

tutorial - 0 - Get data, model, export, simulate, plot, and manipulate data

September 7, 2020

```
[31]: # expand cells to the 95% of the display width
from IPython.core.display import display, HTML
display(HTML("<style>.container { width: 95% !important; }</style>"))
```

<IPython.core.display.HTML object>

1 Tutorial: Automatic rule-based modeling of metabolism, protein-protein interactions, and regulation of gene expression employing *Atlas*

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Notes: This tutorial was created for the manuscript “*Atlas*: Automatic modeling of regulation of bacterial gene expression and metabolism using rule-based languages”, first submitted for peer-review to Bioinformatics on May, 2020.

1.1 Prerequisites

0. The tutorial was prepared and executed on Ubuntu 20.04, PathwayTools version 24, and docker engine version 19.03.8.
1. PathwayTools must be installed and running to obtain data from the EcoCyc database. Please, run `pathway-tools -lisp -python-local-only` before to obtain any data. (Optional) The PathwayTools software could be executed in the background, with help of `nohup` `pathway-tools -lisp -python-local-only > /dev/null 2> /dev/null &`. Please follow instructions at <http://pathwaytools.org/> to obtain a licensed copy of the software from <https://biocyc.org/download-bundle.shtml>. However, data could be manually formatted using a text-based editor or a spreadsheet software.

Note: If you ran into the `pathway-tools/ptools/24.0/exe/aclssl.so: undefined symbol: CRYPTO_set_locking_callback` error, please follow instructions [here](#):

<https://github.com/glucksfall/atlas/tree/master/PTools-v24>. Instructions will guide you to install a docker image that is able to run pathway tools, but does not include it, so you still need to obtain the software with a valid license.

2. (Highly recommended) Install Docker. Please follow instructions for a supported Operating System <https://docs.docker.com/engine/install/>: On Ubuntu, install it with `apt-get install docker.io`. On Win10, install Docker Desktop with WSL2 support <https://docs.docker.com/docker-for-windows/wsl/>. On MacOS, install Docker Desktop <https://docs.docker.com/docker-for-mac/install/>. The Docker `networkbiolab/pleiades` installs the python packages, the jupyter server, and the stochastic simulators.
3. (Recommended) Jupyter notebook. We recommend the use of Anaconda3 <https://www.anaconda.com/products/individual> because of the easier installation of the stochastic simulators from <https://anaconda.org/alubbock>.
4. (Optional) A stochastic simulator, supported by the pySB python package ([BNG2](#), [NFsim](#), [KaSim](#) or [Stochkit](#)). pySB requires BNG2 to simulate models with NFsim.
5. (Optional) Cytoscape to visualize metabolic networks and others.
6. (Optional) A deterministic simulator: pySB supports ODE integration via `scipy.integrate.ode`, BioNetGen ODE integration, and CUDA-accelerated ODE integration with Marco S. Nobile's cupSODA software (<https://github.com/aresio/cupSODA>). If the user feel comfortable with SBML models, pySB could export to SBML and deterministic simulation done with libRoadRunner (<http://libroadrunner.org/>), Tellurium (<http://tellurium.analogmachine.org/>), COPASI (<http://copasi.org/>), etc.

1.2 Installation

0. If you are running the docker image "pleiades", please go directly to the section "Preamble".
1. To install, please follow one of the following steps:
 1. Install the docker image "pleiades" using `docker pull networkbiolab/pleiades`. The container is based on the Anaconda3 software and it installs Atlas, and the stochastic simulators BNG2, NFsim, KaSim, and Stochkit. After building the image, please run the container with `docker run --detach --publish 10000:8888 networkbiolab/pleiades`, and go to `localhost:10000` in your preferred browser. The required password is `pleiades`.
 2. Download or clone the Github repository from <https://github.com/networkbiolab/pleiades> with `git clone https://github.com/networkbiolab/pleiades foo` (where `foo` is an absolute or relative path). Then, you could build the docker image with `docker build foo --tag pleiades` and run it with `docker run --detach --publish 10000:8888 pleiades`. Finally, go to `localhost:10000` in your preferred browser. The required password is `pleiades`.
 3. Install with pip3: `sudo -H python3 -m pip install pleiades` or `python3 -m pip install pleiades --user`. Pleiades is a meta-package that install Atlas (the rule-based modeller), Pleione (a genetic algorithm for parameter calibration of RBMs, compatible with SLURM), Alcyone (to perform identifiability analysis of parameters), and Sterope (to perform sensitivity analysis of parameters in kappa RBMs, compatible with SLURM). You should install, configure, and run the jupyter notebook on your own:

```
example sudo -H pip3 install jupyter && nohup python3 -m jupyter notebook
--port=8888 --no-browser --port-retries=0 > /dev/null 2> /dev/null &
```

4. Download or clone the Github repository from <https://github.com/networkbiolab/atlas> with `git clone https://github.com/networkbiolab/atlas foo` (where `foo` is an absolute or relative path). Requisites must be fulfilled manually with `pip3`: `sudo -H python3 -m pip install pandas pysb pythoncyc jupyter seaborn` or `python3 -m pip install pandas pysb pythoncyc jupyter seaborn --user`.

1.3 Objectives

1. Get metabolic data: enzyme names, substrates, products, and location of enzymes.
2. Basic manipulations of metabolic data: change reversibility, change enzyme location.
3. Get composition of complexes for protein-protein interactions.
4. Basic manipulations of interaction data: add and remove interactions.
5. Perform simulation, plot of variables, and export of models.
6. Get DNA architecture and build a model from a manually written network.
7. Get protein-DNA and TF-metabolite interactions and build a model from manually written networks.
8. Add gene regulatory interactions to transcription rules: Activation of transcription.

1.4 Preamble: load *Atlas*

```
[2]: # testing source code
# required if atlas was cloned from GitHub and this notebook is executed from
# → the tutorial directory.
import sys
sys.path.append("..")

import atlas_rbm.atlas as atlas
import atlas_rbm.utils as utils
import atlas_rbm.export as export
import atlas_rbm.simulation as simulation
```

```
[3]: utils.checkPathwayTools()
```

PathwayTools is running. Available PGDB are: META, ECOLI

```
[3]: True
```

```
[4]: utils.execPToolsDocker('ptools-v24')
```

Doing nothing since PathwayTools is running.

1.5 Getting data to model metabolism

In this tutorial, we will obtain data from EcoCyc to model the lactose degradation that occurs in *Escherichia coli*. We choose the lactose metabolism since it was discovered in the decade of 1960s and it is a common model of gene regulation with more than 50 years of biochemical information. In an side

note, the characterization of the lactose operon and others rewarded their authors the 1965 Nobel Prize in Physiology or Medicine (<https://www.nobelprize.org/prizes/medicine/1965/summary/>)

The lactose operon from *E. coli* consists of three genes: the β -galactosidase gene *lacZ*, the lactose permease gene *lacY* (also known as lactose-proton symporter), and the galactoside O-acetyltransferase gene *lacA*.

Note: The location is informed for all gene products, including the location of complexes formed by the encoded protein. Therefore, the location may reflect the location of the enzyme or its components before complex assembly. Still, there are 154 enzymatic and 3 transport reactions without known gene, and the location will set to unknown.

Note: The location could be changed later with help of an utility function. The function `utils.metabolicNetwork.FromEnzymeList()` produces data in three formats for the **GENE OR COMPLEX** column: 1. The **product** key produces networks with the name of the enzyme, but the model will not be compatible with the modeling of gene expression. **Note:** The **product** key will try to retrieve the location of the enzyme or transporter. If not, the location refer to the location of the monomers of the complex, e.g. "BETAGALACTOSID-CPLX". 2. The **gene** key produces networks with the name of the gene (is the default), but the model will not be compatible with modeling of protein-protein interactions, specially for the modeling of heteromers. **Note:** For enzyme complexes, the **genes** key will decompose the complex into its genes and locations of their products, e.g. the *araFGH* transporter ("ABC-2-CPLX") where the *araG* product locates to the cytoplasm and the inner membrane when interact with *araH*. 3. The **complex** key produces hyper-networks (brackets denote subnetworks, and we will subsequently call it as *hypergraph notation*), and in that way, Atlas produces a model that will be compatible with the modeling of protein-protein interactions and gene expression. **Note:** For enzyme complexes, the **complex** key will decompose the complex into its genes AND write the stoichiometry of the complex in the *hypergraph notation*. If one or more monomers have two or more locations, the output will show as many rows to show all combinations. Be aware that some could be not physiologically possible.

```
[5]: %time network = utils.metabolicNetwork.FromEnzymeList('ECOLI',
↳ ['GALACTOACETYLTRAN-CPLX', 'LACY-MONOMER', 'BETAGALACTOSID-CPLX',
↳ 'ABC-2-CPLX'], fmt = 'product')
network
```

CPU times: user 35.8 ms, sys: 61.9 ms, total: 97.7 ms

Wall time: 911 ms

```
[5]:
```

	GENE OR COMPLEX	ENZYME	LOCATION	REACTION \
0	ABC-2-CPLX		inner membrane	ABC-2-RXN
1	BETAGALACTOSID-CPLX		cytosol	BETAGALACTOSID-RXN
2	BETAGALACTOSID-CPLX		cytosol	RXN0-5363
3	BETAGALACTOSID-CPLX		cytosol	RXN-17726
4	BETAGALACTOSID-CPLX		cytosol	RXN0-7219
5	GALACTOACETYLTRAN-CPLX		cytosol	GALACTOACETYLTRAN-RXN
6	LACY-MONOMER		inner membrane	TRANS-RXN-24
7	LACY-MONOMER		inner membrane	TRANS-RXN-94
8	LACY-MONOMER		inner membrane	RXN0-7215
9	LACY-MONOMER		inner membrane	RXN0-7217

10	LACY-MONOMER	inner membrane	RXN-17755
	SUBSTRATES		PRODUCTS \
0	WATER,ATP,L-ARABINOSE		Pi,ADP,PROTON,L-ARABINOSE
1	CPD-15972,WATER		GALACTOSE,Glucopyranose
2	Alpha-lactose		ALLOLACTOSE
3	CPD-3561,WATER		GALACTOSE,Fructofuranose
4	CPD-3785,WATER		GALACTOSE,D-ARABINOSE
5	Beta-D-Galactosides,ACETYL-COA	6-Acetyl-Beta-D-Galactosides,CO-A	
6	PROTON,Alpha-lactose		PROTON,Alpha-lactose
7	PROTON,MELIBIOSE		PROTON,MELIBIOSE
8	PROTON,CPD-3561		PROTON,CPD-3561
9	PROTON,CPD-3785		PROTON,CPD-3785
10	PROTON,CPD-3801		PROTON,CPD-3801
	FWD_RATE	RVS_RATE	
0	1.0	0.0	
1	1.0	0.0	
2	1.0	1.0	
3	1.0	0.0	
4	1.0	0.0	
5	1.0	0.0	
6	1.0	1.0	
7	1.0	1.0	
8	1.0	1.0	
9	1.0	1.0	
10	1.0	1.0	

Note: Using the `utils.metabolicNetwork.FromEnzymeList()` function with the `all_enzymes()` and `all_transporters()` functions from the `pythoncyc` package, we could obtain a **draft** network and a **draf** rule-based model of metabolism for the considered organism:

```
[6]: import pythoncyc
%time network = utils.metabolicNetwork.FromEnzymeList('ECOLI', pythoncyc.
↳select_organism('ECOLI').all_transporters())
%time utils.metabolicNetwork.expand_network(network, 'ecocyc-v24-tps-cytoscape.
↳txt')
network.to_csv('ecoli-tps-v24.txt', sep = '\t', index = False)
%time atlas.construct_model_from_metabolic_network('ecoli-tps-v24.txt',
↳noObservables=True, noInitials=True, toFile = 'model-transporters.py')
len(network[network['ENZYME LOCATION'].str.match('unknown')]) # how many
↳reactions are not mapped to a gene product
```

```
CPU times: user 2.82 s, sys: 2.83 s, total: 5.65 s
Wall time: 53.7 s
CPU times: user 8.82 ms, sys: 0 ns, total: 8.82 ms
Wall time: 8.78 ms
```

It was found duplicated reaction names in the network.
Please check the conflicting_reactions.txt and correct them if necessary.
CPU times: user 519 ms, sys: 38.7 ms, total: 558 ms
Wall time: 570 ms

[6]: 3

```
[7]: import pythoncyc
%time network = utils.metabolicNetwork.FromEnzymeList('ECOLI', pythoncyc.
↳select_organism('ECOLI').all_enzymes())
%time utils.metabolicNetwork.expand_network(network, 'ecocyc-v24-enz-cytoscape.
↳txt')
network.to_csv('ecoli-enz-v24.txt', sep = '\t', index = False)
%time atlas.construct_model_from_metabolic_network('ecoli-enz-v24.txt',
↳noObservables=True, noInitials=True, toFile = 'model-enzymes.py')
len(network[network['ENZYME LOCATION'].str.match('unknown')]) # how many
↳reactions are not mapped to a gene product
```

Unable to retrieve data for CPLX0-7889. Please, review the information at
<https://biocyc.org/ECOLI/NEW-IMAGE?object=CPLX0-7889> and post an issue at
<https://github.com/networkbiolab/atlas> if you believe it is a software error.
CPU times: user 9.35 s, sys: 9.23 s, total: 18.6 s
Wall time: 2min 50s
CPU times: user 17.6 ms, sys: 2.36 ms, total: 19.9 ms
Wall time: 19.8 ms
It was found duplicated reaction names in the network.
Please check the conflicting_reactions.txt and correct them if necessary.
CPU times: user 1.16 s, sys: 35.9 ms, total: 1.2 s
Wall time: 1.21 s

[7]: 155

However, we must know the exact enzyme ID from the database. As an alternative, we could obtain the metabolic reactions from the ECOLI database from PathwayTools with help of the `utils.metabolicNetwork.FromGeneList()` function. The function produce data in three formats for the GENE OR COMPLEX column: 1. The **product** key produces networks with the name of the enzyme, but the model will not be compatible with the modeling of gene expression. **Note:** The **product** key will find the product that perform the enzymatic reaction, e.g. *lacZ* produces monomers that interact to produce the enzyme. Also, it will write the location of the product(s) of the queried gene. It is a good practice to query only one gene of the complex. 2. The **gene** key produces networks with the name of the gene (is the default), but the model will not be compatible with modeling of protein-protein interactions, specially for the modeling of heteromers. **Note:** The **gene** key will write the location of the product(s) of the queried gene., e.g. *araF*, *araG*, and *araH*. 3. The **complex** key produces hyper-networks, and in that way, Atlas will produce a model that is compatible with the modeling of protein-protein interactions and gene expression. **Note:** The **complex** key will write the stoichiometry of the enzymatic complex of the queried gene, and the mapped location of the complex, all in *hypergraph notation*. It is a good practice query only one gene of the complex.

8	1.0	1.0
9	1.0	1.0
10	1.0	0.0
11	1.0	1.0
12	1.0	0.0
13	1.0	0.0

Note: The function is particularly slow since it needs to build a dataframe to map gene names (e.g. *lacY*) to the internal identification name that PathwayTools understand: EG10526 <-> lacY. With that internal identification, the function looks for enzymes, reactions of those enzymes, the substrates and products of those reactions, and the location of the enzymes. We could accelerate the process (useful for repeated gathering of data) if we execute `utils.returnCommonNames()` and utilize that precalculated dataframe and pass to the utility functions.

```
[9]: # %time utils.getData('ECOLI', 'EG10526')
%time utils.getData('ECOLI', 'EG10526')['common_name'] # for simplicity of the
↳output
```

CPU times: user 903 µs, sys: 716 µs, total: 1.62 ms
Wall time: 14.3 ms

```
[9]: 'lacY'
```

```
[10]: %time df_genes = utils.returnCommonNames('ECOLI')
```

CPU times: user 2.37 s, sys: 2.3 s, total: 4.67 s
Wall time: 48.8 s

```
[11]: %time utils.metabolicNetwork.FromGeneList('ECOLI', ['lacZ', 'lacA', 'lacY',
↳'araF', 'araG', 'araH'], fmt = 'product', precalculated = df_genes)
# ~35 times faster
```

CPU times: user 71.3 ms, sys: 58.4 ms, total: 130 ms
Wall time: 1.38 s

```
[11]:
```

	GENE OR COMPLEX	ENZYME LOCATION	REACTION \
0	ABC-2-CPLX	periplasmic space	ABC-2-RXN
1	ABC-2-CPLX	inner membrane	ABC-2-RXN
2	ABC-2-CPLX	cytosol	ABC-2-RXN
3	ABC-2-CPLX	inner membrane	ABC-2-RXN
4	GALACTOACETYLTRAN-CPLX	cytosol	GALACTOACETYLTRAN-RXN
5	LACY-MONOMER	inner membrane	TRANS-RXN-24
6	LACY-MONOMER	inner membrane	TRANS-RXN-94
7	LACY-MONOMER	inner membrane	RXN0-7215
8	LACY-MONOMER	inner membrane	RXN0-7217
9	LACY-MONOMER	inner membrane	RXN-17755
10	BETAGALACTOSID-CPLX	cytosol	BETAGALACTOSID-RXN
11	BETAGALACTOSID-CPLX	cytosol	RXN0-5363

12	BETAGALACTOSID-CPLX	cytosol	RXN-17726
13	BETAGALACTOSID-CPLX	cytosol	RXN0-7219
	SUBSTRATES		PRODUCTS \
0	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE	
1	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE	
2	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE	
3	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE	
4	Beta-D-Galactosides,ACETYL-COA	6-Acetyl-Beta-D-Galactosides,CO-A	
5	PROTON,Alpha-lactose	PROTON,Alpha-lactose	
6	PROTON,MELIBIOSE	PROTON,MELIBIOSE	
7	PROTON,CPD-3561	PROTON,CPD-3561	
8	PROTON,CPD-3785	PROTON,CPD-3785	
9	PROTON,CPD-3801	PROTON,CPD-3801	
10	CPD-15972,WATER	GALACTOSE,Glucopyranose	
11	Alpha-lactose	ALLOLACTOSE	
12	CPD-3561,WATER	GALACTOSE,Fructofuranose	
13	CPD-3785,WATER	GALACTOSE,D-ARABINOSE	

	FWD_RATE	RVS_RATE
0	1.0	0.0
1	1.0	0.0
2	1.0	0.0
3	1.0	0.0
4	1.0	0.0
5	1.0	1.0
6	1.0	1.0
7	1.0	1.0
8	1.0	1.0
9	1.0	1.0
10	1.0	0.0
11	1.0	1.0
12	1.0	0.0
13	1.0	0.0

```
[12]: %time utils.metabolicNetwork.FromGeneList('ECOLI', ['lacZ', 'lacA', 'lacY',
↳ 'araF', 'araG', 'araH'], fmt = 'genes', precalculated = df_genes)
```

CPU times: user 81.6 ms, sys: 50.6 ms, total: 132 ms
Wall time: 1.42 s

[12]:	GENE OR COMPLEX	ENZYME LOCATION	REACTION \
0	araF	periplasmic space	ABC-2-RXN
1	araG	inner membrane	ABC-2-RXN
2	araG	cytosol	ABC-2-RXN
3	araH	inner membrane	ABC-2-RXN
4	lacA	cytosol	GALACTOACETYLTRAN-RXN

5	lacY	inner membrane	TRANS-RXN-24
6	lacY	inner membrane	TRANS-RXN-94
7	lacY	inner membrane	RXN0-7215
8	lacY	inner membrane	RXN0-7217
9	lacY	inner membrane	RXN-17755
10	lacZ	cytosol	BETAGALACTOSID-RXN
11	lacZ	cytosol	RXN0-5363
12	lacZ	cytosol	RXN-17726
13	lacZ	cytosol	RXN0-7219

	SUBSTRATES	PRODUCTS \
0	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
1	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
2	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
3	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
4	Beta-D-Galactosides,ACETYL-COA	6-Acetyl-Beta-D-Galactosides,CO-A
5	PROTON,Alpha-lactose	PROTON,Alpha-lactose
6	PROTON,MELIBIOSE	PROTON,MELIBIOSE
7	PROTON,CPD-3561	PROTON,CPD-3561
8	PROTON,CPD-3785	PROTON,CPD-3785
9	PROTON,CPD-3801	PROTON,CPD-3801
10	CPD-15972,WATER	GALACTOSE,Glucopyranose
11	Alpha-lactose	ALLOLACTOSE
12	CPD-3561,WATER	GALACTOSE,Fructofuranose
13	CPD-3785,WATER	GALACTOSE,D-ARABINOSE

	FWD_RATE	RVS_RATE
0	1.0	0.0
1	1.0	0.0
2	1.0	0.0
3	1.0	0.0
4	1.0	0.0
5	1.0	1.0
6	1.0	1.0
7	1.0	1.0
8	1.0	1.0
9	1.0	1.0
10	1.0	0.0
11	1.0	1.0
12	1.0	0.0
13	1.0	0.0

```
[13]: %time utils.metabolicNetwork.FromGeneList('ECOLI', ['lacZ', 'lacA', 'lacY',
↪ 'araF', 'araG', 'araH'], fmt = 'complex', precalculated = df_genes)
```

CPU times: user 137 ms, sys: 106 ms, total: 243 ms
Wall time: 2.38 s

[13]: GENE OR COMPLEX \

0	[araG,araG,araH,araH,araF]
1	[araG,araG,araH,araH,araF]
2	[araG,araG,araH,araH,araF]
3	[araG,araG,araH,araH,araF]
4	[araG,araG,araH,araH,araF]
5	[araG,araG,araH,araH,araF]
6	[lacA,lacA,lacA]
7	lacY
8	lacY
9	lacY
10	lacY
11	lacY
12	[lacZ,lacZ,lacZ,lacZ]
13	[lacZ,lacZ,lacZ,lacZ]
14	[lacZ,lacZ,lacZ,lacZ]
15	[lacZ,lacZ,lacZ,lacZ]

	ENZYME LOCATION	REACTION \
0	[inner membrane,inner membrane,inner membrane,...	ABC-2-RXN
1	[cytosol,cytosol,inner membrane,inner membrane...	ABC-2-RXN
2	[inner membrane,inner membrane,inner membrane,...	ABC-2-RXN
3	[cytosol,cytosol,inner membrane,inner membrane...	ABC-2-RXN
4	[inner membrane,inner membrane,inner membrane,...	ABC-2-RXN
5	[cytosol,cytosol,inner membrane,inner membrane...	ABC-2-RXN
6	[cytosol,cytosol,cytosol]	GALACTOACETYLTRAN-RXN
7	inner membrane	TRANS-RXN-24
8	inner membrane	TRANS-RXN-94
9	inner membrane	RXN0-7215
10	inner membrane	RXN0-7217
11	inner membrane	RXN-17755
12	[cytosol,cytosol,cytosol,cytosol]	BETAGALACTOSID-RXN
13	[cytosol,cytosol,cytosol,cytosol]	RXN0-5363
14	[cytosol,cytosol,cytosol,cytosol]	RXN-17726
15	[cytosol,cytosol,cytosol,cytosol]	RXN0-7219

	SUBSTRATES	PRODUCTS \
0	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
1	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
2	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
3	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
4	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
5	WATER,ATP,L-ARABINOSE	Pi,ADP,PROTON,L-ARABINOSE
6	Beta-D-Galactosides,ACETYL-COA	6-Acetyl-Beta-D-Galactosides,CO-A
7	PROTON,Alpha-lactose	PROTON,Alpha-lactose
8	PROTON,MELIBIOSE	PROTON,MELIBIOSE
9	PROTON,CPD-3561	PROTON,CPD-3561

10	PROTON,CPD-3785	PROTON,CPD-3785
11	PROTON,CPD-3801	PROTON,CPD-3801
12	CPD-15972,WATER	GALACTOSE,Glucopyranose
13	Alpha-lactose	ALLOLACTOSE
14	CPD-3561,WATER	GALACTOSE,Fructofuranose
15	CPD-3785,WATER	GALACTOSE,D-ARABINOSE

	FWD_RATE	RVS_RATE
0	1.0	0.0
1	1.0	0.0
2	1.0	0.0
3	1.0	0.0
4	1.0	0.0
5	1.0	0.0
6	1.0	0.0
7	1.0	1.0
8	1.0	1.0
9	1.0	1.0
10	1.0	1.0
11	1.0	1.0
12	1.0	0.0
13	1.0	1.0
14	1.0	0.0
15	1.0	0.0

The output is a pandas dataframe that could be exported with `network.to_csv(path)`, or in a two-columns format that Cytoscape could interpret as a network. The `utils.metabolicNetwork.expand_network` function reorders and exports the dataframe as a text file (in this case to `./tutorial.txt`).

```
[14]: %time network = utils.metabolicNetwork.FromGeneList('ECOLI', ['lacZ', 'lacA', 'lacY'], fmt = 'genes', precalculated = df_genes)
      ↪ %time utils.metabolicNetwork.expand_network(network, './lactose-metabolism-cytoscape-v1.txt')
```

```
CPU times: user 53 ms, sys: 40.8 ms, total: 93.7 ms
Wall time: 970 ms
CPU times: user 916  $\mu$ s, sys: 740  $\mu$ s, total: 1.66 ms
Wall time: 1.41 ms
```

The following image was prepared from the `tutorial.txt` file, and you could reproduce it with Cytoscape: 1. Click on the **Import Network from File System** icon or click on **File -> Import -> Network from File....** 2. Navigate to the file and click on **Open**. 3. **SOURCE**, **TARGET**, and **EDGE ATTRIBUTE** are OK, but the 4th columns must be the **SOURCE NODE ATTRIBUTE** and the 5th column the **TARGET NODE ATTRIBUTE**. Click on the header and change it to the correct attribute. The attributes will help later to filter and to add format to nodes and edges. 4. Click on **Filter** (on the right), then on the **+** icon and finally on **Column Filter**: 1. On the selector, click on **Edge**: **EDGE ATTRIBUTE** and change **contains** to **is**. 1. Write **NO REVERSIBLE**

that will select edges that correspond to irreversible reactions. Click on **Style**, then **Edge** (in the bottom), and click on the 3rd column to bypass the format of the **Target Arrow Shape** and select your favorite arrow shape. 2. Write **REVERSIBLE** and bypass the format of the **Source Arrow Shape AND Target Arrow Shape**, and select your favorite arrow shape. 2. On the selector, click on **Node: SOURCE_NODE_ATTRIBUTE**: 1. Write **RXN** that will select nodes encoding the reactions. Click on **Style**, then on **Node** and bypass the **Fill Color**. In the new window, you could set-up the color, e.g. **#00AA50** 2. Write **GENE_PROD** that will select nodes encoding the gene name, protein name, or the enzyme name. Click on **Style**, then on **Node** and bypass the **Fill Color**. In the new window, you could set-up the color, e.g. **#CC0033** 3. Write **MET** that will select nodes encoding substrate metabolites. Click on **Style**, then on **Node** and bypass the **Fill Color**. In the new window, you could set-up the color, e.g. **#00ABDD**. Also, set a shape for nodes, to differentiate substrates from products. 3. On the selector, click on **Node: TARGET_NODE_ATTRIBUTE**: 1. Write **MET** that will select nodes encoding product metabolites. Click on **Style**, then on **Node** and bypass the **Fill Color**. In the new window, you could set-up the color, e.g. **#00ABDD**

The result will be similar to

If we inspect the network, we could highlight four things: 1. The **lacA** reaction is disconnect from the network formed by the **lacZ** and **lacY** reactions; 2. The **lacY** reactions do not inform the metabolite compartment, so substrates and products refer to the same node; 3. The utilization of *internal codes* for certain compounds; and 4. The impossibility of alpha-lactose degradation into glucose (glucopyranose) and galactose.

We could retrieve information from the reaction and metabolite nodes with the `utils.getData()` function:

```
[15]: # utils.get_data('ECOLI', 'CPD-3561')
      %time utils.getData('ECOLI', 'CPD-3561')['names'] # for simplicity of the output
```

```
CPU times: user 421 µs, sys: 1.34 ms, total: 1.77 ms
Wall time: 13.2 ms
```

```
[15]: ['lactulose', 'CEPHULAC (TN)']
```

We observe that **CPD-3561** correspond to **lactulose**. Therefore, the **EcoCyc** database informs that **LacY** is able to incorporate **alpha-lactose**, **melibiose**, **lactulose (CPD-3561)**, **3-O-galactosylarabinose (CPD-3785)**, and **melibionate (CPD-3801)** into the cell cytoplasm. Interestingly, the common synthetic activator **IPTG** (o-nitrophenyl- -galactoside) is mentioned in the description for the lactose transporter (<https://biocyc.org/gene?orgid=ECOLI&id=EG10526#>), but there is no inclusion of the **IPTG** transport in the reactions of **LacY**.

```
[16]: # utils.get_data('ECOLI', 'CPD-15972')
      %time utils.getData('ECOLI', 'CPD-15972')['names'] # for simplicity of the
      ↪output
```

```
CPU times: user 1.63 ms, sys: 0 ns, total: 1.63 ms
Wall time: 12.5 ms
```

```
[16]: ['lactose',
      '&beta;-D-galactopyranosyl-(1&rarr;4)-D-glucopyranose',
      'D-lactose',
      '&beta;-D-Galp-(1&rarr;4)-D-Glcp']
```

Next, EcoCyc informs that LacZ could metabolize lactose (CPD-15972) into galactose and glucopyranose, lactulose into β -galactose and fructofuranose, and 3-O-galactosylarabinose into β -galactose and arabinose. The difference in metabolite names between the lacY and lacZ reactions is enough to make a Section ???. Specifically, the name lactose refers to the two anomers α - and β -lactose because of EcoCyc uses generic names when enzymes show no stereoselectivity. In addition, galactose has also an anomeric center, but the database refers to it as β -D-galactose without indication the enzyme could produce α -galactose from α -lactose.

```
[17]: %time print(utils.getData('ECOLI', 'GALACTOSE')['common_name'])
print()
%time print(utils.getData('ECOLI', 'GALACTOSE')['synonyms'])
```

```
&beta;-D-galactopyranose
```

```
CPU times: user 1.18 ms, sys: 949  $\mu$ s, total: 2.13 ms
```

```
Wall time: 22.7 ms
```

```
['&beta;-D-galactopyranose', '&beta;-D-galactose', 'cerebrose',
 '6-(hydroxymethyl)tetrahydropyran-2,3,4,5-tetraol']
```

```
CPU times: user 1.98 ms, sys: 0 ns, total: 1.98 ms
```

```
Wall time: 22.7 ms
```

The advantage of the procedure is the ability to modify the data programatically using python functions (<https://pandas.pydata.org/>) or export the data and manipulate it using a text processor or a spreadsheet software. For routinary changes, we included utility functions to make batch manipulation to the data:

```
[18]: # Transport reactions: Add compartments to substrates and/or products
%time network = utils.metabolicNetwork.setTransport(network, geneLst = ␣
␣['lacY'], fromLst = ['PER'], toLst = ['CYT'])
network
```

```
CPU times: user 7.39 ms, sys: 612  $\mu$ s, total: 8 ms
```

```
Wall time: 7.42 ms
```

```
[18]:  GENE OR COMPLEX ENZYME LOCATION      REACTION \
0      lacA      cytosol  GALACTOACETYLTRAN-RXN
1      lacY  inner membrane      TRANS-RXN-24
2      lacY  inner membrane      TRANS-RXN-94
3      lacY  inner membrane      RXN0-7215
4      lacY  inner membrane      RXN0-7217
5      lacY  inner membrane      RXN-17755
6      lacZ      cytosol  BETAGALACTOSID-RXN
7      lacZ      cytosol      RXN0-5363
```

8	lacZ	cytosol	RXN-17726
9	lacZ	cytosol	RXN0-7219

	SUBSTRATES	PRODUCTS \
0	Beta-D-Galactosides, ACETYL-COA	6-Acetyl-Beta-D-Galactosides, CO-A
1	PER-PROTON, PER-Alpha-lactose	PROTON, Alpha-lactose
2	PER-PROTON, PER-MELIBIOSE	PROTON, MELIBIOSE
3	PER-PROTON, PER-CPD-3561	PROTON, CPD-3561
4	PER-PROTON, PER-CPD-3785	PROTON, CPD-3785
5	PER-PROTON, PER-CPD-3801	PROTON, CPD-3801
6	CPD-15972, WATER	GALACTOSE, Glucopyranose
7	Alpha-lactose	ALLOLACTOSE
8	CPD-3561, WATER	GALACTOSE, Fructofuranose
9	CPD-3785, WATER	GALACTOSE, D-ARABINOSE

	FWD_RATE	RVS_RATE
0	1.0	0.0
1	1.0	1.0
2	1.0	1.0
3	1.0	1.0
4	1.0	1.0
5	1.0	1.0
6	1.0	0.0
7	1.0	1.0
8	1.0	0.0
9	1.0	0.0

Note: By default, Atlas interprets the default location of monomers as cytoplasmatic. When setting the location to CYT, the `setTransport()` function will delete a previous compartment or append nothing to the name of the monomer.

```
[19]: # Irreversibility of reactions per gene: Change the Reverse Rate to zero
%time network = utils.metabolicNetwork.setIrreversibility(network, geneLst = ['lacY', 'lacA'])
network
```

CPU times: user 197 μ s, sys: 3.35 ms, total: 3.55 ms
Wall time: 3.22 ms

	GENE OR COMPLEX	ENZYME LOCATION	REACTION \
0	lacA	cytosol	GALACTOACETYLTRAN-RXN
1	lacY	inner membrane	TRANS-RXN-24
2	lacY	inner membrane	TRANS-RXN-94
3	lacY	inner membrane	RXN0-7215
4	lacY	inner membrane	RXN0-7217
5	lacY	inner membrane	RXN-17755
6	lacZ	cytosol	BETAGALACTOSID-RXN

7	lacZ	cytosol	RXN0-5363
8	lacZ	cytosol	RXN-17726
9	lacZ	cytosol	RXN0-7219

	SUBSTRATES	PRODUCTS \
0	Beta-D-Galactosides, ACETYL-COA	6-Acetyl-Beta-D-Galactosides, CO-A
1	PER-PROTON, PER-Alpha-lactose	PROTON, Alpha-lactose
2	PER-PROTON, PER-MELIBIOSE	PROTON, MELIBIOSE
3	PER-PROTON, PER-CPD-3561	PROTON, CPD-3561
4	PER-PROTON, PER-CPD-3785	PROTON, CPD-3785
5	PER-PROTON, PER-CPD-3801	PROTON, CPD-3801
6	CPD-15972, WATER	GALACTOSE, Glucopyranose
7	Alpha-lactose	ALLOLACTOSE
8	CPD-3561, WATER	GALACTOSE, Fructofuranose
9	CPD-3785, WATER	GALACTOSE, D-ARABINOSE

	FWD_RATE	RVS_RATE
0	1.0	0.0
1	1.0	0.0
2	1.0	0.0
3	1.0	0.0
4	1.0	0.0
5	1.0	0.0
6	1.0	0.0
7	1.0	1.0
8	1.0	0.0
9	1.0	0.0

```
[20]: # Irreversibility of reactions per reaction. The beta-galactosidase has also
      ↪ isomerase activity (reversible reaction)
%time network = utils.metabolicNetwork.setIrreversibility(network, rxnLst =
      ↪ ['BETAGALACTOSID-RXN', 'RXN-17726', 'RXN0-7219'])
network
```

CPU times: user 3.87 ms, sys: 0 ns, total: 3.87 ms

Wall time: 3.34 ms

[20]:	GENE OR COMPLEX	ENZYME	LOCATION	REACTION \
0	lacA		cytosol	GALACTOACETYLTRAN-RXN
1	lacY	inner	membrane	TRANS-RXN-24
2	lacY	inner	membrane	TRANS-RXN-94
3	lacY	inner	membrane	RXN0-7215
4	lacY	inner	membrane	RXN0-7217
5	lacY	inner	membrane	RXN-17755
6	lacZ		cytosol	BETAGALACTOSID-RXN
7	lacZ		cytosol	RXN0-5363
8	lacZ		cytosol	RXN-17726

9	lacZ	cytosol	RXN0-7219
	SUBSTRATES		PRODUCTS \
0	Beta-D-Galactosides, ACETYL-COA		6-Acetyl-Beta-D-Galactosides, CO-A
1	PER-PROTON, PER-Alpha-lactose		PROTON, Alpha-lactose
2	PER-PROTON, PER-MELIBIOSE		PROTON, MELIBIOSE
3	PER-PROTON, PER-CPD-3561		PROTON, CPD-3561
4	PER-PROTON, PER-CPD-3785		PROTON, CPD-3785
5	PER-PROTON, PER-CPD-3801		PROTON, CPD-3801
6	CPD-15972, WATER		GALACTOSE, Glucopyranose
7	Alpha-lactose		ALLOLACTOSE
8	CPD-3561, WATER		GALACTOSE, Fructofuranose
9	CPD-3785, WATER		GALACTOSE, D-ARABINOSE

	FWD_RATE	RVS_RATE
0	1.0	0.0
1	1.0	0.0
2	1.0	0.0
3	1.0	0.0
4	1.0	0.0
5	1.0	0.0
6	1.0	0.0
7	1.0	1.0
8	1.0	0.0
9	1.0	0.0

```
[21]: # Compartment of reactions. The lacY gene is a protein located to the inner
      ↪ membrane of E.coli
%time network = utils.metabolicNetwork.setEnzymeLocation(network, geneLst =
      ↪ ['lacY'], compartmentLst = ['iMEM'])
network
```

CPU times: user 1.41 ms, sys: 0 ns, total: 1.41 ms
Wall time: 1.26 ms

[21]:	GENE OR COMPLEX	ENZYME LOCATION	REACTION \
0	lacA	cytosol	GALACTOACETYLTRAN-RXN
1	lacY	inner membrane	TRANS-RXN-24
2	lacY	inner membrane	TRANS-RXN-94
3	lacY	inner membrane	RXN0-7215
4	lacY	inner membrane	RXN0-7217
5	lacY	inner membrane	RXN-17755
6	lacZ	cytosol	BETAGALACTOSID-RXN
7	lacZ	cytosol	RXN0-5363
8	lacZ	cytosol	RXN-17726
9	lacZ	cytosol	RXN0-7219

	SUBSTRATES	PRODUCTS \
0	Beta-D-Galactosides, ACETYL-COA	6-Acetyl-Beta-D-Galactosides, CO-A
1	PER-PROTON, PER-Alpha-lactose	PROTON, Alpha-lactose
2	PER-PROTON, PER-MELIBIOSE	PROTON, MELIBIOSE
3	PER-PROTON, PER-CPD-3561	PROTON, CPD-3561
4	PER-PROTON, PER-CPD-3785	PROTON, CPD-3785
5	PER-PROTON, PER-CPD-3801	PROTON, CPD-3801
6	CPD-15972, WATER	GALACTOSE, Glucopyranose
7	Alpha-lactose	ALLOLACTOSE
8	CPD-3561, WATER	GALACTOSE, Fructofuranose
9	CPD-3785, WATER	GALACTOSE, D-ARABINOSE

	FWD_RATE	RVS_RATE
0	1.0	0.0
1	1.0	0.0
2	1.0	0.0
3	1.0	0.0
4	1.0	0.0
5	1.0	0.0
6	1.0	0.0
7	1.0	1.0
8	1.0	0.0
9	1.0	0.0

```
[22]: %time utils.metabolicNetwork.expand_network(network, './
↳ lactose-metabolism-cytoscape-v2.txt')
```

CPU times: user 962 µs, sys: 772 µs, total: 1.73 ms
Wall time: 1.34 ms

```
[23]: %time model = atlas.construct_model_from_metabolic_network(network, verbose =
↳ False) # verbose = True will print the pySB functions needed to reproduce
↳ the model
model
```

CPU times: user 459 ms, sys: 972 µs, total: 460 ms
Wall time: 458 ms

```
[23]: <Model 'atlas_rbm.construct_model_from_metabolic_network' (monomers: 2, rules:
10, parameters: 40, expressions: 0, compartments: 0) at 0x7f7d69e88e80>
```

```
[24]: utils.analyseConnectivity(model, '/opt/git-repositories/KaSim.Kappa-Dev/KaSa')
# Do not worry, later we will set the initial condition.
# Due to the multiple instances, we create only metabolites at cytoplasm and
↳ the transport reactions will fail.
```

There are some non applicable rules:
rule TRANS_RXN_24 (File "_U076DMIMNC.kappa", line 47, characters 15-282:) will

never be applied.
 rule TRANS_RXN_24_rev (File "_U076DMIMNC.kappa", line 48, characters 19-286:) will never be applied.
 rule TRANS_RXN_94 (File "_U076DMIMNC.kappa", line 49, characters 15-274:) will never be applied.
 rule TRANS_RXN_94_rev (File "_U076DMIMNC.kappa", line 50, characters 19-278:) will never be applied.
 rule RXNO_7215 (File "_U076DMIMNC.kappa", line 51, characters 12-266:) will never be applied.
 rule RXNO_7215_rev (File "_U076DMIMNC.kappa", line 52, characters 16-270:) will never be applied.
 rule RXNO_7217 (File "_U076DMIMNC.kappa", line 53, characters 12-266:) will never be applied.
 rule RXNO_7217_rev (File "_U076DMIMNC.kappa", line 54, characters 16-270:) will never be applied.
 rule RXN_17755 (File "_U076DMIMNC.kappa", line 55, characters 12-266:) will never be applied.
 rule RXN_17755_rev (File "_U076DMIMNC.kappa", line 56, characters 16-270:) will never be applied.

Every monomer and complex of monomers may occur in the model.

Note: Even with an initial condition of zero, KaSA will report reactions as reachable because it analyzes the connectivity of the network. Unreachable rules or agents will those that: first, monomers are not instantiated (missing `%init:` statements in `kappa`), and two, *Rules* not producing the monomers in the required state (e.g. phosphorylation). For sake of completeness, we produced `kappa` files that fail the connectivity analysis:

```
[25]: utils.analyseConnectivity('fail_rules.kappa', path = '/opt/git-repositories/
      ↪KaSim.Kappa-Dev/KaSa')
      # Missing instantiation of the lacA trimer, so impossibility to catalyze its
      ↪reactions
```

There are some non applicable rules:

rule GALACTOACETYLTRAN_RXN_galactose (File "fail_rules.kappa", line 187, characters 34-534:) will never be applied.
 rule GALACTOACETYLTRAN_RXN_galactose_rev (File "fail_rules.kappa", line 188, characters 38-538:) will never be applied.

Every monomer and complex of monomers may occur in the model.

```
[26]: utils.analyseConnectivity('fail_agents.kappa', path = '/opt/git-repositories/
      ↪KaSim.Kappa-Dev/KaSa')
      # Missing instantiation of Acetyl-CoA and CoA, substrate and product of the
      ↪lacA trimer.
      # It does not matter if the rate of the reaction is zero.
      # (by design, Atlas writes reversible reactions even if the rate for the
      ↪reverse reaction is zero)
```

There are some non applicable rules:

rule GALACTOACETYLTRAN_RXN_galactose (File "fail_agents.kappa", line 187, characters 34-534:) will never be applied.

rule GALACTOACETYLTRAN_RXN_galactose_rev (File "fail_agents.kappa", line 188, characters 38-538:) will never be applied.

Every monomer and complex of monomers may occur in the model.

To simulate, we need to set the initial condition:

```
[27]: # initial condition
# for metabolites
simulation.set_initial.met(model, 'Alpha_lactose', 'per', 100)
# Message of "Initial not found" explained: By default, Atlas creates initials
↳for cytosolic metabolites, proteins, and complexes (DNA and RNAs are always
↳cytosolic)
simulation.set_initial.met(model, 'PROTON', 'per', 100) # required for lactose
↳transport
simulation.set_initial.met(model, 'WATER', 'cyt', 100) # required for lactose
↳hydrolysis

# for proteins
simulation.set_initial.prot(model, 'lacY', 'imem', 1) # required for lactose
↳transport.
simulation.set_initial.prot(model, 'lacZ', 'cyt', 1) # required for lactose
↳isomerization
```

Initial not found. Creating Initial...

Initial not found. Creating Initial...

Initial not found. Creating Initial...

```
[27]: <Model 'atlas_rbm.construct_model_from_metabolic_network' (monomers: 2, rules:
10, parameters: 43, expressions: 0, compartments: 0) at 0x7f7d69e88e80>
```

```
[28]: utils.analyseConnectivity(model, '/opt/git-repositories/KaSim.Kappa-Dev/KaSa')
```

Every rule may be applied.

Every monomer and complex of monomers may occur in the model.

Once the model has a suitable initial condition, the user could export it to a variety of formats or simulate directly inside the notebook

```
[29]: # export to
%time export.to_sbml(model, 'export-to-sbml.sbml')
%time export.to_matlab(model, 'export-to-matlab.m')
%time export.to_mathematica(model, 'export-to-mathematica.wl')
%time export.to_potterswheel(model, 'export-to-potterswheel.m')
%time export.to_bngl(model, 'export-to-bngl.bngl')
%time export.to_bngnet(model, 'export-to-bngnet.net')
```

```
%time export.to_kappa(model, 'export-to-kappa.kappa')
%time export.to_python(model, 'export-to-python.py')
%time export.to_pysb(model, 'export-to-pysb.py')
# %time export.to_stochkit(model, 'export-to-stochkit.xml') # pySB error
%time export.to_json(model, 'export-to-json.json')
```

```
CPU times: user 466 ms, sys: 70.5 ms, total: 536 ms
Wall time: 1.34 s
CPU times: user 81 ms, sys: 2.44 ms, total: 83.4 ms
Wall time: 87.9 ms
CPU times: user 173 ms, sys: 1.56 ms, total: 174 ms
Wall time: 179 ms
CPU times: user 157 ms, sys: 0 ns, total: 157 ms
Wall time: 162 ms
CPU times: user 2.79 ms, sys: 0 ns, total: 2.79 ms
Wall time: 4.18 ms
CPU times: user 9.26 ms, sys: 3.17 ms, total: 12.4 ms
Wall time: 518 ms
CPU times: user 4.11 ms, sys: 0 ns, total: 4.11 ms
Wall time: 3.91 ms
CPU times: user 193 ms, sys: 4.57 ms, total: 198 ms
Wall time: 202 ms
CPU times: user 3.67 ms, sys: 0 ns, total: 3.67 ms
Wall time: 5.04 ms
CPU times: user 3.03 ms, sys: 0 ns, total: 3.03 ms
Wall time: 40.7 ms
```

```
[30]: # simulation
bng = '/opt/git-repositories/bionetgen.RuleWorld/bng2/'
kasim = '/opt/git-repositories/KaSim4.Kappa-Dev/'
cupsoda = '/opt/git-repositories/cupSODA.aresio/'
stochkit = '/opt/git-repositories/StochKit.StochSS' # not the bin folder

%time data0 = simulation.scipy(model, start = 0, finish = 10, points = 2000)
%time data1 = simulation.cupsoda(model, start = 0, finish = 10, points = 2000, ↵
↪ path = cupsoda) # only if you have a GPU NVIDIA; comment if not.
%time data2 = simulation.bngODE(model, start = 0, finish = 10, points = 2000, ↵
↪ path = bng)
%time data3 = simulation.bngSSA(model, start = 0, finish = 10, points = 2000, ↵
↪ n_runs = 20, path = bng)
# %time data4 = simulation.bngPLA(model, start = 0, finish = 10, points = 2000, ↵
↪ n_runs = 20, path = bng) # requires refinement of the model
%time data5 = simulation.bngNF(model, start = 0, finish = 10, points = 2000, ↵
↪ n_runs = 20, path = bng)
%time data6 = simulation.kasim(model, start = 0, finish = 10, points = 2000, ↵
↪ n_runs = 20, path = kasim)
```

```
# %time data7 = simulation.stochkit(model, start = 0, finish = 10, points = 2000, n_runs = 20, path = stochkit) # pySB error
```

```
CPU times: user 3.5 s, sys: 1.15 s, total: 4.65 s
Wall time: 3.53 s
CPU times: user 1.44 s, sys: 30.9 ms, total: 1.47 s
Wall time: 2.38 s
CPU times: user 1.62 s, sys: 13.7 ms, total: 1.64 s
Wall time: 2.36 s
CPU times: user 4.07 s, sys: 92.1 ms, total: 4.16 s
Wall time: 6.6 s
CPU times: user 3.11 s, sys: 120 ms, total: 3.23 s
Wall time: 5.37 s
CPU times: user 2.55 s, sys: 177 ms, total: 2.73 s
Wall time: 4.24 s
```

Note: The ODE and SSA simulations require the execution of the network generation, while KaSim and NFsim are network-free simulators.

Finally, we plot the simulation results. The result of the `simulation.scipy()`, `simulation.scipy()` function is a pandas dataframe. In the case of stochastic simulations (SSA, KaSim, NFsim, Stochkit), the function returns a dictionary with a list of dataframes for each simulations (`sims` key), a dataframe with the average (`avrg` key) and a dataframe with the standard deviation (`stdv` key) of those simulations. Currently, we included three kind of plots, although the user could access the dataframes and plot directly with methods in the seaborn package (<https://seaborn.pydata.org/>), in the pandas package (<https://pandas.pydata.org/pandas-docs/stable/reference/api/pandas.DataFrame.plot.html>), or with matplotlib (<https://matplotlib.org/>).

```
[31]: import seaborn
import matplotlib.pyplot as plt

palette = seaborn.color_palette('colorblind')

for kind in ['scatter', 'plot']:
    # first plot, periplasmic concentration
    fig, ax = plt.subplots(1, 2, figsize = (4*2, 3*1), dpi = 100)
    simulation.plot.metabolite(data3['avrg'], 'Alpha_lactose', 'per', ax = ax[0],
    **{'kind' : kind},
    plt_kws = {'s' : 2, 'color' : palette[0], 'label' : r'$\alpha$-lactose', 'alpha' : .5})

    # second plot, cytoplasmic concentration
    simulation.plot.metabolite(data3['avrg'], 'Alpha_lactose', 'cyt', ax = ax[1],
    **{'kind' : kind},
    plt_kws = {'s' : 2, 'color' : palette[1], 'label' : r'$\alpha$-lactose', 'alpha' : .5})
```

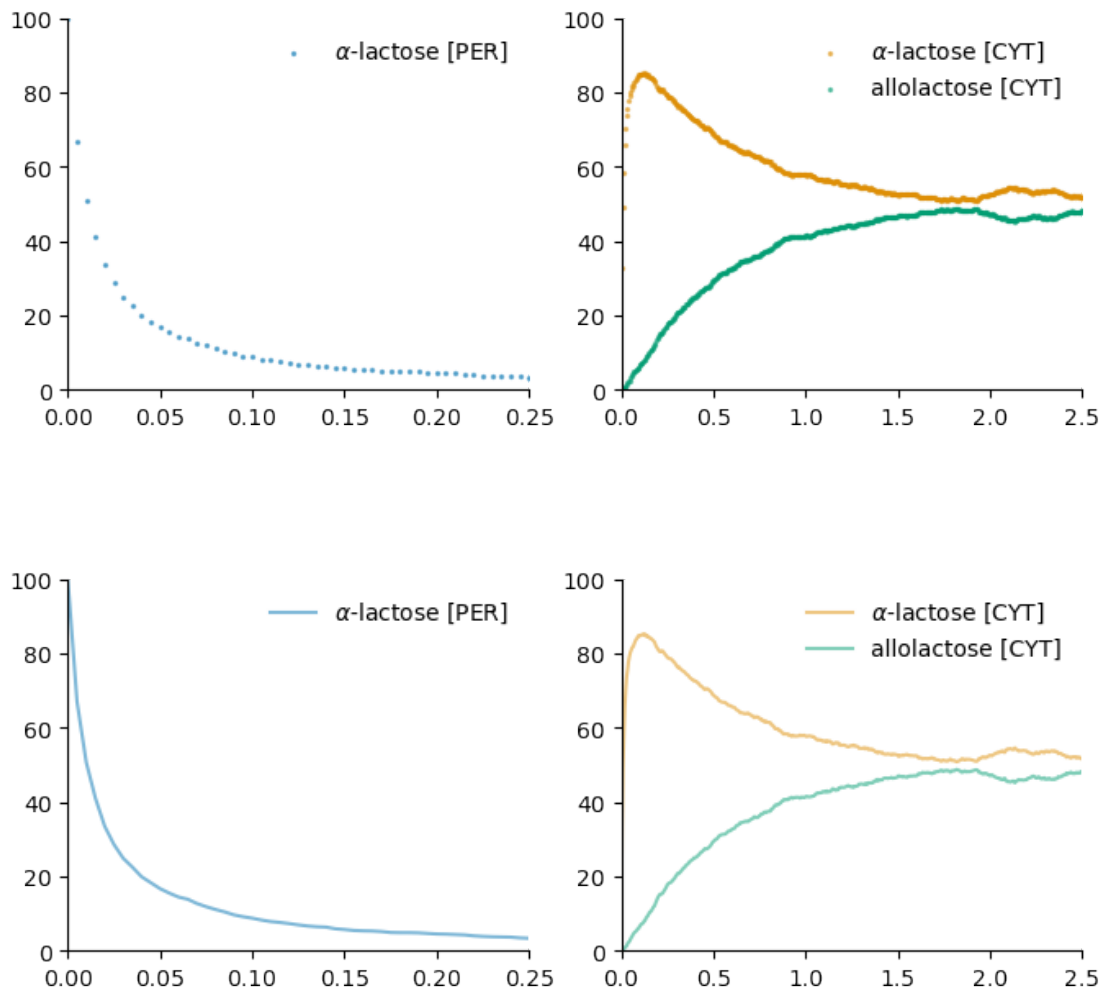
```

# second plot, cytoplasmic concentration
simulation.plot.metabolite(data3['avrg'], 'ALLOLACTOSE', 'cyt', ax = ax[1],
↳**{'kind' : kind},
    plt_kws = {'s' : 2, 'color' : palette[2], 'label' : r'allolactose_
↳[CYT]', 'alpha' : .5})

ax[0].set_xlim((0,.25))
ax[1].set_xlim((0,2.5))
ax[0].set_ylim((0,100))
ax[1].set_ylim((0,100))

seaborn.despine()

```



```

[32]: import seaborn
import matplotlib.pyplot as plt

```

```

palette = seaborn.color_palette('colorblind')

# first plot, periplasmic concentration
fig, ax = plt.subplots(1, 2, figsize = (4*2, 3*1), dpi = 100)
simulation.plot.metabolite(data3, 'Alpha_lactose', 'per', ax = ax[0], **{'kind': '
↳: 'fill_between', 'weight' : .5},
    plt_kws = {'s' : 2, 'color' : palette[0], 'label' : r'$\alpha$-lactose
↳[PER]', 'alpha' : .5})

# second plot, cytoplasmic concentration
simulation.plot.metabolite(data3, 'Alpha_lactose', 'cyt', ax = ax[1], **{'kind': '
↳: 'fill_between', 'weight' : .5},
    plt_kws = {'s' : 2, 'color' : palette[1], 'label' : r'$\alpha$-lactose
↳[CYT]', 'alpha' : .5})

# second plot, cytoplasmic concentration
simulation.plot.metabolite(data3, 'ALLOLACTOSE', 'cyt', ax = ax[1], **{'kind' : '
↳: 'fill_between', 'weight' : .5},
    plt_kws = {'s' : 2, 'color' : palette[2], 'label' : r'allolactose [CYT]',
↳: 'alpha' : .5})

# first plot, periplasmic concentration
simulation.plot.metabolite(data0, 'Alpha_lactose', 'per', ax = ax[0], **{'kind': '
↳: kind},
    plt_kws = {'s' : 2, 'color' : palette[0], 'label' : r'ODE'})

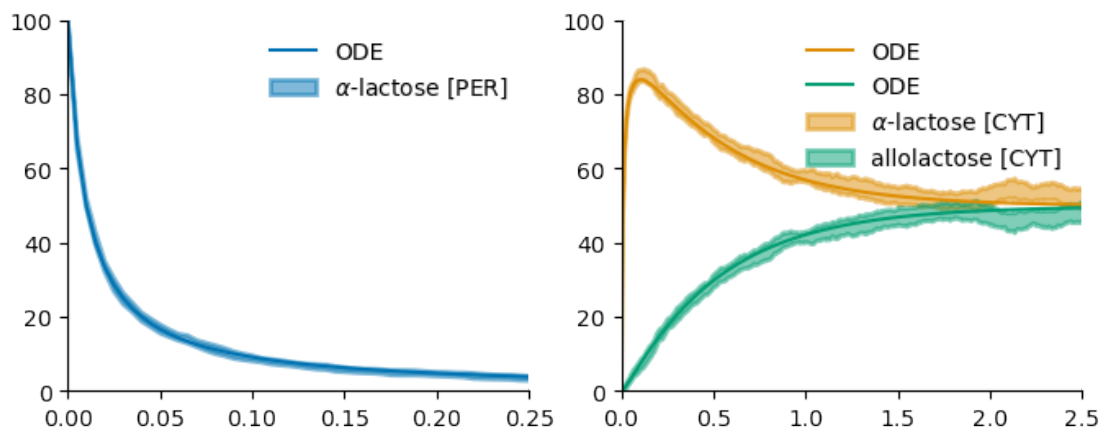
# second plot, cytoplasmic concentration
simulation.plot.metabolite(data0, 'Alpha_lactose', 'cyt', ax = ax[1], **{'kind': '
↳: kind},
    plt_kws = {'s' : 2, 'color' : palette[1], 'label' : r'ODE'})

# second plot, cytoplasmic concentration
simulation.plot.metabolite(data0, 'ALLOLACTOSE', 'cyt', ax = ax[1], **{'kind' : '
↳: kind},
    plt_kws = {'s' : 2, 'color' : palette[2], 'label' : r'ODE'})

ax[0].set_xlim((0,.25))
ax[1].set_xlim((0,2.5))
ax[0].set_ylim((0,100))
ax[1].set_ylim((0,100))

seaborn.despine()

```

As expected, the degradation of lactose into glucose and galactose is impossible due to a Section ???. The disconnected pathway will be manually corrected and explained in a second tutorial. Also, we must note that we considered the enzymatic reactions are performed by the monomers (e.g. one monomer of LacZ catalyze one reaction), although biochemical information informs the monomers are catalytically active only when the complex is assembled (e.g. one LacZ tetramer catalyzes four reactions). This will be considered next, when we will model protein-protein interactions.

1.6 Modeling protein-protein interactions

We could consult the stoichiometry of complexes with PythonCyc, and vice versa, querying which complexes are formed by a certain gene product (although indirectly):

```
[33]: import pythoncyc
print(pythoncyc.select_organism('ECOLI').
      ↪monomers_of_protein('BETAGALACTOSID-CPLX'))
# we obtained the code "BETAGALACTOSID-CPLX" with the gathering of the
      ↪metabolic data in "product" format.
print(pythoncyc.select_organism('ECOLI').
      ↪monomers_of_protein('GALACTOACETYLTRAN-CPLX'))
# Similarly, the code "GALACTOACETYLTRAN-CPLX".

print(pythoncyc.select_organism('ECOLI').
      ↪genes_of_protein('BETAGALACTOSID-CPLX'))
print(pythoncyc.select_organism('ECOLI').
      ↪genes_of_protein('GALACTOACETYLTRAN-CPLX'))

print(pythoncyc.select_organism('ECOLI').all_products_of_gene('EG10527'))
print(pythoncyc.select_organism('ECOLI').all_products_of_gene('EG10524'))
```

```
[['|BETAGALACTOSID-MONOMER|'], [4]]
[['|GALACTOACETYLTRAN-MONOMER|'], [3]]
[['|EG10527|']]
```

```
['|EG10524|']
['|BETAGALACTOSID-MONOMER|', '|BETAGALACTOSID-CPLX|']
['|GALACTOACETYLTRAN-MONOMER|', '|GALACTOACETYLTRAN-CPLX|']
```

We included the function `utils.interactionNetwork.FromGeneList()` to write possible mechanism of assembly from the stoichiometry data of complexes.

```
[34]: %time network = utils.interactionNetwork.FromGeneList('ECOLI', genes = ['lacZ',
↳ 'lacA', 'lacY'], precalculated = df_genes)
network
```

CPU times: user 8.44 ms, sys: 4.21 ms, total: 12.7 ms

Wall time: 60.3 ms

```
[34]:
```

	SOURCE	TARGET	LOCATION	FWD_RATE	\
0	lacA	lacA	[cytosol,cytosol]	1.0	
1	lacA	[lacA,lacA]	[cytosol,cytosol,cytosol]	1.0	
2	lacZ	lacZ	[cytosol,cytosol]	1.0	
3	lacZ	[lacZ,lacZ]	[cytosol,cytosol,cytosol]	1.0	
4	lacZ	[lacZ,lacZ,lacZ]	[cytosol,cytosol,cytosol,cytosol]	1.0	

	RVS_RATE
0	1.0
1	1.0
2	1.0
3	1.0
4	1.0

```
[35]: atlas.read_network('network-lac-ProtProt.tsv')
```

```
[35]:
```

	SOURCE	TARGET	FWD_RATE	RVS_RATE	LOCATION
0	lacZ	lacZ	1.0	0.0	cytosol
1	[lacZ,lacZ]	[lacZ,lacZ]	1.0	0.0	cytosol
2	lacA	lacA	1.0	0.0	cytosol
3	lacA	[lacA,lacA]	1.0	0.0	cytosol

Despite the effort, the function could retrieve interactions that would be hard to observe in nature (parsimony). Also, because we obtain from BioCyc the protein composition of complexes and not intermediates, we could miss interactions or add interactions that never occur because we programmed the complete enumeration of possible ordered mechanisms. From the network, we could remove interaction by its indexes, and add interactions defining its participants:

```
[36]: network = utils.interactionNetwork.removeInteraction(network, index = [3,4])
network = utils.interactionNetwork.addInteraction(network, source = ['lacZ',
↳ 'lacZ'], target = ['lacZ', 'lacZ'])
# we supposed a dimer-dimer interaction is more plausible than a trimer-monomer
↳ interaction to form a tetramer
network
```

	SOURCE	TARGET	LOCATION	FWD_RATE	RVS_RATE
0	lacA	lacA	[cytosol, cytosol]	1.0	1.0
1	lacA	[lacA, lacA]	[cytosol, cytosol, cytosol]	1.0	1.0
2	lacZ	lacZ	[cytosol, cytosol]	1.0	1.0
3	[lacZ, lacZ]	[lacZ, lacZ]	cytosol	1.0	1.0

Note: If the location of complexes do not match the number of components, Atlas will suppose the location (or the first defined location) is valid for all components of the final complex.

```
[37]: %time model = atlas.construct_model_from_interaction_network(network, verbose =   
      ↪False)  
model
```

CPU times: user 198 ms, sys: 0 ns, total: 198 ms
Wall time: 196 ms

```
[37]: <Model 'atlas_rbm.construct_model_from_interaction_network' (monomers: 1, rules:  
      4, parameters: 14, expressions: 0, compartments: 0) at 0x7f7d5a00d7f0>
```

```
[38]: utils.analyzeConnectivity(model)
```

Every rule may be applied.

Every monomer and complex of monomers may occur in the model.

```
[39]: # initial condition  
      # for proteins  
simulation.set_initial.prot(model, 'lacZ', 'cyt', 12)  
simulation.set_initial.prot(model, 'lacA', 'cyt', 12)  
  
# simulation  
bng = '/opt/git-repositories/bionetgen.RuleWorld/bng2/'  
kasim = '/opt/git-repositories/KaSim4.Kappa-Dev/'  
cupsoda = '/opt/git-repositories/cupSODA.aresio/'  
stochkit = '/opt/git-repositories/StochKit.StochSS' # not the bin folder  
  
%time data0 = simulation.scipy(model, start = 0, finish = 10, points = 2000)  
%time data1 = simulation.cupsoda(model, start = 0, finish = 10, points = 2000,   
      ↪path = cupsoda)  
%time data2 = simulation.bngODE(model, start = 0, finish = 10, points = 2000,   
      ↪path = bng)  
%time data3 = simulation.bngSSA(model, start = 0, finish = 10, points = 2000,   
      ↪n_runs = 20, path = bng)  
# %time data4 = simulation.bngPLA(model, start = 0, finish = 10, points = 2000,   
      ↪n_runs = 20, path = bng)  
%time data5 = simulation.bngNF(model, start = 0, finish = 10, points = 2000,   
      ↪n_runs = 20, path = bng)
```

```
%time data6 = simulation.kasim(model, start = 0, finish = 10, points = 2000,
↳n_runs = 20, path = kasim)
# %time data7 = simulation.stochkit(model, start = 0, finish = 10, points =
↳2000, n_runs = 20, path = stochkit) # pySB error
```

```
CPU times: user 336 ms, sys: 11 ms, total: 347 ms
Wall time: 759 ms
CPU times: user 233 ms, sys: 13.5 ms, total: 247 ms
Wall time: 717 ms
CPU times: user 283 ms, sys: 5.43 ms, total: 289 ms
Wall time: 747 ms
CPU times: user 1.41 s, sys: 34.2 ms, total: 1.44 s
Wall time: 2.72 s
CPU times: user 971 ms, sys: 81.6 ms, total: 1.05 s
Wall time: 2.32 s
CPU times: user 935 ms, sys: 159 ms, total: 1.09 s
Wall time: 1.79 s
```

```
[40]: import seaborn
import matplotlib.pyplot as plt

palette = seaborn.color_palette('colorblind')

# first plot, periplasmic concentration
fig, ax = plt.subplots(1, 2, figsize = (4*2, 3*1), dpi = 100)
simulation.plot.protein(data3, 'lacZ', 'cyt', ax = ax[0], **{'kind' :
↳'fill_between', 'weight' : .5},
    plt_kws = {'s' : 2, 'color' : palette[0], 'label' : 'LacZ monomers',
↳'alpha' : .5})

simulation.plot.protein(data3, 'lacA', 'cyt', ax = ax[0], **{'kind' :
↳'fill_between', 'weight' : .5},
    plt_kws = {'s' : 2, 'color' : palette[1], 'label' : 'LacA monomers',
↳'alpha' : .5})

simulation.plot.cplx(data3, 'lacAx3', 'cyt', ax = ax[1], **{'kind' :
↳'fill_between', 'weight' : .5},
    plt_kws = {'s' : 2, 'color' : palette[2], 'label' : r'LacA trimers [CYT]',
↳'alpha' : .5})

simulation.plot.cplx(data3, 'lacZx4', 'cyt', ax = ax[1], **{'kind' :
↳'fill_between', 'weight' : .5},
    plt_kws = {'s' : 2, 'color' : palette[3], 'label' : r'LacZ tetramers [CYT]',
↳'alpha' : .5})

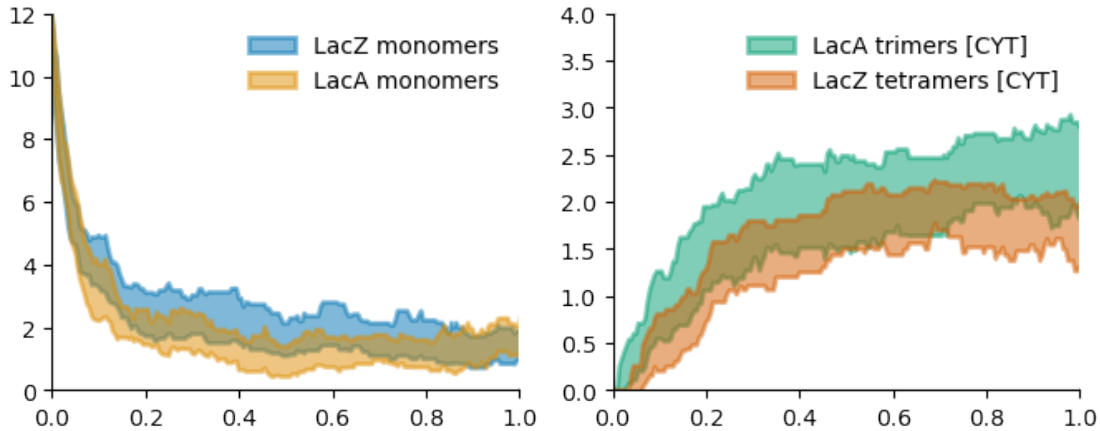
ax[0].set_xlim((0,1))
```

```

ax[1].set_xlim((0,1))
ax[0].set_ylim((0,12))
ax[1].set_ylim((0,4))

seaborn.despine()

```



1.7 Modeling gene expression

To model gene expression (transcription and translation) we must first retrieve data for the genomic architecture of the considered genes. For this purpose, we could use the `operon_of_gene` method from the `pythoncyc` package to obtain the operons of a gene:

```

[41]: import pythoncyc
      TUs = pythoncyc.select_organism('ECOLI').operon_of_gene(df_genes['gene_
      ↪name']['lacZ'])
      TUs

```

```

[41]: ['|TU0-4703|', '|TU00036|', '|TU0-4701|', '|TU0-4702|', '|TU0-14521|']

```

```

[42]: components = []
      for TU in TUs:
          x = utils.getData('ECOLI', TU)['components']
          print(TU, '=', x)
          components.append(x)

```

```

|TU0-4703| = ['|TERM0223|', '|TERM0222|', '|EG10524|', '|EG10526|', '|EG10527|',
'|PM0-4943|']
|TU00036| = ['|BS0-5469|', '|BS0-5468|', '|BS0-5467|', '|BS0-5463|',
'|BS0-5462|', '|BS0-5461|', '|TERM0223|', '|TERM0222|', '|PM00045|',
'|BS0-3622|', '|BS00104|', '|BS00105|', '|BS00106|', '|EG10524|', '|EG10526|',
'|EG10527|']
|TU0-4701| = ['|TERM0223|', '|TERM0222|', '|BS0-3622|', '|EG10524|',

```

```
'|EG10526|', '|EG10527|', '|PM0-4941|']
|TU0-4702| = ['|TERM0223|', '|TERM0222|', '|BS0-3622|', '|EG10524|',
'|EG10526|', '|EG10527|', '|PM0-4942|']
|TU0-14521| = ['|TERM0223|', '|TERM0222|', '|EG10524|', '|EG10526|',
'|PM0-9804|']
```

Note: An operon is a functional unit of DNA under the control of a promoter. In bacteria, operons could be polycistronic (two or more genes under the control of a unique promoter) or monocistronic (only one gene under the control of a promoter). However, the information from the pythoncyc could led to misinterpretation of the genomic architecture, and we do not provide a function to cover all possibilities. Please, look at <https://ecocyc.org/gene?orgid=ECOLI&id=EG10527#tab=TU> and similar webpages to construct by yourself the network using a spreadsheet or text editor software.

Here is an example depicting the DNA architecture of the lactose operons (see above note an output of `operon_of_gene`). Using brackets, Atlas understand that DNA promoters, ribosome binding sites (RBS), coding DNA sequences (CDS), and terminators are part of the same architectural unit, and therefore, Atlas defines the RNA products and protein products. From the example, Atlas will write rules for the synthesis of 10 molecules of RNA (five promoters times two terminators), which produce three proteins:

```
[43]: utils.read_network('network-lac-operon-arq.tsv')
```

```
[43]:
```

	UPSTREAM	DOWNSTREAM	RNAP_FWD_DOCK_RATE	RNAP_RVS_DOCK_RATE	\
0	[lacZ-pro4	lacZ-pro3	1.0	1.0	
1	lacZ-pro3	lacZ-pro2	1.0	1.0	
2	lacZ-pro2	lacZ-pro1	1.0	1.0	
3	lacZ-pro1	lacZ-rbs	1.0	1.0	
4	lacZ-rbs	lacZ-cds	NaN	NaN	
5	lacZ-cds	lacY-pro1	NaN	NaN	
6	lacY-pro1	lacY-rbs	1.0	1.0	
7	lacY-rbs	lacY-cds	NaN	NaN	
8	lacY-cds	lacA-rbs	NaN	NaN	
9	lacA-rbs	lacA-cds	NaN	NaN	
10	lacA-cds	lacA-ter1	NaN	NaN	
11	lacA-ter1	lacA-ter2]	NaN	NaN	

	RNAP_FWD_SLIDE_RATE	RNAP_FWD_FALL_RATE	RIB_FWD_DOCK_RATE	\
0	1.0	NaN	NaN	
1	1.0	NaN	NaN	
2	1.0	NaN	NaN	
3	1.0	NaN	NaN	
4	1.0	NaN	1.0	
5	1.0	NaN	NaN	
6	1.0	NaN	NaN	
7	1.0	NaN	1.0	
8	1.0	NaN	NaN	
9	1.0	NaN	1.0	
10	1.0	1.0	NaN	

11	1.0	1.0	NaN
----	-----	-----	-----

	RIB_RVS_DOCK_RATE	RIB_FWD_SLIDE_RATE	RIB_FWD_FALL_RATE
0	NaN	NaN	NaN
1	NaN	NaN	NaN
2	NaN	NaN	NaN
3	NaN	NaN	NaN
4	1.0	1.0	1.0
5	NaN	NaN	NaN
6	NaN	NaN	NaN
7	1.0	1.0	1.0
8	NaN	NaN	NaN
9	1.0	1.0	1.0
10	NaN	NaN	NaN
11	NaN	NaN	NaN

Notes:

- RNAP_FWD_DOCK_RATES and RNAP_RVS_DOCK_RATES are valid values only for the description of the reversible interaction of the RNA Polymerase to the promoters.
- RNAP_FWD_SLIDE_RATES are valid values for the transition of the RNA Polymerase from the UPSTREAM to the DOWNSTREAM DNA parts. We describe rules where the RNA Polymerase could not move back.
- RNAP_FWD_FALL_RATES are valid values only for the description of the unbinding of the RNA Polymerase and the UPSTREAM identifying a DNA terminator.
- RIB_FWD_DOCK_RATES and RIB_RVS_DOCK_RATES are valid values only for the description of the reversible interaction of the bacterial Ribosome to the RBS.
- RIB_FWD_SLIDE_RATES are valid values for the transition of the Ribosome from the UPSTREAM to the DOWNSTREAM RNA parts. We describe rules where the Ribosome could not move back.
- RIB_FWD_FALL_RATES are valid values only for the description of the unbinding of the Ribosome and the UPSTREAM identifying a CDS.
- Encode DNA parts as: “name of the gene”-“type of DNA part” (**pro#**: Promoter, **rbs**: Ribosome Binding Site, **cds**: Coding DNA Sequence, **ter#**: (transcriptional) Terminator). Multiple promoters and terminators must have a numeric identifier (to replace #). For instance, **lacZ-pro1** identifies the most proximal promoter to the **lacZ** gene.
- In the case of Binding Sites (BS) for the interaction of Transcriptional Factors with DNA, please use the notation: BS-“name of the gene or regulator”-“upstream genomic coordinate”-“downstream genomic coordinate”. The coordinates could be relatives or absolutes. For instance **BS-araC-56-72** describe the binding site located upstream the transcription start site of the **araC** gene, from the -56 nucleotide to the -72 nucleotide.

We provide two functions to reconstruction transcription and translation from the network of DNA parts (known as a “genome graph”). Here, we will use the first of these two functions. The `construct_model_from_genome_graph()` function is an automatization of the Kappa Bio-Brick Framework (<https://www.sciencedirect.com/science/article/pii/S1571066111001289>). The

function writes rules for transcription of DNA mediated by a RNAP-CPLX agent (to keep notation with BioCyc) and translation of RNA mediated by a RIBOSOME-CPLX agent:

```
[44]: %time model = atlas.construct_model_from_genome_graph('network-lac-operon-arq.
      ↪tsv')
model
```

CPU times: user 718 ms, sys: 31.7 ms, total: 750 ms

Wall time: 750 ms

```
[44]: <Model 'atlas_rbm.construct_model_from_genome_graph' (monomers: 4, rules: 28,
parameters: 50, expressions: 0, compartments: 0) at 0x7f7daa5f7d90>
```

```
[45]: %time model = atlas.construct_model_from_genome_graph('network-ara-operons-arq.
      ↪tsv')
model
```

CPU times: user 1.03 s, sys: 0 ns, total: 1.03 s

Wall time: 1.05 s

```
[45]: <Model 'atlas_rbm.construct_model_from_genome_graph' (monomers: 4, rules: 50,
parameters: 78, expressions: 0, compartments: 0) at 0x7f7d52f40280>
```

1.8 Modeling regulation of gene expression

The second function that describe gene transcription and translation also consider the specificity of sigma factors for bacterial promoters. In bacteria, there is only one RNA polymerase that synthesize mRNA, rRNA, and tRNA. To control which genes are in active transcription, promoters show differential specificity for one or several sigma factors. In the case of *E. coli*, the bacteria has seven of those sigma factors: rpoD (exponential growth phase), rpoE (heat and protein misfolding stress), rpoH (heat stress), rpoN (nitrogen homeostasis), rpoS (stationary phase), fecI (iron starvation), and fliA (flagella synthesis) that interact with the core RNA polymerase (rpoAABC) and form an interaction interface for the holoenzyme with promoters.

```
[46]: utils.read_network('network-lac-sigma-specificity.tsv')
```

```
[46]:
```

	SOURCE	TARGET	FWD_DOCK_RATE	RVS_DOCK_RATE	\
0	[rpoA,rpoA,rpoB,rpoC,rpoD]	lacZ-pro4	1.0	1.0	
1	[rpoA,rpoA,rpoB,rpoC,rpoD]	lacZ-pro3	1.0	1.0	
2	[rpoA,rpoA,rpoB,rpoC,rpoD]	lacZ-pro2	1.0	1.0	
3	[rpoA,rpoA,rpoB,rpoC,rpoD]	lacZ-pro1	1.0	1.0	
4	[rpoA,rpoA,rpoB,rpoC,rpoD]	lacY-pro1	1.0	1.0	

	FWD_SLIDE_RATE
0	1.0
1	1.0
2	1.0
3	1.0

Note: The TARGET name must match the UPSTREAM name in the network that describe the genomic architecture:

```
[47]: network = utils.read_network('network-lac+rpoABCD-operons-arq.tsv')
network[['UPSTREAM', 'DOWNSTREAM']][network['UPSTREAM'].str.contains('pro')] #
↳ shows the UPSTREAM and DOWNSTREAM columns where the UPSTREAM column contains
↳ the 'pro' string
```

```
[47]:      UPSTREAM DOWNSTREAM
0    [lacZ-pro4  lacZ-pro3
1     lacZ-pro3  lacZ-pro2
2     lacZ-pro2  lacZ-pro1
3     lacZ-pro1  lacZ-rbs
6     lacY-pro1  lacY-rbs
12   [rpoA-pro1  rpoA-rbs
15   [rpoB-pro1  rpoB-rbs
20   [rpoD-pro1  rpoD-rbs
```

```
[48]: %time model = atlas.
↳ construct_model_from_sigma_specificity_network('network-lac-sigma-specificity.
↳ tsv', 'network-lac+rpoABCD-operons-arq.tsv')
model
```

CPU times: user 1.41 s, sys: 7.98 ms, total: 1.41 s
Wall time: 1.43 s

```
[48]: <Model 'atlas_rbm.construct_model_from_sigma_specificity_network' (monomers: 4,
rules: 55, parameters: 91, expressions: 0, compartments: 0) at 0x7f7daa6b9be0>
```

```
[49]: %time model = atlas.
↳ construct_model_from_sigma_specificity_network('network-ara-sigma-specificity.
↳ tsv', 'network-ara+rpoABCD-operons-arq.tsv')
```

CPU times: user 1.7 s, sys: 6.96 ms, total: 1.71 s
Wall time: 1.91 s

However, sigma factors is only one of the many mechanism a bacteria has to control gene expression. Canonically speaking, Transcription Factors (TFs) drive the control of gene expression. TFs are proteins that bind to DNA or unbind from DNA at speciliazed sequences. Most TFs are sensors of metabolites: lacI senses allolactose, araC senses arabinose, crp senses cyclic AMP, etc. We employ an interaction network to model such information:

```
[50]: net1 = utils.read_network('network-lac+ara-TFs+DNA.tsv')
# net1 # for better visualization
net1[0:1]
```

```
[50]:
```

	SOURCE	TARGET	FWD_RATE	RVS_RATE	LOCATION
0	[crp,SMALL-CAMP,crp,SMALL-CAMP]	BS-crp-51-72	1.0	1.0	cytosol

Note: We appended the prefix `SMALL-` to tell Atlas that the interaction partner is a metabolite and not a protein (an interaction network is interpreted by default as a protein-protein interaction network). We employ the *hypergraph notation* to denote a complex of agents.

```
[51]: net2 = utils.read_network('network-lac+ara-ProtMet.tsv')
      net3 = utils.read_network('network-lac-ProtProt.tsv')
```

Networks of the same kind (except metabolic networks) must be concatenated to produce valid models. Atlas numbers the rules and there is a probability to produce non unique rule names. In the case of metabolic networks, we use the reaction name to name rules (and check for unique names prior to write rules).

```
[52]: import pandas
      network = pandas.concat([net1, net2, net3]).reset_index(drop = True)
      # rules are numbered and networks of the same kind cannot be processed
      # → separately to then combine them into one bigger model (uniqueness of rule
      # → names)
      # reset index is optional, but we used it to show the model produce all 30
      # → interactions (index start at zero)
      network[-5:]
```

```
[52]:
```

	SOURCE	TARGET \
30	[araC,SMALL-alpha-L-arabinopyranose,araC]	SMALL-alpha-L-arabinopyranose
31	lacZ	lacZ
32	[lacZ,lacZ]	[lacZ,lacZ]
33	lacA	lacA
34	lacA	[lacA,lacA]

	FWD_RATE	RVS_RATE	LOCATION
30	1.0	1.0	cytosol
31	1.0	0.0	cytosol
32	1.0	0.0	cytosol
33	1.0	0.0	cytosol
34	1.0	0.0	cytosol

```
[53]: %time model = atlas.construct_model_from_interaction_network(network)
      model
```

```
CPU times: user 988 ms, sys: 88 µs, total: 988 ms
Wall time: 1.01 s
```

```
[53]: <Model 'atlas_rbm.construct_model_from_interaction_network' (monomers: 3, rules:
      35, parameters: 89, expressions: 0, compartments: 0) at 0x7f7daa93d370>
```

Data from the BioCyc database concerning regulation of gene expression could be obtained with

help of the `utils.getData()` function. After obtaining the components of an operon, we could obtain the genomic coordinates of *most* of the DNA parts, and the center position of DNA binding sites:

```
[54]: for DNA_part in components[1]:
        data = utils.getData('ECOLI', DNA_part)
        print(DNA_part, data['left_end_position'], data['abs_center_pos'],
              ↪data['right_end_position'])
```

```
|BS0-5469| None 366410 None
|BS0-5468| None 366397 None
|BS0-5467| None 366374 None
|BS0-5463| None 366348 None
|BS0-5462| None 366338 None
|BS0-5461| None 366320 None
|TERM0223| 361140 None 361179
|TERM0222| 361212 None 361236
|PM00045| None None None
|BS0-3622| None 366404.5 None
|BS00104| None 365932 None
|BS00105| None 366425 None
|BS00106| None 366333 None
|EG10524| 361249 None 361860
|EG10526| 361926 None 363179
|EG10527| 363231 None 366305
```

```
[55]: print(TUs[1])
for DNA_part in components[1]:
    data = utils.getData('ECOLI', DNA_part)
    if data['involved_in_regulation'] != None:
        # print(data['involved_in_regulation'][0])
        regulation = utils.getData('ECOLI', data['involved_in_regulation'][0])
        print(DNA_part, regulation['regulator'], regulation['mode'])
print()

print(TUs[2])
for DNA_part in components[2]:
    data = utils.getData('ECOLI', DNA_part)
    if data['involved_in_regulation'] != None:
        # print(data['involved_in_regulation'][0])
        regulation = utils.getData('ECOLI', data['involved_in_regulation'][0])
        print(DNA_part, regulation['regulator'], regulation['mode'])
print()

print(TUs[3])
for DNA_part in components[3]:
    data = utils.getData('ECOLI', DNA_part)
    if data['involved_in_regulation'] != None:
```

```
#         print(data['involved_in_regulation'][0])
regulation = utils.getData('ECOLI', data['involved_in_regulation'][0])
print(DNA_part, regulation['regulator'], regulation['mode'])
```

```
|TU00036|
|BS0-5469| |PD00288| ['-']
|BS0-5468| |PD00288| ['-']
|BS0-5467| |PD00288| ['-']
|BS0-5463| |PD00288| ['-']
|BS0-5462| |PD00288| ['-']
|BS0-5461| |PD00288| ['-']
|BS0-3622| |CPLX0-226| ['-']
|BS00104| |PD00763| ['-']
|BS00105| |PD00763| ['-']
|BS00106| |PD00763| ['-']
```

```
|TU0-4701|
|BS0-3622| |CPLX0-226| ['-']
```

```
|TU0-4702|
|BS0-3622| |CPLX0-226| ['-']
```

We hope you could format data by hand, as many interactions with a Transcription factor and a small metabolite are described as metabolic reactions, meanwhile we formatted that data as interaction networks:

```
[56]: print(utils.getData('ECOLI', 'CPLX0-226')['components']) # CPLX0-226 is the
      ↪ CRP-cAMP complex
print(utils.getData('ECOLI', 'CPLX0-226')['consensus_sequence']) # we could
      ↪ obtain the consensus DNA sequence if known
print(utils.getData('ECOLI', 'PC00004')['appears_in_left_side_of']) # in
      ↪ contrast to the "catalyzes" slot of gene products
print(utils.getData('ECOLI', 'BETAGALACTOSID-CPLX')['catalyzes'])
print(utils.getData('ECOLI', 'CPLX0-226')['appears_in_right_side_of']) #
      ↪ equivalent output
print(' + '.join(utils.getData('ECOLI', 'RXN0-269')['left']) + '->' + ' + ' +
      ↪ join(utils.getData('ECOLI', 'RXN0-269')['right']))
```

```
[ '|PC00004|', '|CAMP|' ]
[ 'AAATGTGAtctagaTCACATTT' ]
[ '|RXN0-269|' ]
[ '|BETAGALACTOSID-ENZRXN|', '|ENZRXN0-6526|', '|ENZRXN0-8150|',
  '|ENZRXN0-8152|' ]
[ '|RXN0-269|' ]
|PC00004| + |CAMP| -> |CPLX0-226|
```

Finally, we will incorporate regulatory relationships through the copy of rules describing the docking or sliding of the Ribosome through the DNA. For the inactivation of transcription by competition

of a TF and a RNA polymerase for the same or overlapping binding site, we simply add interactions of the TF and copy the docking rule of the RNA polymerase with a condition:

```
[57]: %time network = utils.read_network('network-lac+ara-TFs+DNA.tsv')
      network[0:5]
```

CPU times: user 3.89 ms, sys: 0 ns, total: 3.89 ms

Wall time: 3.3 ms

```
[57]:
```

	SOURCE	TARGET	FWD_RATE	RVS_RATE	\
0	[crp,SMALL-CAMP,crp,SMALL-CAMP]	BS-crp-51-72	1.0	1.0	
1	[crp,SMALL-CAMP,crp,SMALL-CAMP]	BS-crp-22-1	1.0	1.0	
2	lacI BS-lacI-422-402		1.0	1.0	
3	lacI BS-lacI-21-1		1.0	1.0	
4	lacI BS-lacI-72-92		1.0	1.0	


```

LOCATION
0  cytosol
1  cytosol
2  cytosol
3  cytosol
4  cytosol
```

```
[58]: %time model1 = atlas.
      ↪construct_model_from_interaction_network('network-lac+ara-TFs+DNA.tsv')
      model1
```

CPU times: user 545 ms, sys: 0 ns, total: 545 ms

Wall time: 544 ms

```
[58]: <Model 'atlas_rbm.construct_model_from_interaction_network' (monomers: 3, rules:
      24, parameters: 53, expressions: 0, compartments: 0) at 0x7f7daa5ba280>
```

```
[59]: %time model2 = atlas.
      ↪construct_model_from_sigma_specificity_network('network-lac-sigma-specificity.
      ↪tsv', 'network-lac+rpoABCD-operons-arq.tsv')
      model2
```

CPU times: user 1.35 s, sys: 0 ns, total: 1.35 s

Wall time: 1.35 s

```
[59]: <Model 'atlas_rbm.construct_model_from_sigma_specificity_network' (monomers: 4,
      rules: 55, parameters: 91, expressions: 0, compartments: 0) at 0x7f7daa6b9d00>
```

```
[60]: %time model = atlas.combine_models([model1, model2], verbose = False)
      model
```

CPU times: user 1.9 s, sys: 0 ns, total: 1.9 s

Wall time: 1.91 s

[60]: <Model 'atlas_rbm.atlas' (monomers: 5, rules: 79, parameters: 144, expressions: 0, compartments: 0) at 0x7f7daa650a90>

```
[61]: atlas.add_regulation(model, name = 'docking_2_lacZ_pro3',  
                           conditions = ['BS-crp-51-72', 'BS-crp-22-1'], replace = True)
```

[61]: <Model 'atlas_rbm.atlas' (monomers: 5, rules: 79, parameters: 146, expressions: 0, compartments: 0) at 0x7f7daa650a90>

```
[62]: atlas.add_regulation(model, name = 'docking_3_lacZ_pro2',  
                           conditions = ['BS-crp-51-72', 'BS-crp-22-1'], replace = True)
```

[62]: <Model 'atlas_rbm.atlas' (monomers: 5, rules: 79, parameters: 148, expressions: 0, compartments: 0) at 0x7f7daa650a90>

```
[63]: atlas.add_regulation(model, name = 'docking_4_lacZ_pro1',  
                           conditions = ['[crp,SMALL-CAMP,crp,SMALL-CAMP,BS-crp-51-72]',  
                                         '[crp,SMALL-CAMP,crp,SMALL-CAMP,BS-crp-22-1]'],  
                           ↪replace = True)
```

[63]: <Model 'atlas_rbm.atlas' (monomers: 5, rules: 79, parameters: 150, expressions: 0, compartments: 0) at 0x7f7daa650a90>