]) are displayed in the command window.
 - *Reaction participation*: Determines the absolute (displayed in the command window) and relative (in %, displayed in the network maps) frequency of the appearance of each reaction in the selected MCSs.

- *Cut set size histogram*: Displays a histogram representing the frequency distribution of the MCSs sizes. Quite useful to assess the fragility of the system.
- *Cut set with lowest side-effects*: As for the feature “*Select non-affected modes*” this functionality computes a relationship between the currently selected sets of elementary modes and MCSs, respectively: from all selected MCSs the one which hits the lowest number of selected modes is determined. This MCS (or several MCSs in case that several minima exist) becomes the new selection. This feature is useful to find an optimal intervention strategy (which hits all undesired modes but spares most of the others).
- *Save cut sets*: All MCSs (not only the selection) will be saved in a file whose name has to be given. The extension of this file must be “mat” (MATLAB-format).
- *Export cut sets*: All SELECTED MCSs can be exported as a (binary) matrix to an ASCII file (columns: reactions, rows: MCSs). The first row contains the names of the reactions.

Analysis/Minimal cut sets/Compute (via EFM/EFV) ...: Calculation of minimal cut sets (MCSs) via elementary (flux) modes /elementary flux vectors. In contrast to the duality-based MCS computation procedure described above, this function requires the EFMs/EFVs to be calculated in a predecesing step.

A dialog box opens in which several checkboxes allow one to specify certain parameters for the computation of MCS from a given set of EFMs. The calculation of a new set of MCSs is *always related to the current set of elementary modes* (or *elementary vectors*; in the following we will always refer to elementary modes). This means, as mentioned above, that you cannot compute MCSs with this function before you have not calculated or loaded a set of modes. Furthermore, at least one mode must be selected. The currently selected set of modes will be considered as all possible realizations of a certain function or phenotype (e.g., modes with a low product yield) which are to be “destroyed”, i.e. the current selection of modes will be interpreted as the set of “target modes” (or “undesired modes”). If all modes are selected, the complete network will become inoperable by the cut sets computed. The MCSs calculated are those minimal sets of reactions, whose removal from the network will guarantee that none of the selected modes will “survive”, hence, that these pathways are destroyed and non-functional anymore. The “minimal” property means, that no subset of the MCSs would be (structurally) sufficient for this goal. For example, in Fig. 1, {R1,R6} would be an MCS for the complete network. Each essential reaction is an MCS having a size of one. (For further details see [9].) As a new feature, CNA now also allows the computation of *constrained MCSs* [21]: one can define up to two sets of elementary modes from which a minimum number (to be specified) must not be hit by the MCSs to be computed. In this way, one may compute MCSs that, for instance, disable the synthesis of a certain product while preserving the ability to produce other components (including biomass)[21]. In principle, this constrained set of MCSs would be contained in the complete set of MCSs without the side constraint (and one could select them after the computation) but it can be highly beneficial to consider these constraints already during the algorithm since it can drastically reduce the number of admissible MCSs. The “desired modes” (modes that should not be hit by the MCSs) are specified with the help of the two clipboards available for elementary modes (see above).

Before starting the calculation of MCSs, the user has the several options to specify the MCSs to be computed:

- *Exclude reactions with 0 value*: All reactions having a “0” in their associated text box are considered as reactions which *must not* be contained in the MCSs to be calculated. This option can be used to compute only a subset of all MCSs which do not involve the

respective reactions.

- *Prefiltering duplicate/superset vectors:* When elementary flux vectors of inhomogeneous systems have been calculated it is sometimes useful to identify vectors whose support pattern is a duplicate or even a superset of another and then to eliminate those before calculation. The result with/without prefiltering is always the same but sometimes prefiltering accelerates computation. Note that elementary flux modes (of the homogeneous system) are per definition support-minimal; so here a prefiltering will not be useful (except if multifunctional enzymes are specified; see next item).
- *Request multifunctional enzymes:* If the network contains reactions which are catalyzed by the same enzyme (e.g. transketolase) then this can be considered properly during the calculation of MCSs (since both reactions will be eliminated simultaneously in mutants unable to synthesize the respective enzyme). If this checkbox is selected then – after starting the calculation procedure – the user will be requested to define (arbitrary many) sets of reactions each related to a particular multifunctional enzyme (input of reaction IDs is supported with autocomplete; see section 0.6). Each reaction set of a multifunctional enzyme can be defined by entering the respective names of the reactions in a provided window. After choosing ‘OK’ a new set can be defined. One can exit this loop by letting the input field empty and clicking on ‘OK’.
- *Save at least ... modes from clipboard 1:* As described above, using this option one can impose constraints to the MCSs to be computed. In an initial step, one selects a set of modes to be preserved (they should not be hit by the MCSs) and copies them to the elementary-modes clipboard 1 (see clipboard tools for elementary modes). Then, by selecting this checkbox and by specifying a minimum number of modes to be preserved, CNA will compute only those MCSs that hit all target modes and satisfy the additional constraint of saving the required minimum number of modes (the “desired modes” being specified by clipboard 1).
- *Save at least ... modes from clipboard 2:* The same tool as the previous one but in combination with clipboard 2. One may also combine both clipboards allowing one to define more complex constraints: one may specify up two sets of “desired modes” (one is clipboard 1, the other in clipboard 2) and for each of these sets a minimum number of modes that must be preserved. A complex example: using this facility we may compute MCSs that will destroy all modes but save (i) at least one mode that will enable biomass synthesis (out of a set of biomass-producing modes specified in clipboard 1) and (ii) at least one mode out of a set of modes (in clipboard 2) that produce synthesis of a product of interest.
- *Late check (during postprocessing):* An algorithmic flag meaningful only in combination with the two previous options: The fulfillment of the side constraints (minimum number of modes to be saved) is normally checked during the algorithm which is especially advantageous if the set of “desired modes” is relatively small. However, if the set of desired modes specified in the clipboard is relatively high (compared to the set of target modes) it might be better to discard non-admissible MCSs in a post-processing step which can be enforced by selecting this checkbox.
- *Maximal cut set size:* Here, the user can specify the maximal size of the MCSs to be calculated (= number of reactions an MCS contains). Especially in large networks, the number of MCSs can grow rapidly. Therefore, by limiting the size of the MCSs, the user may focus only on the subset of smaller MCS, which are usually the most important ones.

When the computation has been started the progress can be seen in the command window. The overall computation time depends on your computer, the selected set of modes, the number of involved reactions and the algorithm used. As in the case of elementary modes it can take a few

seconds or several days (or even longer ...).

After successful MCS computation, a panel comes up that can be used to view and analyze the MCS as was described above under item ‘Analysis/Minimal cut sets/Compute (directly via duality)’; see there the subsection “Exploring computed Minimal Cut Sets via the MCS panel”.

Analysis/Minimal cut sets/Load ...: Allows one to load minimal cut sets that were saved in earlier sessions. Select a proper file (usually, the modes are saved in MATLAB files with extensions “mat”). Warning: cut sets should be newly calculated if they have been computed and saved before the network structure has been modified. Otherwise, an error message could appear.

Analysis/Minimal cut sets/Show ...: minimal cut sets, which are still in the memory (loaded or calculated earlier in the current session) will be displayed again.

The submenu **Analysis/Flux balance analysis** provides three functions related for flux balance analysis / flux optimization.

Analysis/Flux Balance Analysis/Flux optimization (FBA): This function optimizes (here: minimizes!) a linear objective function. The following parameters are involved in this procedure:

- the currently defined rates in text boxes of reactions are considered as constraints
- the valid range (constraints) of the reaction rates (min/max) as well as the coefficient in the linear objective function are basic properties of each reaction and can be defined by using the menu item „Network composer“. The currently defined objective function can be displayed via menu item “Analysis/Flux Balance Analysis/Show objective function”.

One classical FBA optimization example would be „Maximize growth rate“ for an uptake of one mmol of a certain substrate. For this scenario, you may set the value 1 in the text box associated with the uptake rate and set the coefficients for the linear objective function for each reaction on 0, except for „mue“ which gets the coefficient -1 (keep always in mind: the objective function will be minimized!).

Note: This function **needs at least one working LP solver (see section 0.5)**. When starting this routine, the user may choose one solver from a list of available solvers detected by CNA.

Analysis/Flux Balance Analysis/Parsimonious FBA”: This function does exactly the same as the previous one (FBA) but ensures that (in case that the optimal flux distribution is not unique) among all optimal flux vectors the one with the minimal sum of all (absolute) fluxes will be selected. Note that even with this additional secondary objective, the optimal flux distribution is not necessarily unique. Note also that FBA and parsimonious FBA may deliver the same solution, dependent on the used solver and (non-)uniqueness of optimal solutions.

Note: This function **needs at least one working LP solver (see section 0.5)**. When starting this routine, the user may choose one solver from a list of available solvers detected by CNA.

It may happen that, during the calculation of an FBA (see above) or FVA (see below) scenario the solver returns an infeasibility message, for example, due to inconsistent flux values. To check for and resolve those infeasibilities the following function can be used.

Analysis/Flux Balance Analysis/Show objective function: Shows the linear objective function as currently defined by the reaction properties (via “the coefficients in the linear objective function”; see section 3.1). This objective function will be used in the following function.

Analysis/ Check/Resolve infeasibility of FBA scenario: This function checks whether an FBA scenario (consisting of the given fluxes in the text boxes and the predefined flux bounds) is feasible or not (i.e. whether a flux distribution fulfilling all constraints exists), resolves occurring infeasibilities by correcting the given flux values and performs afterwards, with the possibly corrected scenario, an FBA or FVA computation (as specified by the user) (see also Ref [33]). When starting this function, a panel comes up in which the user has to make four different specifications (Fig. 14):

- 1) Whether a linear (LP) or quadratic (QP) optimization has to be performed when checking for infeasibility and resolving potential infeasibilities. The LP corresponds to a minimization of weighted absolute changes, while the QP delivers a weighted least-squares solution [33]. The LP-based approach needs an LP solver (which can be chosen in the next step as usual; see below), while the QP approach needs either MATLAB’s quadprog solver contained in the optimization toolbox or CPLEX’s cplexqp solver (CPLEX installation required)). The available QP solvers are already indicated in the first drop-down menu in the control panel in Fig. 14 and if no QP solver is detected then the QP-based approach cannot be chosen.
- 2) Corrections for given fluxes needed to make the system feasible can be penalized via four different weighting schemes. For example all corrections can be equally weighted or relative to their flux values (i.e. larger fluxes will be more likely changed when resolving infeasibilities). One option also allows the use of the measurement variances (a property of each reaction; see Table 1 and section 3.1).
- 3) For given zero fluxes one may specify a particular weight (this will be normally large (>1000) to ensure that corrections of zero fluxes are unlikely).
- 4) As fourth specification, one needs to select one of five possible actions to be done once the scenario has been corrected (or was feasible anyway). For example, one may perform an FBA, parsimonious FBA or FVA with the corrected system.

The result of the correction procedure and of the specified calculations will be displayed in the text boxes. Additional information regarding the feasibilities, redundancies and the introduced corrections are shown in the command window.

As an alternative, one may also use “Classical (algebraic) metabolic flux analysis/Flux analysis” (see below), which can also balance inconsistent flux scenarios (however, that function cannot deal with flux bound constraints and thus also neglects reaction reversibilities; but it can deal with measurement variances).

Note: This function **needs at least one working LP solver (see section 0.5); also when using the QP-based approach.** When starting this routine, the user may choose one LP solver from a list of available solvers detected by CNA. As mentioned above, for the QP-based approach, MATLAB’s or CPLEX’s QP solver must be available to solve the quadratic programming problem.

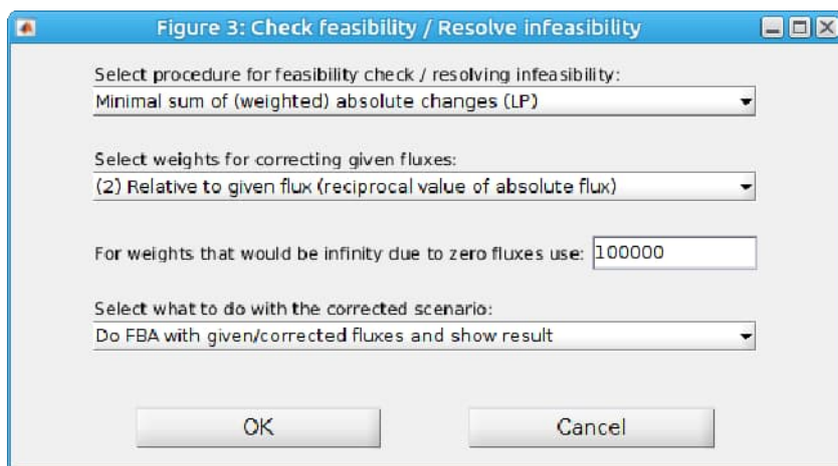


Fig. 14: Window for checking/resolving infeasibility in FBA scenarios

Analysis/Flux variability analysis: Useful for checking the *feasible* range (upper and lower bounds) of each reaction rate consistent with a given set of predefined fluxes and consistent with the min/max reaction values defined for each reaction. Obviously, the computed upper and lower boundaries will lie within the min/max range defined by the reaction properties. Given values in the reaction text boxes are considered as predefined reaction rates (the macromolecule concentrations are also read; if applicable). Linear optimization is then used to determine for each reaction the feasible lower and upper boundary and the result is displayed in the command window. The flux ranges are also displayed in the text boxes and different colors are used to indicate (i) predefined fluxes (fixed prior to computation), (ii) fluxes that are uniquely determined (upper and lower boundary are identical), (iii) reactions that have no fixed flux but are essential (i.e, either the minimum flux is positive or the maximum flux is negative), and (iv) all other flux ranges that contain the zero flux as possible value. This function can also be used as a substitute for “Flux Analysis ...” (see below) as long as the system is feasible and non-redundant. It even has the advantage that the min/max constraints of the reactions (and thus the reversibility of the reactions) is explicitly taken into account; in contrast to classical flux analysis. It may therefore result in a larger number of uniquely determined rates. However, it may take longer since several linear optimization problems must be solved. Moreover, if the defined scenario is overly stringent and thus not feasible (function ‘Check feasibility’ can be used to test this) an error message will be shown and one should then try “Flux Analysis ...” to treat redundant systems.

Note: This function **needs at least one working LP solver (see section 0.5)**. When starting this routine, the user may choose one solver from a list of available solvers detected by CNA.

Analysis/Phase plane analysis ...: This function projects the solution space on a two-dimensional plane of two selected reaction rates. This is frequently used, for example, to study the dependency between growth rate and synthesis rate of a product of interest (then also called production envelope) but is also useful for many other applications. The user has to specify the identifiers of the selected reactions (input of reaction IDs is supported with autocomplete; see section 0.6). If more than two reactions are defined (say the number is x), then all $x*(x-1)/2$ phase planes of all pairs of reactions will be drawn as subplots in one figure. The phase plane is approximated by sampling the reaction rates in n equidistant steps, this number n must also be defined. Usually, $n=20$ is sufficient (also in large networks) but it can be adapted to get finer approximations. The defined flux boundes and the currently defined rates in text boxes of

reactions are considered as constraints. The two reaction rates must be bounded by these constraints; an error message will otherwise come up.

Note: This function **needs at least one working LP solver (see section 0.5)**. When starting this routine, the user may choose one solver from a list of available solvers detected by CNA.

The submenu “Yield analysis” provides two functions:

Analysis/Yield analysis/Maximize yield ...: This function can be used to maximize a yield Y defined by a linear fractional (yield) function of the type $Y = \mathbf{c}^T \mathbf{r} / \mathbf{d}^T \mathbf{r}$, where \mathbf{r} is the (steady state) flux vector and \mathbf{c} and \mathbf{d} are vectors indicating weighting factors for the reaction rates. In typical applications, \mathbf{c} contains a single non-zero entry for the product (P) excretion reaction and \mathbf{d} contains a single non-zero entry for the substrate (S) uptake reaction, hence, the yield of P/S would be maximized. However, more general yield functions can be optimized as well by choosing appropriate factors in \mathbf{c} and \mathbf{d} . The user will be asked to specify the reaction identifiers (input of reaction IDs is supported with autocomplete; see section 0.6) with non-zero values in \mathbf{c} and \mathbf{d} , respectively, together with the respective weighting factor (only weights different from “1” need to be defined; negative weights can also be used, e.g., if a substrate uptake reaction is defined in the direction of excretion thus having negative fluxes for substrate uptake). For example, specifying “R1 2” for \mathbf{c} and “R2 R3 2” for \mathbf{d} would maximize the function $Y=2*R1/(R2 + 2* R3)$. The optimal yield is calculated and (one) optimal solution resulting in the optimal yield is displayed. Note that

- the defined flux boundes and the currently defined rates in text boxes of reactions are considered as constraints.
- optimal yield solutions often coincide with optimal solutions obtained by FBA, but this is not generally true.
- the yield function must be bounded; an error message will otherwise come up. In particular, there should be no flux vector in the network where the denominator becomes zero AND the nominator is non-zero.

Note: This function **needs at least one working LP solver (see section 0.5)**. When starting this routine, the user may choose one solver from a list of available solvers detected by CNA.

Analysis/Yield analysis/2D yield space plot...: This function generates a two-dimensional yield space plot displaying the dependency between two yields (yield functions). Yield space plots are useful, for example, to study the trade-off between growth (biomass) yield and the yield of a certain product of interest. The two yield functions $Y1=\mathbf{c1}^T \mathbf{r} / \mathbf{d1}^T \mathbf{r}$ and $Y2 = \mathbf{c2}^T \mathbf{r} / \mathbf{d2}^T \mathbf{r}$ must be defined in the same way as explained above for “Maximize yield ...”. The yield space is approximated by sampling the yields in n equidistant steps, this number n must also be defined. Usually, $n=20$ is sufficient (also in large networks) but it can be adapted to get finer approximations. As for maximizing yields, note that:

- the defined flux boundes and the currently defined rates in text boxes of reactions are considered as constraints.
- the two yield functions must be bounded; an error message will otherwise come up. In particular, there should be no flux vector in the network where the denominator of a yield becomes zero while the nominator is non-zero.

Note: This function **needs at least one working LP solver (see section 0.5)**. When starting this routine, the user may choose one solver from a list of available solvers detected by CNA.

The “Classical (algebraic) metabolic flux analysis” submenu provides three functions related to classical metabolic flux analysis.

Analysis/Classical (algebraic) metabolic flux analysis/Classify flux scenario and rates (algebraic redundancy/determinacy): The currently specified scenario (characterized by the currently defined and non-defined values of reaction rates) is investigated to classify non-defined rates to be (uniquely) calculable (C) or non-calculable (###) and to classify defined (“measured”) rates to be redundant (balanceable) (B) or non-redundant (M). The classification of rates also allows for a classification of the complete system: If redundant (balanceable) rates occur then the system is redundant and the number of degrees of redundancy is shown. Furthermore, if any unknown rate is non-calculable then the system is underdetermined (otherwise: determined). The classification result is shown in the network maps (using the above introduced symbols #, B, C, M and the respective box colors as defined in section 2) as well as in the command window of MATLAB. This function is also useful for planning experiments: the user can test whether a certain unknown rate is calculable by the use of a certain set of measured rates. For this purpose the user must set a numerical value of all reaction rates which are thought to be measured on an arbitrary value and all other rates on undefined (“#”) and then start this function. (Note: The real values of the defined rates have no influence on the classification. Note also that this classification does not consider reaction reversibilities. To take this into account use FBA/FVA-related functions).

Analysis/Classical (algebraic) metabolic flux analysis/Flux analysis ...: This menu function provides methods for (classical) metabolic flux analysis (MFA), i.e. for calculating unknown fluxes from measured or known rates (specified by the user in the text boxes). Note that this classical (algebraic) MFA used only the steady-state constraint together with the given fluxes to compute unknown fluxes. It does not consider the flux bounds (and therefore also no reaction reversibilities). As alternative, one may alternatively either use “Flux Variability Analysis” or “Check / Resolove infeasibility in FBA scenario” (see also Ref [33]).

Two basic types of MFA scenarios can occur, depending on the redundancy of the defined scenario (see also menu function “Classify rates” (above) and ref. [1]):

Case 1) The scenario is non-redundant: In this case, no redundancies (and related inconsistencies) in the given rates occur. In a non-redundant scenario, on the basis of the currently defined values, all (uniquely) calculable rates are determined. The results are shown in the network maps and the color of each text box indicates whether the respective reaction rate was defined or calculated or non-calculable (for colors see section 2). Some further information is displayed in the command window. In particular, warnings will appear when a reaction rate has left its admissible range (by checking the rate minimum and maximum).

Again, note that Case 1 can also be well handled via “Flux variability analysis” (see above) with the additional advantage that reaction reversibilities are explicitly accounted for.

Case 2) The scenario is redundant: In this case, some of the defined rates carry redundant information. The scenario can now be either inconsistent or consistent, usually they are inconsistent (non-redundant scenarios are always consistent). An example for a redundant scenario is shown in Fig. 2 where the growth rate (μ) and the rate for uptake of A (R1) as well as the following conversion to B (R2) and then to D (R3) have been defined. It is obvious that R2 and R3 must have the same value and defining both rates results in a redundant system where R2 and R3 are redundant/balanceable.

Whether a system is redundant and (if so) which of the defined rates are redundant can also be easily checked by the function “Classify rates ...” described above.

If a redundant scenario was defined, CNA provides basically three different methods to treat this scenario and to deal with potential inconsistencies. Accordingly, a panel comes up (which is in parts identical to Fig. 14) where the user has to specify the method to be used:

- *Minimize (least-squares) deviation from steady state*: This function determines the least squares solution of the unknown rates by using the pseudoinverse. For those unknown rates, for which the solution is unique, the values are displayed in the text boxes (with color “calculated”). Importantly, no balancing/corrections of the redundant/balanceable rates will be carried out, i.e. inconsistent systems will remain inconsistent and the solution will then violate the steady state condition for at least one metabolites (but the least-squares solution ensures that these deviations are minimized). The resulting residuals are displayed in the command window. Note also that the weighting scheme that can be defined in the panel that comes up will not be used and will thus have no effect.
- *Balance inconsistencies via weighted least squares*: This is the standard procedure for handling redundant systems in classical MFA (see e.g. [5,6]): First, the redundant/balanceable rates are identified and finds minimal corrections in these rates to make the scenario consistent. Here, a weighted least-squares approach is used to minimize the (weighted and squared) changes. The weights define the penalties for the change in the given corrections and four different weighting schemes can be used (see also “Analysis/Check/Resolve infeasibility in FBA scenario”). For example all corrections can be equally weighted or relative to their flux values (i.e. larger fluxes will more likely be changed when resolving inconsistencies). The classical approach is to use the measurement variances (a property of each reaction; see Table 1 and section 3.1) to set large weights (penalties) for rates with low measurement variances. For given zero fluxes one may specify a particular weight (this will normally be large (>1000) to ensure that not or only small corrections are applied to zero fluxes since the latter are often well-known).
Once the corrections for the redundant rates have been computed to obtain a feasible system, the unknown rates are computed from the corrected scenario from which then the (uniquely) calculable rates are displayed (with color “calculable”). Redundant rates (with their corrected value) are displayed (with color “special”). Given rates that are not redundant are displayed as usual in color “defined” and non-calculable rates remain in color “standard”. Further information about the correction procedure are displayed in the command window. If the weighting scheme with measurement variances was used, a consistency check based on the residuals will be performed. If the (variances-weighted) residual h is higher than the chi-squares value for confidence interval 0.95, then modeling or measurement errors are expected with a probability of at least 95 %. Additionally, the variances of the calculated (actually: estimated) rates are also displayed in the command window (see also ref. [6]).
- *Gross measurement error detection*: This function can be applied only in redundant systems with at least 2 degrees of redundancies. The goal is the identification of the redundant rate that causes the largest deviation from the steady state assumption. For this purpose for each redundant rate a separate balance of inconsistencies via the above described weighted least squares estimation is performed where the respective rate is considered as non-defined (only the third weighting scheme, which uses the measurement variances, can be used). Thus, for each balanceable rate one gets the variances-weighted residual h if the rate is not involved (these h 's are shown in the command window). The

reaction, for which the h decreases at most, causes the largest error. This result is also displayed in the command window. No results are displayed in the network maps – every value remains unchanged. (Note: if one would like to calculate the flux distribution with the smallest error one has to set the rate with the largest error to unknown (“#”) and to then to use the above described method *Balance inconsistencies via weighted least squares*.

Again: note that “classical flux analysis” cannot consider reaction reversibilities and min/max reaction rates. As long as the system is not redundant, one may also use “Flux variability analysis” (FVA) which takes min/max reaction rates into account and may therefore result in a larger number of uniquely determined fluxes. Another option to deal with inconsistent scenarios is the “Check feasibility of flux scenario” function (see above).

Analysis/Classical (algebraic) metabolic flux analysis//Sensitivity analysis...: This function calculates the sensitivities for all currently unknown but calculable rates with respect to a certain currently defined rate. For this purpose, one of the known (entered) rate (here denoted by r_m) can be chosen from a list box. After clicking on the OK-button the linear dependency $\partial r_c / \partial r_m$ of each unknown but calculable rate r_m with respect to the chosen rate r_m is computed and displayed in the text boxes of r_m . Calculated sensitivities get the background color “calculated”. Defined rates get the color “defined”, except the rate for which the sensitivities have been calculated (gets color “special”). Non-calculable rates remain in color “standard”, because their sensitivity is non-calculable, too. In redundant systems the sensitivity is computed with respect to a simple (non-weighted) and non-balancing least squares estimation (see “.../Flux Analysis ...”).

Analysis/Net conversion of external metabolites ...: Based on the currently defined flux scenario, the net conversion of external metabolites (see section 3.1) is calculated and displayed (input of the species IDs is supported with autocomplete; see section 0.6).

Analysis/In-Out fluxes at a metabolite ...: Shows a (stacked) bar chart with all influxes and effluxes around a specified metabolite. The fluxes are computed by multiplying the reaction rates (given in the text boxes) with the stoichiometric coefficient of the selected metabolite in these reactions. This feature is useful, for example, to identify the reactions with the highest production / consumption of a particular metabolite. You may apply this function also in the case where only a subset of all reaction rates (producing/consuming the respective metabolite) is given in the textboxes.

3.3.6 Menu “Info”

Info ...: Shows info about *CellNetAnalyzer*.

4. Constructing and analyzing signal-flow (signaling, regulatory) networks

4.1 How signal-flow networks are represented in CellNetAnalyzer

In CNA, the construction and analysis of models of signal-flow networks is done in a similar way as for mass-flow (stoichiometric) networks. However, whereas stoichiometric networks describe the flow of material in a (stoichiometrically) exact manner, signal-flow networks may capture signal (information) and, to a certain extent, also material flows in a more abstract way by describing causal and logical (Boolean) dependencies. This leads to a different network representation. In CNA, a signal-flow network is a Boolean network represented as a signed directed hypergraph. The latter can easily be converted to an interaction graph (see our paper [11]). Before confusing you too much, let's see how signaling networks can be set-up in CNA in practice.

Signal-flow networks are composed of elements of two different types only, namely of *species* and of *reactions*, here also referred to as *interactions*. The species are the network nodes, equivalent to the metabolites in stoichiometric networks, and interactions are directed hyperarcs in the network similar but not fully equivalent to reactions in stoichiometric models. In signaling networks, each reaction represents one possibility how (i.e. under which conditions) the end node of this reaction can be activated (or produced; see below). The element types *biomass constituents* and *assembly routes* are not available in interaction networks.

Attributes of species and reactions in signal-flow networks are listed in Table 2.

Element Type	Species	Reaction (Interaction)
	network nodes	interactions (causal relationships, e.g. activations, inhibitions etc.)
Text Box	logical state	signal flow
Attributes	<ul style="list-style-type: none"> • <i>full name</i> • <i>identifier</i> • <i>default value</i> • <i>text box parameters</i> • <i>comments/notes</i> 	<ul style="list-style-type: none"> • <i>identifier</i> • <i>reaction (interaction) equation</i> • <i>default value</i> • <i>time scale</i> • <i>flag 1: whether this interaction is to be excluded in logical computations</i> • <i>flag 2: whether the logical interaction is treated with incomplete truth table</i> • <i>flag 3: whether the interaction is monotone</i> • <i>text box parameters</i> • <i>comments/notes</i>

Table 2: Network elements in signal-flow networks and their attributes.

As indicated in Table 2 and in contrast to metabolites in stoichiometric networks, all species have an associated text box usually displaying information about the current or computed (logical or discrete) state of that species. Consequently, as for reactions, a default value can be defined for species as well.

Interactions have some similar and but also different attributes compared to reactions in stoichiometric networks. First, the associated text box usually displays the *signal flow* along this interac-

tion (hyper-)arc (instead of a reaction rate). The reaction or “interaction equation” can be defined symbolically by using the species identifiers. It defines *which* species participate (with positive or negative influence) in this interaction and *which* species can be activated along this interaction. Syntax and semantics of interaction equations are similar as in stoichiometric reaction equations but there are some fundamental differences. In a two-level (binary) network an interaction equation looks as follows

$$[!]<Spec1> + [!]<Spec2> + [!]<Spec2> \dots = <SpecR>$$

<Spec1>, <Spec2>, and <SpecR> are species identifiers. On the right-hand side of the equation only one species identifier (<SpecR>) can appear. Each equation represents one possibility to activate the “target species” SpecR and the “+” sign on the left hand-side denotes a logical AND while the (optional) exclamation marks “!” indicate logical NOT operations. (Note that the meaning of the “+” in CNA differs thus from the symbolic representation of Boolean functions where “+” represents normally an OR operation - but in CNA it is an AND).

An example: the interaction equation

$$A + !B + C = D$$

states that D can be activated if A AND C are active AND B NOT. Note that no space between the NOT operator „!“ and B is allowed whereas space is mandatory between species identifiers and „+“ and „=“. Note that all the AND activation rules (i.e. all reaction equations) by which a certain species (here D) can be activated are implicitly OR-connected. Thus, if one would define additionally to the interaction above a second interaction such as

$$A + !C = D$$

the total Boolean function for D resulting from these two interactions would read: D is active if (A AND NOT(B) AND C) OR (A AND NOT(C)). Thus, all the interaction equations pointing in one and the same species define implicitly the so-called (minimal) disjunctive normal form (DNF) of the Boolean function describing how the species (here D) can be activated.

Metabolic (stoichiometric) reactions can be represented in this framework up to a certain (qualitative) degree. For example, $A + B = C$ might represent a metabolic reaction where A and B are required (are converted) to produce C. However, stoichiometric coefficients (except “1”) cannot be captured because coefficients in interaction equations have a different meaning as explained in the following: In binary logical networks, only on / off (0 and 1) states are considered for each species. CNA also allows the construction of multi-valued logical networks in which a species may have arbitrary many discrete states. In the interaction equation, this is indicated by coefficients in front of the species identifiers (separated with blanks from the identifiers):

$$[Coeff1] [!]<Spec1> + [Coeff2] [!]<Spec2> + [Coeff3] [!]<Spec2> \dots = [Coeff3] <SpecR>$$

For example, the equation

$$2 A + !C + 2 !G = 2 F$$

expresses: If A is has at least level 2 AND C is NOT active (level is 0) AND G is NOT at level 2 (or higher) then F will reach level 2. As indicated by “at least” and “not higher”, CNA assumes

monotone behavior. That means, in the example above, F gets level 2 if $C < 1$ (i.e. $C=0$), $G < 2$ $A \geq 2$. If you want to have an interaction to be non-monotone, i.e. to be active only if the species states obey exactly the coefficients, you may deactivate flag 3 (Table 2). So if the above example is non-monotone, then F gets level 2 if and only if $A=2$, $C=0$, and $G \neq 2$ (\neq denotes “unequal”). Hence, if $A=3$, $C=0$, $G=1$, then F gets level 2, only if the interaction is monotone. On the other hand, if $A=2$, $C=0$, $G=3$ then F gets level 2 only in the non-monotone case (because of $G=3$).

As shown in this example, the level (or coefficient) „1“ needs not to be written explicitly (as done here for !C). Therefore, the general interaction equation as given above for binary networks, is just the special case where each species in the equation is associated with a coefficient of “1”.

Note that the exclamation mark is not allowed on the right-hand side and that every species may occur only once in each equation.

Regarding this general principle of constructing multi-level networks there is one important rule: If a species is activated by several different interactions, always the highest possible activation level will be set. For example, having the following two interaction equations

$$\begin{aligned} 2 A &= 3 G \\ !B + 2 C &= 2 G \end{aligned}$$

and assuming that the left-hand side of both equations is fulfilled (A is at level 2, B is zero and C is 2), then G will be assigned level 3 (the highest one). The signal flow along the first interaction will be “3” and along the second one “2”.

Furthermore, input and output arcs may be defined, such as

$$\begin{aligned} = A & \text{ (if the signal flow along this interaction is set on “1”, then A will be activated)} \\ B = & \text{ (if B has state “1” then this interaction will have an (output) signal flow of “1”)} \end{aligned}$$

As already mentioned above, the logical network representation as used herein can be transformed into an interaction graph (the hyperarcs are split into arcs and all species entering an interaction equation with NOT (“!”) are the starting points of a negative (inhibiting) arc in the graph model). However, in certain case one knows that a species, let’s say A, has a positive influence on B but not in such a strong way that it can activate B alone. Then, A would not appear in the logical model. To indicate, that there is nevertheless an influence, one can declare an interaction

$$A = B$$

and set the *flag* (1) that this interaction is to be *excluded in logical computations* (see Fig. 18).

In realistic networks, one is sometimes not sure whether the influence of some species on another species has to be combined with a logical AND or a logical OR. An example: assume A activates C and B activates C but you don’t know whether both are required for (full) activation of C. Here one might define an interaction

$$A + B = C$$

and set another (the second) *flag*, indicating that this interaction has to be treated with an *incomplete truth table* (see Fig. 18). This means that C becomes zero if $A=B=0$ and becomes active if

$A=B=1$. These are the identical entries for the AND and OR truth tables. For the other two possible cases ($A=1/B=0$ and vice versa), the activation level remains undefined. Note that the interaction

$$A + !B = C ,$$

where this flag has been set, has a defined value for C when $A=1/B=0$ ($C=1$) or $A=0/B=1$ ($C=0$) and undefined otherwise.

Finally, another parameter of interactions is the *time scale*. Sometimes, one knows that a certain interaction becomes active significantly later than others (e.g. due to the involvement of gene expression). Then it makes sense to assign a larger time scale value (e.g. “1”) to this particular interaction, whereas all others get a time scale of 0. In principle, arbitrary time scale values can be used; a lower value indicates that an interaction is earlier available. This parameter is used for setting quickly time scale scenarios (see section 4.3).

Fig. 16 depicts the interactive network map of a simple signal-flow network in CNA. Notice that the default colour (grey) and the size of the reaction text boxes have been configured differently compared to species (yellow text box; see section 2) ensuring that species and reactions can easily be distinguished. Connections of species entering an interaction with a negative (NOT) operation were indicated in the network map by a red colour in the map. Hence, the interaction equation of arc 7 is $!D = A$ and for 2&3 it reads $!I1 + I2 = E$.

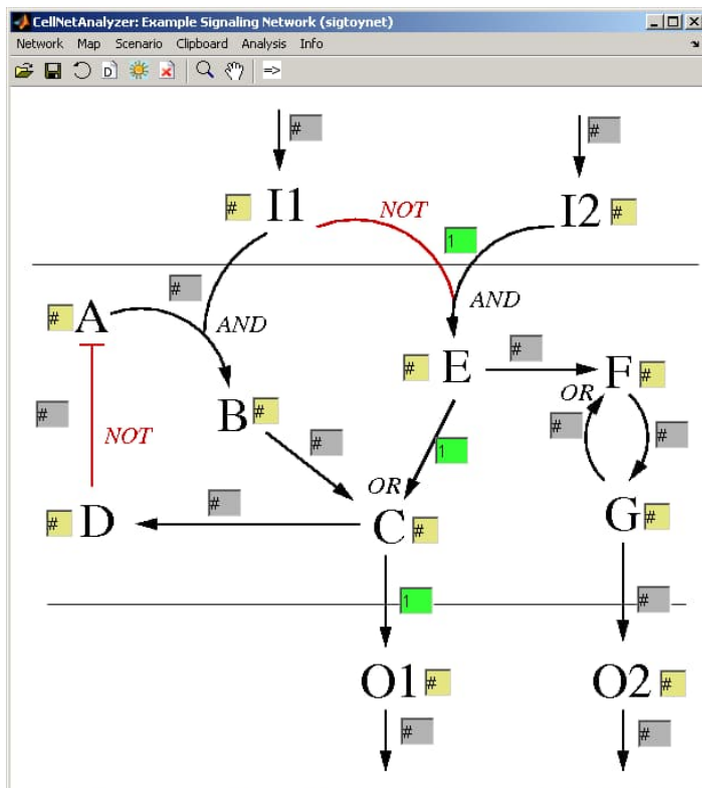


Fig. 16: Interactive network map of a simple signal-flow network that could represent a signaling or regulatory network. The green text boxes indicate a signaling path from $I1$ to $O1$ (or from $I2$ to $O1$; regarding this ambiguity see remarks in section 4.3.).

Note: As for mass-flow networks, using dedicated API functions (see section 7.2.1), the species and reaction notes can also be used to store generic data for species and reactions in a structured

way. Thus, any parameter or textual information (e.g., KEGGID), which are not covered by CNA's reserved fields, can be stored under user-specified field names. The data are serialized in the reactions or species notes and can later be retrieved again.

4.2 Composing and editing the structure of signal-flow networks

Bearing the technical differences and the new attributes of elements in mind, composing signal-flow networks with the masks provided by CNA is analogous to mass-flow networks (section 3.2). The network composer appears similar as in Fig. 5 (but only with species and reactions as possible network elements) and can be used in the same way as described in section 3.2. Again, the masks for existing species and interactions can be opened by right-clicking on the associated text boxes on the network map. It is not possible to open multiple editing masks at the same time. Therefore if you open a new editing mask any previously opened mask will be deleted and possible changes discarded.

The masks for declaring/editing species and reactions/interactions are depicted in Figs. 17 and 18. The text box parameters can be defined in the same way as described in section 3.2 (and all text boxes can be moved simultaneously using the respective button in the network composer window). There is only one difference: Text boxes for species and reactions can be of type „Editable” and „Non-editable” (as in mass-flow networks) and additionally of type „Non-visible”. The latter is helpful in large networks for hiding text boxes of certain (not that important) network elements. This helps to clean up crowded interactive network maps. Note that you can, irrespective of the set default visibility, always hide and display all species /reactions independently during a network session (see menu functions). Furthermore, be aware of that the values currently contained in hidden text boxes are considered as usual when performing computations.

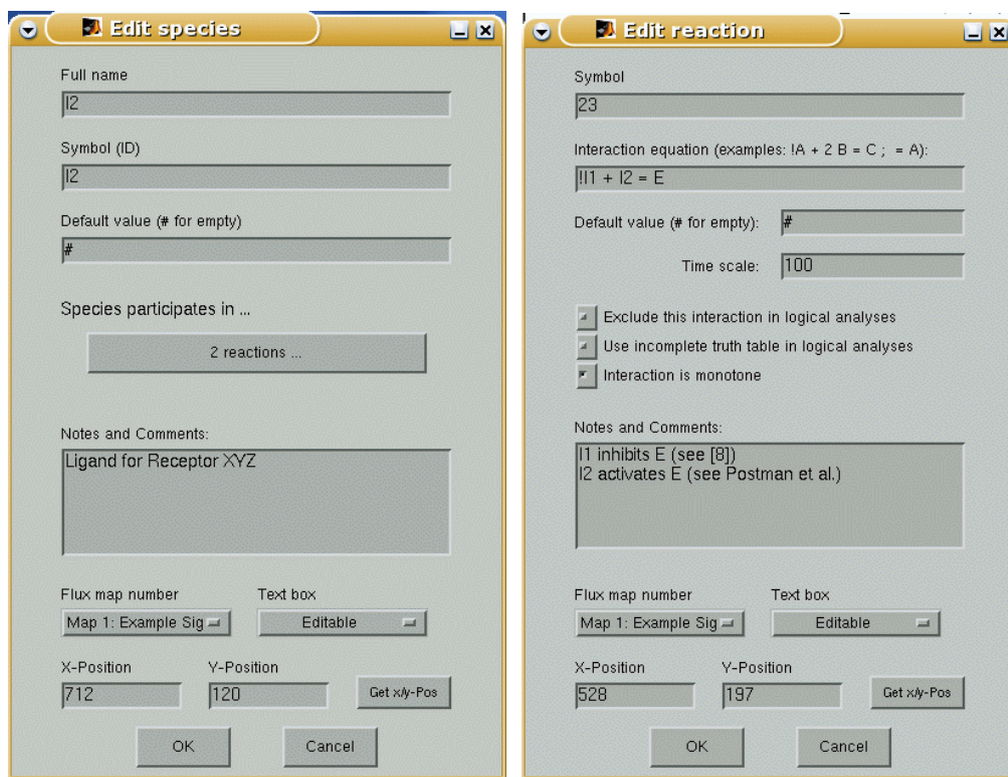


Fig. 17 (left): Mask for editing/declaring species in signal-flow networks.

Fig. 18 (right): Mask for editing/declaring reactions (interactions) in signal-flow networks.

Searching for elements in the network (element selector): See corresponding section (3.2) for mass-flow networks.

4.3 Toolbox (menu functions) for analyzing signal-flow networks

CNA provides a number of functions for studying signaling networks and the menu (Fig. 19) is similar as the menu for mass-flow networks (Fig. 11): it has a similar set-up, it is partitioned into several groups of functions and can be called via the menu item “CellNetAnalyzer” within the interactive maps (Fig. 19).

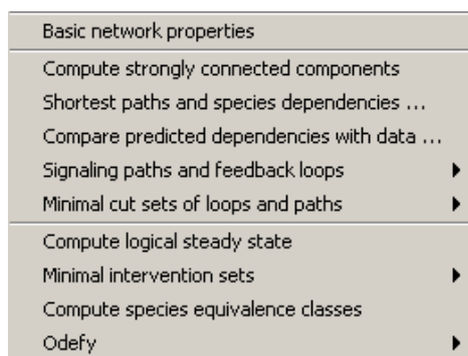


Fig. 19: The CNA submenu “Analysis” for signal-flow networks.

The menu is partitioned into the following groups, with (*) and (**) indicating the core functions for exploring the functional properties of the interaction network:

The menu bar for signal-flow networks has six entries (see also Fig. 16):

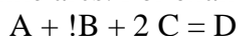
- **Network:** Editing/Saving the network model
- **Map:** Zoom tools and saving a map with current values.
- **Scenario:** Saving/Setting/Loading scenarios
- **Clipboard:** Clipboard and arithmetic operations
- **Analysis:** functions for analyzing the network (see Fig. 19).
- **Info:** info about CellNetAnalyzer

Many menu items are analogous to their counterparts in mass-flow networks (cf. section 3.3).

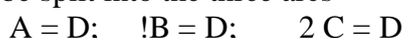
4.3.1 Menu “Network: Editing/Saving the network model

Network/Network composer ... : This opens the main window for editing and saving the network structure as it was described in detail in the previous sections.

Network/Convert to interaction graph: Each interaction with AND connections (i.e. hyperarcs with at least two species on the left-hand side of the interaction equation) are split into single graph-like arcs. For example, an interaction equation named ‘ieq1’ that reads



would be split into the three arcs



The original interaction (‘ieq1’) is removed whereas the three new arcs represent now new

interactions. Each of them gets a new name (composed of the original name of the split interaction and the name of the source species of the arc; in the example above: 'ieq1_A', 'ieq1_B', 'ieq1_C') and an associated text box, which is positioned closely to the original text box. For certain computations, this explicit graph representation is more appropriate. For example, the computation of signaling or influence paths (see below) is internally always performed within the underlying graph model and then mapped back to the hypergraph model which might sometimes lead to non-unique mappings. For example, Fig. 16 shows a signaling path computed with CNA (see below) in which the interaction (hyperarc) 2&3 participates – it is actually not clear whether the path starts at the left (I1) or the right (I2) branch of 2&3. After changing the network into an explicit graph representation, the display of this path would unambiguously indicate that either I1 or I2 is involved. Certain computations (e.g. minimal species cut sets of paths) even require the graph representation.

Before the network is converted to an interaction graph, the user is asked whether he really wants to perform the conversion since the original hypergraph model will be lost in this session (functions for logical analysis are usually not sensible anymore, e.g. splitting 2&3 in Fig. 16 would mean that either $I1=1$ OR $I2=0$ would suffice to activate E which is not equivalent to the original hypergraph model). Hence, changes made in the hypergraph model before should first be saved (whereas the new (graph-like) network structure may also be saved separately.). Furthermore the user is asked whether CNA should remove parallel (duplicate) arcs which may arise during splitting. For example, assume you have two interactions that may activate a species Z, namely $A + B = Z$ and $A + C = Z$. Splitting these two hyperarcs would lead to four new arcs: $A = Z$, $B = Z$, $A = Z$, $C = Z$. Thus, the connection $A = Z$ would be represented by two equivalent (parallel) arcs. This may lead to undesired effects. For the example above: computing signaling paths or feedback loops, all paths and loops that lead over A and then Z would be computed twice as there are two parallel arcs from A to Z. This can be avoided by letting CNA remove parallel arcs when converting the network to an interaction graph.

Network/Export provides two methods for exporting the network model.

Network/Export/Export interaction matrix ...: Exports interaction matrix, NOT flags (how interaction matrix and NOT flags are stored, see chapter 7), reactions and species identifiers for further analysis outside of *CellNetAnalyzer*. The user can choose whether the matrix is to be exported as ASCII or MATLAB file. In ASCII format the network is saved as follows:

```

first row: <reaction names>
second row <name of first species> <first row of interaction matrix>
third row: <name of second species> <second row of interaction matrix>
....<and so on>

```

If the matrix is stored in MATLAB format, 4 variables are saved which can be reloaded in MATLAB later on:

```

species_names: names of the m metabolites (char array)
reaction_names: names of the q reactions (char array)
NOTflags: a zero indicates a NOT operation in the respective interaction equation
intermat: interaction matrix (m x q), where the m rows correspond to the
species in species_names and the q columns to the reactions in reaction_names

```

Network/Export/Export in SBML qual format ...: After specifying a file location, the logical model is exported in SBML qual format (qual = qualitative / logical models).

Network/Import SBML qual format: Imports simple SBML qual models (from an .xml file)

via the libSBML/SBMLToolbox parser (the required files of the SBMLToolbox are included in the CNA distribution; see also <http://www.sbtoolbox.org/>) and python scripts. Note that only simple SBML qual models can be imported (for example, only AND, OR and NOT operators are allowed; only 0 and 1 are allowed for the discrete state variables) and that this function requires MATLAB >= 2014.b and Python installed.

Generally, to create a new CNA signal-flow project from an SBML qual file proceed as follows:

- 1.) Create a new CNA project with the project manager (section 2). There, you can also specify which network map(s) you want to use.
- 2.) Load the network, which is empty at this stage.
- 3.) Start “Network/Import SBML qual model ...” and select the SBML file. After conversion, the network will be reloaded and the text boxes for the reactions are arranged (by default) in the upper right corner. You now have to move the text boxes to their respective position on the network map (via Network composer; see section 3.2).
- 4.) Save the network and you are done.

SBML qual models can also be imported via an API function (see section 7.4).

Network/Visibility of text boxes ...: This sub-menu allows one to hide/make visible the text boxes of reactions and/or species. This is often useful to make crowded network maps easier to read. The default visibility of all text boxes may be set back whenever required.

Network/Element Selector ...: Tool for searching network elements fulfilling a given set of properties. Explained in detail in section 3.2 (Fig. 10).

Network/Display interaction equations: Displays all interaction equations in the command window.

Network/Show names of network elements: In each text box the name of the associated network element (reaction or species) is shown. This function is especially useful when moving text boxes (“Move Text Boxes ...” in network composer window; section 3.2).

The names of reactions and species, respectively, are also shown as tooltip strings when moving the mouse over the text boxes.

4.3.2 Menu “Map”

Map/Set original map size: Each network map is resized to its original resolution (pixel-based graphics) or the original size (MATLAB figures). The maps should then appear well-displayed. Ideally, the network graphics should a resolution not larger larger than the monitor resolution.

Map/Zoom tools on/off: Displays the zoom tools in the network map in which this menu item has been selected (or the zoom tools disappear if they are currently visible). The zoom tools are always mounted in the upper right corner of the map. The scrollbars allow one to scroll the map left/right and up/down, respectively (note: sometimes, the scrollbars are not correctly displayed by MATLAB; you should then click somewhere in the map or just minimize and then maximize the window). The ‘+’ button enables to zoom in. Press the ‘+’ button and then select two points on the map using the cross-hair. The rectangle spanned by these two points will be the new region displayed in the window. The text boxes are also zoomed with the respective factor. By pressing the ‘-’ button one can zoom out (factor of 2). Finally, by pressing the ‘R’ (reset) button

the entire map is displayed again. Note that the assignment of text box positions can also be done in the zoomed state.

Pan (select from toolbar): Can be used to activate the pan mode of the figure. When the pan mode is switched on you can click and drag the network map around inside the window. This will only have an effect if a part of the map is currently displayed (e.g. after zooming in). Due to limitations of the MATLAB GUI the textboxes are hidden while dragging the maps. Pan can be used in conjunction with zoom but then zoom must be activated first.

Map/Save map in graphics file ... : The map within which this menu item is selected can be saved as a graphics in a selectable format (e.g., EPS, bitmap graphics). The map will be saved together with the text boxes and the currently displayed values. You should not save the map as a MATLAB figure!

4.3.3 Menu “Scenario”: Saving/Setting/Loading scenarios

The **Scenario** menu provides several functions for saving, setting, loading and displaying scenarios. A scenario is the set of numeric values of reactions and species that are currently displayed in the respective text boxes.

Scenario/Reset last scenario: The values defined prior the last calculation are displayed in the text boxes.

Scenario/Set default scenario: The default values are displayed in the text boxes. For network elements without a default value, “#” is displayed indicating an unknown/undefined value.

Scenario/Save scenario: Opens a dialog box for saving the current text boxes values in a file. By default, the file has extension “val” (“values”).

Scenario/Save species values for ProMoT: Opens a dialog box for saving the current species values in a text file that can be loaded by ProMoT (the values can then be displayed in the context of the network using ProMoT’s Visual Explorer). The user also has to define an “attribute name” that will be required by ProMoT (see ProMoT manual).

Scenario/Save reaction values for ProMoT: Opens a dialog box for saving the current reaction values in a text file that can be loaded by ProMoT (the values can then be displayed in the context of the network using ProMoT’s Visual Explorer). The user also has to define an “attribute name” that will be required by ProMoT (see ProMoT manual).

Scenario/Load scenario: For loading a scenario (formerly saved by “Save scenario”). A window opens for selecting a scenario file (by default with extension “val”). In case that some numeric values are already given in the text boxes, the user will be asked whether the given values are to be overwritten (current text box values of all those elements are retained for which no new value is assigned in the loaded scenario) or whether all text box values are set to be undefined (“#”) before loading the scenario. After loading, the text box values are highlighted in the same way as done by Scenario.../Highlight values (see below).

Attention has to be paid if the scenario had been saved prior the network structure has been modified. Then, problems may occur if names of network element have been changed. Network elements that have been added will not cause problems (they get value “#”).

Scenario/Clear all values: The string “#” (meaning “undefined”) is displayed in all text boxes.

Scenario/Clear all reaction values: The string “#” (meaning “undefined”) is displayed in all reaction text boxes, current text box values for species remain unchanged.

Scenario/Clear all species values: The string “#” (meaning “undefined”) is displayed in all species text boxes, current text box values for reactions remain unchanged.

Scenario/Set all input/output gates to zero: A zero value is displayed in all text boxes

(i) of input and output species, i.e. of species without predecessor or without successor

(ii) and of all interactions representing an input or output arc.

Other text box values may be retained or not (the user is requested). This prevents (activating) signal input crossing the boundaries and is useful, for example, for computing feedback loops.

Scenario/Set time scale scenario...: The user is requested to enter a time scale. Then, all reactions having a time-scale larger than the entered value will be assigned a zero value (indicated also by color “special” assigned for the respective text boxes). The user is asked whether the values of all other text boxes should be retained or cleared (set to “#”, i.e. undefined).

Scenario/Highlight values: This function highlights all text boxes with a given (known) value; zero values are indicated by color “special” and non-zero values by color “computed” (see section 2). Undefined values get standard color. This is especially useful in large networks.

Scenario/Highlight values (on/off): This function colors the reaction and species text boxes according to their displayed value. Text boxes with a zero value are indicated by color “special” and those with non-zero values by color “computed”. Boxes with non-numerical values remain in standard color. This function is useful to quickly identify on/off values of the current scenario.

Scenario/Highlight values (heatmap): This function is similar as the previous one but uses the heatmap style for highlighting the text boxes. The user has to define two threshold values; one for reaction text boxes and one for species text boxes. These thresholds define the ‘zero’ or ‘neutral’ value. All text boxes having a larger value than the threshold (separately for species and reactions) will become green; the saturation of the green hue is determined by the ratio

$$(\text{text box value})/(\text{max_value_of_all text_boxes} - \text{threshold value}).$$

Analogously, for all text boxes having a smaller value than the threshold value, the text box color becomes red; the saturation of the red hue is determined by the ratio:

$$(\text{text box value})/(\text{threshold value} - \text{min_value_of_all text_boxes}).$$

(Boxes with non-numerical values have standard color.) One may use this function, for instance, to display and highlight measured data of gene expression or protein phosphorylation (activity) within a network context.

Scenario/Show current values in bar chart: Displays currently given reaction and species values in (two) bar charts. Useful for comparisons, e.g. participation levels of reactions in signaling paths

4.3.4 Menu “Clipboard”

Clipboard/Values to clipboard: Works like a usual clipboard: The currently given numeric

values in the text boxes are copied to a clipboard. The following two functions can be applied to the clipboard.

Clipboard/Paste values from clipboard: Takes the values from the clipboard and pastes them back into the text boxes.

Clipboard/Arithmetic operations: Useful for arithmetic operations on reaction and species values of two different scenarios (e.g. computing and highlighting differences). The four basic calculation operations are available for linking the values currently displayed in the text boxes with the values from the clipboard. Alternatively, arithmetic operations can also be performed with a user-defined numerical value. The results are displayed in the text boxes.

4.3.5 Menu “Analysis”: Analyzing the network

Analysis/Basic network properties: Displays some basic topological properties in the command window. This function is particularly useful to detect errors in the network structure that might have been inserted during network construction.

- *Species participating in no interaction* are detected
- *Sources and Sinks:* Species that are not influenced by others (=sources: no predecessor) or that do not influence others (=sinks: no successor) are detected.
- *Input Arcs and Output Arcs:* Interactions with an empty left-hand side (INPUT) or with an empty right-hand side (OUTPUT) in their interaction equations are identified. These arcs typically represent signal flows across system boundaries (e.g. *I_{in}* and *O_{out}* in Fig. 16).
- *All non-monotone interactions* and those with incomplete truth table or to be excluded in logical computations (see Table 2) are displayed.
- *Parallel Reactions:* Reactions with the same participating species are detected.
- *Number of different time scales.*
- *Basic Properties* of the interaction matrix.

Analysis/Compute strongly connected components: A strongly connected component (SCC) is a maximal subgraph of the interaction graph in which paths between all pairs of nodes exist. Consequently, when searching for cycles that involve a certain node it is only necessary to search in the SCC that contains this node. After calculation, SCCs that comprise more than two nodes are visualized on the network map: The species that belong to the same SCC are labeled with the same number and in addition their text boxes are colored with the same color. Because colors (currently four) are used repetitively, species belonging to different SCCs can have the same color but their labels are unique to the SCC.

Analysis/Shortest paths and species dependencies ...: Similar function as “Graph-theoretical path lengths ...” in mass-flow networks but even more powerful. Allows for the calculation of *graph-theoretical* (shortest) path lengths between all species, separately for negative and positive paths from which then (by taking the minimum of both) the unsigned shortest path is also computed. Additionally, the average path length and the network diameter are computed (for the unsigned shortest path length). Furthermore, a dependency matrix is determined (see below) which is extremely helpful for detecting functional dependencies between all pairs of species.

Note that a signal-flow network in CNA is a Boolean network represented as a *hypergraph*. Therefore, for applying graph-theoretical methods, this hypergraph is temporarily transformed into its underlying interaction graph (see also “Convert to interaction graph”). Regarding the interpretation it is important to realize that the shortest paths which are computed with this

function indicate whether there is a positive/negative/any influence of a species A on B, without a guarantee that the path is active under a given scenario in the logical model. Note that the shortest paths algorithms work well also in large networks.

Prior to the computation, the user can choose several options (Figure 20):

- *Exclude species with given zero value:* This option can be used to exclude certain species and, hence, certain possible paths in the network while computing the shortest paths (enables, for example, to simulate a certain failure or knock-out). Just set a zero value in the text boxes of species to be excluded and select this option.
- *Exclude reactions with given zero value:* This option allows one to exclude certain interactions and, hence, certain possible paths in the network while computing the shortest paths. Set a zero value in the respective reaction text box before computing the paths and select this option.
- *Display distance matrices:* Displays in three separate windows the shortest positive/negative/unsigned path lengths (distances) as colored arrays. The array element e_{AB} represents the shortest path length from A to B. The larger the path length is the brighter the array element. A blue cell marks species pairs between which no path exists.
- *Display dependency matrix:* Finally, a dependency matrix is determined which characterizes the functional dependency between each pair of species: The color of the matrix element e_{AB} indicates one of 6 possible types of dependency: (1) black: no influence of A on B, (2) yellow: A has activating and inhibiting effect on B, (3) light red: A is pure inhibitor of B, (4) light green: A is pure activator of B, (5) dark red: A is total inhibitor of B, (6) dark green: A is total activator of B. Dependencies with respect to one single species may also be displayed directly within the network maps after the computation has been finished (see below).

In very large networks, the creation of distance and dependency matrices might fail in MATLAB. Use then instead the option:

- *Export distance + dependency matrix:* This allows for an export of the calculated distance and dependency matrices into an ASCII file. The user may then further study the path lengths matrix by its own algorithms. Each distance matrix (total, negative, positive path lengths) and the dependency matrix will be exported in separate files (the user specifies the files upon finishing the computation). The dependency matrix contains in each cell one of the classification number 1-6 as described above.
- *Algorithm:* In contrast to standard shortest path algorithms in unsigned graphs, the computation of shortest positive / shortest negative paths and cycles is not trivial; the algorithm is not polynomial-time and computing the pos./neg. paths may thus take a long time (dependent on network size and structure). CNA provides three algorithms: (i) a simple exhaustive search algorithm which delivers exact results and which – although not polynomial – proved to be efficient enough also for larger networks with several hundreds of nodes (ii) for larger networks or those with very complicated structure (e.g. many negative feedback loops), CNA provides an approximative algorithm which is faster (polynomial complexity; seconds or minutes in large and complicated networks) and has, to our experience, a high probability to compute the correct distances [14]. The approximative algorithm is definitely correct if no negative feedback circuit is contained in the network. However, for special configurations it may happen that negative (positive) paths between a pair of nodes A and B are overlooked if a positive (negative) path from A to B exists and the network contains negative feedback loops. (iii) Another exact algorithm which turned out to be favorable (compared to the first one) in certain classes of networks is the Two-

Step-Algorithm [14]. In general, try first the exact algorithms (i) or (iii), if they are too slow take (ii).

- *Maximal Path Length*: You are allowed to restrict the paths to be considered by setting an upper bound for the path length (longer distances will not be considered decreasing also the computation time).

After computation, depending on the chosen options, the distance matrices and/or dependency matrix are shown and exported if desired. Besides, average path length, network diameter (longest shortest path length) and the number of connected components in the graph are displayed in the command window.

Finally, a small panel comes up allowing a detailed influence/dependency analysis (relying on the computed dependency matrix): Enter a species identifier in the respective input field, select (case 1) “Dependency on all species” or (case 2) “Influence on all species” and click then on “Compute”. In the first case, each compound is classified into one of the following categories: (total) activator (only positive paths to the selected species exist), (total) inhibitor (only negative paths to the selected species exist), ambivalent factor (positive and negative paths to the selected species exist) or non-affecting factor (no path to the specified species exist at all). The classification is displayed in the species text boxes, also illustrated by different box colors. In the second case, the converse relationships is determined (for which compounds is the specified species an (total) activator, (total) inhibitor, ambivalent factor or non-influencing) and then displayed in the species text boxes. Regarding the classifications, in particular for *total* activators and inhibitors, see [11].

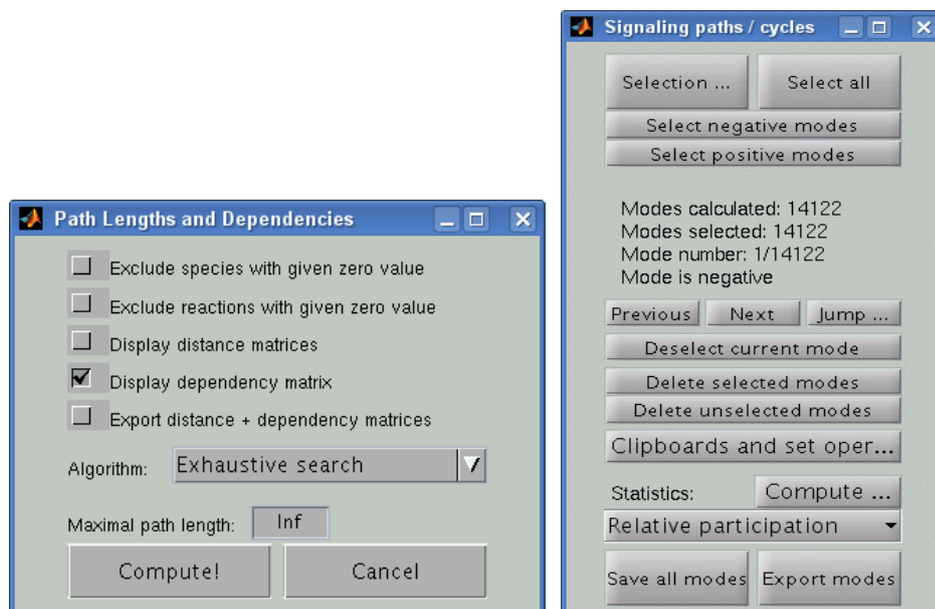


Fig. 20 (left): Options for computing shortest paths and species dependencies

Fig. 21: Control panel for displaying and analyzing signaling paths and cycles.

Analysis/Compare predicted dependencies with data ...: With the help of the dependency matrix, causal dependencies between each pair of species in the model can be predicted. Having data available where the activation level of certain species is measured in response to different stimuli (e.g. adding ligands and/or inhibitors), one can detect whether an increase or decrease in the activation levels is in accordance with the predicted causal dependencies (see [17]). For example (Fig. 16), assume species O2 shows a higher activation level when it is measured in

absence of input I2 than it does in presence of I2. Since I2 is a (strong) activator for O2 in this network, this leads to a contradiction between the model and the experimental data: an increase in I2 cannot induce a decrease in O2, as there are only positive paths connecting I2 and O2. (Hint: this function assumes that the data reflects the initial responses of the species, i.e. weak activators and weak inhibitors contained in the dependency matrix are handled in the same way as strong activators and strong inhibitors, respectively.)

The information on the experimental data must be stored in a structure array named *CNADEPProject* that contains the following fields:

CNADEPProject.InputMatrix: Matrix that contains in the rows the different experimental scenarios (i.e. adding a ligand, inhibiting a certain species ...) and in the columns the species. It is not necessary that all metabolites of the current model have a column in the *InputMatrix* (however, the column order must fit with the *MetaboliteNames* field and the columns in *OutputMatrix* (see below)). A positive entry corresponds to adding a ligand or activating a certain species (a larger value indicates a higher concentration) whereas negative entries are used for inhibitors (a smaller value (i.e. larger absolute value) indicates a stronger inhibitor). Also important, a species having an undefined value is indicated with 'NaN'.

CNADEPProject.OutputMatrix: Matrix that contains observed species values (measured or from literature). The rows (scenarios) and columns (species) must fit with the *InputMatrix*.

CNADEPProject.MetaboliteNames: Cell array of strings or character array that contains the metabolite (species) names (the first entry corresponding to the first column of the *InputMatrix* or *OutputMatrix*, second entry to second column and so on). The metabolite names have to fit with the species names in the model (although not all species of the model must be present here, e.g. if they have not been measured).

CNADEPProject.NegativeStates (optional): Cell array of strings or character array that contains species that are assumed to be constitutive active at time 0 and are inactivated when the signal proceeds. If the field does not exist, an empty field is created.

CNADEPProject.ScenariosToCompare (optional): Matrix that contains in each row one pair of scenarios the user wants to compare. For example to analyze the influence of a ligand, one compares a scenario x where the ligand was added with a scenario y where it was not added. The respective row in the matrix *ScenariosToCompare* would store the indices of these two scenarios.

If this field does not exist, all scenarios are compared (pairwise) to each other.

When starting this routine, similar to the computation of shortest paths and dependencies, the user can choose several options prior to the computation. For the comparison of the predicted dependencies with experimental data the model's dependency matrix is computed. Therefore, the first three options are already known from the previous function (see figure 19):

- *Exclude species with given zero value*
- *Exclude reactions with given zero value*
- *Compute approximative dependency matrix*

Additional options are:

- *Display comparison results*: The result of the comparison is displayed as colored matrix, where the rows correspond to the compared scenarios and the columns to the measured species. Row labels indicate which ligands/inhibitors were added in both scenarios with the same concentration (green dot for added ligands, red dot for added inhibitors) and how the stimuli of the two scenarios differ (e.g. ligand A↑). For example, the label I1 •, A •, I2 ↑ refers to the comparison of two scenarios that were

both stimulated with the same concentrations of ligand I1 and of an inhibitor for A, but with different concentrations of ligand I2 (possibly I2 was only added in one of both scenarios), i.e. the influence of an increased level of I2 is analyzed. The color of a matrix element indicates one of 10 possible cases that are explained in the legend displayed in another window (e.g. light green: treatment causes increase as expected; orange: no effect expected, but treatment causes decrease).

- *Export comparison results*: Saves the structure *depmat_compare_results* to file. The structure contains the following fields:
 - depmat_compare_results.ExpectedDependencies*: Matrix that describes for each comparison the expected up/down behavior (resulting from the model structure).
 - depmat_compare_results.FoldChange*: Matrix that contains the fold changes in the compared scenarios.
 - depmat_compare_results.DiscChange*: Matrix that contains discretized values that one obtains by comparing the FoldChange with a fixed threshold (see below). 1: significant increase (FoldChange > threshold); -1: significant decrease (FoldChange < 1/threshold); 0: no significant change; 5: states that are not compared, because an external intervention on the measured species was made.
 - depmat_compare_results.ComparisonResult*: Matrix that contains the result of the comparison, coded with 1, 2, ..., 10. 1: no significant change, positive or negative effect expected; 2: increase although no change expected; 3: decrease although no change expected; 4: increase although negative effect expected; 5: decrease although positive effect expected; 6: influence on a species is not considered if an external intervention on this species was made; 7: decrease as expected; 8: increase as expected; 9: no significant change as expected; 10: ambivalent in the model.
 - depmat_compare_results.Label*: Row labels, indicating which scenarios are compared and how the stimuli differ.

Finally, the fields *Ratio threshold* and *Data file* should be specified:

- *Ratio threshold*: threshold to define when the FoldChange is significant. FoldChange > threshold: significant increase; FoldChange < 1/threshold: significant decrease. Default value is 1.5.
- *Data file*: Enter the path of the MATLAB (mat) file where you saved the *CNADEPProject* data structure. Alternatively, you can use the *Browse* button.

Start the computation by clicking the *Start!* button.

To test this function an example data file (containing a fictitious dataset) for the Signaling Toy model is provided ('testdepmatcomp.mat' in the 'sigtoynet' directory).

Analysis/Signaling paths and feedback loops: This sub-menu allows one to compute, display and analyze feedback loops and signaling paths in the underlying interaction graph of the logical network model. In general, it is recommended to first convert the logical (hypergraph) model into an interaction graph (see above) when using this function, however, it can also directly be performed in the logical network. The sub-menu provides two main functions:

Feedback loops ...: Computes the internal feedback loops (= circuits or (elementary) cycles) in the underlying interaction graph of the network. For example, in Fig. 16, there are two internal circuits (loops, directed cycles). Although the elementary modes algorithm can be used for computing the cycles (see [11]), CNA uses a backtracking algorithm for enumerating the cycles. The

following three options can be chosen in a dialog box appearing after selecting this menu item:

(i) *Enforce/Exclude reactions by given values*: if selected, the numerical values given in the text boxes of the reactions (interactions) confine the set of cycles to be computed (similar as for elementary modes in metabolic networks): all interactions with a zero value in their text box are blocked during circuit computation and if an interaction has a non-zero value in its text box then CNA will compute only those cycles that involve this interaction. For example, to compute only the cycles involving interaction 6, one would set a “1” in the associated text box (one circuit would be found). Note that it is not possible to enforce a “hypergraphical” interaction where several species are AND-connected. Thus, interactions “1&4” und “2&3” can not be enforced in Fig. 16. For those cases, you should first convert the network into an interaction graph (see above). Another important issue is, that multi-level interactions (coefficients larger than 1 in an interaction equation) are treated as binary interactions (all non-zero coefficients are set to 1).

(ii) *Enforce/Exclude species by given values*: as for reactions, when selecting this checkbox, the numerical values given in the text boxes of the reactions (interactions) confine the set of cycles to be computed: only those cycles are computed, that run over all the nodes having a non-zero value in their box and that do not visit nodes with a zero-value in their box. For example, to compute the (only) circuit where A is involved, write a “1” into the box of A, select this checkbox and start the calculation. As for reactions, multi-level interactions (coefficients larger than 1 in an interaction equation) are treated as binary interactions (all non-zero coefficients are set to 1).

(iii) *Consider unsigned graph*: if selected, CNA ignores the direction of the arcs (i.e. an undirected graph model is evaluated). This can be useful e.g. to compute also feedforward loops (there are two such feedforward loops in Fig. 16 (additionally to the two feedback loops). (Note that option (ii) is similar to option “Check reversibility” for elementary modes in mass-flow networks).

Signaling paths...: This function works similar to the previous one: Instead of feedback loops it computes all paths connecting one species from a specified set of start nodes with another species from a specified set of end (target) nodes. In the upcoming panel, the user may specify a set of start and a set of target nodes by entering the respective species identifiers. Additionally (as a short cut), he may optionally include all input nodes (= nodes that have no predecessor or are connected to an input arc) to the set of start nodes or/and all output nodes (= nodes that have no successor or are connected to an output arc) to the set of target nodes. As explained above for feedback loops, one may optionally exclude or enforce certain interactions and/or nodes by setting zero/non-zero values into the respective text boxes prior to computation. An example: five signaling paths would be found, if one selects the input nodes (I1, I2) as start nodes and the output nodes (O1, O2) as target nodes. Only two of them would remain if one sets a 1 in the text box of interaction 9 and a zero in the text box of reaction 8 (and enables the checkbox ‘*Enforce/Exclude reactions by given values*’).

When the computation of cycles or signaling paths has finished, a control panel comes up (Fig. 20) which is very similar to the panel for managing the computed set of elementary modes in mass-flow networks (Fig. 12, paths and cycles are also referred to as modes in Fig. 21). The usage is completely analogous as explained for elementary modes in section 3.3. Paths and loops can be displayed subsequently in the network by using the “Next” and “Previous” button. Involved reactions on a path are indicated by value “1” and the text box color is “calculated“. Note again that paths and cycles are computed in the interaction graph and then mapped back to

the logical hypergraph (see remarks for menu item “Convert to interaction graph”). Specific selections of modes can be made by using the ‘Selection tool’ (input of reaction / species IDs is supported with autocomplete; see section 0.6). The same statistical computations can be made as for elementary modes; exceptions are the features “(Product) Yields and rate ratios” and “Control-effective fluxes” which are for obvious reasons not available in signal-flow networks. Regarding Minimal Cut Sets see below.

There is one major difference regarding elementary modes in stoichiometric networks and paths and cycles in signaling networks: a reversibility property is not specified in signal-flow networks. On the other hand, paths and cycles can be positive (activating influence) or negative (inhibitory influence), depending on the parity of negative signs (NOTs) on the path. The sign of the currently displayed mode is therefore shown in the control panel. Furthermore, two additional buttons allow for a selection of all negative or all positive modes of the current selection. With the clipboard functions and the buttons ‘Delete selected modes’, ‘Delete unselected modes’ and ‘Deselect current modes’ one may assemble more complicated selections of the set of paths/loops (as explained for elementary modes in section 3.3.). Finally the signaling paths and feedback loops may be saved or exported to an ASCII file.

The other two items of the “*Signaling paths and feedback loops...*” sub-menu are *Load ...* and *Show ...* which allow one to load a set of paths/loops saved in an earlier session or to show a set of paths/loops computed or loaded earlier in the current session (and still being in memory, see also section 3.3. for the same menu items for elementary modes).

Analysis/Minimal cut sets of loops and paths: As the minimal cut sets in mass-flow networks attack a selected set of elementary modes (section 3.3), the minimal cut sets (MCSs) computed with this function attack (cut, interrupt) the currently selected set of paths or/and loops in a signal-flow network. Hence, before computing MCSs for paths/loops the user has to compute these modes by one of the two methods described above.

In signaling networks one may determine MCSs removing (only) *reactions* or MCSs removing (only) *species*. (From an experimental point of view, the removal of species is realistic in signaling networks, while practically infeasible in metabolic networks). Therefore, when choosing a function from this sub-menu, the user has to specify whether species MCS or reaction MCSs are to be computed/loaded/displayed.

Computing, analyzing and displaying reaction MCSs is analogous as for elementary modes (see section 3.3):

- (i) the user selects a (sub)set of modes (paths/cycles) that is to be attacked (interrupted) – the current selection defines this set of target paths/cycles.
- (ii) As for MCS for elementary modes, using the paths/cycles clipboards one may define up to two sets of paths/cycles for which a minimum number (to be specified) must not be hit by the MCSs to be computed.
- (iii) choose from menu “Minimal cut sets for paths .../Compute...” and choose *reaction MCSs*
- (iv) specify options for calculation (the same as for “metabolic MCSs” (described in section 3.3), except that multifunctional enzymes are not supported)
- (v) when the MCSs have been computed, a control panel comes up allowing to walk through the set of MCSs (while displaying them in the interactive maps), to select certain subsets of MCSs, to compute statistical properties and to save the MCSs (these functionalities are all described in detail in section 3.3)

Note that negative/positive effects of removing reactions are not considered when computing the MCSs (see also item (b) for species MCSs).

Computing MCSs for species, some particularities have to be taken into account:

- (a) Species MCSs can only be computed in interaction *graphs* (this is also recommended for reaction MCSs), Therefore, if you want to compute species MCSs for paths, first apply “Convert to interaction graph” from the menu, compute the paths in the interaction graph and then the species MCSs for these paths.
- (b) CNA assumes that you want to interrupt a signal flow along a path or within a feedback loop. Whereas this procedure is straightforward for feedback loops and in graphs with exclusively activating (+) arcs, it might have ambiguous effects for signaling paths with negative arcs. For example, assume in Fig. 22 the negative path starting in I1 and leading to O1 via E should be attacked. Each species (I1, E, C, O1) on this path would represent one Minimal Cut Set. However, whereas removing I1 would have a positive effect on O1, removing E would have the opposite effect: it would indeed correctly disrupt the path starting in I1 but the inhibition along this path would virtually still be active since E is an activator of O1 (removing E is structurally equivalent to activate I1 inhibiting E). These effects are not taken into account by the MCS approach – the computed MCSs will only ensure, that no signal can flow from the start to the end node in any of the target paths. Such effects can be much better taken into account when computing Minimal Interventions Sets (see below) in the logical model, possibly with incomplete truth tables.

The rest is analogous as for reaction MCSs. Options for the computational procedure may be specified (same options as offered for reaction MCSs). Then, a control panel comes up which is similar as the one for reaction MCSs and analogous in usage. Of course, the main difference is, that for a displayed MCSs not reactions but species are marked in the network maps. Inactivated (removed) species are indicated by a zero value (the box color is of type “special”) in the interactive maps. Specific selections of MCSs can be made, some statistical features can be computed (again analogous as for MCSs in metabolic networks) and all MCSs can be saved or exported.

Again, with the *Load ...* item in the minimal cut sets sub-menu one may load a saved set of (reactions or species) MCSs.

The remaining functions of the” **Analysis**” menu analyze explicitly the logical network.

Analysis/Compute logical steady state: Takes the currently defined reaction values (=fixed signal flows) and species values (= fixed states) and computes from this information – as far as uniquely possible – the resulting logical steady state of the species as it follows from the Boolean network structure. This function is in a certain sense similar to metabolic flux analysis (section 3.3).

First, the user may fix some states of species (which is particularly relevant for INPUT species and for indicating knock-outs or knock-ins in the network). Additionally, the user may enter values for signal flows along selected interactions. This is particularly relevant for INPUT arcs (see Figure 21), but it might also be useful to deactivate certain interactions by setting the signal flow along this interaction to zero. Note that only INPUT arcs are allowed to have a zero or non-zero value whereas all other interactions may only be assigned a zero (non-zero values will be ignored in the latter case). The order of evaluation of given species/reaction values is as follows:

- if a species value has been defined, then the logical state of this species will be fixed to that value; any given value for a reaction that points into that species will be ignored
- a reaction will be considered to be off (i.e. temporarily removed) if a signal flow of zero has been defined for that reaction (note: if all reactions pointing into a species have been set to zero, this species will be assigned a zero in steady state, i.e. this is analogous to setting the respective species value to zero)

- if a non-zero value has been defined for an INPUT arc, then the state of the node connected to this INPUT arc will be fixed to that value. If non-zero values for several INPUT arcs pointing into one and the same species have been defined then the one with the highest activation level will be taken to fix the state of the species. If a non-zero value has been defined for at least one INPUT arc all other signal flows pointing into that reaction will have no effect on the logical steady state of this species. Thus, if you have two interactions “= A” and “B = 2 A” then setting a “1” for the first interaction and letting the second undefined means that A will be fixed to 1 irrespective whether B will be assigned a value of 1 or not. Furthermore, for an INPUT arc of type “= 2 A”, setting any value different to “2” will be automatically converted to “2” in order to be consistent with the level of the signal level flowing along this reaction. Generally, a simpler way than using INPUT arcs is to set the respective node value directly in the associated species box.
- setting a non-zero value for reactions that are not INPUT arcs will be ignored

After defining fixed species and reaction values and starting this function, the logical steady states resulting from the given scenario are computed. The algorithm tries iteratively to derive logical steady states for species until no further update is possible (see reference [11]). Fig. 22 shows an example: Setting the signal flows for the input arc *I1in* to zero and for *I2in* to 1 (given signal flows of input arcs are colored as “defined” – here in blue) and letting all other species and reaction values undefined, the logical steady state as displayed in Fig. 22 would result. For example, since I1 is inactive and I2 active, E becomes activated by hyperarc 2&3 and will remain in this active state (so the logical (qualitative) steady state of E will be 1). Then E will activate C, O1, D, F, G and O2. B becomes inactive because I1 is inactive and, since D is active, A becomes inactive. Resulting zero values for states (species) and signal flows (reactions) are indicated by color “special” (here red), non-zero values (the activation level) are indicated in color “calculated (here green) and unknown values in standard color. The signal flow values indicate whether an activation of the connected target species occurs (non-zero value: the level of activation reached) or not (zero value) along this interaction.

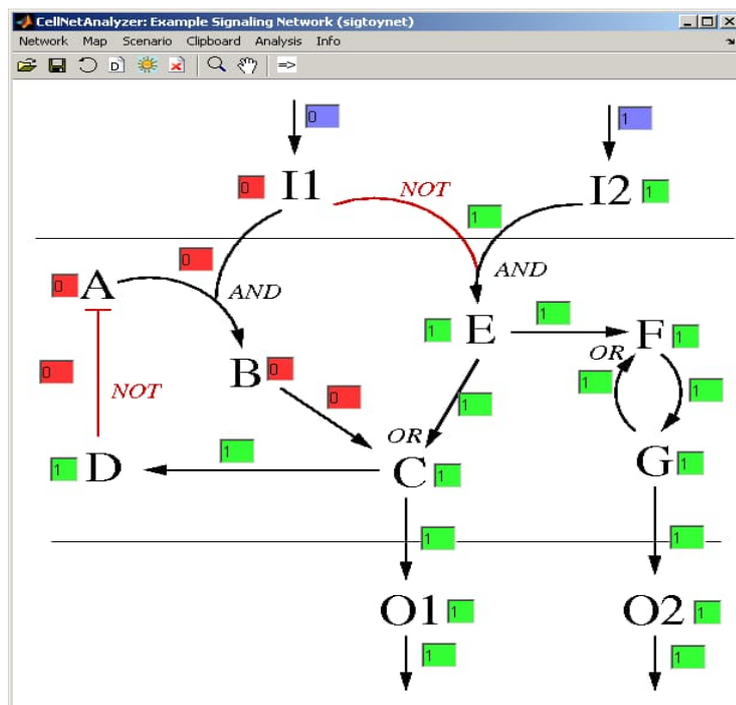


Fig. 22: A scenario of logical steady state analysis (smaller boxes: species; larger boxes: reactions). Predefined values are indicated in blue. Green values indicate calculated “on” state, red values “off” state.

Important remarks:

- The logical steady state resulting from a scenario depends heavily on the fixed signal flows and species states. It is based on an algorithm for three-valued logic (see ref. [20]).
- Note that a (global) logical steady state might not exist at all for a given scenario or that it cannot be resolved completely. For example, setting only the signal flow of input arc at I2 to 1 and everything else as unknown, nothing could be concluded. Setting instead the incoming signal flow at I1 to 1, we could derive at least a (partial) logical steady state, namely $E = 0$.
- There are a few cases where CNA does not identify existing logical steady state values. CNA does not (yet) check whether positive feedback loops are self-sustaining by initial state values of participating species (for example, in a dynamic Boolean network model, assuming initial values of “1” for F and G would result in partial logical steady states $F=G=1$ due to the positive feedback loop between F and G). Positive feedback loops can, however, be identified by the algorithms described above and therefore be checked by the user himself. There might be some other, for cellular networks very unlikely situations, where existing logical steady states are not identified. Certainly, all steady state values that have been found by CNA are definitely correct.
- As described in section 4.1, if the “Exclude in logical computations” – flag is set for a reaction, then it will not be considered as a possible activation route for the logical steady state identification procedure.
- Note also the particularity of interactions with incomplete truth table (section 4.1)

Analysis/Minimal intervention sets ...: As explained above, minimal cut sets (MCSs) computed for a set of paths/cycles refer to the underlying interaction graph of the logical signal-flow network. In contrast, this function computes *minimal intervention sets* (of species) directly from the logical network, i.e. the more constrained logical behavior of the network is taken into account (see [11,12]). Minimal intervention sets (MISs) are a set of interventions that fulfill a user-defined intervention goal. An intervention can be a permanent deactivation (knock-out/knock-down) or a permanent/constitutive activation (e.g. induced by overexpression of certain genes as occurring in certain diseases like cancer) of a certain node. The computation of logical MISs requires three steps guided by CNA:

- (i) The user defines an intervention goal. This can be done by setting the *desired levels* of reactions or/and species in the respective text boxes. For example, in Fig. 22 we might set a “0” for species O1 and a “1” for the output arc O2out (which is equivalent to setting a “1” for O2). This intervention goal demands that O2 has to be active in logical steady state and O1 inactive.
- (ii) The user may a priori fix certain states (for species) and signal flows (for reactions). [Regarding the conventions for fixing species and reaction values, see the explanations given under “Compute logical steady state”]. Setting such a particular scenario is useful to compute MISs valid only under certain conditions (of interest). In our example, we might assume that I2 is active, i.e. we set a zero for the input arc I2in. The MISs to be computed will then have to ensure that O2 is active and O1 inactive (in logical steady state) under the condition that I1 is inactive.
- (iii) First of all, certain species may be marked as non-removable/non-inactivatable (set a “-1” in the respective text box) or as non-activatable (set a “-2”) or as neither inactivatable nor activatable (set a “-3”). Since the computation of logical MCSs is intricate, this may reduce the computation time considerably. In addition, two options can be used to globally restrict the activation/inactivation (cf. Fig 23): (a) Species may be knocked-out

(removed/inactivated). (b) Permanent activation of species. Deselecting this option and enabling knock-outs (option (a) is on) is equivalent to set a “-2” in all species text boxes. In contrast, selecting this option and disabling knock-outs (option (a) is off) is equivalent to set a “-1” in all species text boxes. A third option (c) allows to exclude species that have a fixed value (due to the values set in second step) from interventions. This often makes sense because e.g. the value of boundary species is fixed due to external constraints. In our example, we assume that all species are removable and activatable. If we would select option (c), then I2 would be excluded no MIS containing I2 would be computed.

A second set of options can be used to define the maximum number of interventions, the maximum number of violated goals and whether or not to report the minimal number of discrepancies. As for MCSs of elementary modes and paths/cycles, setting an upper boundary for the cardinality of MISs may drastically reduce the computational demand. From a practical point of view, the small MISs are the most relevant ones. If you want to have all leave the default “Inf” which we use also in our example. If the maximum number of violated goals is greater than zero then also intervention sets are accepted that do not fulfill all goals as long as they do not contain more errors than the specified value. When the option to report the minimal number of discrepancies is activated then during calculation the minimum number of errors of all tested intervention sets is tracked. In case intervention sets were found this value simply corresponds to the minimum number of errors of these sets. When no intervention sets were found this value is the minimum number of errors that can be obtained by any intervention set under the specified conditions (in particular the maximum number of interventions). The actual value is printed to the MATLAB console. Note that the determination of the minimum number of discrepancies may increase the computation time compared to the case where this flag is off (in particular when multiple scenarios are used which is currently only possible via the corresponding API function; cf. sec. 7.6).

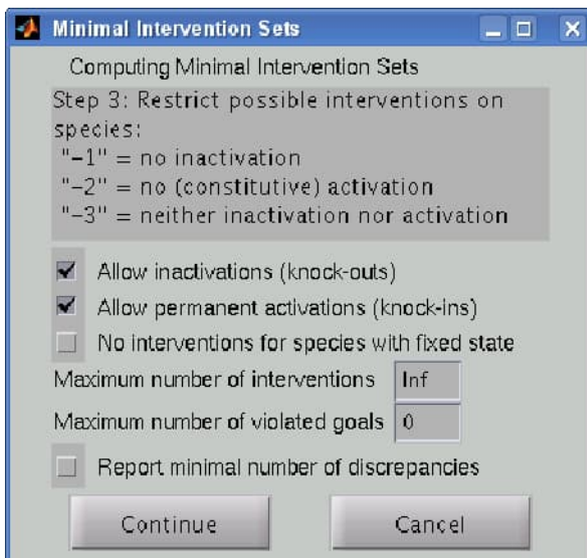


Fig. 23: Dialog for the third step when formulating a problem for minimal intervention sets.

Then, CNA computes the logical MISs which is, as already mentioned, another complex (combinatorial) problem. Upon finishing the computation, a similar control panel as for species MCSs of paths/cycles comes up by which the user can step through the set of MISs and display each MIS in the interactive maps. Inactivated (removed) species are indicated by a zero value (the box color is of type “special”) and activated species by a “1” (and box color “calculable”).

Specific selections of MISs can be made. Two statistical features can be calculated for the current selection: a histogram of the size of the MIS can be computed or a histogram with the relative participation of each species (separately for permanent activation/deactivation) can be displayed. Finally, the set of computed MISs can be saved or exported.

For our accompanying example, we would get 43 (!) MISs. Thereof two comprise only inactivations representing “logical cut sets” (as only cuts are contained): {!I1, !C} and {!I1, O1}. Three MISs are built up exclusively by activations: {I1, D, F}, {I1, D, G}, {I1, D, O2}. The remaining 38 MISs comprise at least one inactivation and at least one activation. One example is {!C, E}. (The exclamation mark indicates here a cut i.e. knock-out or inactivation).

Computing MIS as solutions to an integer linear program (ILP):

With this feature it is possible to more efficiently calculate the MIS for networks that only use monotone interactions and have no multi-value nodes. Furthermore, multiple scenarios are not supported nor may any goals be violated in the MIS. This is an experimental feature and can be invoked by setting the ‘Maximum number of violated goals’ (see Fig. 23) to a negative value. In order to calculate the MIS as solutions to an ILP a CPLEX installation with working JAVA interface is required (see instructions in section 0.5).

All other options described above have the same meaning and can be used as usual.

Analysis/Compute species equivalence classes: This function computes equivalence classes of species useful to identify synchronized network regions. All species of an equivalence classes are strictly correlated (positively or negatively) in every logical steady state that can be reached by the network, i.e. there synchronization holds for all possible inputs. For example, A and B are in an equivalence class if it follows from A being on (off) that B is also on (off) and vice versa. Species A and B are also in an equivalence class if they are negatively correlated, i.e. if it follows from A being off (on) that B is on (off) and vice versa. In the latter case the equivalence class would be written as {A, !B} or {!A, B}. You may also clamp values for species/arcs prior computation; in this case the equivalence is computed with respect to this scenario (for example, A and B might be in an equivalence class only if an input, say C, is set to be off). You should also keep in mind that the equivalences are only valid if the network reaches a logical steady state.

The computed equivalence classes are displayed in the text boxes (class number; with alternating text box colors for each class) and in the command window and may also be saved in a text file. Importantly, this function can only be used in binary-level networks.

Analysis/Odefy ...: *Odefy* is a plugin for CNA developed (with the help of CNA’s API) by Jan Krumsiek and Dominik Wittmann in the group of Fabian Theis at the Helmholtz Zentrum Munich. The *Odefy* homepage can be found here: <http://www.helmholtz-muenchen.de/cmb/odefy>. *Odefy* uses a new technique for transforming a discrete Boolean network into a continuous ODE (ordinary differential equations) model using multivariate polynomial interpolation ([16]). The ODE model can then be exported (e.g. as MATLAB or SBML file) or be simulated on-the-fly in CNA. *Odefy* also allows the simulation of a Boolean network (with synchronous or asynchronous update rules) and it can also run independently of CNA from MATLAB command line (see *Odefy* documentation). Importantly, so far *Odefy* works only with Boolean, not with multi-valued logical models.

It follows a brief explanation of the two main features (copied from the *Odefy* documentation which can be found in *CellNetAnalyzer/code/ext/odefy/doc/index.html*).

Odefy.../Export ODE model: *Odefy* can export the generated ODE model as MATLAB scripts, in SB Toolbox format, GNA or SQUAD format. In the dialog which pops up you select a filename

where the model will be stored. For MATLAB script files or SB Toolbox format you need to specify whether to create a HillCube ODE model, a normalized HillCube ODE model or a BooleCube ODE model (see [16]).

Odefy.../ Simulate Boolean / ODE model: Simulates Boolean models (only binary logic allowed) – either with synch/asynch. Boolean simulation or using the ODEs generated from the Boolean model. In the Simulation dialog you can specify all settings required for the continuous/Boolean simulation of the logical model:

- *Time units to simulate* - The number of time units to be calculated. For continuous simulations, *Odefy* uses the MATLAB `ode15s` ODE solver function with adaptive approximation intervals.
- *Edit parameters* - Here you can edit the Hill kinetics parameters for each interaction. Only required when simulating with HillCube / normalized HillCube (see [16]).
- *Edit initial values:* *Odefy* reads the initial values of the species directly from the species text boxes; so initial values cannot be edited when using *Odefy* with CNA. Reactions with a zero in their textbox are temporarily removed from the network before starting the simulation (see also remark below).
- *Simulation type* - determines whether a HillCube, a normalized HillCube (reaches $y=1$ at $x=1$) or a BooleCube transformation to an ODE model is chosen (see [16]). Alternatively, discrete Boolean simulations (with synchronous, asynchronous (with fixed order) or randomly asynchronous update rules) will be performed. Note that the BooleCube simulation ignores the n and k Hill parameters whereas the Boolean simulation ignores all parameters. For Boolean simulations with synchronous (all nodes fire at once) or randomly asynchronous (one randomly selected node fires) update rules, no parameters are required. For (non-random) asynchronous update (in each time step the state of only one node is updated and after n time steps each of the n nodes has been updated once), you must specify an order in which the species are updated. In *Odefy* you have three options: you can (i) load a permutation from the current workspace, (ii) update the species in sequential order or (iii) generate a random permutation.
Note: When performing asynchronous Boolean simulation, one time step corresponds to an update step of one species!
- *Plot type:* You can choose “Regular line diagram” or “Heatmap style”.
- *Save results into workspace* - If this box is checked, the simulation time vector and the corresponding concentration matrix will be stored in the MATLAB workspace as `simt` and `simy`, respectively. If you are performing a Boolean simulation, no time vector is created.

By clicking on *Simulate*, the simulation is started. Results will be displayed in a plot (of the chosen style).

Important: before the actual simulation starts, *Odefy* reads the values from the species and reaction text boxes. Values set in species text boxes are used *as initial values* for the respective species (should be between 0 and 1). Species for which no value is defined will be assigned an initial value of 0. Regarding the reaction text box values: all reactions with a zero value in their text box are (temporarily) removed from the network (non-zero values are ignored). In this way, a knock-out of a node can be simulated by setting all reactions pointing into that node to zero.

4.3.6 Menu “Info”

Info ...: Shows info about *CellNetAnalyzer*.

5. Editing ASCII files of network projects

In *CellNetAnalyzer* there are four possible ways to define/import the structure of a network (project):

- Use the GUI masks to define your network (as described in section 3.2 and 4.2).
- One may manually create or edit the network-specific files which are read when loading a project in CNA. In some cases, this might be more convenient than using the masks, especially when constructing large networks. This chapter describes how a network project is saved in CNA and which network-specific files are required.
- Mass-flow (stoichiometric) models can be imported and exported in SBML and METATOOL format (detailed in section 3.3 and chapter 6).
- One may use the CNA's Application Programming Interface (API) to construct a new network project (via MATLAB's command line). Existing (possibly partial) information of the network structure (such as the stoichiometric matrix) can be used in the API-based generation of a new project. This solution is described in detail in chapter 7 and is recommended if some parts of the network structure are available/can be loaded into in the MATLAB workspace.

Additionally, with dedicated API functions (see section 7.2.1), the reaction and species notes can be used to store generic data for reactions and species/metabolites in a structured way which can later be retrieved again.

This chapter describes how CNA network projects are stored on disk.

5.1 Declaring a network project

Project-specific files are stored in the associated project directory. All network projects and their associated directories are registered in the file "networks" in the CNA main directory. A new project can be manually registered or an existing one can be modified by editing this file. Each row in this file registers a network project as follows:

```
<Name of the network project>    <directory>    <network type>
```

The directory should be a sub-directory of the CNA main directory and the relative path instead of the absolute one should be given. Network type can be "1" (mass-flow) or "2" (signal-flow). An example, defining a project of a mass-flow network named "Stoichiometric Network Example" with project directory "SmallExample":

```
Stoichiometric;Network;Example SmallExample 1
```

Semicolons can be used to mark blanks, hence, the String "Stoich.;Network;Example" will be translated to "Stoich. Network Example".

5.2 Network project files

First, for both mass-flow and signal-flow networks, the project directory should contain the

graphics serving as CNA network maps (see chapter 1 and 2). If no network map is available you may use the provided dummy picture “dummy.pcx” located in the CNA main directory.

Several ASCII files stored in the project directory are used by CNA for representing mass-flow (MF) and signal flow (SF) networks. These files are in a special format interpretable by CNA. The first three files (*app_para.m*, *metabolites*, *reactions*) are required for both MF and SF:

app_para.m

This file contains general information on the network project, including text box colors, text box size, in which graphic files the network maps are stored etc. This file should always be created by using the Project Manager (e.g. when initializing and registering a new project).

reactions

This file contains the reactions. Each row defines a reaction and its properties (see section 3.1 and 4.1) as follows:

<identifier> <reaction equation> | <default value> <Par1> <Par2> <Par3> <x-Pos> <y-Pos> <map number> <text box type> <Par4>

The (symbolic) reaction (in SF: interaction) equation can be defined as described in section 3.1 and 4.1, respectively. All equations must be finished by the pipe symbol as indicated above. For all reactions, except the biomass synthesis reaction (whose identifier is “mue”), a reaction equation must be given.

<default value>: If the default value is unknown write “#” or “NaN”.

Par1: MF: rate minimum; SF: flag for incomplete truth table (0 = off, 1 = on)

Par2: MF: rate maximum; SF: time scale

Par3: MF: coefficient in linear objective function (for optimization problems); SF: flag for “Exclude in logical computations” (1: exclude; 0: include)

<x-Pos> and <y-Pos> are the pixel coordinates of the lower left corner of the text box associated with this reaction. <map number> is the network map where the box has to be appear. <text box type> is “1” for editable, “2” for non-editable and “3” for non-visible. The non-visible type is not allowed in mass-flow networks.

Par4: MF: Variance of measurements; SF: flag for monotony (0 = non-monotone)

Example: reaction R1::A in Fig. 1 (stoichiometric network):

R1::A = 1 A | # -Inf Inf 0 73.5 97.5 1 1 0.01

Example: biomass synthesis (“mue”) in Fig. 1 (stoichiometric network):

mue | # 0 10 0 394.5 448.5 1 1 0.01

Example: reaction 2&3 in Fig. 22 (signal-flow network):

2&3 !I1 + I2 = E | # 0 1 0 653 148 1 1 1

metabolites

Stores the metabolites (MF) or species (SF), respectively, including their attributes. Each row defines a metabolite/species with its parameters as described in section 3.1 and 4.1, respectively:

<identifier> <full name> <Par1> <Par2> <x-Pos> <y-Pos> <map number> <textbox type>

Par1: MF: not used; SF: default value (if no default value is known write “#” or “NaN”).

Par2: MF: External flag (is “0” if the metabolite is internal and “1” if external); SF: not used

<x-Pos>, <y-Pos>, <map number> and <text box type> define the position and type of the associated text box as explained for reactions. Note that this information is only relevant and used for SF (since only there species have an associated text box).

Example: species A in Fig. 22:

```
A    SpeciesA    NaN    0    269    155    1    1
```

The two files *reactions_notes* and *metabolites_notes* are optional for both MF and SF. They contain comments and notes for the reactions / species defined in the files *reactions* and *metabolites*, respectively. If the respective file does not exist, the comments are set to be empty. The set-up of both files is simple:

```
<identifier1> <Notes1>
<identifier2> <Notes2>
.....
```

<identifier> is a identifier of a species/reaction as defined in the *reactions* and *metabolite* files
 <Notes> is a string containing the notes/comments for the respective network element. Spaces are allowed in the String but line breaks are not. You may encode a line break by “;:;”. This string will be replaced when displaying the notes in *CellNetAnalyzer*. Note that the IDs need not be in the same order as in the *reactions/metabolites* files and not for all species and reactions a comment has to be included.

The following three files are additionally required (at least as empty files) for mass-flow networks:

macromolecules

Defines the biomass components including their attributes and parameters:

```
<identifier> <full name> <default value> <x-Pos> <y-Pos> <map number> <text box type>
```

<x-Pos>, <y-Pos>, <text box type> and <map number> describe the coordinates and style of the associated text box as for reactions. The (cumulative) synthesis equation for each biomass constituent is defined in file *macromolecule_synthesis* (see below).

Example: biomass constituent BC1 in Fig.1:

```
BC1    Biomass_Component_1    0.4    225.5    423.5    1    1
```

macromolecule_synthesis

Each row describes the cumulative synthesis equation for one biomass constituent (see section 3.1). It expresses the amount of metabolites (unit: [mmol Metabolite / g biomass component]) required for the synthesis of 1 gram of this biomass component. The equation can be formulated by using the metabolite identifiers, similar as for reaction equations:

```
<BiomassConst.> = <coeff1> <Met1> + <coeff2> <Met2> + ..... + <coeffx> <Metx>
```

Example: Synthesis equation for BC1 in Fig.1:

```
BC1 = 2 A + 1 C
```

assembly

Each row describes an assembly route and its properties (see section 3.1) as follows:

<Metabolite> <Biomass Constituent> <x-Pos> <y-Pos> <map number>

<x-Pos>, <y-Pos> and <map number> define the location of the associated text box as for reactions. <Metabolite> is the identifier of the metabolite consumed for the biomass constituent given in the second identifier.

Example: assembly of A into BC1

A	BC1	181.5	331.5	1
---	-----	-------	-------	---

6. Import/Export of models in SBML, METATOOL, and COBRA format

For stoichiometric (mass-flow / metabolic) networks, *CellNetAnalyzer* supports

- import and export of SBML models (Level 2 and 3, now also supporting the flux balance constraints (fbc) package 2 for level 3)
- export of the model in METATOOL format and
- import/export of COBRA models.

These conversions can be done either via the menu in the GUI (see section 3.3) or via dedicated API functions (see section 7.4). Note that several model features contained in imported SBML model files (e.g., compartment, charge and chemical formula of a metabolite; name and gene product association of a reaction) are stored in the generic data fields of CNA which can later be extracted and used for certain calculations (see section 7.2.1). The following fields will be saved in the generic data structures if the corresponding information is provided in the SBML model:

Reactions: reaction_id; reaction_name; geneProductAssociation

Species: constant; hasOnlySubstanceUnits; compartment; compartment:constant; fbc:charge; fbc:chemicalFormula

Within the GUI, structured generic data of mass-flow networks can also be edited by a dedicated editor for structured data (see section 3.2).

For signaling/regulatory (signal-flow) networks, CNA supports export and import of SBML qual models either via the GUI menu (see 4.3.1) or via a dedicated API function (see section 7.4). Note that only simple SBML qual models can be imported (for example, only AND, OR and NOT operators are allowed; only binary states (0 and 1) are allowed for the discrete state variables) into a CNA signal-flow network (SFN) project.

7. Application Programming Interface (API)

The API (Application Programming Interface) of CNA allows interested users and developers

- to read/write/edit or import/export the network structure of a project
- to read values from text boxes (GUIs), then to perform own calculations and finally to display the results in the interactive network maps
- to call selected functions of CNA (such as the computation of elementary modes or signaling paths) without the CNA GUI (some functions can only be called via API)
- to integrate user-developed functions as an entry in CNA's menu bar – you can thus construct and integrate plugins for CNA
- to change network/project attributes directly from MATLAB's command line (only for advanced users; not recommended for beginners)

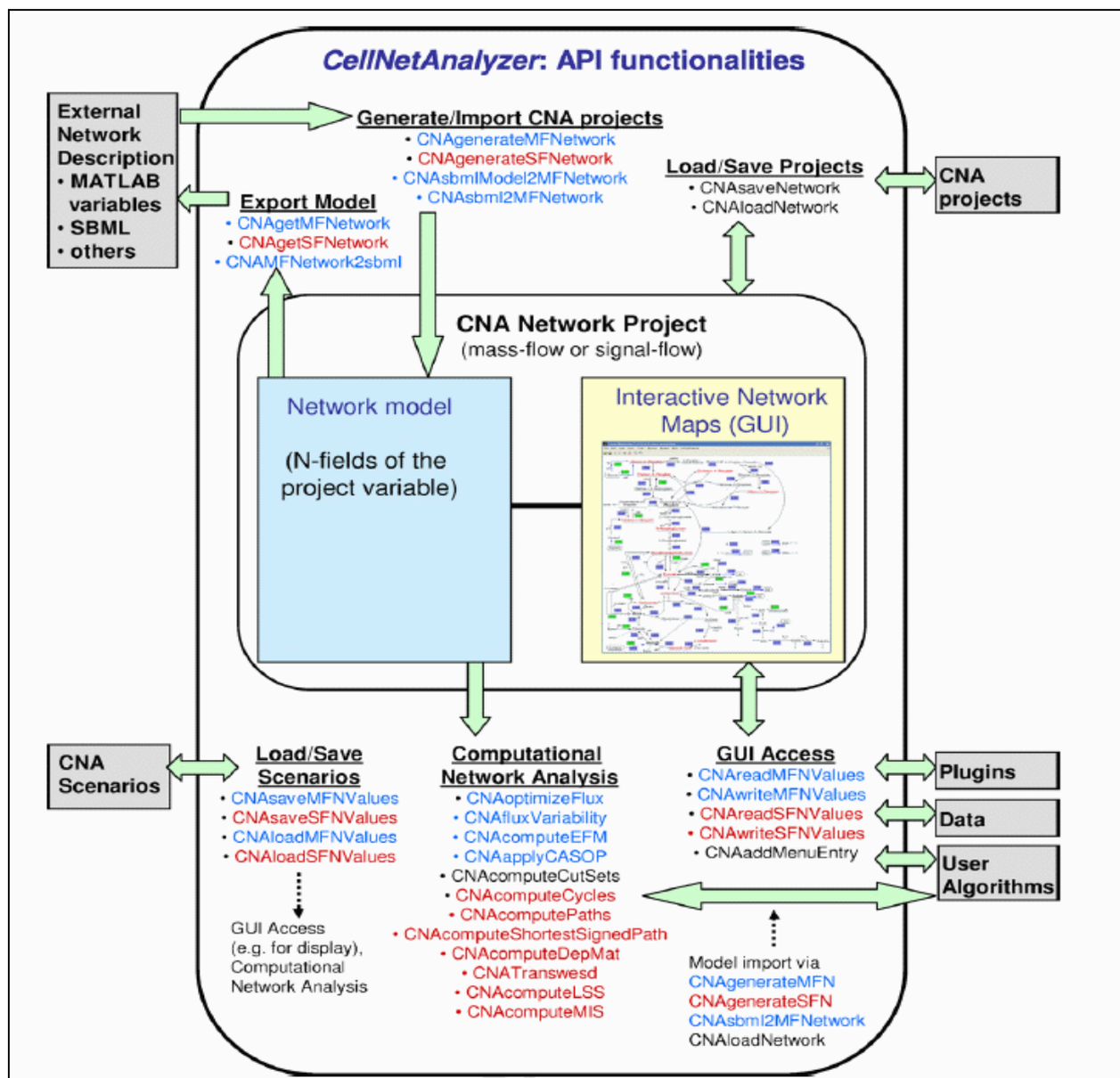


Fig. 24: Overview of (selected) API functions in CellNetAnalyzer.

CNA comprises several functions that have specific API functionalities (the name of these

functions starts with prefix ‘CNA’) which will be explained below. Furthermore, we will also document the internal variable structure of CNA, which should make interfacing CNA with other applications easier.

Before using any of the functionality presented in this chapter make sure that CNA is initialized. This can be done by calling the **startcna** function (if this function is not available go into the *CellNetAnalyzer* directory first). Calling **startcna** adds all relevant CNA directories to the MATLAB path, initializes the global variable **cnan** and launches the project manager window. You can suppress the project manager (e.g. if you want to operate in batch-mode or if you are using octave) by calling **startcna(1)** (see also below).

7.1. The internal variable structure of CellNetAnalyzer

In order to encapsulate the data belonging to one project and to allow working with multiple projects, (network) project variables were introduced in CNA version 9.0.

In CNA every network project is represented by a project variable which is a MATLAB structure (type struct) containing several fields for the different types of data associated with a project. Moreover, some of the fields are structs themselves to allow grouping of variables that belong together. Basically this data model follows the idea of object orientation as it was introduced in MATLAB in version 7. However, none of MATLAB's object orientation functionality is currently used and therefore features like inheritance do not work for CNA network variables.

As already mentioned, once loaded, a CNA network project is represented in the MATLAB workspace as a struct variable (which we will denote in the following with **cnap**). In general, the fields of a CNA project struct variable can be roughly classified into

- i) **N-fields** (= network fields): fields that define mathematically the network structure and its attributes (e.g. stoichiometric matrix).
- ii) **M-fields** (= meta fields): are fields that are additionally required to define a CNA network project (mainly GUI-related data such as text box size or text box color and other meta data such as the path of the project directory).
- iii) **L-fields** (= local fields): the field ‘**cnap.local**’ is itself a struct and its fields are temporary variables that are changed during actions or calculations. In particular, fields of ‘**cnap.local**’ are used to pass/return parameters to/from functions. For using the API functionality, users do not have to care about these fields.

Several fields are identical for mass-flow and signal-flow networks but there are also specific fields for each network type. In the following we document the N- and P- fields of mass-flow and signal-flow networks. Note that not all of the listed fields need to exist in a project variable when performing certain calculations. Furthermore, from version 2018.1 on, CNA also provides the option to store additional data sets (not covered by the standard data fields of a CNA project variable) in a generic way (see section 7.2.1 below).

Mass-flow networks

(we assume in the following that the network has n species, q reactions and m macromolecules; w is a placeholder for an appropriate non-zero number)

N-fields of mass-flow projects (compare with sections 3.1 and 5):

`cnap.type`: type of the network (`cnap.type = 1` for mass-flow and `cnap.type=2` for signal-flow)
`cnap.specID`: ($n \times w$) char array containing the identifiers of the species
`cnap.specLongName`: ($n \times w$) char array containing the long (full) names of the species
`cnap.specNotes`: ($1 \times n$) cell array containing the notes for the species
`cnap.specExternal`: ($1 \times n$) vector indicating which species are external (1) and which not (0)
`cnap.specInternal`: ($1 \times w$) vector with the indices of the internal species; it holds
that `cnap.specInternal=find(~cnap.specExternal);`
`cnap.numS`: number of species in the network (`cnap.numS=n`)
`cnap.numI`: number of internal species (`cnap.numI=size(cnap.specInternal,2)`)
`cnap.numR`: number of reactions in the network (`cnap.numR=q`)
`cnap.reacID`: ($q \times w$) char array containing the identifiers of the reactions
`cnap.reacNotes`: ($1 \times q$) cell array containing the notes for the reactions
`cnap.mue`: is the index of the 'mue' (biosynthesis) reaction; if 'mue' is not contained in the
`reacID` list, then `cnap.mue` is empty
`cnap.objFunc`: ($q \times 1$) vector containing the coefficients of the reactions in the linear objective
function (to be used for "Flux optimization")
`cnap.reacMin`: ($q \times 1$) vector containing the lower boundaries of the reaction rates (e.g. 0 for an
irreversible reactions or `-Inf` for irreversible reactions)
`cnap.reacMax`: ($q \times 1$) vector containing the upper boundaries of the reaction rates (e.g. `Inf` for
arbitrary large rates)
`cnap.reacDefault`: ($q \times 1$) vector containing the default (reaction rate) values to be displayed in
the reaction text boxes (`NaN` indicates an empty value)
`cnap.reacVariance`: ($q \times 1$) vector containing the variances of (measured) reaction rates (used for
some metabolic flux analyses; see section 3.3)
`cnap.stoichMat`: the ($n \times q$) stoichiometric matrix of the network
`cnap.macroID`: ($m \times w$) char array containing the identifiers of the macromolecules
`cnap.macroLongName`: ($m \times w$) char array containing the full names of the macromolecules
`cnap.macroDefault`: ($m \times 1$) vector containing the default values of the macromolecules (to be
displayed in the macromolecule text boxes; `NaN` is not allowed here)
`cnap.numMac`: number of macromolecules (`cnap.numMac=m`)
`cnap.macroComposition`: ($m \times w$) matrix defining the stoichiometry of the macromolecules with
respect to the metabolites (species): matrix element `macroComposition(i,j)` stores how much
of metabolite *i* (in mmol) is required to synthesize 1 gram of macromolecule *j*

M-fields of mass-flow projects (compare with section 2, 3.1 and 5):

`cnap.has_gui`: flag that indicates whether this project variable is associated with an active GUI
`cnap.netnum`: index of the network in the project manager.
`cnap.path`: gives the (relative or absolute) path of the project directory (in which the project files
are located)
`cnap.net_var_name`: the name of the network project (identical with the name of the project
variable; here denoted with the placeholder 'cnap:') This field contains thus the name of the
variable as it is called in the base workspace. When loading a network a name for the network
variable is automatically generated based on the directory name of the network folder. It is
essential that `net_var_name` matches the name of the workspace network variable name for
the proper operation of the graphical user interface (see technical aspects). This has to be
taken into account if you are running a project with a GUI and want to modify its data from
the command prompt. Neither the variable name nor the string in `cnap.net_var_name` must be

changed while the network is used because this would result in errors when using the GUI elements.

`cnap.unsaved_changes`: indicates whether there are unsaved changes in the network structure (1) or not (0) (made by using the network composer)

`cnap.epsilon`: CNA epsilon (smallest number greater than zero) to be used in CNA calculations

`cnap.reacBoxes`: a ($q \times 6$) array defining the position and appearance of the reaction text boxes; `cnap.reacBoxes(:,1)` not in use currently; `cnap.reacBoxes(:,2)` defines the x-position of each box; `cnap.reacBoxes(:,3)` defines the y-position, `cnap.reacBoxes(:,4)` contains the MATLAB GUI handles of the text boxes, `cnap.reacBoxes(:,5)` contains the map numbers (where the boxes are to be displayed) and `cnap.reacBoxes(:,6)` defines the style of each text box (1=editable; 2=non-editable)

`cnap.macroBoxes`: a ($m \times 6$) array defining the position and appearance of the macromolecule text boxes; the columns of this array are analogous as for `cnap.reacBoxes`

`cnap.macroSynthBoxes`: a ($w \times 6$) array defining the assembly routes including the parameters of their text boxes (w = number of defined assembly routes): `cnap.macroSynthBoxes(:,1)` the index of the metabolite involved in this assembly route, `cnap.macroSynthBoxes(:,2)` the index of the macromolecule in whose assembly the metabolite is involved; `cnap.macroSynthBoxes(:,3)` defines the x-position of the text boxes of the assembly routes; `cnap.macroSynthBoxes(:,4)` defines the y-position of the text boxes; `cnap.macroSynthBoxes(:,5)` contains the MATLAB GUI handles of the text boxes; `cnap.macroSynthBoxes(:,6)` stores the map numbers (where the boxes are to be displayed)

`cnap.nummacsynth`: number of assembly routes (= `size(cnap.macroSynthBoxes,1)`)

`cnap.maps`: is a ($w \times 2$) cell array of strings (w = number of interactive maps in the project). `cnap.maps{i,1}` is the name (title) of the i -th map; `cnap.maps{i,2}` is the graphic file (optionally with path) of the i -th map.

`cnap.nummaps`: is the number of interactive maps in this project (`cnap.nummaps=size(cnap.maps,1)`)

`cnap.figs`: is a ($w \times 9$) array (w = number of interactive maps) defining some figure properties: `cnap.figs(:,1)` stores the handles of the figures; `cnap.figs(:,2:3)` stores the resolution of the figure; `cnap.figs(:,4)` indicates whether the resource (graphic file) of the figure is a pixel-based graphic (0) or a MATLAB figure (1); `cnap.figs(:,5:9)` stores (temporarily) some parameters required for installing zoom tools on the maps

`cnap.color1`: (1×3) vector with the RGB values defining reaction text box color 'standard'

`cnap.color2`: (1×3) vector with the RGB values defining reaction text box color 'calculated'

`cnap.color3`: (1×3) vector with the RGB values defining reaction text box color 'special'

`cnap.color4`: (1×3) vector with the RGB values defining reaction text box color 'defined'

`cnap.specBoxColor`: (1×3) vector with the RGB values of the color of macromolecule text boxes

`cnap.textColor`: (1×3) vector with the RGB values defining the text color in reaction and macromolecule text boxes

`cnap.macroSynthColor`: (1×3) vector with the RGB values defining the text color in assembly text boxes

`cnap.reacBoxWidth`: ($1 \times w$) vector (w = number of interactive maps) defining the width of reaction text boxes (separately for each map)

`cnap.reacBoxHeight`: ($1 \times w$) vector (w = number of interactive maps) defining the height of reaction text boxes (separately for each map)

`cnap.specBoxWidth`: ($1 \times w$) vector (w = number of interactive maps) defining the width of macromolecule text boxes (separately for each map)

`cnap.specBoxHeight`: ($1 \times w$) vector (w = number of interactive maps) defining the height of

macromolecule text boxes (separately for each map)
 cnap.reacFontSize: (1 × w) vector (w = number of interactive maps) defining the font size of the text in the reaction text boxes (separately for each map)
 cnap.specFontSize: (1 × w) vector (w = number of interactive maps) defining the font size of the text in the macromolecule text boxes (separately for each map)
 cnap.has_gui: flag that indicates whether this project variable is associated with an active GUI
 cnap.pan_icon_handles (only version 7.3 and higher): stores GUI handles
 cnap.figmenu: stores GUI handles

Signal-flow networks

(we assume in the following that the network has n species, q reactions/interactions; w is a placeholder for an appropriate non-zero number)

N-fields of signal-flow projects (compare with sections 4.1 and 5):

cnap.type: type of the network (cnap.type = 2 for signal-flow)
 cnap.numS: number of species in the network (cnap.numS=n)
 cnap.numR: number of reactions in the network (cnap.numR=q)
 cnap.specID: (n × w) char array containing the identifiers of the species
 cnap.specLongName: (n × w) char array containing the long (full) names of the species
 cnap.specNotes: (1 × n) cell array containing the notes for the species
 cnap.specDefault: (n × 1) vector containing the default values to be displayed in the species text boxes (NaN indicates an “empty”)
 cnap.reacID: (q × w) char array containing the identifiers of the interactions
 cnap.reacNotes: (1 × q) cell array containing the notes for the interactions
 cnap.reacDefault: (q × 1) vector containing the default values to be displayed in the interaction text boxes (NaN indicates an empty value)
 cnap.monotony: (1 × q) vector containing the monotony flag for the interactions (0 if not monotone; non-zero if monotone)
 cnap.excludeInLogical: (1 × q) vector containing the “exclude in logical calculations” flag of the interactions (1 if to be excluded; 0 otherwise)
 cnap.incTruthTable: (1 × q) vector containing the “incomplete truth table” flag of the interactions (1 for incomplete truth table, 0 otherwise)
 cnap.timeScale: (q × 1) vector containing the time scales of the interactions
 cnap.interMat: (n × q) interaction matrix of the network (rows: species; columns: interactions); defines together with cnap.notMat the logic of each interaction (each interaction is an AND clause; see also section 4.1 and reference [11])
 cnap.notMat: (n × q) matrix (rows: species; columns: interactions); stores the NOTs appearing in the interactions; if species i participates with its inverted value (by a NOT operation) in the j-th interaction then cnap.notMat(i,j) = 0 (otherwise 1)

M-fields of signal-flow projects (compare with section 2, 4.1 and 5):

cnap.has_gui: flag that indicates whether this project variable is associated with an active GUI
 cnap.netnum: index of the network in the project manager.
 cnap.path: gives the (relative or absolute) path of the project directory (in which the project files

are located

- `cnap.net_var_name`: the name of the network project (identical with the name of the project variable; here denoted with the placeholder 'cnap:') This field contains thus the name of the variable as it is called in the base workspace. When loading a network a name for the network variable is automatically generated based on the directory name of the network folder. It is essential that `net_var_name` matches the name of the workspace network variable name for the proper operation of the graphical user interface (see technical aspects). This has to be taken into account if you are running a project with a GUI and want to modify its data from the command prompt.
- `cnap.unsigned_changes`: indicates whether there are unsigned changes in the network structure (1) or not (0) (made by using the network composer)
- `cnap.epsilon`: CNA epsilon (smallest number greater than zero) to be used in CNA calculations
- `cnap.reacBoxes`: a ($q \times 6$) array defining the position and appearance of the reaction text boxes; `cnap.reacBoxes(:,1)` not in use currently; `cnap.reacBoxes(:,2)` defines the x-position of each box; `cnap.reacBoxes(:,3)` defines the y-position, `cnap.reacBoxes(:,4)` contains the MATLAB GUI handles of the text boxes, `cnap.reacBoxes(:,5)` contains the map numbers (where the boxes are to be displayed) and `cnap.reacBoxes(:,6)` defines the style of each text box (1=edit-able; 2=non-editable)
- `cnap.specBoxes`: a ($n \times 6$) array defining the position and appearance of the species text boxes; the columns of this array are analogous as for `cnap.reacBoxes`
- `cnap.maps`: is a ($w \times 2$) cell array of strings (w = number of interactive maps in the project). `cnap.maps{i,1}` is the name (title) of the i-th map; `cnap.maps{i,2}` is the graphic file (optionally with path) of the i-th map.
- `cnap.nummaps`: is the number of interactive maps in this project (`cnap.nummaps=size(cnap.maps,1)`)
- `cnap.figs`: is a ($w \times 9$) array (w = number of interactive maps) defining some figure properties: `cnap.figs(:,1)` stores the handles of the figures; `cnap.figs(:,2:3)` stores the resolution of the figure; `cnap.figs(:,4)` indicates whether the resource (graphic file) of the figure is a pixel-based graphic (0) or a MATLAB figure (1); `cnap.figs(:,5:9)` stores (temporarily) some parameters required for installing zoom tools on the maps
- `cnap.color1`: (1×3) vector with the RGB values defining text box color 'standard'
- `cnap.color2`: (1×3) vector with the RGB values defining text box color 'calculated'
- `cnap.color3`: (1×3) vector with the RGB values defining text box color 'special'
- `cnap.color4`: (1×3) vector with the RGB values defining text box color 'defined'
- `cnap.specBoxColor`: (1×3) vector with the RGB values defining the color of species text boxes
- `cnap.textColor`: (1×3) vector containing the RGB values of the text color to be used for both species and reaction text boxes
- `cnap.macroSynthColor`: (1×3) vector: (currently unused in signal-flow networks)
- `cnap.reacBoxWidth`: ($1 \times w$) vector (w = number of interactive maps) defining the width of reaction text boxes (separately for each map)
- `cnap.reacBoxHeight`: ($1 \times w$) vector (w = number of interactive maps) defining the height of reaction text boxes (separately for each map)
- `cnap.specBoxWidth`: ($1 \times w$) vector (w = number of interactive maps) defining the width of species text boxes (separately for each map)
- `cnap.specBoxHeight`: ($1 \times w$) vector (w = number of interactive maps) defining the height of species text boxes (separately for each map)
- `cnap.reacFontSize`: ($1 \times w$) vector (w = number of interactive maps) defining the font size of the text in the reaction text boxes (separately for each map)
- `cnap.specFontSize`: ($1 \times w$) vector (w = number of interactive maps) defining the font size of the

text in the species text boxes (separately for each map)
cnap.pan_icon_handles (only version 7.3 and higher): stores GUI handles
cnap.figmenu: stores GUI handles

Knowing the structure of the CNA project variable, you may easily export selected data from a project (e.g. the stoichiometric Matrix). You may also manually change some entries, however, it is then up to you to ensure that the network remains consistent. For example, if you add a reaction (manually) to the stoichiometric matrix, you also have to update all fields that contain reaction specific information (such as reacID, reacMin, reacMax etc.).

In the following, the API functions of CNA are briefly described (in a few cases there are extended descriptions). A detailed description of each function and its parameters can be obtained by entering “help <funcname>” in MATLAB (CNA must have been started).

7.2. API functions for creating, copying and editing network projects

CNAgenerateMFNetwork

Assume that you have a stoichiometric reaction network that you would like to analyze with CNA (or some CNA functionalities). Usually, you will have the stoichiometric matrix and the reversibilities of the reactions. However, a CNA mass flow project requires many more variables (fields) to be defined. Here, it would be desirable to have a function, which generates a CNA mass flow project from such partial information. This is exactly one purpose of the API function

Usage: `[cnap, errval] = CNAgenerateMFNetwork(cnap,nodisp)`

The project variable ‘cnap’ is usually a structure that contains some (N-)fields of a CNA mass-flow project whose existing fields are checked for consistency and to which missing fields are added and initialized with default values. Display of detailed messages can be repressed (nodisp=1; default: 0). The output errval indicates whether consistency errors have been found (errval=1) or not (errval=0).

If cnap has no N-field name (or no field cnap.stoichMat) then **CNAgenerateMFNetwork** will generate and initialize an empty CNA mass-flow project structure. If cnap has already some N-fields of a CNA mass-flow project, all missing N-fields will be generated and filled with default values and then returned. Additionally, the M-fields reacBoxes, macroBoxes, nummacsynth, macroSynthBoxes and epsilon will be appended with default values as well. For example, the text box position of reactions (cnap.reacBoxes) and macromolecules (cnap.macroBoxes) will be set to the upper left corner. These M-fields are required to save the project using **CNAsaveNetwork** (see below). Upon saving the network, the project can later be registered via the project manager where also all other required M-fields (text box size and style etc.) can then be defined.

Even though the returned project variable ‘cnap’ lacks most M-fields, it can already be used for calling several CNA functions (see below) which do not access GUI elements.

Importantly, if the passed argument cnap contains already several N-fields (e.g. cnap.stoichMat, cnap.numms, cnap.numr) this function will check the consistency of the predefined fields and return a nonzero errval if an inconsistency is found. The procedure will also be stopped at this point, so the creation of the network variable will then be incomplete.

Here is an example: assume you have (only) the stoichiometric matrix of a network in the MATLAB workspace. Let's call this matrix N. Create now a new structure z having the (only) field "stoichMat":

```
z.stoichMat=N;
```

Entering now

```
[z, errval] = CNAgenerateMFNetwork(z);
```

will create a CNA mass-flow project structure in the variable ‘z’ containing all N-fields and the few M-fields mentioned above. Non-existing fields are initialized with default values, for example z.reacMin, z.objFunc will be zero vectors, reacMax an Infinity vector. With this variable you can already call functions that do not access the

GUI (see below). You may then save the generated network as a CNA project with **CNAsaveNetwork**(see below)

```
z.path= 'path/to/project/directory';  
z=CNAsaveNetwork(z);
```

Note that it is necessary to specify the save directory in z.path before calling **CNAsaveNetwork**.

If you then register the project in the project manager you will automatically set the rest of the M-fields of this project (including a network map, if you want). You can then reload the project later either with the "Load w/o GUI" button or (if you provided a network map) as usual with 'Start'.

Note that in the above example **CNAgenerateMFNetwork** will first check whether z has a field "stoichMat". If not, an empty field stoichMat will be added to z and all other required fields will be initialized accordingly. As mentioned, the correct dimensions of existing fields will be checked. For example, if your input variable z has the two fields "stoichMat" and "reacMin" and if `size(cnap.stoichMat,2)~=size(cnap.reacMin,1)` you will get an error message and (errval will be 1). The function will then stop and the project variable remains incomplete.

Another Example

Here is another, more explicit example how to use **CNAgenerateMFNetwork** to create a new metabolic network model. If not done yet, start CNA by calling **startcna** (we here do that without launching the project manager; see section 7.8):

```
startcna(1);
```

Consider the following reaction scheme:

```
R1: A = 2 B + C
```

```
R2: B + D = E
```

```
R3: C + E = 3 F
```

which gives the stoichiometric matrix (rows: metabolites, columns: reactions):

```
simple_net.stoichMat= [-1 0 0; 2 -1 0; 1 0 -1; 0 -1 0; 0 1 -1; 0 0 3];
```

The metabolites B, C, E are considered internal (and thus A, D and F external):

```
simple_net.specExternal= [1 0 0 1 0 1];
```

If all reactions are irreversible and you don't want to assign your own metabolite or reaction names you can already call

```
simple_net= CNAgenerateMFNetwork(simple_net);
```

to set up the project variable. If you want to assign e.g. metabolite names you can use

```
simple_net.specID= strvcat('A', 'B', 'C', 'D', 'E', 'F');
```

to do so (assigning reacID works analogously). Reaction reversibility can be controlled with the reacMin and reacMax fields. Typically all reactions have +Inf as reacMax and the distinction between reversible and irreversible reactions is made by setting reacMin to -Inf or 0 respectively. In order to make reactions two and three reversible you can e.g. execute

```
simple_net.reacMin= zeros(size(simple_net.stoichMat, 2), 1);
```

```
simple_net.reacMin([2 3])= -Inf;
```

You can now call

```
simple_net= CNAgenerateMFNetwork(simple_net);
```

which sets reacMax to +Inf for all reactions and configures the remaining fields of the project variable.

To save the network create a new directory (here called simple_path like the project variable), set up the path field to point to it and lastly call **CNAsaveNetwork** (as explained in section 7.3):

```
simple_net.path= './simple_net';
```

```
simple_net= CNAsaveNetwork(simple_net);
```

You can later register the network in the project manager (cf. chapter 2 and 7.3 below) and now continue working with the project through the API functions described below. When you are finished, you can call

```
close_cna;
```

which closes all projects running with a GUI, removes the CNA directories from the MATLAB path and deletes the **cna** variable (section 7.9).

CNAgenerateSFNetwork

Analogous as for mass-flow projects, there is also a function

Usage: `[cnap, errval] = CNAgenerateSFNetwork(cnap)`

for creating a signal-flow project (variable) from partial information. Again, the fields in 'cnap' will be checked for consistency and all missing N-fields and the M-fields reacBoxes, specBoxes and epsilon will be added. Here,

CNAgenerateSFNetwork starts with looking whether the field 'cnap.interMat' exists. If not, an empty project will be created, otherwise, all other signal-flow N-fields will be added (if they do not exist yet) or checked for consistency (if already present in cnap).

An example: assume you have (only) the interaction matrix and the NOT matrix of a Boolean network somewhere in the MATLAB workspace (for the definition of the interaction and NOT matrix see reference [11]). Let's call these matrices I and N, respectively. Create now a new structure z having the fields interMat and notMat:

```
z.interMat=I;  
z.notMat=N;
```

Entering now

```
[z,errval] = CNAgenerateSFNetwork(z);
```

will create a CNA signal-flow project variable 'z' that has all N-fields (defining the network topology) of a signal-flow network and the M-fields reacBoxes, specBoxes and epsilon. Non-existing fields are initialized with default values. With this variable you can already call functions that do not access the GUI (see below). You may save the generated network project as a CNA SFN project via **CNASaveNetwork** (see below):

```
z.path= 'path/to/project/directory';  
znew=CNASaveNetwork(znew);
```

Note that it is necessary to specify the save directory in z.path before calling **CNASaveNetwork**.

If you then register then the project in the project manager (or by editing the file "networks") you will automatically set the rest of the M-fields of this project (including a network map, if you want). You can then reload the project later either with the "Load w/o GUI" button or (if you provided a network map) as usual with 'Start'.

Again, the correct dimensions of existing fields will be checked. For example, if $\text{any}(\text{size}(z.\text{interMat}) \sim \text{size}(z.\text{notMat}))$ you will get an error message and errval will be 1.

The following two functions facilitate to export/copy a CNA network project into a new struct variable. Only the (N-) fields defining the network structure are copied whereas all GUI-related fields are not transferred.

CNAgetMFNetwork

Usage: `mfn = CNAgetMFNetwork(cnap,c_macro);`

cnap is a CNA mass-flow project structure and c_macro is a (cnap.nummac,1) vector defining the biomass composition. If macromolecules are not defined for this project set c_macro=[].

All those fields of the project cnap will be copied into the output mfn that define the network topology and related parameters. GUI-related (P-) and temporary (L-) fields are not copied. Hence, mfn will contain all N-fields of cnap, except epsilon, has_gui, reacBoxes, macroBoxes, macroSynthBoxes and nummacsynth.

If the project cnap has macromolecules (and uses thus CNA's biomass synthesis reaction 'mue'), the vector biocomp will be used together with cnap.macroComposition to calculate the stoichiometry of reaction 'mue' which is then inserted into the column with index cnap.mue.

This function provides thus a convenient way to copy/export the network structure of a CNA mass-flow project into a new variable, thereby getting rid of GUI and temporary variables of the project. The new struct variable can then be used independently of CNA.

CNAgetSFNetwork

Usage: `sfn = CNAgetSFNetwork(cnap);`

cnap is a CNA signal-flow project structure. All those fields of the project cnap will be copied into the output sfn that define the network topology and related parameters. GUI-related (P-) and temporary (L-) fields are not copied. Hence, sfn will contain all N-fields of cnap, except epsilon, has_gui, reacBoxes and specBoxes.

This function provides thus a convenient way to copy/export the network structure of a CNA signal-flow project into a new variable, thereby getting rid of GUI and temporary variables of the project. The new struct variable can then be used independently of CNA.

CNAdeleteReaction

Usage: [cnap] = **CNAdelateReaction**(cnap,delreac)

This function deletes all the reactions specified in the 'delreac' index vector from the project 'cnap' and returns the modified project 'cnap'. All related parameters of the deleted reactions (including text boxes (if with GUI)) will be removed from the project.

CNAdelateSpecies

Usage: [cnap] = **CNAdelateSpecies**(cnap,delspec)

This function deletes all the species (metabolites) specified in the 'delspec' index vector from the project 'cnap' and returns the modified project 'cnap'. All related parameters of the deleted species (including GUI elements) will be removed from the project.

CNAaddReactionMFN

Adds a reaction to a CNA mass-flow project.

Usage: cnap = **CNAaddReactionMFN**(cnap, reac, equation, lb, ub, objCoeff, ...
defaultRate, measVar, notes, flxMapNo, editable, xpos, ypos)

CNAaddSpeciesMFN

Adds a reaction to a CNA mass-flow project.

Usage: cnap = **CNAaddSpeciesMFN**(cnap, spec, spec_ext, spec_name, spec_notes)

CNAgenerateGPRrules

Generates a struct array that contains the information on gene-enzyme-reaction-associations. Gene-information can either be provided in the cnap-structure as the value of the field 'geneProductAssociation' (see CNAsetGenericReactionData), or be provided separately as a cell array. Syntax follows Cobra's manner ('and', 'or', curved brackets)

Usage: [cnap, enzymes, genes, gpr_rules] = **CNAgenerateGPRrules**(cnap, grRules, bool_enz_only)

CNAintegrateGPRrules

Extends a given mass-flow network with pseudo-reactions and pseudo-metabolites for genes and enzymes which can later be used, for example, to compute gene cut sets (this function is also used in CNAgeneMCSEnumerator2).

Usage: [gcnap, rmap, gmap, gkoCost, gkiCost, gT, gD, rType, sType, gpr_rules] = ...
CNAintegrateGPRrules(cnap, gpr_rules, koCost, kiCost, T, D, gkoCost, gkiCost, expand_notknockable)

CNAgenerateMaps

Creates maps for a given CNA mass-flow project listing all reaction names of the project (and positioning the reaction text boxes behind the respective reaction name). These maps can be used as a surrogate for network graphics for a GUI-based analysis of the project.

Usage: [cnap,err] = **CNAgenerateMap**(cnap,show_eq)

7.2.1. API functions for managing generic data in CNA projects

CNA provides API functions to use the reaction and species notes to store generic data for reactions and species/metabolites in a structured way. Thus, any parameter (e.g., ΔG_R^0 of a reaction or chemical formula of a metabolite) or textual information (e.g., KEGGID), which are not covered by CNA's default parameters, can be stored under user-specified field names. In particular, certain species/reaction information obtained when importing an SBML file are stored in the generic data (see section 6). The generic data are serialized in the reactions or species notes (cnap.reacNotes/cnap.specNotes) and can later be retrieved for further processing.

The functions to save data in the reaction notes are `CNAsetGenericReactionData()` or `CNAsetGenericReactionData_with_array()`. To save data in the species notes the functions are `CNAsetGenericSpeciesData()` or `CNAsetGenericSpeciesData_with_array()` and can be used analogously. You can save numeric and textual data under a user-defined keyword.

CNAsetGenericReactionData

This function saves a textual or numeric value for a reaction under a specified keyword in the reaction notes of a CNA project. If data already exist with the specified keyword, then it will be overwritten. If no reaction exists with the given index then the function returns a nonzero errval.

```
[cnap, errval]=CNAsetGenericReactionData(cnap,idx,key1,value1,key2,value2, ...)
```

For example to save the KEGG id of reaction number 5 under the keyword 'kegg_id' use the function as follows:

```
[cnap, errval] = CNAsetGenericReactionData(cnap,5,'kegg_id','R12345')
```

CNAsetGenericReactionData_with_array

Sometimes it is more convenient to store an array of data for all reactions in one go using the following function:

```
[cnap, errval] = CNAsetGenericReactionData_with_array(cnap,key,values)
```

For example to save all the KEGG ids of your reactions under the keyword 'kegg_id' use the function as follows:

```
[cnap,errval] = CNAsetGenericReactionData_with_array(cnap,...  
                                                    'kegg_id',{ 'R12345', 'R12346', ...})
```

Again, already existing data with this key will be overwritten. If the key is no valid field name or no the length of the values array differs from the number of reactions in the cna project then the function returns a nonzero errval.

CNAsetGenericSpeciesData ... analogous to `CNAsetReactionData`

CNAsetGenericSpeciesData_with_array ... analogous to `CNAsetSpeciesData_with_array`

Once generic data have been saved in a project, they can later be accessed again. The functions to retrieve the data from the reaction notes are `CNAgetGenericReactionData()` or `CNAgetGenericReactionData_as_array()`. To retrieve the data in the species notes the functions are `CNAgetGenericSpeciesData()` or `CNAgetGenericSpeciesData_as_array()` and can be used analogously.

CNAgetGenericReactionData

This function retrieves a value saved under a specified keyword in the reaction notes of a reaction in a CNA project.

```
Usage: [val1,val2,...,errval] =  
       CNAgetGenericReactionData(cnap,idx,key1,key2,...);
```

If no data is stored under the given key(s) the function returns a nonzero errval.

For example, to read the 'geneProductAssociation' for the reaction with index 5:

```
[data, errval] = CNAgetGenericReactionData(coli,5,'geneProductAssociation')
```

CNAgetGenericReactionData_as_array

To retrieve the data stored under a given keyword for all reactions at once use the following function:

```
[data, errval] = CNAgetGenericReactionData_as_array(cnap,key):
```

If no data is stored under the given key the array is filled with a default value (NaN for numeric data and the empty string for textual data). For example to read all the formula of the reactions that are stored under the keyword 'formula' use the function as follows:

```
[data,errval]= CNAgetGenericReactionData_as_array(coli,'formula')
```

CNAgetGenericSpeciesData ... analogous to CNAgetGenericReactionData

CNAgetGenericSpeciesData_as_array .. analogous to CNAgetGenericReactionData_as_array

There are two functions to receive all keywords that are used in the generic data of the reactions or species:

```
[keywords, err] = CNAgetKeywordsGenericReactionData(cnap)
```

```
[keywords, err] = CNAgetKeywordsGenericSpeciesData(cnap)
```

7.3. Saving and loading CNA projects

CNA saves a network project via several ASCII files (section 5.2). The following functions can be used to save and load network projects to/from these project files.

CNAsaveNetwork

```
Usage: cnap = CNAsaveNetwork(cnap);
```

This function stores the network structure of a mass-flow project variable cnap. It creates/overwrites all the files of a CNA network project (see section 5.2.). cnap must have all N-fields of a CNA mass-flow project and additionally the M-fields cnap.reacBoxes, cnap.macroBoxes, cnap.nummacsynth and cnap.macroSynthBoxes. For example, a CNA project variable created with **CNAgenerateMFNetwork** or **CNAgenerateSFNetwork** can directly be saved with this function. Note that the cnap.path field must be set to an appropriate, absolute path where the project files will be stored. Existing project files will be overwritten! If the directory does not exist then it will be created. The function returns the mass-flow project.

Once you have saved a network project, you can register it in the CNA network list as follows (cf. end of section 2):

- 1) Click on "New" in the "Network Project Manager".
- 2) Enter the (preferably relative) path to the project directory where the project files have been stored. Define the name of the project. Additionally, you may declare source files for the network maps – if none is available choose the “dummy” network map as inserted by default.
- 3) Click on "Save" and answer the next question with "Yes".
- 4) The project now appears at the bottom of the project list in the project manager.

A saved network project can also be reloaded via **CNAloadNetwork** described below. However, unless the network has been registered in the CNA network list it can only be loaded without GUI (see *no_gui* option below). The reason for this is that registration creates an `app_para.m` file, which contains GUI configuration options, in the project directory. If a suitable `app_para.m` file (e.g. taken from some other directory) is added to a project directory, then its project can also be loaded with GUI without having been registered in the project manager.

CNAloadNetwork

Usage: `result = CNAloadNetwork(network, no_gui, return_cnap)`

This function loads a CNA (mass-flow or signal-flow) network project with or without GUI.

7.4 Import and Export of SBML, COBRA, and METATOOL Models

The following API functions serve for import/export of CNA mass flow projects from/to SBML, COBRA toolbox, and METATOOL. In addition, import/export of signal flow networks from/to SBML is possible. Import functions create a CNA mass flow project which can afterwards be saved via **CNAsaveNetwork** and then, optionally, be registered in the project manager's list of networks (see above). We start with import / export of METATOOL models.

CNAmetatool2MFNetwork

Usage: `[cnap, errval] = CNAmetatool2MFNetwork(fname)`

For importing stoichiometric networks defined in METATOOL format. This function takes the name of the input file as argument and uses a parser for METATOOL input files together with the **CNAgenerateMFNetwork** function (see above) to create a new project variable which is then returned by the function. The new project variable can then be used with the CNA API functions or saved as a CNA project (via **CNAsaveNetwork**) and finally be registered as a new project (as described in section 7.3).

CNAMFNetwork2metatool

Usage: `err = CNAMFNetwork2metatool(cnap, fname, macromol)`

Exports a CNA mass-flow project to a METATOOL-compatible file.

The following two methods allow import and export of COBRA toolbox models:

CNAcobra2cna

This function creates a CNA project from a COBRA model.

Usage: `cnap = CNAcobra2cna(cbmodel)`

Basically, the CNA mass flow project fields *specID*, *specLongName*, *stoichMat*, *reacMin*, *reacMax*, *reacID* and *objFunc* are filled from their counterparts in the COBRA model.

CNAcna2cobra

This function creates a COBRA model from a CNA mass flow.

Usage: `cbmodel = CNAcna2cobra(cnap)`

Basically, the COBRA model fields *description*, *rxns*, *metS*, *S*, *lb*, *ub*, *c*, *osense*, *rxnNames*, *metNames* are filled from their counterparts in the CNA model.

CNASbmlModel2MFNetwork

Usage: `[cnap, errval] = CNASbmlModel2MFNetwork(filename, ext_comparts)`

This function uses the *TranslateSBML* function of the libSBML MATLAB interface (see: <http://sbml.org/Software/SBMLToolbox>) to convert an SBML file into a CNA mass-flow network (MFN) project. The required libSBML files for several operating systems are distributed with *CellNetAnalyzer*. The fbc (flux balance constraints) package 2 for SBML level 3 is supported. Note that several model features often contained in imported SBML model files (e.g., compartment, charge and chemical formula of a metabolite; name and gene product association of a reaction) are stored in the generic data fields of CNA which can later be extracted and used for certain calculations (see also section 6 and 7.2.1).

CNASbmlModel2SFNetwork

Usage: `[cnap, errval] = CNASbmlModel2SFNetwork(filename)`

Converts simple qualitative SBML model (for example, only AND, OR and NOT operators are allowed; only 0 and 1 are allowed for the discrete state variables) into a CNA signal-flow network (SFN) project. Uses the *TranslateSBML* function of the libSBML Matlab interface for parsing an SBML (xml) file. Requires MATLAB >= 2014.b and python installed.

Two functions allow export of CNA models:

CNAMFNetwork2sbml

Usage: `err = CNAMFNetwork2sbml(cnap,fname,macromol,fbc)`

Exports CNA mass-flow project to an SBML file (with path specified in *fname*). Note that the model will be exported in SBML Level 3 version 1 (w/o fbc package (fbc = flux balance constraints)).

CNASFNetwork2sbmlQual

Usage: **CNASFNetwork2sbmlQual**(cnap, fname)

Exports CNA signal-flow project to an SBML file complying with the ‘SBML qual’ specifications.

The logical model is interpreted as a binary model and treats ITT as AND gates; half-edges are ignored. All interactions affecting one node are collected into one transition to fulfill the coherence recommendation of the SBML qual standard.

7.5 Accessing the GUI of CNA

Most API functions of CNA do not access (and do not explicitly need) the GUI of CNA, i.e. they access only N-fields of a CNA project variable. The API functions described in this section, allow you to access the GUI (i.e. the interactive maps and text boxes) of a network project loaded with GUI. CNA itself uses internally different functions for doing the same, however, the API functions described in this section simplify this procedure for the user.

The following four functions allow you to read/write numerical values from/into the text boxes of a network project.

CNAreadMFNValues

Usage: `[reacval, macroval] = CNAreadMFNValues(cnap);`

Given a mass-flow project that has been loaded with GUI (i.e. all N- and M-fields must exist in cnap), this function returns the numerical values currently set in the text boxes of reactions ($q \times 1$ vector reacval) and macromolecules ($m \times 1$ vector macroval), respectively. If the text box of the i -th reaction does not contain a numerical value, then reacval(i)=NaN. Nothing is changed in the project variable cnap; it is therefore not returned.

CNAwriteMFNValues

Usage: `cnap = CNAwriteMFNValues(cnap,reacval,reacol,macroval);`

Given a mass-flow project that has been loaded with GUI (all N- and M-fields must exist in cnap), this function writes numerical values into the text boxes. The $q \times 1$ vector **reacval** contains the values to be written into the reaction text boxes, i.e. reacval(i) appears in the text box of the i -th reaction. An unknown/undefined value is indicated by value NaN which will then be represented in the box as ‘###’. The $q \times 1$ vector reaccol specifies the color of each reaction text box. Each element in reaccol is an integer value from the set {1,2,3,4} defining the color index of the associated reaction text box. The indices have the following meaning (for colors in CNA see chapter 2): 1=’standard’, 2=’calculated’, 3=’defined’, 4=’special’.

The $m \times 1$ vector macroval contains the numerical values to be written in the text boxes of the macromolecules. They usually define the biomass composition, i.e. the percentages of the biomass constituents given by g macromolecule/gDW (see chapter 3). If you don’t have macromolecules in your model simply use an empty matrix []. The color of macromolecule text boxes is fixed and can thus not be changed. The function returns the network project with a few updated L-fields. For example, the values displayed in the text boxes before using this function are saved so that they can be recalled by the ‘Reset last scenario’ function.

CNAwriteMFNText

Usage: `cnap = CNAwriteMFNText(cnap,reactext,reacol,macrottext)`

Given a mass-flow project that has been loaded with a GUI this function writes strings into the text boxes of reactions and (if existent) of macromolecules. As for `CNAwriteMFNValues` (see above), text boxes of reactions can be colored using different color indices.

There are two analogous functions for reading/writing text box values in signal-flow networks:

CNAreadSFNValues

Usage: `[reacval, specval] = CNAreadSFNValues(cnap);`

Given a signal-flow project that has been loaded with GUI (i.e. all signal-flow N- and M-fields must exist in `cnap`), this function returns the numerical values currently set in the text boxes of reactions ($q \times 1$ vector `reacval`) and species ($n \times 1$ vector `specval`), respectively. If the text box of the i -th reaction does not contain a numerical value, then `reacval(i)=NaN`; the same is done for species. Nothing is changed in the project variable `cnap`; it is therefore not returned.

CNAwriteSFNValues

Usage: `cnap = CNAwriteSFNValues(cnap,reacval,reacol,specval,speccol);`

Given a signal-flow project that has been loaded with GUI (all signal-flow N- and M-fields must exist in `cnap`), this function writes numerical values into the reaction and species text boxes. The $q \times 1$ vector `reacval` contains the values to be written into the reaction text boxes, i.e. `reacval(i)` appears in the text box of the i -th reaction. Analogously, the $n \times 1$ vector `specval` contains the values to be written into the species text boxes, i.e. `specval(i)` appears in the text box of the i -th species. An unknown/undefined value can be indicated by value `NaN` which will then be represented in the respective reaction/species box as `###`. The text box colors are specified by the $q \times 1$ vector `reacol` (reaction text boxes) and by the $m \times 1$ vector `speccol` (species text boxes). Each element in `reacol` and `speccol` is an integer value from the set $\{1,2,3,4\}$ defining the color index of the associated reaction/species text box. The indices have the following meaning: 1="standard", 2="calculated", 3="defined", 4="special". Note that "standard" is different for species and reactions; the other three color indices refer to the same color (see also chapter 2). The function returns the network project with a few updated L-fields. For example, the values displayed in the text boxes before using this function are saved so that they can be recalled by the 'Reset last scenario' function.

The following function can be applied to both types of projects and can be used to integrate own entries (and callbacks) into the CNA pull-down menu.

CNAaddMenuEntry

Usage: `menhandles = CNAaddMenuEntry(cnap,mname,fname);`

`cnap` is a signal- or mass-flow project variable. This function adds a new entry to the CNA pulldown menu "Analysis". `mname` and `fname` are strings defining the name of the menu entry and the associated callback function, respectively. The added menu item will read:

"User function: <mname>"

and is appended to the CNA menu "Analysis" in all interactive maps associated with the project. If the inserted menu entry is selected, the (user-created) function `fname` will be called. `fname` will usually correspond to a m- or MEX-file. Note that the extensions (.m, .dll, .mexglx and so on) must not be contained in `fname`.

The function returns the handles of the created menu entries (if your network project consists of w network maps, then w handles will be returned). The handles may be used to delete inserted menu entries at a later stage. Arbitrary menu entries can be inserted by the user. The project variable `cnap` is not returned as it remains unchanged.

Example

The following example illustrates the usage of the API functions described in this section and also how to access fields of a CNA project structure. Assume you have a mass flow network in which you have computed a flux distribution. You might want to normalize the fluxes by a certain reaction rate (e.g. by a substrate uptake reaction). In addition, by using the text box colors you want to distinguish between reactions carrying no flux, positive flux (running forward) and negative flux (running backward). A MATLAB function (`normalize_flux.m`) which interfaces CNA and performs this task could look like this (you find this function in the directory `CellNetAnalyzer/code/api`):

```

%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
function cnap = normalize_flux(cnap, reacname)

index=[];for z = 1:cnap.numr      if(strcmp(reacname,deblank(cnap.reacID(z,:)))
index=z;                        break;      endend
if isempty(index)  warndlg(['Reaction name ',reacname,' does not exist'],'Error');
return;
end

[rr,mm]=CNAreadMFNValues(cnap);

if(rr(index)==0 | isnan(rr(index)))
warndlg('Normalization cannot be performed','Error');
return;
end

rr=rr/rr(index);
colidx=ones(length(rr),1);
colidx(find(rr==0))=2;
colidx(find(rr>0))=3;
colidx(find(rr<0))=4;

cnap=CNAwriteMFNValues(cnap,rr,colidx,mm);
%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%

```

In case you want to make this function callable from the CNA menu (“Analysis”) of a project `MetNet` (i.e. there is a project variable `MetNet`) you should write the following two lines in the MATLAB command window:

```

rname='Glucose_uptake'; %example for setting name of the reaction
menhandles=CNAaddMenuEntry(MetNet,'Flux normalization',...
                           'MetNet=normalize_flux(MetNet, rname);');

```

7.6 API functions for analyzing a network project

The following API functions give you the opportunity to call CNA’s algorithms for computing and analyzing complex network properties, such as paths, cycles, elementary modes and so forth. Most of these functions (exceptions are, for example, **CNAcomputeCutsets** and **CNATranswedd**) require a CNA network project from which they usually access some selected N-fields (i.e. no access to GUI). You may thus call these functions with projects that have been loaded with or without GUI and also projects with “partial” information generated e.g. with **CNAGenerateMFNetwork** or **CNAGenerateSFNetwork** as described above. The results are usually returned as MATLAB arrays or matrices. The network project itself is not changed or modified.

Here we document the functions and also indicate which N-fields in the network project are required. In general, the documentation for all API functions can also be obtained online in MATLAB by entering: `help <function name>`.

7.6.1 Analysis of Signal-flow networks

The following API functions are useful for analyzing signal-flow networks. This includes functions for exploring interaction graphs and for studying the behavior of logical networks.

CNAbasicNetworkProp

This function computes basic topological and structural properties of a CNA model.

Usage: **CNAbasicNetworkProp**(cnap)

CNAcomputeCycles

This function enumerates cycles (feedback loops) in signal flow networks.

Usage: [cycles, csigs] = **CNAcomputeCycles**(cnap, edge_constr, node_constr, undirected)

CNAcomputePaths

This function enumerates paths in signal flow networks.

Usage: [paths, psigs] = **CNAcomputePaths**(cnap, edge_constr, ...
node_constr, add_in, addout, specstart, specend)

CNAcomputeShortestSignedPaths

This function calculates the shortest signed paths in signal flow networks.

Usage: [distpos, distneg, idx, dist] = **CNAcomputeShortestSignedPaths**(cnap, maxdist, edge_constr, ...
node_constr, algo)

CNAcomputeDepMat

Calculates the dependency matrix of signal flow networks.

Usage: [depmat,idx, distpos,distneg] = **CNAcomputeDepMat**(cnap, maxdist, edge_constr, node_constr, algo)

CNATranswedd

Computes the transitive reduction of a weighted signed (interaction) graph. Useful as false positive reduction method in reverse engineering of cellular interaction graphs (see ref. [19]). A CNA project variable is not required.

Usage: [graph] = **CNATranswedd**(graph,epsilon,fullcheck,pathexact,acyclic,maxpathl)

CNAcomputeLSS

Calculates the logical steady state in a signal-flow network for a given set of input stimuli or fixed signal/species values.

Usage: [spec_lass, inter_lass] = **CNAcomputeLSS**(cnap, spec_vals, inter_vals)

CNAcomputeMIS

Calculates (logical) minimal intervention sets in signal flow networks fulfilling a set of goals defined by a scenario. It is possible to define multiple scenarios whose goals all have to be fulfilled by the MIS. The goals and fixed species/interactions are specific for each scenario while the remaining parameters apply to all scenarios.

Usage: [mis, idx, ec, stat] = **CNAcomputeMIS**(cnap, scen, max_mis_size, err_tol, count_minerr,...
spec_restr, allow_inact, allow_actv, excl_fix_spec , dispval, fpath, fname)

7.6.2 Analysis of Mass-flow networks

The following API functions can be used for analyzing mass-flow networks.

CNAbasicNetworkProp

This function computes basic topological and structural properties of a CNA model.

Usage: **CNAbasicNetworkProp**(cnap)

CNAoptimizeFlux

Performs flux optimization in mass-flow networks: ojective function `cnap.objFunc` is **minimized** by linear programming (LP). Parsimonious flux optimization can be performed and additional inequality constraints can be specified.

Usage: [optFlux, success, status, optval] = **CNAoptimizeFlux**(cnap, fixedFluxes, c_macro, solver, dispval, pFBA, A_ieq, b_ieq)

CNAoptimizeYield

This function maximizes the yield function c^*r/d^*r for flux vectors r within the flux space defined by `cnap` and the 'fixedFluxes' vector.

Usage: function [maxyield,flux_vec,success, status] =
CNAoptimizeYield(cnap, c, d, fixedFluxes, c_macro, solver, dispval)

CNAfluxVariability

Flux variability analysis in mass-flow networks. Given a CNA mass-flow project (with or without GUI) this function determines the range of feasible fluxes for each reaction by solving linear optimization problems. Some fluxes may be fixed and additional inequality constraints can be defined.

Usage: [minFlux,maxFlux,success,status] =
CNAfluxVariability(cnap,fixedFluxes,c_macro,solver,reactdx,A_ieq,b_ieq)

CNAcomputeElemConsRel

Computation of elementary conservation relations in a mass-flow network.

Usage: CR = **CNAcomputeElemConsRel**(cnap,solver,positive,c_macro)

CNAremoveConsRel

This API function removes successively conservation relations from a mass-flow project by deleting metabolites (or by switching internal metabolites to external) until no further conservation relation remains in the system. The user can iteratively select a species from a list of metabolites (being part of at least one conservation relation) which is then deleted to break a conservation relation. This is repeated until no further conservation relations remains. Alternatively, the metabolites to be deleted / to be set to external are automatically chosen by the function (no user input).

Usage: [cnap,delspec] = **CNAremoveConsRel**(cnap,rational,setToExt,auto)

CNAplotPhasePlane

Usage: [success,status] = **CNAplotPhasePlane**(cnap, plotrxns, fixedFluxes, c_macro, solver, numdisc)

Given a mass-flow project (with or without GUI), some predefined fluxes (fixedFluxes), and a list of n reactions (plotrxns), this function displays $n*(n-1)/2$ many 2D projections of the flux space onto the reaction rates (one projection for each possible pair of the n reactions in plotrxns).

CNAplotYieldSpace

Generates 2D yield space plot for two user-specified yields through sampling in discrete steps

Usage: [yieldspace,success] = **CNAplotYieldSpace**(cnap,c,d,c2,d2,numdisc,fixedFluxes,c_macro,solver,dispval)

Given a mass-flow project (cnap, with or without GUI), possibly some predefined fluxes (fixedFluxes), and a predefined biomass composition (c_macro; relevant for projects with variable biomass composition), this function uses (numdisc many) discrete steps to approximate and plot the bounds of the two-dimensional yield space defined by the two yield functions $Y1=c*r/d*r$ and $Y2=c2*r/d2*r$ (r: feasible flux vector). The calculated bounds of the yield space will be returned and the convex hull of yield space bounds be plotted.

CNAcomputeEFM

Computes elementary (flux) modes / elementary (flux) vectors or a minimal generating set (convex basis) of flux cones or flux polyhedra associated with mass-flow networks. Two different scenarios can be considered:

(i) In the homogeneous case, the solution space defined by the steady state assumption and reversibility constraints form a polyhedral (flux) cone. Here, elementary modes can be computed which are particular flux vectors of this cone with minimal support (irreducible set of reactions with non-zero rate). The set of elementary modes includes all extreme rays of the flux cone, but usually many more vectors. CNAcomputeEFM can also compute a convex basis of the flux cone. In contrast to elementary modes, the convex basis is a minimal set of vectors sufficient to generate all flux vectors within the flux cone by non-negative linear combinations of convex basis vectors. As implemented herein, the convex basis will always represent a subset of the elementary modes. Note that the convex basis is only unique if there is no reversible flux vector.

Specifying inhomogeneous constraints (e.g., fixing a reaction rate to a non-zero value or introducing non-zero upper and/or lower boundaries for the reaction rates) leads to the inhomogeneous case where the solution space becomes a more general flux polyhedron. CNAcomputeEFM computes then either the elementary vectors of the flux polyhedron (a generalization of elementary modes; the set of elementary vectors will, for example, include all extreme rays and extreme points (if existent) of the resulting solution space). Again, alternatively, a convex basis (= minimal set of generators spanning the resulting flux polyhedron) of the flux polyhedron can also be calculated. Note that the zero point will not be delivered, even if it is an extreme point of the flux polyhedron.

For more information see section 3.3.5 and reference [29].

Usage: [efm, irrev, idx, ray] = **CNAcomputeEFM**(cnap, constraints, solver, irrev_flag, ...
convbasis_flag, iso_flag, c_macro, display, efmtool_options)

CNAcomputeCutsets

This function uses the Berge algorithm to compute minimal cut sets (or minimal hitting sets) for a given set of binary vectors (which may represent the support of certain paths/cycles/elementary modes/elementary vectors / sets or hyperedges). The operation is equivalent to hypergraph transversal.

A CNA project variable is not required.

Usage: cutsets = **CNAcomputeCutsets**(targets, mcsmax, names, sets2save, earlycheck, prefilter)

CNAMCSEnumerator

CNAMCSEnumerator2

CNAregMCSEnumerator

CNAfindRegMCS

CNAgeneMCSEnumerator

CNAgeneMCSEnumerator2

All these API functions are different variants to compute minimal cut sets (MCS) in mass-flow networks via the dual algorithm presented in [24]. This algorithmic approach does not need to calculate the elementary modes in a preprocessing step (in contrast to CNAcomputeCutsets computing the MCS from a given set of EFMs).

Generally, given a mass-flow project *cnap* and a set of 'undesired' (target) flux vectors (defined by matrix T and vector t as shown below) and (optionally) a set of 'desired' flux vectors (defined by matrix D and vector d) MCSs are computed fulfilling the following properties: Knocking out the reactions (or genes in case of gene MCS) of an MCS guarantees that all (target) flux vectors v obeying

$$\begin{aligned} \text{cnap.stoichimat} * v &= 0 \\ \text{reac.Min} &\leq v \leq v.\text{reacMax} \\ T*v &\leq t \end{aligned}$$

will be blocked (are infeasible) whereas at least one flux vector r fulfilling

$$\begin{aligned} \text{cnap.stoichimat} * r &= 0 \\ \text{cnap.reacMin} &\leq r \leq \text{cnap.reacMax} \\ D*r &\leq d \end{aligned}$$

will be kept functional.

If D and d are non-empty (i.e., if desired flux vectors have been defined) the MCSs computed represent CONSTRAINED MCSs (cMCSs), otherwise unconstrained MCSs. CNAMCSEnumerator2 is the latest and most up to date version with many new features (e.g., multiple target and desired regions, reaction deletions and additions etc.).

In order to run these functions it is necessary that the MATLAB CPLEX and, for some functions, also the Java CPLEX interfaces work (see section 0.5). Note that the JAVA CPLEX is not mandatory anymore for CNAMCSEnumerator2.

Here is a brief description of the different variants of these functions:

CNAMCSEnumerator

This is the original algorithm used in [24]

Usage: [mcs] = **CNAMCSEnumerator**(cnap, T, t, D, d, notknockable, maxMCS, maxMCSsize, filename)

CNAMCSEnumerator2

This is an extended and revised version of CNAMCSEnumerator. For example, it allows to also consider reaction additions (knock-ins) and variable costs of the interventions. In this way, regulatory MCS as explicitly calculated by CNAMCSEnumerator (see below) can also be computed via CNAMCSEnumerator2. Also single MCS as computed by CNAfindRegMCS can be found by this algorithm [32]. CPLEX, intlinprog and glpk are supported as MILP solvers.

Usage: [mcs, status, cmp_mcs, cmp_cnap, mcs_idx_cmp_full, obj] = ...
CNAMCSEnumerator2(cnap,T,t,D,d,koCost,kiCost,maxSolutions,maxCost,options,verbose)

CNAMCSEnumerator3

This is an extended and revised version of CNAMCSEnumerator2 and allows the use of more general descriptions of undesired (target) and desired regions (here called modules). CNAMCSEnumerator3 also supports CPLEX, intlinprog, glpk and additionally Gurobi as MILP solver.

Usage: function [mcs, status, cmp_mcs, cmp_cnap, mcs_idx_cmp_full, obj] = ...
CNAMCSEnumerator3(cnap,modules,koCost,kiCost,maxSolutions,maxCost,options,verbose)

CNAregMCSEnumerator

This API function extends CNAMCSEnumerator to enumerate not only smallest constrained Minimal Cut Sets (cMCS; combinations of reaction knockouts fulfilling a given intervention goal) but also smallest *regulatory* constrained Minimal Cut Sets (which are combinations of reaction knockouts and flux up- and down-regulations) in large-scale networks [Ref. 25].

Usage: [mcs, reacNames] = **CNAregMCSEnumerator**(cnap,T,t,D,d,notknockable,maxMCSnum,maxMCSsize, ...
filename,useIntegratedMILP,react_off,regulation,time_limit,default_flux_limit)

CNAfindRegMCS

Similar to 'CNAregMCSEnumerator', but only attempts to find a single (not necessarily smallest) constrained minimal cut set in mass-flow networks. Optionally, up- and down-regulation of certain reaction fluxes may be considered as intervention strategies in combination with reaction cuts (knockouts) yielding then 'regulatory cMCSs'.

Usage: [mcs, reacNames, reg, sys, obj, cs] = **CNAfindRegMCS**(cnap,T,t,D,d,notknockable,
maxMCSsize,filename,react_off,regulation,time_limit,EpGap)

CNAgeneMCSEnumerator

This function is similar to CNAMCSEnumerator but instead of calculating minimal reaction cut sets, this function determines minimal gene cut sets based on gene-enzyme-reaction associations. "Regulatory cuts" as in CNAregMCSEnumerator are not allowed. However, the algorithm uses the integrated approach that is also employed by CNAregMCSEnumerator. Before the core cut set algorithm starts, the gene-enzyme-reaction associations are integrated into the metabolic network similar as described by Machado et al. (2016), PLoS Comput Biol 12: e1005140.

Usage: [mcs, gene_idx]= **CNAgeneMCSEnumerator**(cnap, T, t, D, d, notknockable, maxMCSnum,
maxMCSsize, react_off, time_limit, default_flux_limit, enzymes, rgm, erm, has_gn_ass)

CNAgeneMCSEnumerator2

This is a revised version of CNAgeneMCSEnumerator to compute gene MCS. It uses internally the functions CNAgenerateGPRrules and CNAintegrateGPRrules to integrate the GPR rules as pseudo-reactions and then CNAMCSEnumerator2 to calculate the (gene) MCS. [32]

Usage: [rmcs, full_mcs, full_cnap, cmp_mcs, cmp_cnap, mcs_idx_cmp_full, status, obj] =
CNAgeneMCSEnumerator2(cnap,T,t,D,d,koCost,kiCost,maxSolutions,maxCost,timeLimit,
use_bigM,enum_method,gkoCost,gkiCost,gpr_rules,useCompression,verbose)

CNAcharacterizeIS

Characterizes a set of intervention strategies (IS) at reaction level, e.g. MCS computed via MCSEnumerator(2), by up to 10 different criteria, groups it into equivalence classes and generates a ranking. [31]

Usage: [IS_rankingStruct, IS_rankingTable] = **CNAcharacterizeIS** cnapRef , IS_lb, IS_ub, IS_to_rank, idx,
cytMet, D, d, T, t, mdParam, lbCore, ubCore, crit, wCoeff,solver)

CNAcharacterizeGeneMCS

Characterizes a set of gene MCS, e.g., computed via MCSgeneEnumerator(2), by up to 10 different criteria, groups it into equivalence classes and generates a ranking.

Usage: [MCS_rankingStruct, MCS_rankingTable] = **CNAcharacterizeGeneMCS**(cnapRef , MCS_lb, MCS_ub, MCS_to_rank, idx, cytMet, D, d, T, t, mdParam, lbCore, ubCore, gmcs, intvCost, gene_and_reac_names , gmcs_rmcs_map, crit, wCoeff, solver)

CNAapplyCASOP

Calculation of reaction weights and reaction ranking for identification of knockout or overexpression candidates according to our recently proposed method CASOP (= Computational Approach for Strain Optimization aiming at high Productivities) [18]. Applicable in mass-flow networks. The target metabolite (desired product) has to be defined as an external metabolite. The biomass reaction can be defined either by the standard biomass synthesis reaction "mue" (see Section 3) or by a user defined biomass synthesis reaction. In the latter case, an external biomass metabolite has to be defined (which must be produced in the user-defined biomass synthesis reaction) and the names of the biomass metabolite and synthesis reaction are additional arguments for CNAapplyCASOP.

Usage: [ReacWeightMatrix, RatingMatrix]= **CNAapplyCASOP**(cnap, gamma_vec, k_vec, product_name, ... molm_product, uptake_reaction_names, rating_boundaries_vec, plot_reaction_ids, ... legendID, EMoptions, BMreaction, BMmetabolite)

For arguments see also the original CASOP publication [18].

CNAreduceMFNetwork

Reduces a mass-flow networks to a sub- (or core) network based on user-given specifications. The function starts with a pruning step followed by an (optional) compression step (via CNAcompressMFNetwork). See ref. [26].

Usage: [cnapred, delreactions, delmetabolites] = **CNAreduceMFNetwork**(cnap, FG, protect_func, ... protect_met, protect_reac, feas, rmin, solver, comp, rational)

CNAcompressMFNetwork

Loss-free network compression of mass-flow networks. Based on the stoichiometric matrix and reaction reversibilities this function compresses the reaction network by

- (i) removing blocked reactions
- (ii) lumping enzyme (reaction) subsets
- (iii) lumping choke points (= metabolites produced / consumed by one unique reaction and consumed/produced by many reactions)
- (iv) removing conservation relations (= dependent metabolites).

Compression based on these rules is loss-free meaning, for example, that the compressed system has the equivalent set of elementary modes etc. Certain reactions or/and metabolites can be protected against compression. Compression based on efmtool's rational arithmetic can be used optionally.

[Note: this function is routinely used in CNA, for example before calculating elementary modes.]

Usage: [redsmat, irrev, reacidx, metidx, cnapcomp]=**CNAcompressMFNetwork**(cnap, protect_reac, protect_spec, ... rmCR, rmChoke, rational, blocked_reac, nodisp)

CNAcomputeOptMDFpathway

This function calculates a pathway (flux vector v) with the maximal Max-min driving force (MDF), i.e., a pathway with associated metabolite concentrations where the minimum driving force of all participating reactions is maximal. (For MDF see Noor et al. (2014), PLOS Comp Biol, 10:e1003483)

Usage: [mdf, v, conc, dfs] = **CNAcomputeOptMDFpathway**(cnap, RT, G0, Cmin, Cmax, ...
D, d, fixed_ratios, stoichMat)

CNAmake Feasible

This function finds minimal (weighted) changes in a set of given (fixed) fluxes to make a (possibly) infeasible flux scenario feasible.

Usage: [FeasFlux, cor, success] = **CNAmakeFeasible**(cnap, fluxes, c_macro, w, linsolver, useqp, verbose)

7.7 API functions for loading and saving scenarios

CNAloadMFNValues

Load scenarios (reaction and macromolecule values) of mass-flow networks from a (CNA) val-file.

Usage: [reacval, macroval, err, reacMin, reacMax, objFunc] = **CNAloadMFNValues**(cnap, fname)

CNASaveMFNValues

Save scenarios (reaction and macromolecule values) of a mass-flow project to a (CNA) val-file.

Usage: err = **CNASaveMFNValues**(cnap, fname, reacval, macroval, fullscenario, numformat)

CNAloadSFNValues

Load interaction and species values from a (CNA) val-file in signal-flow networks.

Usage: [reacval, specval] = **CNAloadSFNValues**(cnap, fname)

CNASaveSFNValues

Saves interaction and species values into to a (CNA) val-file.

Usage: err = **CNASaveSFNValues**(cnap, fname, reacval, specval)

7.8 Starting a network project w/o GUI

It is possible to work with a project without using the GUI (at least to some extent) which might be useful for certain applications. You can load the network without the GUI by clicking on

“Start w/o GUI” in the project manager.

You may even suppress the project manager GUI when starting CNA, e.g. if you want to operate in batch-mode, if you call CNA only for calculating some network features, or if you are using octave. You can do this by calling **startcna(1)**. As described in a previous section, the API function **CNALoadNetwork** allows one to load network projects (optionally) without GUI, i.e., without interactive maps. Note again that you must definitely initialize CNA (with **startcna**) before using any of the API functionality presented in this chapter. Calling **startcna** - which is located in *CellNetAnalyzer*'s main directory - adds all relevant CNA directories to the MATLAB path, initializes the global variable **cnan** and launches the project manager window (unless called with **startcna(1)**). To finish CNA enter **close_cna**, which closes all projects that are running with a GUI, removes the CNA directories from the MATLAB path and deletes the **cnan** variable.

7.9 The global *cnan* variable and CNA options

When starting the CNA with **startcna**, the global variable *cnan* is initialized and appears in the base workspace. *cnan* is a structure variable whose fields are used to store data for the project manager, certain GUI handles of projects with an active GUI and information about the CNA configuration. For the user the most relevant field is *cnan.options* which controls the behavior of the CNA and its GUI to some degree. The current options and their possible values are:

cnan.options.ask_when_closing_project: controls whether a confirmation dialog comes up when closing one of the network maps. Possible values are 0: never ask; 1: ask when there are unsaved changes; 2: ask always.

cnan.options.ask_when_saving_net: controls whether a confirmation dialog comes up when saving the network structure. Possible values are 0: never ask; 1: ask if the project has an active GUI.

cnan.options.ask_when_closing_composer: controls whether the user is asked to save changes when closing the network composer and unsaved changes exist. Possible values are 0: never ask; 1: ask if unsaved changes exist.

7.10 Example for using the CNA API from another application

Suppose you have an application that provides a stoichiometric matrix and information about reaction reversibilities. Let's assume that the stoichiometric matrix is called *st* and that there is a vector *irrev* that contains 0 for reversible reactions and 1 for irreversible ones. The length of *irrev* must be the same as the number of columns of *st*. Furthermore, let *st* contain only internal metabolites. The following example illustrates how you can use the API of CNA to calculate the elementary modes and the corresponding minimal cut sets from the latter:

```
...
startcna(1); % initialize CNA without project manager
cnap.stoichMat= st;
cnap.reacMin= zeros(length(irrev), 1); % must be a column vector
cnap.reacMin(irrev == 0)= -Inf; % set up reversibility
%all other fields of cnap will be set (with default values) by the following command
cnap= CNAgenerateMFNetwork(cnap);
[ems, irrev_ems, ems_idx] = CNAcomputeEFM(cnap);
mcs=CNAcomputeCutsets(ems);
close_cna; % close CNA and clean up
...
```

Of course you could also encapsulate these commands in a separate function. In any case, you have to make sure that the *CellNetAnalyzer* directory is in the MATLAB path because otherwise *startcna* will not be found (and the CNA paths would not be set).

7.11 CNA plugins

The API functions of *CellNetAnalyzer* allow interested developers to build their own applications interfacing CNA. As demonstrated, it can also be used to build CNA plugins such as ODEfy (which is even a permanent part of CNA). On CNA's website you can find a list of downloadable plugins developed by other users.

8. References

Articles with “+” are open access and thus freely available at the journal’s website. Let us know if you would like to have a copy of any of the cited journal articles.

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Online manual: http://www.mpi-magdeburg.mpg.de/projects/cna/manual_cellnetanalyzer.pdf

Online tutorial: http://www.mpi-magdeburg.mpg.de/projects/cna/CNA_tutorial.pdf

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