



A software platform for genome-scale metabolic models simulation, reconstruction and visualization

User manual

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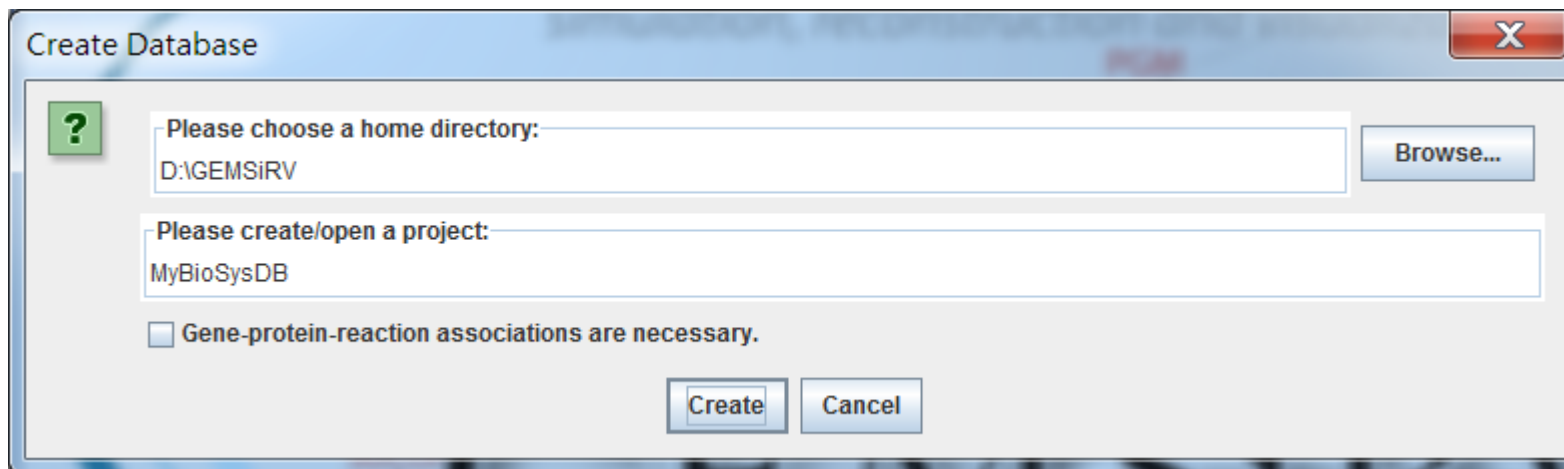
Mar 06, 2012

GEMSiRV

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Basically, a metabolic network is an assembly of biochemical reactions. While information about reactions is sufficient for modeling the network, the more information on associated genes or proteins, the more useful for the investigation of cellular responses in gene or protein level. Gene-protein-reaction (GPR) associations can be described in two-layer relations: “gene and protein” and “protein and reaction”, which are usually saved in spreadsheet format, the required information and available models are summarized in http://sb.nhri.org.tw/GEMSiRV/en/Metabolic_Models. On the other hand, published genome-scale metabolic models are commonly exchanged in Systems Biology Markup Language (SBML) format, but the protein information is lost or can not be recovered to the two-layer relation. Therefore, GEMSiRV provides different schema for these two types of metabolic reconstructions. If you want to create a project with clear two-layer relations of GPR associations, please check the checkbox of Gene-protein-reaction associations are necessary to generate the three-index schema (gene, protein and reaction indices) for reconstruction. Otherwise, GEMSiRV will generate the two-index schema (gene and reaction indices) automatically.



Create Database

?

Please choose a home directory:

D:\GEMSiRV Browse...

Please create/open a project:

MyBioSysDB

☐ Gene-protein-reaction associations are necessary.

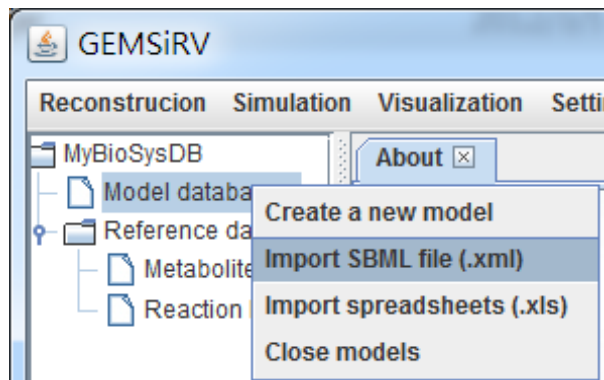
Create Cancel

Reconstruction

Click on Reconstruction in the menu bar to open **Model databases** and **Reference databases**.

Model importing and editing

Right click on **Model databases** to Import SBML file (.xml) or to Import spreadsheets (.xls), you can import a metabolic model in SBML/spreadsheet format. Some existing metabolic models can be found and downloaded from http://sb.nhri.org.tw/GEMSiRV/en/Metabolic_Models



You can directly edit/update the content of the imported model by right clicking on a cell.

In table of Reaction Index:

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB

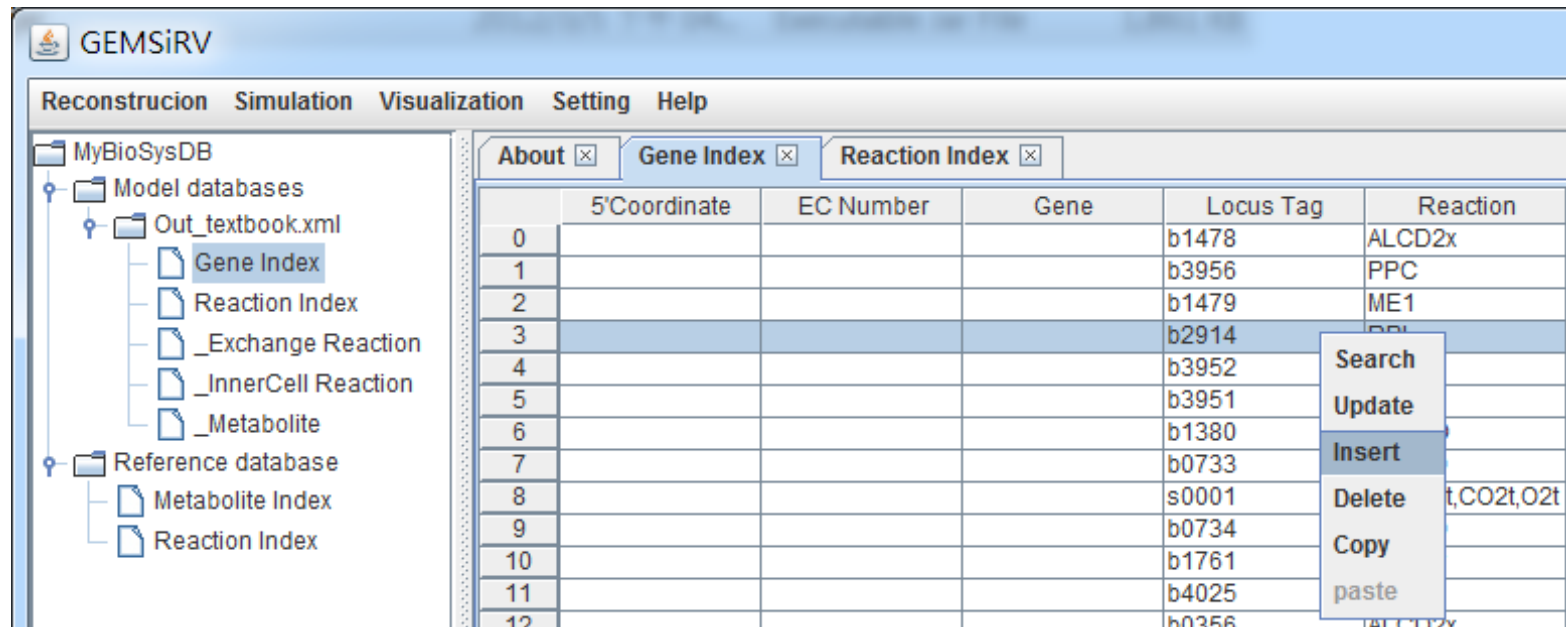
- Model databases
 - Out_textbook.xml
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
- Reference database
 - Metabolite Index
 - Reaction Index

About [x] Gene Index [x] Reaction Index [x]

	Abbreviation	Confidence	Equation	Gene	Name
0	ADK1		[c] : amp + atp <=...	(b0474)	adenylate kinase
1	GLUSy		[c] : akg + gln-L +...	(b3212+b3213)	glutamate syntha...
2	GLUDy		[c] : glu-L + h2o +...	(b1761)	glutamate dehyd...
3	FORt2		for[e] + h[e] --> fo...	(b0904),(b2492)	formate transport...
4	FUM		[c] : fum + h2o <=...	(b1612),(b1611)...	fumarase
5	PDH		[c] : coa + nad + ...	(b0114)+(b0116...	pyruvate dehydro...
6	GLUt2r		glu-L[e] + h[e] <=...	(b4077)	L-glutamate tran...
7	ALCD2x		[c] : etoh + nad <...	(b0356),(b1241)...	alcohol dehydrog...
8	ICDHyr		[c] : icit + nadp <=...	(b1136)	isocitrate dehydr...
9	PYRt2r		h[e] + pyr[e] <==> ...		pyruvate reversib...
10	SUCDi		[c] : q8 + succ --> ...	(b0722+b0721+...	succinate dehydr...
11	FRD7		[c] : fum + q8h2 --...	(b4151+b4153+...	fumarate reducta...
12	NADH16		(4.0) h[c] + nadh[...	(b2285+b2284+...	NADH dehydrog...
13	TPI		[c] : dha + h2o <--> g...	(b3010)	triose-phosphate

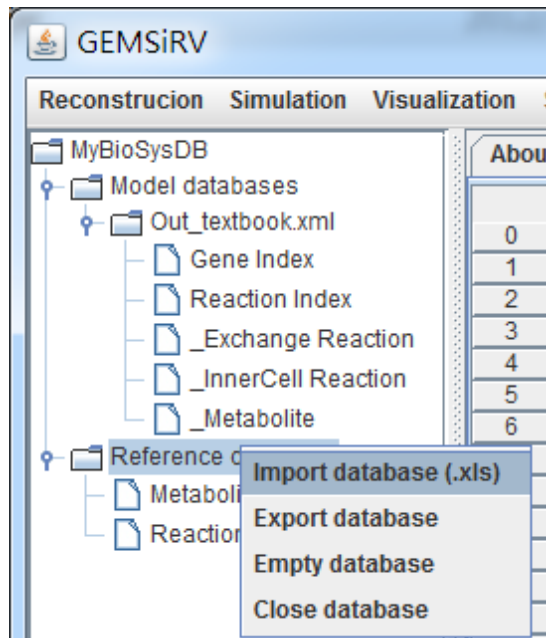
Search
Update
Insert
Delete
Copy
paste

In table of Gene Index:

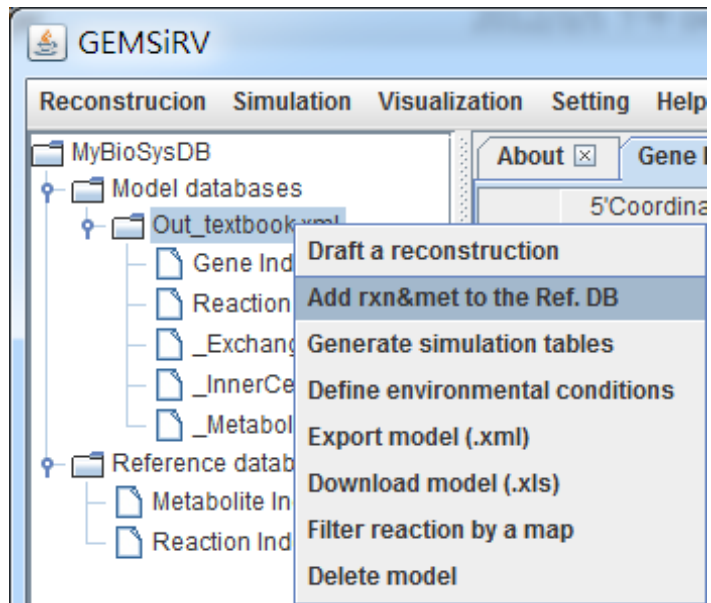


Reference database construction

Right click on **Reference database** to Import database (.xls), you can import a reference database to construct your own reference database. Available reference databases including BiGG, KEGG, MetaCyc and Model SEED databases can be found and downloaded from http://sb.nhri.org.tw/GEMSiRV/en/Reference_Databases.



Or, you can right click on the model you imported to Add rxn&met to the Ref. DB, so that you can add the information about metabolites and reaction described in the model to the reference database that you have created.



You can right click on the Reaction Index of **Reference database** to evaluate charge/mass balance of equation.

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB

- Model databases
 - Out_textbook.xml
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
- Reference database
 - Metabolite Index
 - Reaction Index

Reaction Index

	ABBREVIATION	EC_NUMBER	EQUATION
0	ADK1		[c] : amp + atp <==>...
1	GLUSy		[c] : akg + gln-L + h ...
2	GLUDy		[c] : glu-L + h2o + n...
3	FORT2		for[e] + h[e] --> for[c]...
4	FUM		[c] : fum + h2o <==>...
5	PDH		[c] : coa + nad + pyr ...
6	GLUt2r		glu-L[e] + h[e] <==> ...
7	ALCD2x		[c] : etoh + nad <== ...
8	ICDHyr		[c] : icit + nadp <==> ...
9	pyrD2r		h[e] + pyr[e] <==> h[...
10	q8h2		[c] : q8 + succ --> fu...
11	FRD7		[c] : fum + q8h2 --> ...
12	NADH16		(4.0) h[c] + nadh[c] ...

Evaluate charge/formula balance

You can add/edit the equation of reaction by using equation dialog or type directly. For example, for reaction PGK (phosphoglycerate kinase), you can type "[c] : 3pg + atp <==> 13dpg + adp" in its equation or you can enter the EQUATION dialog to edit its content.

☒ \rightleftharpoons
☐ \rightarrow

Left Column:
 (1) 3pg[c]
 (1) atp[c]
 Stoichiometry: 1
 Compartment: c
 Reactant:
 Add Delete

Right Column:
 (1) 13dpg[c]
 (1) adp[c]
 Stoichiometry: 1
 Compartment: c
 Product:
 Add Delete

Evaluate Submit Cancel

Draft reconstruction generation

Firstly, you need to have a close related model organism whose metabolic reconstruction has been built already.

Then you need to prepare a blank reconstruction containing gene information of your interest strain. This file can be generated by GBKParser (<http://sb.nhri.org.tw/GEMSiRV/en/GBKParser>). However, **you need to add the corresponding orthologous genes to the column of Ref-BLAST.**

For example, we want to draft a reconstruction of *Salmonella enteric subsp. Enteric serovar Typhimurium str. LT2 (SLT2)* by mapping to the reconstruction model *iAF1260 of Escherichia coli str. K-12 substr. MG1655 (ECO)*.

Therefore, we download the gbk files of these two strains from RefSeq (<http://www.ncbi.nlm.nih.gov/RefSeq>). With available

NC_003197.gbk and NC_000913.gbk files for SLT2 and ECO respectively, we then use GBKParser to parse basic gene information and amino acid sequences. In addition, we download the metabolic model *iAF1260* from BiGG (<http://bigg.ucsd.edu/>) and modify it with TextReplacer (<http://sb.nhri.org.tw/GEMSiRV/en/TextReplacer>). The ready-to-use model can be found and downloaded from http://sb.nhri.org.tw/GEMSiRV/en/Metabolic_Models.

The amino acid sequence files for SLT2 and ECO can be used to generate the reciprocal orthologous-gene pairs by BLASTP or other available software. For example, MrBac (<http://sb.nhri.org.tw/MrBac>) can be used to generate the needed file. However, the detailed procedure is not described here.

The basic gene information parsed from the gbk file is outputted to a spreadsheet file, e.g. NC_003197.gbk.xls, which can be imported into GEMSiRV directly. Right click on **Model Databases** to Import spreadsheet (.xls).

Original spreadsheet file:

Microsoft Excel - NC_003197.gbk-blast.xls

檔案(F) 編輯(E) 檢視(V) 插入(I) 格式(O) 工具(T) 資料(D) 視窗(W) RExcel 說明(H) Adobe PDF(B)

100%

B I U

回復變更(C)... 結束檢閱(N)...

F1 Ref-BLAST

	A	B	C	D	E	F	G	H	I	J	K
	Locus Tag	Gene	5'Coordinate	Product	EC Number	Ref-BLAST					
1	STM0001	thrL	190	thr operon	--						
2	STM0002	thrA	337	bifunctional	2.7.2.4,;1.1.1	b0002					
3	STM0003	thrB	2801	homoserin	2.7.1.39	b0003					
4	STM0004	thrC	3734	threonine s	4.2.3.1	b0004					
5	STM0005	yaaA	5887	hypothetic	--	b0006					
6	STM0006	yaaJ	7396	putative al	--	b0007					
7	STM0007	talB	7665	transaldol	2.2.1.2	b0008					
8	STM0008	mogA	8729	molybdenu	--	b0009					
9	STM0009	yaaH	9942	hypothetic	--	b0010					
10	STM0010	htgA	10805	hypothetic	--	b0011					
11	STM0011	yaal	11245	hypothetic	--	b0013					
12	STM0012	dnaK	11593	molecular	--	b0014					
13	STM0013	dnaJ	13595	chaperone	--	b0015					
14	STM0014	--	15014	putative tra	--						
15	STM0015	--	16088	putative ba	--						
16	STM0016	--	17026	hypothetic	--						
17	STM0017	--	17486	hypothetic	--						
18	STM0018	--	17867	putative ex	--						
19	STM0019	--	20058	putative hy	--						
20	STM0020	--	23335	putative cy	--						
21	STM0021	bcfA	24469	fimbrial sul	--						
22	STM0022	hcrB	25112	fimbrial ch	--						

Gene Index

就緒

Required fields

Gene Index table of the imported blank reconstruction (NC_003197.gbk-blast.xls):

Reconstrucion Simulation Visualization Setting

Project

- Model databases
 - SBML_export_E.coli textbook_out.xml
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - SBML_export_E. coli iAF1260_out.xml
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - NC_003197_gbk-blast.xls
 - Gene Index
 - Reaction Index
- Reference database
 - Metabolite Index
 - Reaction Index
- Maps
 - E.coli_core.cartomap

Gene Index

	5'Coordinate	Locus Tag	Gene	EC Number	Product	Ref-BLAST
0	190	STM0001	thrL	--	thr operon lead...	
1	337	STM0002	thrA	2.7.2.4;;1.1....	bifunctional asp...	b0002
2	2801	STM0003	thrB	2.7.1.39	homoserine kin...	b0003
3	3734	STM0004	thrC	4.2.3.1	threonine synth...	b0004
4	5887	STM0005	yaaA	--	hypothetical pro...	b0006
5	7396	STM0006	yaaJ	--	putative alanine...	b0007
6	7665	STM0007	talB	2.2.1.2	transaldolase B	b0008
7	8729	STM0008	mogA	--	molybdenum co...	b0009
8	9942	STM0009	yaaH	--	hypothetical pro...	b0010
9	10805	STM0010	htgA	--	hypothetical pro...	b0011
10	11245	STM0011	yaal	--	hypothetical pro...	b0013
11	11593	STM0012	dnaK	--	molecular chap...	b0014
12	13595	STM0013	dnaJ	--	chaperone prot...	b0015
13	15014	STM0014	--	--	putative transcri...	
14	16088	STM0015	--	--	putative bacteri...	
15	17026	STM0016	--	--	hypothetical pro...	
16	17486	STM0017	--	--	hypothetical pro...	
17	17867	STM0018	--	--	putative exochiti...	
18	20058	STM0019	--	--	putative hydroxy...	
19	23335	STM0020	--	--	putative cytopla...	
20	24469	STM0021	bcfA	--	fimbrial subunit	
21	25112	STM0022	bcfB	--	fimbrial chapar...	
22	25803	STM0023	bcfC	--	fimbrial usher	b0940
23	28425	STM0024	bcfD	--	fimbrial subunit	
24	29433	STM0025	bcfE	--	fimbrial subunit	
25	29994	STM0026	bcfF	--	fimbrial subunit	
26	30478	STM0027	bcfG	--	fimbrial chapar...	
27	31274	STM0028	bcfH	--	putative thiol-di...	
28	32116	STM0028...	--	--	hypothetical pro...	

Empty Reaction Index table of the imported blank reconstruction:

The screenshot shows a software window with a blue title bar and standard Windows window controls. The interface is divided into two main sections. On the left is a 'Project' tree view showing a hierarchy of folders and files. On the right is a table titled 'Reaction Index' which is currently empty.

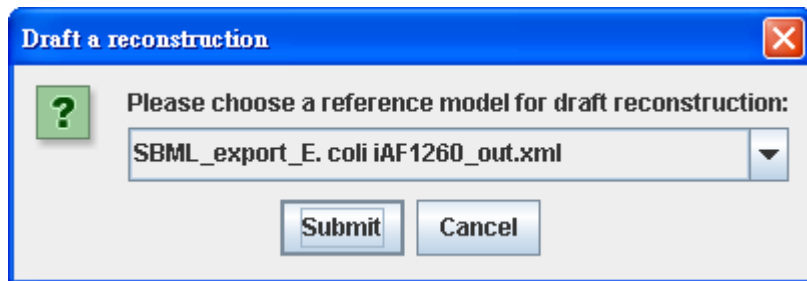
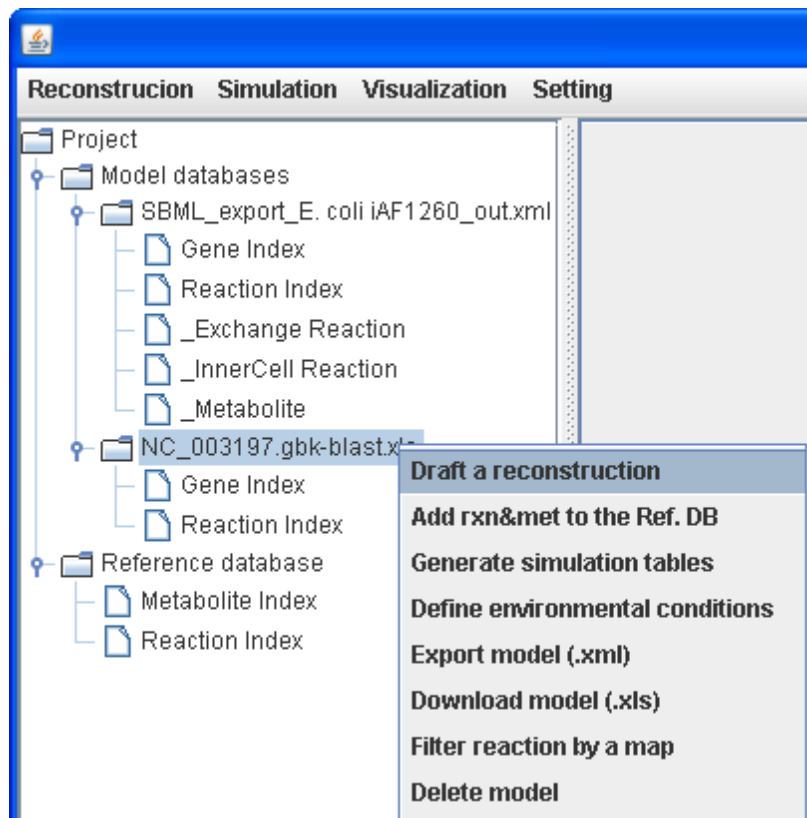
Project Tree:

- Project
 - Model databases
 - SBML_export_E.coli textbook_out.xml
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - SBML_export_E. coli iAF1260_out.xml
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - NC_003197.gbk-blast.xls
 - Gene Index
 - Reaction Index
 - Reference database
 - Metabolite Index
 - Reaction Index
 - Maps
 - E.coli_core.cartomap

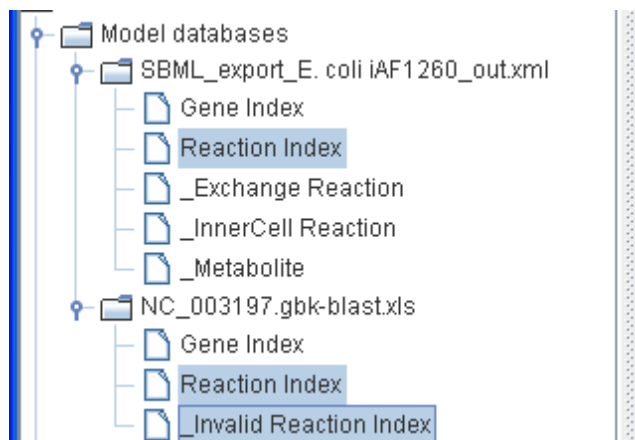
Reaction Index Table:

	Abbreviation	Confidence	Equation	Gene	Name
0					
1					
2					
3					
4					
5					
6					
7					
8					
9					
10					
11					
12					
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					
23					
24					
25					
26					
27					
28					

Right click on the blank reconstruction to Draft a reconstruction by choosing SBML_export_E.coli iAF1260_out.xml as the reference model.

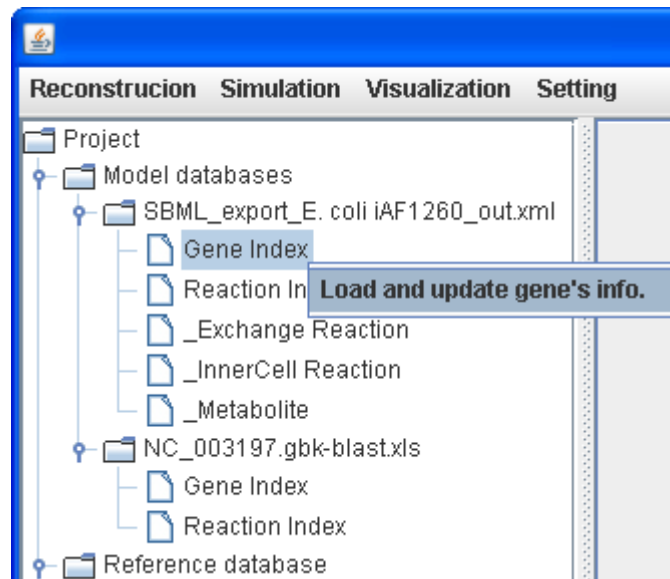


The reactions in the reference reconstruction are classified into two indices (Reaction Index and _Invalid Reaction Index) for the draft reconstruction: one list containing reactions whose associated orthologous genes are present in the blank reconstruction and conform to Boolean statements as described in the reference reconstruction, the other containing those reactions with unknown gene-reaction associations or reactions whose orthologous genes are absent and let to disagree Boolean statements.



Model refinement

Based on the draft reconstruction generated from Model SEED (<http://seed-viewer.theseed.org/seedviewer.cgi?page=ModelView>) or GEMSiRV, users can curate and refine the reconstruction in GEMSiRV. However, the lack of gene information in imported models may hinder the progress. We, therefore, provide a function to load and update the gene information in GEMSiRV. You can right click on the Gene Index of a model to Load and update gene's info., and upload the spreadsheet file generated by GBKParser (<http://sb.nhri.org.tw/GEMSiRV/en/GBKParser>), e.g. NC_000913.gbk.xls for ECO.

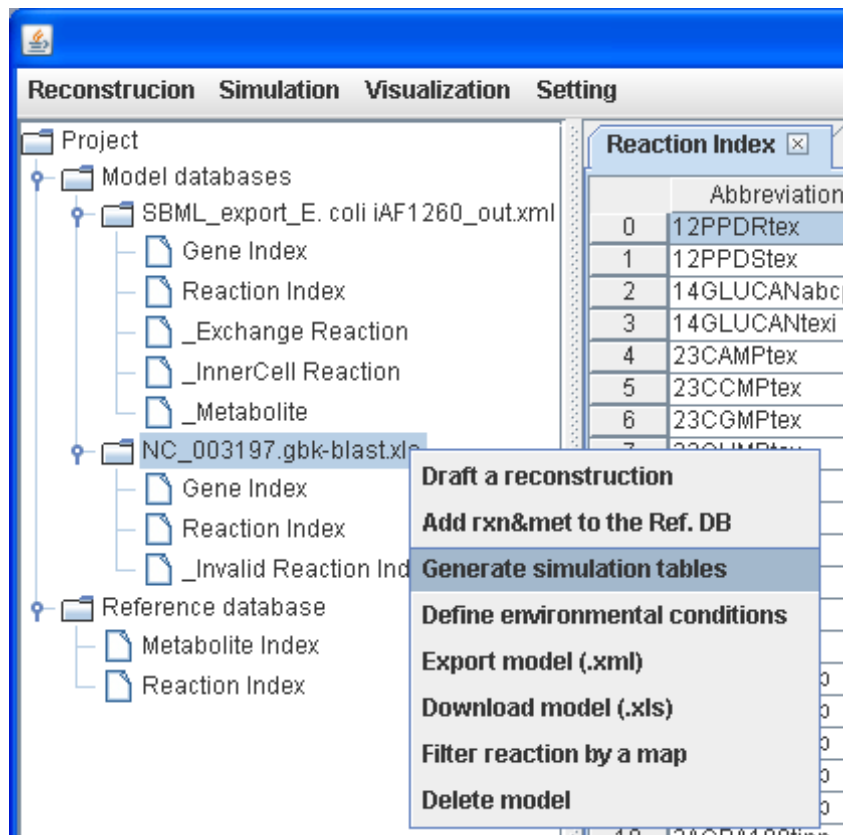


Gene information can be loaded and updated accordingly:

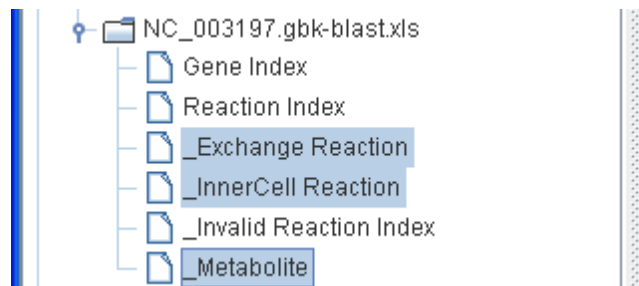
Reconstrucion Simulation Visualization Setting						
<div>Project</div> <div> <div>Model databases</div> <div> <div>SBML_export_E. coli iAF1260_out.xml</div> <div> <div>Gene Index</div> <div>Reaction Index</div> <div>_Exchange Reaction</div> <div>_InnerCell Reaction</div> <div>_Metabolite</div> </div> </div> <div>NC_003197.gbk-blast.xls</div> <div> <div>Gene Index</div> <div>Reaction Index</div> </div> <div>Reference database</div> <div> <div>Metabolite Index</div> <div>Reaction Index</div> </div> <div>Maps</div> </div>						
Gene Index						
	5'Coordinate	EC Number	Gene	Locus Tag	Reaction	
0				s0001	GLYctp,FEEN...	
1	190	--	thrL	b0001		thr
2	337	2.7.2.4;;1.1.1.3	thrA	b0002	ASPK,HSDy	fus
3	2801	2.7.1.39	thrB	b0003	HSK	ho
4	3734	4.2.3.1	thrC	b0004	THRS,4HTRS	thr
5	5234	--	yaaX	b0005		pre
6	6459	--	yaaA	b0006		co
7	7959	--	yaaJ	b0007	GLYt4pp,ALAt4pp	pre
8	8238	2.2.1.2	talB	b0008	TALA	tra
9	9306	--	mog	b0009		pre
10	10494	--	yaaH	b0010		co
11	11356	--	yaaW	b0011		co
12	11786	--	yaal	b0013		pre
13	12163	--	dnaK	b0014		cha
14	14168	--	dnaJ	b0015		cha
15	15445	--	insL	b0016		IS1
16	16903	--	hokC	b4412		tox
17	17489	--	nhaA	b0019	NAt3_2pp	so
18	18715	--	nhaR	b0020		DN
19	20314	--	insB	b0021		IS1
20	20508	--	insA	b0022		Kp
21	21078	--	rpsT	b0023		30
22	21181	--	yaaY	b0024		pre
23	21407	2.7.1.26;;2.7.7.2	ribF	b0025	FMNAT,RBFK	bif
24	22391	6.1.1.5	ileS	b0026	ILETRS	isc
25	25207	3.4.23.36	lspA	b0027		pro
26	25826	5.2.1.8	fkpB	b0028		FK
27	26277	1.17.1.2	ispH	b0029	DMPPS,IPDPS	1-P
28	27293	3.2.2.-	rihC	b0030	ADNUC,URIH,C...	rib

With the aids of simulation and visualization, users can readily identify dead-end metabolites and blocked reactions in the models. Prior to

perform simulation, users need to convert the reconstruction into a mathematical model. Therefore, you can right click on a model to Generate simulation tables to generate a model containing a stoichiometric matrix as well as default systems boundaries.



After clicking on Generate simulation tables, three tables including InnerCell Reaction, Exchange Reaction and Metabolite are generated. The prefix “_” used in these three tables for easily distinguishing from the tables required for reconstruction, e.g. Gene Index, Protein Index (optional) and Reaction Index.



_InnerCell Reaction:

Reconstruction Simulation Visualization Setting

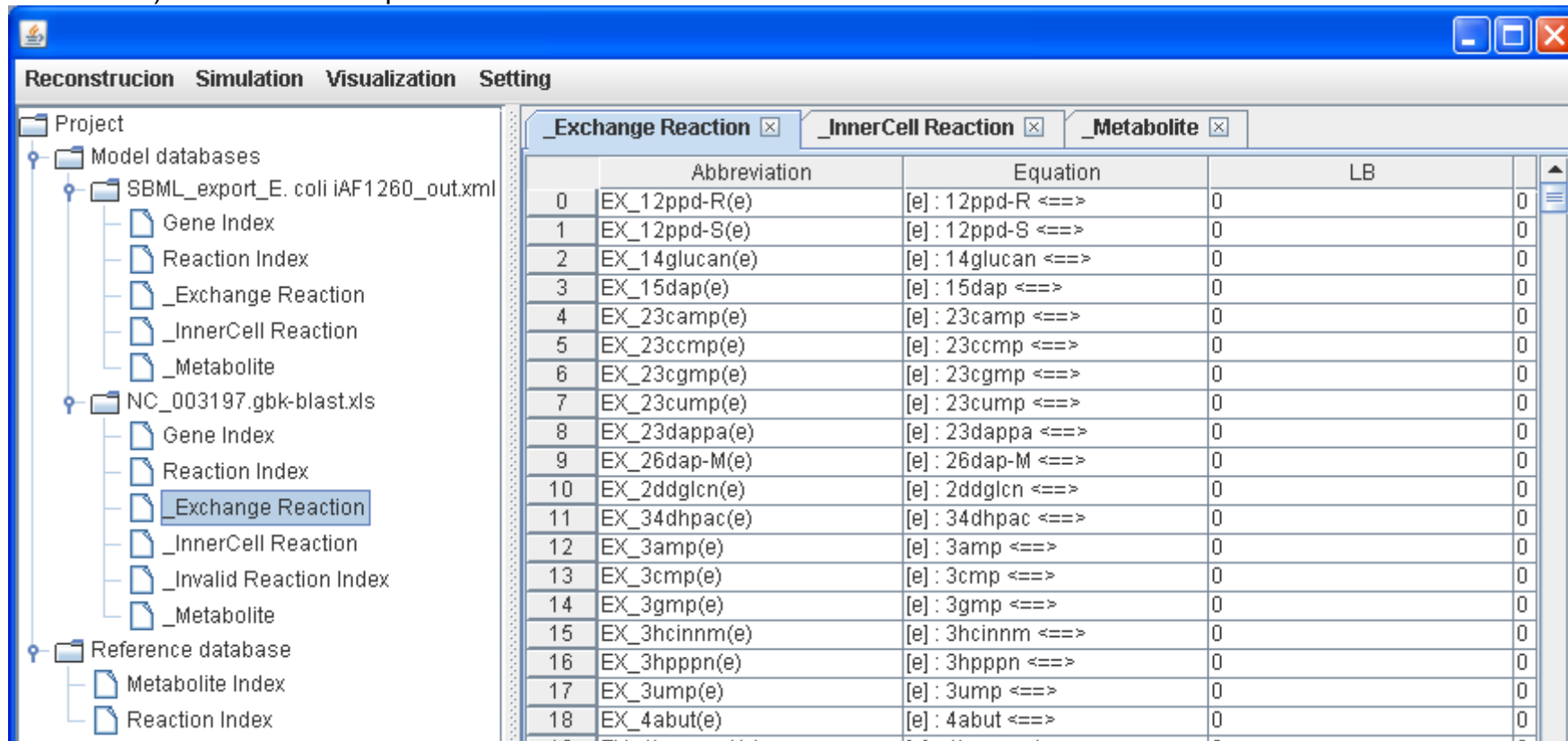
Project

- Model databases
 - SBML_export_E. coli iAF1260_out.xml
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - NC_003197.gbk-blast.xls
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction**
 - _Invalid Reaction Index
 - _Metabolite
- Reference database
 - Metabolite Index
 - Reaction Index

	Abbreviation	Association	Equation	LB	Name
0	12PPDRtex	STM2267,STM03...	12ppd-R[e] <==> ...	-1000.0	(R)-Propane-1,2-
1	12PPDStex	STM2267,STM03...	12ppd-S[e] <==> ...	-1000.0	(S)-Propane-1,2-
2	14GLUCANabcpp	STM4230+STM4...	14glucan[p] + atp...	0	1,4-alpha-D-gluc..
3	14GLUCANtexi	STM4231	14glucan[e] --> 1...	0	1,4-alpha-D-gluc..
4	23CAMPtex	STM2267,STM03...	23camp[e] <==> ...	-1000.0	23cAMP transpor.
5	23CCMPtex	STM2267,STM03...	23ccmp[e] <==> ...	-1000.0	23cCMP transpo..
6	23CGMPtex	STM2267,STM03...	23cgmp[e] <==> ...	-1000.0	23cGMP transpo..
7	23CUMPtex	STM2267,STM03...	23cump[e] <==> ...	-1000.0	23cUMP transpo..
8	23DAPPAtex	STM2267,STM03...	23dappa[e] <==> ...	-1000.0	2,3-diaminopropi.
9	23PDE2pp	STM4403	[p] : 23cump + h2...	0	2',3'-cyclic-nucle...
10	23PDE4pp	STM4403	[p] : 23ccmp + h2...	0	2',3'-cyclic-nucle...
11	23PDE7pp	STM4403	[p] : 23camp + h2...	0	2',3'-cyclic-nucle...
12	23PDE9pp	STM4403	[p] : 23cgmp + h2...	0	2',3'-cyclic-nucle...
13	26DAHtex	STM1473,STM22...	26dap-M[e] <==> ...	-1000.0	meso-2,6-Diami..
14	2AGPA120tipp	STM3009	2ddecg3p[p] --> ...	0	2-Acyl-sn-glycero.
15	2AGPA140tipp	STM3009	2tdecg3p[p] --> 2...	0	2-Acyl-sn-glycero.
16	2AGPA141tipp	STM3009	2tdec7eg3p[p] --...	0	2-Acyl-sn-glycero.
17	2AGPA160tipp	STM3009	2hdecg3p[p] --> ...	0	2-Acyl-sn-glycero.
18	2AGPA161tipp	STM3009	2hdec9eg3p[p] --...	0	2-Acyl-sn-glycero.
19	2AGPA180tinn	STM3009	2ndecg3p[p] --> ...	0	2-Acyl-sn-glycero.

_Exchange Reaction:

Please note that the _Exchange Reaction table will be generated only when you have exchanging metabolites (i.e. extracellular metabolites) in the reaction equations.



	Abbreviation	Equation	LB
0	EX_12ppd-R(e)	[e] : 12ppd-R <==>	0
1	EX_12ppd-S(e)	[e] : 12ppd-S <==>	0
2	EX_14glucan(e)	[e] : 14glucan <==>	0
3	EX_15dap(e)	[e] : 15dap <==>	0
4	EX_23camp(e)	[e] : 23camp <==>	0
5	EX_23ccmp(e)	[e] : 23ccmp <==>	0
6	EX_23cgmp(e)	[e] : 23cgmp <==>	0
7	EX_23cump(e)	[e] : 23cump <==>	0
8	EX_23dappa(e)	[e] : 23dappa <==>	0
9	EX_26dap-M(e)	[e] : 26dap-M <==>	0
10	EX_2ddgln(e)	[e] : 2ddgln <==>	0
11	EX_34dhpac(e)	[e] : 34dhpac <==>	0
12	EX_3amp(e)	[e] : 3amp <==>	0
13	EX_3cmp(e)	[e] : 3cmp <==>	0
14	EX_3gmp(e)	[e] : 3gmp <==>	0
15	EX_3hcinnm(e)	[e] : 3hcinnm <==>	0
16	EX_3hpppn(e)	[e] : 3hpppn <==>	0
17	EX_3ump(e)	[e] : 3ump <==>	0
18	EX_4abut(e)	[e] : 4abut <==>	0

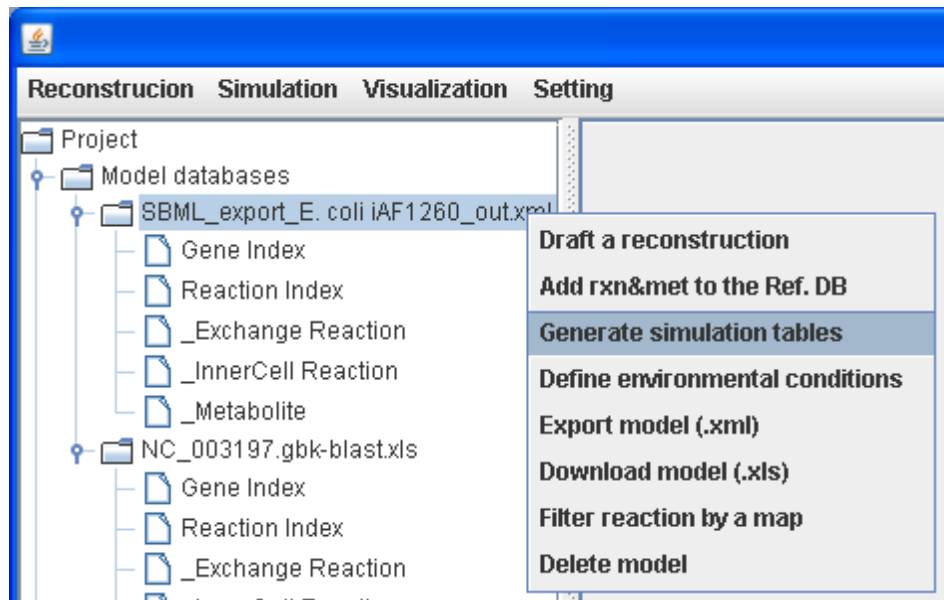
_Metabolite:

The screenshot shows the GEMSiRV software interface. On the left is a 'Project' tree with folders for 'Model databases' (containing 'SBML_export_E. coli iAF1260_out.xml' and 'NC_003197.gbk-blast.xls') and 'Reference database'. The 'Metabolite' file under 'SBML_export_E. coli iAF1260_out.xml' is selected. The main window displays a table of metabolites under the '_Metabolite' tab. The table has columns for Index, Abbreviation, Formula, Name, and Compartment. The data is as follows:

	Abbreviation	Formula	Name	Compartment
0	10fthf	C20H21N7O7	10-Formyltetrahydrofolate	cytoplasm
1	12dgr120	C27H52O5	1,2-Diacyl-sn-glycerol (didod...	cytoplasm
2	12dgr120	C27H52O5	1,2-Diacyl-sn-glycerol (didod...	periplasm
3	12dgr140	C31H60O5	1,2-Diacyl-sn-glycerol (ditetr...	cytoplasm
4	12dgr140	C31H60O5	1,2-Diacyl-sn-glycerol (ditetr...	periplasm
5	12dgr141	C31H56O5	1,2-Diacyl-sn-glycerol (ditetr...	cytoplasm
6	12dgr141	C31H56O5	1,2-Diacyl-sn-glycerol (ditetr...	periplasm
7	12dgr160	C35H68O5	1,2-Diacyl-sn-glycerol (dihex...	cytoplasm
8	12dgr160	C35H68O5	1,2-Diacyl-sn-glycerol (dihex...	periplasm
9	12dgr161	C35H64O5	1,2-Diacyl-sn-glycerol (dihex...	cytoplasm
10	12dgr161	C35H64O5	1,2-Diacyl-sn-glycerol (dihex...	periplasm
11	12dgr180	C39H76O5	1,2-Diacyl-sn-glycerol (diocta...	cytoplasm
12	12dgr180	C39H76O5	1,2-Diacyl-sn-glycerol (diocta...	periplasm
13	12dgr181	C39H72O5	1,2-Diacyl-sn-glycerol (diocta...	cytoplasm
14	12dgr181	C39H72O5	1,2-Diacyl-sn-glycerol (diocta...	periplasm
15	12ppd-R	C3H8O2	(R)-Propane-1,2-diol	cytoplasm
16	12ppd-R	C3H8O2	(R)-Propane-1,2-diol	extracellular
17	12ppd-R	C3H8O2	(R)-Propane-1,2-diol	periplasm
18	12ppd-S	C3H8O2	(S)-Propane-1,2-diol	cytoplasm
19	12ppd-S	C3H8O2	(S)-Propane-1,2-diol	extracellular

Because growth media for modeled organisms may be similar, an environmental condition can be easily set to a model by right clicking on the model to Define environmental conditions.

Here we use the *in silico* (computational) minimal media for the model *iAF1260* as an example (the text file can be downloaded in <http://sb.nhri.org.tw/GEMSiRV/en/Manual>). In order to set the system boundaries to the default values, we right click on the model to Generate simulation tables.



The new simulation tables are generated and replace the previous tables. We set a growth medium for modeling the model. We prepare a text file containing the user-defined boundaries and objective, and then right click on the model to Define environmental conditions.

In silico minimal media for the model iAF1260.

M9 medium_comp.TXT

```

1 [REACTIONS]
2 EX_ca2(e):LB=-1000
3 EX_cl(e):LB=-1000
4 EX_co2(e):LB=-1000
5 EX_cobalt2(e):LB=-1000
6 EX_cu2(e):LB=-1000
7 EX_fe2(e):LB=-1000
8 EX_fe3(e):LB=-1000
9 EX_h2o(e):LB=-1000
10 EX_h(e):LB=-1000
11 EX_k(e):LB=-1000
12 EX_mg2(e):LB=-1000
13 EX_mn2(e):LB=-1000
14 EX_mobd(e):LB=-1000
15 EX_na1(e):LB=-1000
16 EX_nh4(e):LB=-1000
17 EX_pi(e):LB=-1000
18 EX_so4(e):LB=-1000
19 EX_tungs(e):LB=-1000
20 EX_zn2(e):LB=-1000
21 EX_o2(e):LB=-18.5
22 ATPM:LB=8.39,UB=9.39
23 Biomass_Ecoli_core_N(w/GAM)_Nmet2:Objective Coefficient=1
24 EX_glc-D(e):LB=-8

```

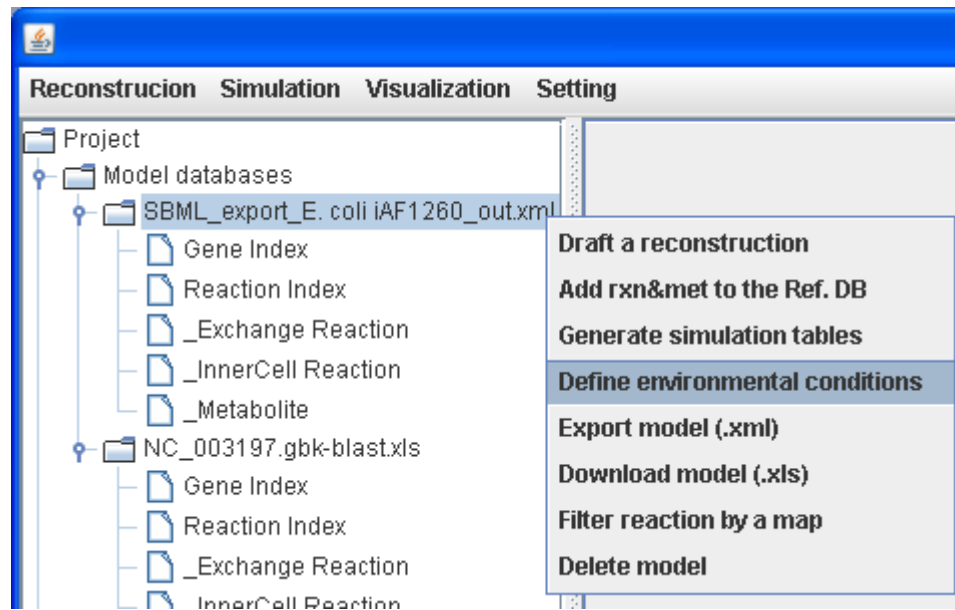
A complete medium to simulate all extracellular metabolites can enter/exit the cell freely.

complete medium.TXT

```

1 [ALL EXCHANGE REACTIONS]
2 LB=-1000
3 UB=1000
4 Objective Coefficient=0
5

```



The user-defined system boundaries and the objective are set in the reconstruction model accordingly.

Reconstrucion Simulation Visualization Setting

Project

- Model databases
 - SBML_export_E. coli iAF1260_out.xml
 - Gene Index
 - Reaction Index
 - Exchange Reaction**
 - _InnerCell Reaction
 - _Metabolite
 - NC_003197.gbk-blast.xls
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Invalid Reaction Index
 - _Metabolite
- Reference database
 - Metabolite Index
 - Reaction Index

_Exchange Reaction

	Abbreviation	Equation	LB	UB	Objective C
0	EX_zn2(e)	[e] : zn2 <==>	-1000	1000.0	0
1	EX_tungs(e)	[e] : tungs <==>	-1000	1000.0	0
2	EX_so4(e)	[e] : so4 <==>	-1000	1000.0	0
3	EX_pi(e)	[e] : pi <==>	-1000	1000.0	0
4	EX_nh4(e)	[e] : nh4 <==>	-1000	1000.0	0
5	EX_na1(e)	[e] : na1 <==>	-1000	1000.0	0
6	EX_mobd(e)	[e] : mobd <==>	-1000	1000.0	0
7	EX_mn2(e)	[e] : mn2 <==>	-1000	1000.0	0
8	EX_mg2(e)	[e] : mg2 <==>	-1000	1000.0	0
9	EX_k(e)	[e] : k <==>	-1000	1000.0	0
10	EX_h2o(e)	[e] : h2o <==>	-1000	1000.0	0
11	EX_h(e)	[e] : h <==>	-1000	1000.0	0
12	EX_fe3(e)	[e] : fe3 <==>	-1000	1000.0	0
13	EX_fe2(e)	[e] : fe2 <==>	-1000	1000.0	0
14	EX_cu2(e)	[e] : cu2 <==>	-1000	1000.0	0
15	EX_cobalt2(e)	[e] : cobalt2 <==>	-1000	1000.0	0
16	EX_co2(e)	[e] : co2 <==>	-1000	1000.0	0
17	EX_cl(e)	[e] : cl <==>	-1000	1000.0	0
18	EX_ca2(e)	[e] : ca2 <==>	-1000	1000.0	0
19	EX_o2(e)	[e] : o2 <==>	-18.5	1000.0	0
20	EX_glc-D(e)	[e] : glc-D <==>	-8	1000.0	0
21	EX_xyly-L(e)	[e] : xyly-L <==>	0	1000.0	0

Or you can simply right click on the reaction to update the lower bound (LB), upper bound (UB) or objective coefficient.

About [x] Reaction Index [x] _Exchange Reaction [x]					
	Abbreviation	Equation	LB	UB	Objective C...
58	EX_butso3(e)	[e] : butso3 <==>	0	999999	0
59	EX_ca2(e)	[e] : ca2 <==>	-999999	999999	0
60	EX_cbi(e)	[e] : cbi <==>	0	999999	0
61	EX_cbl1(e)	[e] : cbl1 <==>	-.01	999999	0
62	EX_cd2(e)	[e] : cd2 <==>	0	999999	0
63	EX_cgly(e)	[e] : cgly <==>	0	999999	0
64	EX_chol(e)	[e] : chol <==>	0	999999	0
65	EX_cit(e)	[e] : cit <==>	0	999999	0
66	EX_cl(e)	[e] : cl <==>	-999999	999999	0

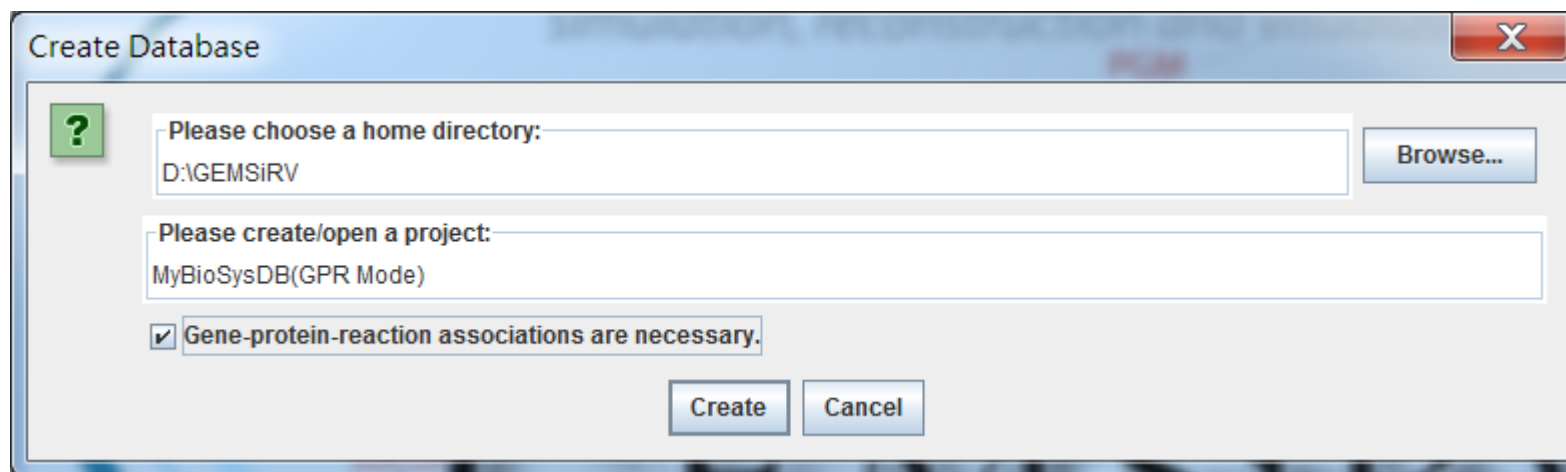
Update					
<div> Click Submit to modify this record, or Cancel </div>					
Abbreviation:		EX_ca2(e)		Equation: [e] : ca2 <==>	
LB:		-999999		Objective Coefficient: 0	
UB:		999999			
<input type="button" value="Submit"/> <input type="button" value="Cancel"/>					

78	EX_cynt(e)	[e] : cynt <==>	0	999999	0
79	EX_cys_D(e)	[e] : cys-D <==>	0	999999	0
80	EX_cys_L(e)	[e] : cys-L <==>	0	999999	0
81	EX_cytd(e)	[e] : cytd <==>	0	999999	0
82	EX_dad_2(e)	[e] : dad-2 <==>	0	999999	0
83	EX_damp(e)	[e] : damp <==>	0	999999	0
84	EX_dca(e)	[e] : dca <==>	0	999999	0
85	EX_dcmp(e)	[e] : dcmp <==>	0	999999	0
86	EX_dcyt(e)	[e] : dcyt <==>	0	999999	0
87	EX_dcca(e)	[e] : dcca <==>	0	999999	0

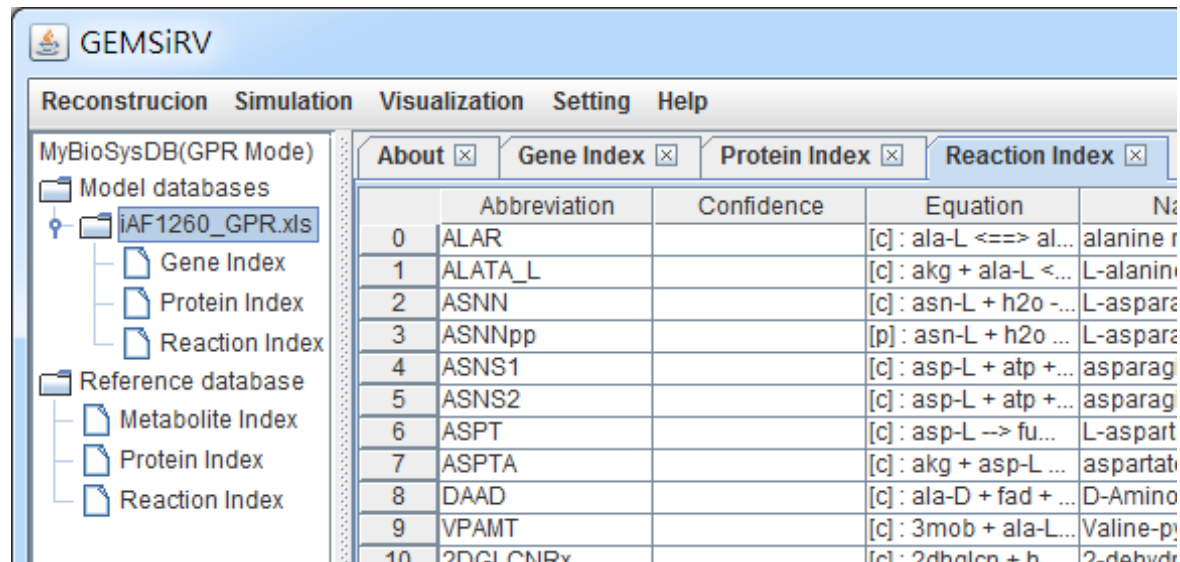
You can freely export or save a metabolic model in SBML format or in spreadsheet format by right clicking on a model to [Export model \(.xml\)](#) or to [Download model \(.xls\)](#). Such models generated by GEMSiRV are fully compatible to GEMSiRV for later importing and simulation.

In addition to the metabolic models saved in SBML format, metabolic reconstructions can be stored in spreadsheet format. The spreadsheet format can store the two-layer relation for gene-protein and protein-reaction associations in network reconstructions. We provide available reconstruction models (GPR) in http://sb.nhri.org.tw/GEMSiRV/en/Metabolic_Models and demonstrate how we use GEMSiRV to reconstruct metabolic networks with GPR relationships.

From reconstruction to model



After clicking on Reconstruction in the menu bar, right click on the **Model databases** to Import spreadsheets (.xls) for importing the reconstruction file of iAF1260_GPR.xls (download from http://sb.nhri.org.tw/GEMSiRV/en/Metabolic_Models). This reconstruction contains three indices: Gene, Protein and Reaction Index.



Then right click on **Reference database** to Import database (.xls) for importing the reference database file Ref_BiGG_GPR.xls which is provided in http://sb.nhri.org.tw/GEMSiRV/en/Reference_Databases.

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - IAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

	ABBREVIATION	EC_NUMBER	EQUATION	KEGGID	NAME	NOTE
0	10FTHF5GLUtl		10fthf5glu[c] --> 10f...		5-glutamyl-10FTHF ...	
1	10FTHF5GLUtm		10fthf5glu[m] --> 10f...		5-glutamyl-10FTHF ...	
2	10FTHF6GLUtl		10fthf6glu[c] --> 10f...		6-glutamyl-10FTHF ...	
3	10FTHF6GLUtm		10fthf6glu[m] --> 10f...		6-glutamyl-10FTHF ...	
4	10FTHF7GLUtl		10fthf7glu[c] --> 10f...		7-glutamyl-10FTHF ...	
5	10FTHF7GLUtm		10fthf7glu[m] --> 10f...		7-glutamyl-10FTHF ...	
6	10FTHFtl		10fthf[c] <==> 10fthf[l]		10-Formyltetrahydro...	
7	10FTHFtm		10fthf[c] <==> 10fthf[...]		10-Formyltetrahydro...	
8	11DOCRTSLtm		11docrts[c] <==> 1...		11-deoxycortisol intr...	
9	11DOCRTSLtr		11docrts[c] <==> 1...		11-deoxycortisol intr...	
10	11DOCRTSTRNtm		11docrtstrn[c] <==> ...		11-deoxycorticoster...	
11	11DOCRTSTRNtr		11docrtstrn[c] <==> ...		11-deoxycorticoster...	
12	12DCR120tl		12dcr120tl --> 12d...		1,2-dicarbonyl...	

A biomass for E. coli, Biomass_Ecoli_core_N (w/ GAM)-Nmet2, is available in the reference database, you can add the reaction to the reconstruction by right clicking on the main window of Reaction Index to Insert. After submitting the abbreviation of reaction "Biomass_Ecoli_core_N (w/ GAM)-Nmet2", the related information including reaction name and equation will be conveyed to the reconstruction from the reference database.

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - IAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

About | Gene Index | Protein Index | **Reaction Index** | Metabolite Index

	Abbreviation	Confidence	Equation	Name
0	ALAR		[c] : ala-L <=> al...	alanine racemase
1	ALATA_L		[c] : akg + ala-L <...	L-alanine transa...
2	ASNN		[c] : asn-L + h2o -...	L-asparaginase
3	ASNNpp		[p] : asn-L + h2o ...	L-asparaginase
4	ASNS1		[c] : asp-L + atp +...	asparagine synt...
5	ASNS2		[c] : asp-L + atp +...	asparagine synt...
6	ASPT		[c] : asp-L -> fu...	L-aspartase
7	ASPTA		[c] : akg + asp-L ...	aspartate transa...
8	DAAD		[c] : ala-D + fad + ...	D-Amino acid de...
9	VPAMT		[c] : 3mob + ala-L...	Valine-pyruvate a...
10	2DGLCNRx		[c] : 2dhgln + h ...	2-dehydro-D-glu...
11	2DGLCNRy		[c] : 2dhgln + h ...	2-dehydro-D-glu...
12	2DGULRx		[c] : 2dhgln + h ...	2-dehydro-L-gulo...
13	2DGULRy		[c] : 2dhgln + h ...	2-dehydro-L-gulo...
14	3HCINNMH		[c] : 3hcinm + h ...	3-hydroxycinnam...
15	3HPPPNH		[c] : 3hpppn + h +...	3-(3-hydroxy-phe...

Search
Update
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Insert [X]

? Click Submit to insert this record, or Cancel

Abbreviation: Confidence:

Note: Other:

Protein: Subsystem:

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - IAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

	Abbreviation	Confidence	Equation	Name	Note	Other	Protein
0	ALAR		[c] : ala-L <=> al...	alanine racemase			Alrec, DadX
1	ALATA_L		[c] : akg + ala-L <...	L-alanine transa...			
2	ASNN		[c] : asn-L + h2o -...	L-asparaginase			AnsA, YbiK
3	ASNNpp		[p] : asn-L + h2o ...	L-asparaginase			AnsB
4	ASNS1		[c] : asp-L + atp +...	asparagine synt...			AsnB
5	ASNS2		[c] : asp-L + atp +...	asparagine synt...			AsnA
6	Biomass_Ecoli_...		[c] : (1.496) 3pg +...	core E. coli biom...			
7	ASPT		[c] : asp-L --> fu...	L-aspartase			AspA
8	ASPTA		[c] : akg + asp-L ...	aspartate transa...			AspC
9	DAAD		[c] : ala-D + fad + ...	D-Amino acid de...			DadA
10	VPAMT		[c] : 3mob + ala-L...	Valine-pyruvate a...			AvtA
11	2DOL GMD...		[c] : 2dhol + h...	2-dehydro D-ala...			ViaF

Or you can search Biomass in the Reaction Index of Reference database:

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - iAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

Index	ABBREVIATION	EC_NUMBER	EQUATION	KEGGID	NAME	NOTE
1221	Biomass_Ecoli_cor		[c] : (1.496) 3pg + (3...		core E. coli biomas...	
1222	biomass_SA_2a		[c] : (0.42) 12dgr_E...		SA biomass 2 witho...	
1223	biomass_SA_2b		[c] : (0.42) 12dgr_E...		SA biomass 2 witho...	
1224	biomass_SA_3a		[c] : (0.42) 12dgr_E...		SA biomass 3 witho...	
1225	biomass_SA_3b		[c] : (0.42) 12dgr_E...		SA biomass 3 witho...	
1226	biomass_SA_4a		[c] : (0.42) 12dgr_S...		SA biomass 4 witho...	
1227	biomass_SA_5a		[c] : (0.42) 12dgr_S...		SA biomass 5 witho...	
1228	biomass_SA_6a		[c] : (0.42) 12dgr_S...		SA biomass 6 witho...	
1229	biomass_SA_6b		[c] : (0.42) 12dgr_S...		SA biomass 6 witho...	
1230	biomass_SA_7a		[c] : (0.01) 10fthf + (...)		SA biomass 7 witho...	
1231	biomass_SA_7b		[c] : (0.01) 10fthf + (...)		SA biomass 7 witho...	
1232	biomass_SA_8a		[c] : (0.01) 10fthf + (...)		SA biomass 8 witho...	
1233	biomass_SA_lipids...		[c] : (0.42) 12dgr_E...		SA biomass 2 lipid...	

Search
Update
Insert
Delete
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paste

then, directly copy and paste to the Reaction Index of iAF1260_GPR.xls

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - iAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

	Abbreviation	Confidence	Equation	Name	Note	Other	Protein
0	ALAR		[c] : ala-L <==> al...	alanine racemase			Alrec, DadX
1	ALATA_L		[c] : akg + ala-L <...	L-alanine transa...			
2	ASNN		[c] : asn-L + h2o -...	L-asparaginase			AnsA, YbiK
3	ASNNpp		[p] : asn-L + h2o ...	L-asparaginase			AnsB
4	ASNS1		[c] : asp-L + atp +...	asparagine synt...			AsnB
5	ASNS2		[c] : asp-L + atp +...	asparagine synt...			AsnA
6	ASPT		[c] : asp-L --> fu...	L-aspartase			AspA
7	ASPTA		[c] : akg + asp-L ...	aspartate transa...			AspC
8	DAAD		[c] : ala-D + fad + ...	D-Amino acid de...			DadA
9	VPAMT		[c] : 3mob + ala-L...	Valine-pyruvate a...			AvtA
10	2DGLCNRx		[c] : 2dhgln + h ...	2-dehydro-D-glu...			YiaE
11	2DGLCNRy		[c] : 2dhgln + h ...	2-dehydro-D-glu...			YiaE
12	2DGULRx		[c] : 2dhgln + h ...	2-dehydro-L-gulo...			YiaE
13	2DGULRy		[c] : 2dhgln + h ...	2-dehydro-L-gulo...			YiaE
14	3HCINNMH		[c] : 3hcinnm + h ...	3-hydroxycinnam...			MhpA
15	3HPPPNH		[c] : 3hpppn + h +...	3-(3-hydroxy-phe...			MhpA
16	3KGK		[c] : 3dhgln + at...	3-keto-L-gulonat...			LyxK

Search
Update
Insert
Delete
Copy
paste

Likewise, you can add a new reaction into the reference.

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - iAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

	ABBREVIATION	EC_NUMBER	EQUATION	KEGGID	NAME	NOTE
1221	Biomass_Ecoli_cor...		[c] : (1.496) 3pg + (3...		core E. coli biomas...	
1222	biomass_SA_2a		[c] : (0.42) 12dgr_E...		SA biomass 2 witho...	
1223	biomass_SA_2b		[c] : (0.42) 12dgr_E...		SA biomass 2 witho...	
1224	biomass_SA_3a		[c] : (0.42) 12dgr_E...		SA biomass 3 witho...	
1225	biomass_SA_3b		[c] : (0.42) 12dgr_E...		SA biomass 3 witho...	
1226	biomass_SA_4a		[c] : (0.42) 12dgr_S...		SA biomass 4 witho...	
1227	biomass_SA_5a		[c] : (0.42) 12dgr_S...		SA biomass 5 witho...	
1228	biomass_SA_6a		[c] : (0.42) 12dgr_S...		SA biomass 6 witho...	
1229	biomass_SA_6b		[c] : (0.42) 12dgr_S...		SA biomass 6 witho...	
1230	biomass_SA_7a		[c] : (0.01) 10fthf + (...)		SA biomass 7 witho...	
1231	biomass_SA_7b		[c] : (0.01) 10fthf + (...)		SA biomass 7 witho...	
1232	biomass_SA_8a		[c] : (0.01) 10fthf + (...)		SA biomass 8 witho...	

Search
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
The reaction of Ec_biomass_iAF1260_core_59p81M can be added into the reference database.

Abbreviation: Ec_biomass_iAF1260_core_59p81M


Equation:

(0.000223) 10fthf[c] + (0.000223) 2ohph[c] + (0.5137) ala-L[c] + (0.000223) amet[c] + (0.2958) arg-L[c] + (0.2411) asn-L[c] + (0.2411) asp-L[c] + (59.984) atp[c] + (0.004737) ca2[c] + (0.004737) cl[c] + (0.000576) coa[c] + (0.003158) cobalt2[c] + (0.1335) ctp[c] + (0.003158) cu2[c] + (0.09158) cys-L[c] + (0.02617) datp[c] + (0.02702) dctp[c] + (0.02702) dgtp[c] + (0.02617) dttp[c] + (0.000223) fad[c] + (0.007106) fe2[c] + (0.007106) fe3[c] + (0.2632) gln-L[c] + (0.2632) glu-L[c] + (0.6126) gly[c] + (0.2151) gtp[c] + (54.462) h2o[c] + (0.09474) his-L[c] + (0.2905) ile-L[c] + (0.1776) k[c] + (0.01945) kdo2lipid4[e] + (0.4505) leu-L[c] + (0.3432) lys-L[c] + (0.1537) met-L[c] + (0.007895) mg2[c] + (0.000223) mlthf[c] + (0.003158) mn2[c] + (0.003158) mobd[c] + (0.01389) murein5px4p[p] + (0.001831) nad[c] + (0.000447) nadp[c] + (0.011843) nh4[c] + (0.04148) pe160[p] + (0.02233) pe160[c] + (0.02632) pe161[c] + (0.04889) pe161[p] + (0.1759) phe-L[c] + (0.000223) pheme[c] + (0.2211) pro-L[c] + (0.000223) pydx5p[c] + (0.000223) ribflv[c] + (0.2158) ser-L[c] + (0.000223) sheme[c] + (0.003948) so4[c] + (0.000223) thf[c] + (0.000223) thmpp[c] + (0.2537) thr-L[c] + (0.05684) trp-L[c] + (0.1379) tyr-L[c] + (0.000055) udcpd[c] + (0.1441) utp[c] + (0.4232) val-L[c] + (0.003158) zn2[c] --> (59.81) adp[c] + (59.81) h[c] + (59.806) pi[c] + (0.7739) ppi[c]

Insert X

 Click Submit to insert this record, or Cancel

ABBREVIATION:	<input type="text" value="F1260_core_59p81M"/>	EC_NUMBER:	<input type="text"/>
EQUATION	<input type="text" value="pi[c] + (0.7739) ppi[c]"/>	KEGGID:	<input type="text"/>
NAME:	<input type="text"/>	NOTE:	<input type="text"/>
VALIDATION	<input type="text"/>		

 GEMSiRV

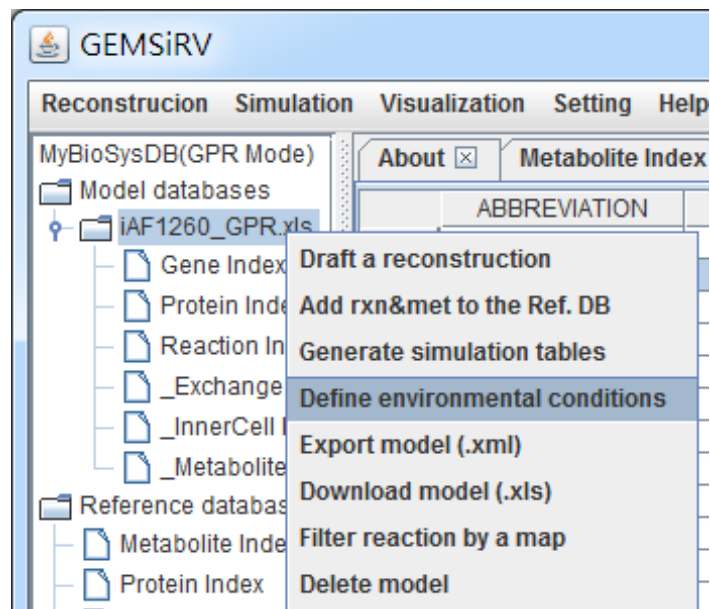
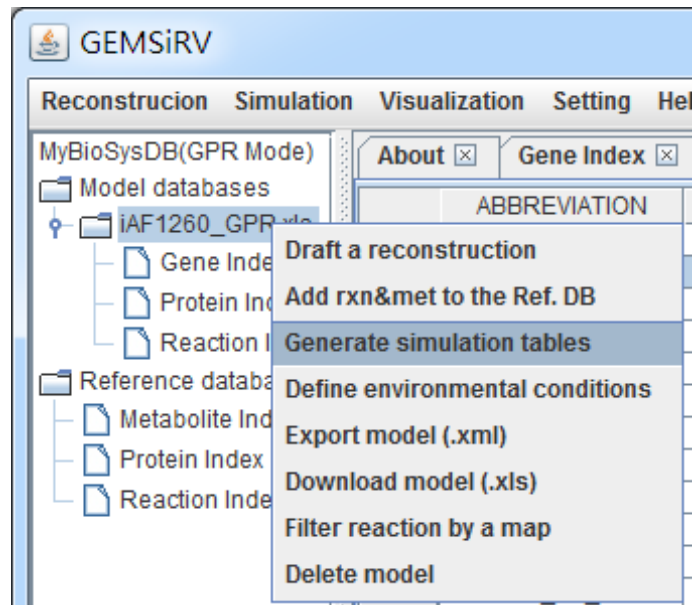
Reconstruction Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - iAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

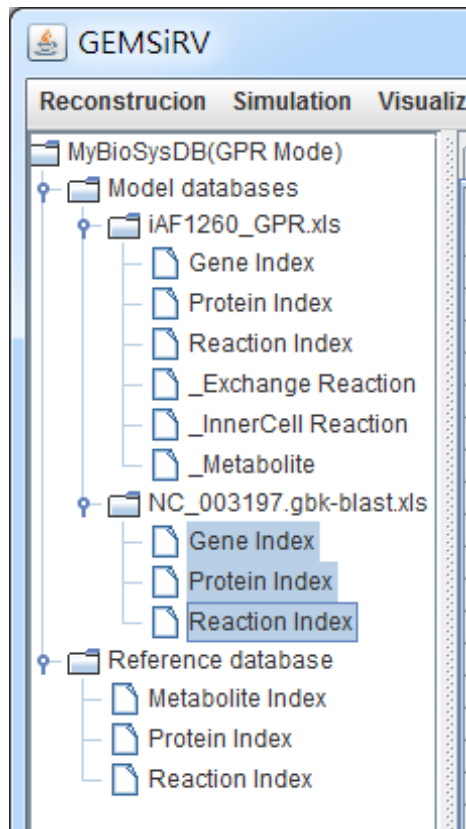
	ABBREVIATION	EC_NUMBER	EQUATION	KEGGID	NAME	NOTE
1221	Biomass_Ecoli_cor...		[c] : (1.496) 3pg + (3...		core E. coli biomas...	
1222	Ec_biomass_iAF12...		(0.000223) 10fthf[c] ...			M
1223	biomass_SA_2a		[c] : (0.42) 12dgr_E...		SA biomass 2 witho...	
1224	biomass_SA_2b		[c] : (0.42) 12dgr_E...		SA biomass 2 witho...	
1225	biomass_SA_3a		[c] : (0.42) 12dgr_E...		SA biomass 3 witho...	
1226	biomass_SA_3b		[c] : (0.42) 12dgr_E...		SA biomass 3 witho...	
1227	biomass_SA_4a		[c] : (0.42) 12dgr_S...		SA biomass 4 witho...	
1228	biomass_SA_5a		[c] : (0.42) 12dgr_S...		SA biomass 5 witho...	
1229	biomass_SA_6a		[c] : (0.42) 12dgr_S...		SA biomass 6 witho...	
1230	biomass_SA_6b		[c] : (0.42) 12dgr_S...		SA biomass 6 witho...	
1231	biomass_SA_7a		[c] : (0.01) 10fthf + (...)		SA biomass 7 witho...	
1232	biomass_SA_7b		[c] : (0.01) 10fthf + (...)		SA biomass 7 witho...	
1233	biomass_SA_8a		[c] : (0.01) 10fthf + (...)		SA biomass 8 witho...	

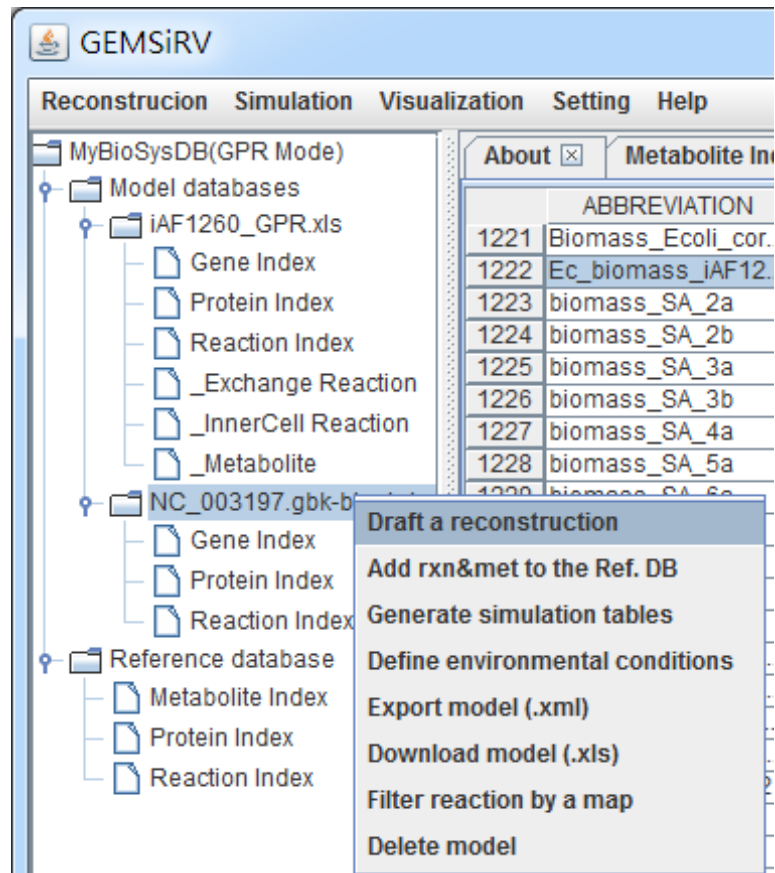
Right click on a reconstruction to Generate simulation tables can convert the reconstruction to a model. Then you can set the system boundaries for simulation.



Draft reconstruction and network refinement

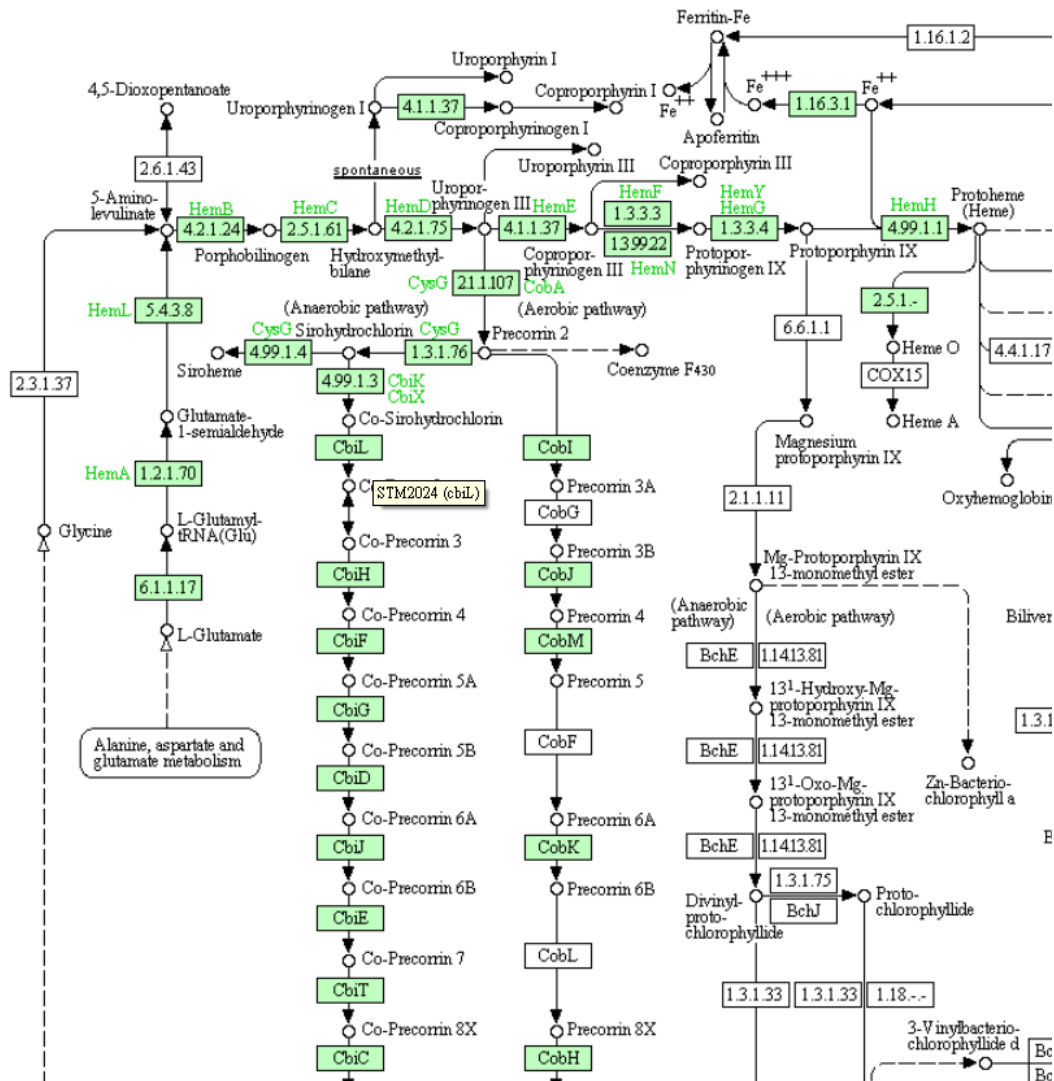
As described previously, we can draft a reconstruction for a genetically related species (e.g. *Salmonella*) with the existing *E. coli* model in GEMSiRV. Therefore, we import the file NC_003197.gbk-blast.xls and draft a reconstruction with reference to iAF1260_GPR.





Then we can refine the draft reconstruction by adding metabolic reactions with gene-protein-reaction associations, some existing reactions in the reference database can be conveyed to the reconstruction. For example, *Salmonella* is reported to be able to synthesize cobalamin due to its metabolic genes (operon) STM2016-STM2035. Therefore, we can manually add those associated reactions and proteins to the draft reconstruction.

PORPHYRIN AND CHLOROPHYLL METABOLISM



Locus

STM2024

Gene

cblL

Protein

CblL

Reaction

CPC2MT

Reaction CPC2MT

Name precorrin-2 C20-methyltransferase

Equation [c] : amet + copre2 --> ahcys + copre3 + h

After clicking into the Protein Index of NC_003197.gbk-blast.xls, right click on the main window of protein index to insert the protein

abbreviation CbiL, the associated gene STM2024 and a note Added to synthesize cobalamin.

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - iAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - NC_003197.gbk-blast.xls
 - Gene Index
 - Protein Index
 - Reaction Index
 - _Invalid Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

	Abbreviation	Gene	Name	Note	Other
0	Aas	STM3010	acyl-ACp synthase	Draft from iAF1260_GP...	AAC
1	Acc	STM0232+STM2366+S...	AcetylCoA carboxylase	Draft from iAF1260_GP...	ACC
2	AceA	STM4184	isocitrate lyase	Draft from iAF1260_GP...	ICL
3	AceB	STM4183	malate synthase A	Draft from iAF1260_GP...	MAL
4	AceEec	STM0152	Pyruvate Dehydrogenas...	Draft from iAF1260_GP...	PDF
5	AceFec	STM0153	Pyruvate dehydrogenas...	Draft from iAF1260_GP...	PDF
6	AckA	STM2337	Acetate kinase	Draft from iAF1260_GP...	ACK
7	AcnA	STM1712	aconitase A	Draft from iAF1260_GP...	ACC
8	AcnB	STM0158	aconitase B	Draft from iAF1260_GP...	MICI
9	AcpH	STM0403	ACP phosphodiesterase	Draft from iAF1260_GP...	FA1I
10	AcpP	STM1196	acyl carrier protein (ACP)	Draft from iAF1260_GP...	AAC
11	AcpS	STM2577	Holo-acp synthase	Draft from iAF1260_GP...	ACP
12	AcpT	STM3583	holo-(acyl carrier protei...	Draft from iAF1260_GP...	ACP
13	AcrEF	STM3390+STM3391	multidrug transport prot...	Draft from iAF1260_GP...	INDI
14	Acs	STM4275	acetyl-CoA synthetase	Draft from iAF1260_GP...	ACS
15	ActP	STM4273	Acetate Permease (Na...	Draft from iAF1260_GP...	ACTP
16	Add	STM1463	adenosine deaminase	Draft from iAF1260_GP...	ADA
17	AdhE	STM1749	Acetaldehyde-CoA dehy...	Draft from iAF1260_GP...	ACA
18	AdhP	STM1567	alcohol dehydrogenase	Draft from iAF1260_GP...	ALC
19	AdiA	STM4296	arginine decarboxylase	Draft from iAF1260_GP...	ARG
20	AdiC	STM4294	Arginine/agmatine anti...	Draft from iAF1260_GP...	ARG
21	Adk	STM0488	Adenylate kinase	Draft from iAF1260_GP...	ADK

Search
Update
Insert
Delete
Copy
paste

Update

Click Submit to modify this record, or Cancel

Abbreviation: CbiL

Note: synthesize cobalamin

Gene: STM2024

Other:

Submit **Cancel**

The gene-protein association will be automatically brought into the Gene Index table.

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

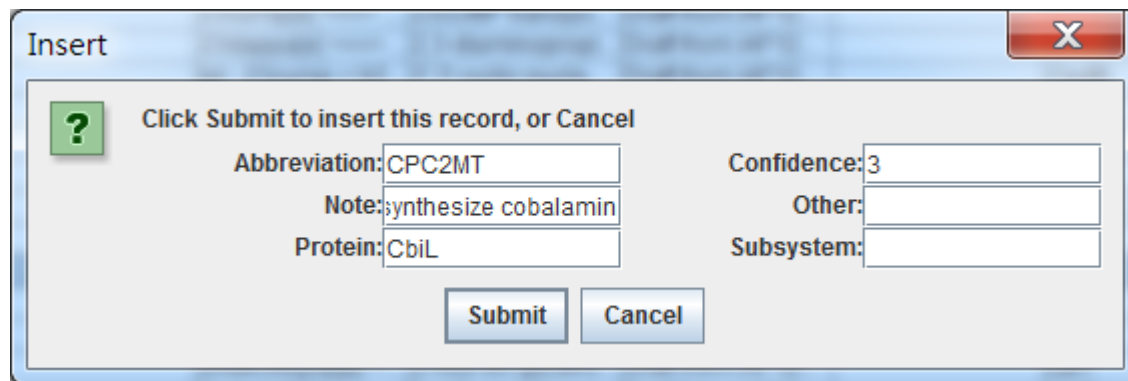
MyBioSysDB(GPR Mode)

- Model databases
 - iAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - NC_003197.gbk-blast.xls
 - Gene Index**
 - Protein Index
 - Reaction Index
 - _Invalid Reaction Index

	5'Coordinate	EC Number	Gene	Locus Tag	Note	Other	Protein
1946	2104417	2.1.1.130	cbiL	STM2024			CbiL
1947	2105208	--	cbiK	STM2025			
1948	2106002	1.3.1.54	cbiJ	STM2026			
1949	2106724	2.1.1.131	cbiH	STM2027			
1950	2107779	--	cbiG	STM2028			
1951	2108533	2.1.1.133	cbiF	STM2029			
1952	2109095	--	cbiT	STM2030			
1953	2109690	2.1.1.132	cbiE	STM2031			
1954	2110823	--	cbiD	STM2032			
1955	2111455	5.4.1.2	cbiC	STM2033			
1956	2112425	--	cobD	STM2034			
1957	2113801	--	cbiA	STM2035			
1958	2115310	--	pocR	STM2036			
1959	2116321	--	pduF	STM2037			
1960	2116846	--	pduA	STM2038			

After clicking into the Reaction Index of NC_003197.gbk-blast.xls, right click on the main window of reaction index to insert the reaction abbreviation CPC2MT, the associated protein CbiL, a note Added to synthesize cobalamin and the confidence score 3 for genetic

evidence.



The image shows a software dialog box titled "Insert" with a red close button (X) in the top right corner. Inside the dialog, there is a green square icon with a white question mark. To the right of the icon is the text "Click Submit to insert this record, or Cancel". Below this text are four input fields arranged in two columns. The left column contains "Abbreviation: CPC2MT", "Note: synthesize cobalamin", and "Protein: CbiL". The right column contains "Confidence: 3", "Other:", and "Subsystem:". At the bottom of the dialog are two buttons: "Submit" and "Cancel".

Abbreviation:	CPC2MT	Confidence:	3
Note:	synthesize cobalamin	Other:	
Protein:	CbiL	Subsystem:	

Submit Cancel

The reaction information including name and equation will be automatically brought into the Reaction Index table.

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - iAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - NC_003197.gbk-blast.xls
 - Gene Index
 - Protein Index
 - Reaction Index
 - _Invalid Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

	Abbreviation	Confidence	Equation	Name	Note	Other	Protein
0	12PPDRtex		12ppd-R[e] <==> ...	(R)-Propane-1,2-...	Draft from iAF12...		OmpC, PhoE, O...
1	12PPDStex		12ppd-S[e] <==> ...	(S)-Propane-1,2-...	Draft from iAF12...		OmpN, PhoE, O...
2	14GLUCANabcpp		14glucan[p] + atp...	1,4-alpha-D-gluc...	Draft from iAF12...		MalG+MalE+Mal...
3	14GLUCANtexi		14glucan[e] --> 1...	1,4-alpha-D-gluc...	Draft from iAF12...		LamB
4	23CAMPtex		23camp[e] <==> ...	23cAMP transpor...	Draft from iAF12...		OmpC, PhoE, O...
5	23CCMPtex		23ccmp[e] <==> ...	23cCMP transpo...	Draft from iAF12...		OmpC, OmpN, P...
6	23CGMPtex		23cgmp[e] <==> ...	23cGMP transpo...	Draft from iAF12...		PhoE, OmpC, O...
7	23CUMPtex		23cump[e] <==> ...	23cUMP transpo...	Draft from iAF12...		OmpN, PhoE, O...
8	23DAPPAtex		23dappa[e] <==> ...	2,3-diaminopropi...	Draft from iAF12...		OmpC, PhoE, O...
9	23PDE2pp		[p] : 23cump + h2...	2',3'-cyclic-nucle...	Draft from iAF12...		CpdB
10	23PDE4pp		[p] : 23ccmp + h2...	2',3'-cyclic-nucle...	Draft from iAF12...		CpdB
11	23PDE7pp		[p] : 23camp + h2...	2',3'-cyclic-nucle...	Draft from iAF12...		CpdB
12	23PDE9pp		[p] : 23cgmp + h2...	2',3'-cyclic-nucle...	Draft from iAF12...		CpdB
13	26DAHtex		26dap-M[e] <==> ...	meso-2,6-Diami...	Draft from iAF12...		OmpC, OmpN, P...
14	2AGPA120tipp		2ddecg3p[p] --> ...	2-Acyl-sn-glycero...	Draft from iAF12...		LpIT
15	CPC2MT	3	[c] : amet + copre...	precorrin-2 C20-...	Added to synthes...		CbiL
16	2AGPA140tipp		2tdecg3p[p] --> 2...	2-Acyl-sn-glycero...	Draft from iAF12...		LpIT
17	2AGPA141tipp		2tdec7eg3p[p] --...	2-Acyl-sn-glycero...	Draft from iAF12...		LpIT
18	2AGPA160tipp		2hdecg3p[p] --> ...	2-Acyl-sn-glycero...	Draft from iAF12...		LpIT
19	2AGPA161tipp		2hdec9eg3p[p] --...	2-Acyl-sn-glycero...	Draft from iAF12...		LpIT
20	2AGPA180tipp		2odecg3p[p] --> ...	2-Acyl-sn-glycero...	Draft from iAF12...		LpIT
21	2AGPA181tipp		2odec11eq3p[p] ...	2-Acyl-sn-glycero...	Draft from iAF12...		LpIT

Likewise, the protein-reaction association will be automatically brought into the Reaction Index table.

GEMSiRV

Reconstrucion Simulation Visualization Setting Help

MyBioSysDB(GPR Mode)

- Model databases
 - iAF1260_GPR.xls
 - Gene Index
 - Protein Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
 - NC_003197.gbk-blast.xls
 - Gene Index
 - Protein Index**
 - Reaction Index
 - _Invalid Reaction Index
- Reference database
 - Metabolite Index
 - Protein Index
 - Reaction Index

	Abbreviation	Gene	Name	Note	Reaction
0	Aas	STM3010	acyl-ACp synthase	Draft from iAF1260_GP...	AACPS1,AACPS2,AACP...
1	Acc	STM0232+STM2366+S...	AcetylCoA carboxylase	Draft from iAF1260_GP...	ACCOAC
2	AceA	STM4184	isocitrate lyase	Draft from iAF1260_GP...	ICL
3	AceB	STM4183	malate synthase A	Draft from iAF1260_GP...	MALS
4	AceEec	STM0152	Pyruvate Dehdrogenas...	Draft from iAF1260_GP...	PDH
5	AceFec	STM0153	Pyruvate dehydrogenas...	Draft from iAF1260_GP...	PDH
6	AckA	STM2337	Acetate kinase	Draft from iAF1260_GP...	ACKr
7	AcnA	STM1712	aconitase A	Draft from iAF1260_GP...	ACONTa,ACONTb
8	AcnB	STM0158	aconitase B	Draft from iAF1260_GP...	MICITD,ACONTa,ACON...
9	AcpH	STM0403	ACP phosphodiesterase	Draft from iAF1260_GP...	FA100ACPHi,FA120AC...
10	AcpP	STM1196	acyl carrier protein (ACP)	Draft from iAF1260_GP...	AACPS1,AACPS2,AACP...
11	AcpS	STM2577	Holo-acp synthase	Draft from iAF1260_GP...	ACPS1
12	CbiL	STM2024	precorrin-2 C20-methyl...	Added to synthesize co...	CPC2MT
13	AcpT	STM3583	holo-(acyl carrier protei...	Draft from iAF1260_GP...	ACPS1
14	AcrEF	STM3390+STM3391	multidrug transport prot...	Draft from iAF1260_GP...	INDOLEt2pp
15	Acs	STM4275	acetyl-CoA synthetase	Draft from iAF1260_GP...	ACS
16	ActP	STM4273	Acetate Permease (Na...	Draft from iAF1260_GP...	Act4pp,GLYCLTt4pp
17	Add	STM1463	adenosine deaminase	Draft from iAF1260_GP...	ADA,DADA
18	AdhE	STM1749	Acetaldehyde-CoA dehy...	Draft from iAF1260_GP...	ACALD,ALCD2x
19	AdhP	STM1567	alcohol dehydrogenase	Draft from iAF1260_GP...	ALCD2x

Simulation

Before simulation, make sure you have set the path of linear programming solver.

To download GNU Linear Programming Kit (GLPK).

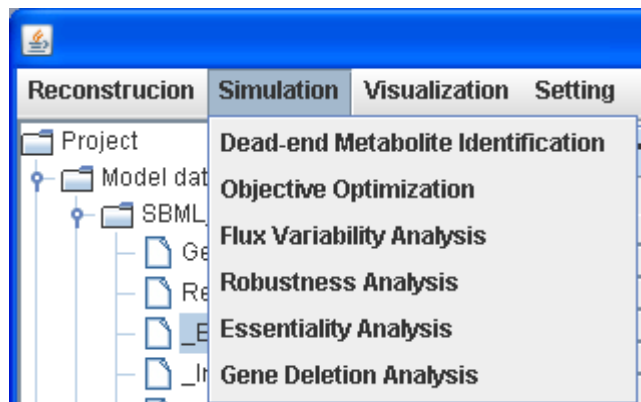
<http://sourceforge.net/projects/winglpk/> (for windows) or

<http://www.gnu.org/software/glpk/> (for Linux/Mac).

After extracting the file you downloaded (e.g. winglpk-4.45.zip), please add the path of glpsol.exe to your Environment variables.

Open the Control Panel -> Click System -> Click Advanced system setting -> Open Environment variables -> Edit Path -> Add variable value ";the path where glpsol.exe locate" (e.g. ;D:\winglpk-4.45\w64)

Click on Simulation in the menu bar to choose which analysis you want to perform.

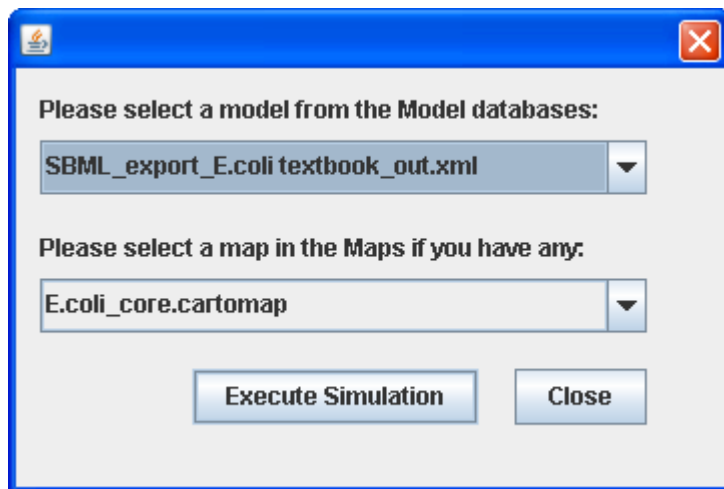
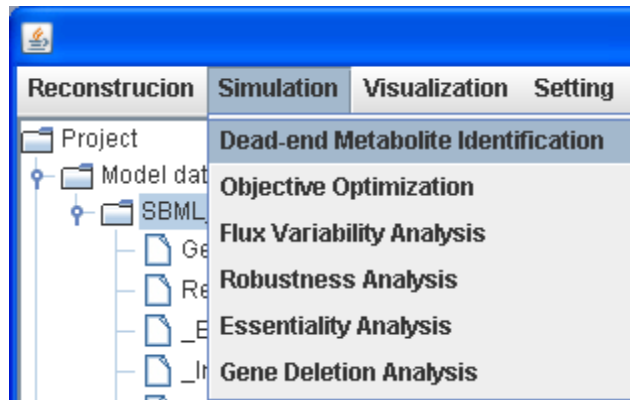


As a case study for demonstration of simulation, we import the E.coli textbook model which was exported from the BiGG into GEMSiRV and use a customized map E.coli_core.cartomap for visualization. You can find and download the model and the map from http://sb.nhri.org.tw/GEMSiRV/en/Metabolic_Models and http://sb.nhri.org.tw/GEMSiRV/en/Metabolic_Maps, respectively.

Dead-end metabolite identification

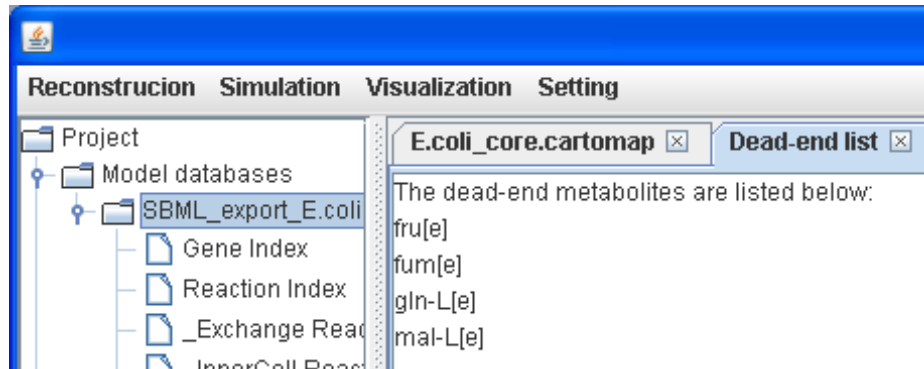
A network reconstruction is converted into a mathematical model including a stoichiometric matrix which describes the connectivity feature of the network and defined systems boundaries before simulation. GEMSiRV can examine the connectivity of all metabolites in a network for dead-end metabolite identification and tag such metabolic dead ends with crosses in the map.

You can select a model and a map (if you have) to perform the examination of network connectivity for dead-end metabolite identification.

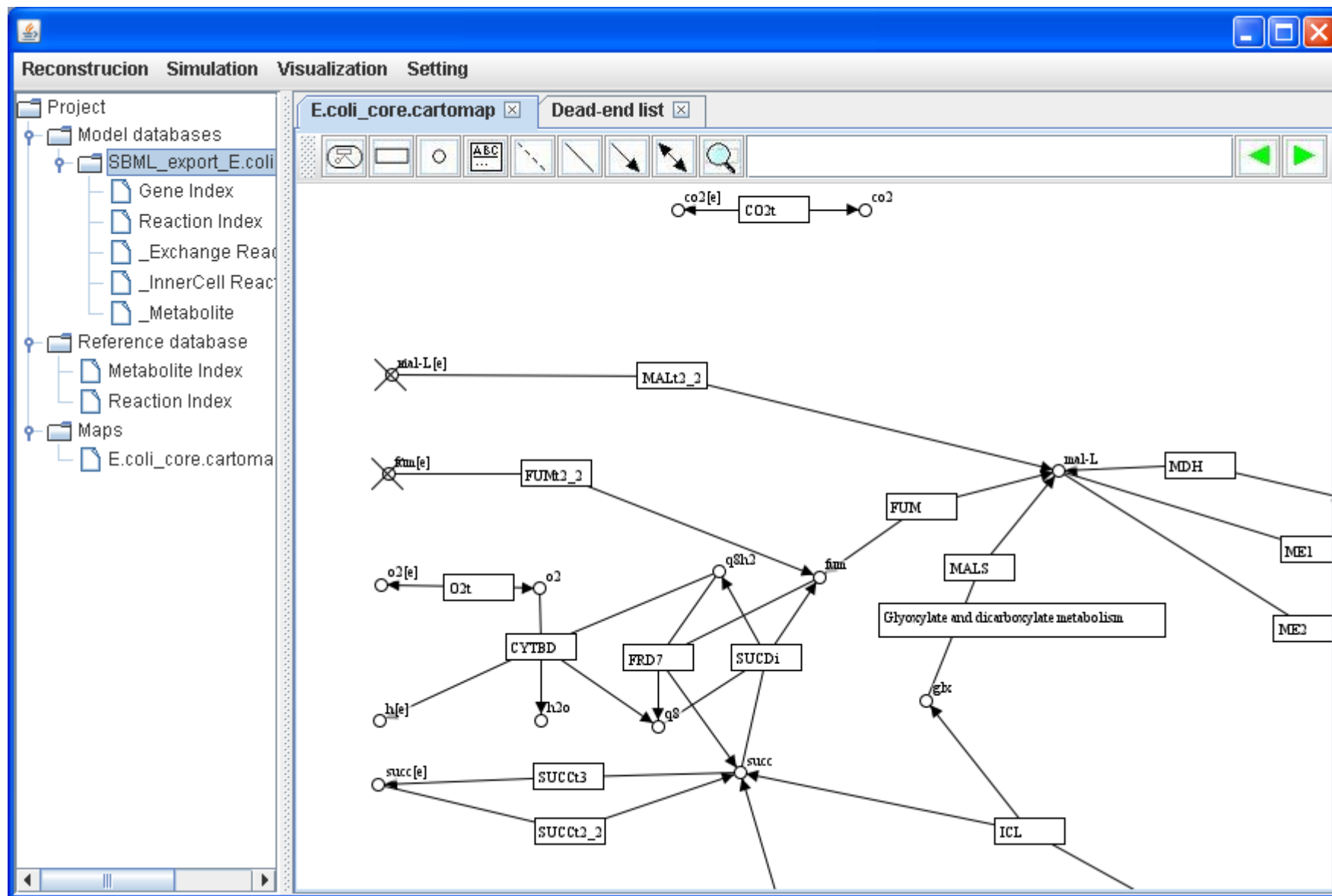


A dead-end metabolite list is generated and those metabolites are tagged with crosses in the map.

A list for dead-end metabolites:



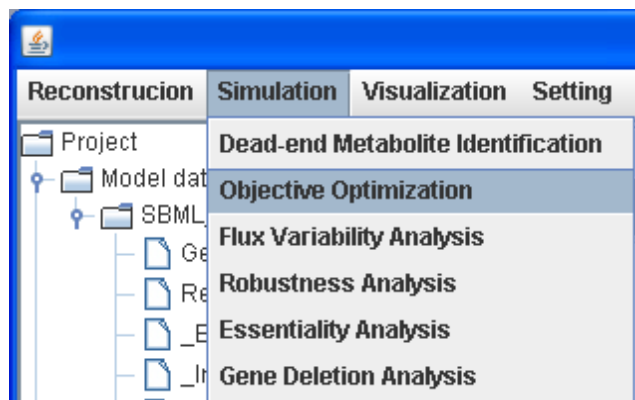
A visualization map with dead-end metabolites:

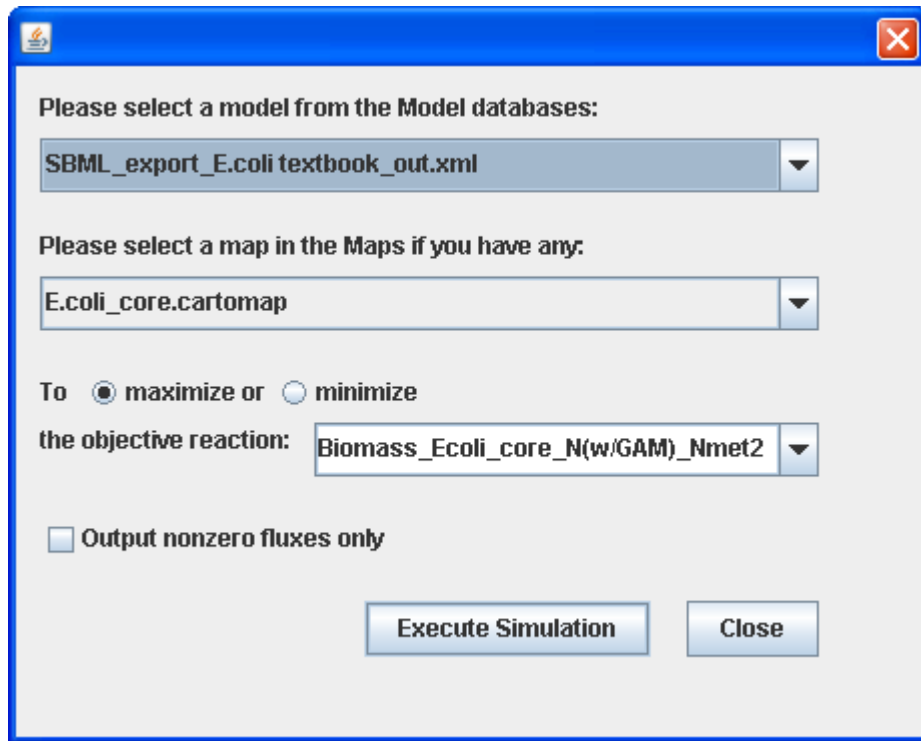


Objective optimization

With a set linear programming solver (e.g. glpk), GEMSiRV can be used to simulate the imported metabolic network model. Please refer to <http://sb.nhri.org.tw/GEMSiRV/en/Installation> for setting up GEMSiRV. Given proper constraints and objective function, the flux results of all reactions in the model will be estimated.

You can select a model and a map (if you have) for objective optimization. The flux results can be visualized in the map.





Please select a model from the Model databases:

SBML_export_E.coli_textbook_out.xml

Please select a map in the Maps if you have any:

E.coli_core.cartomap

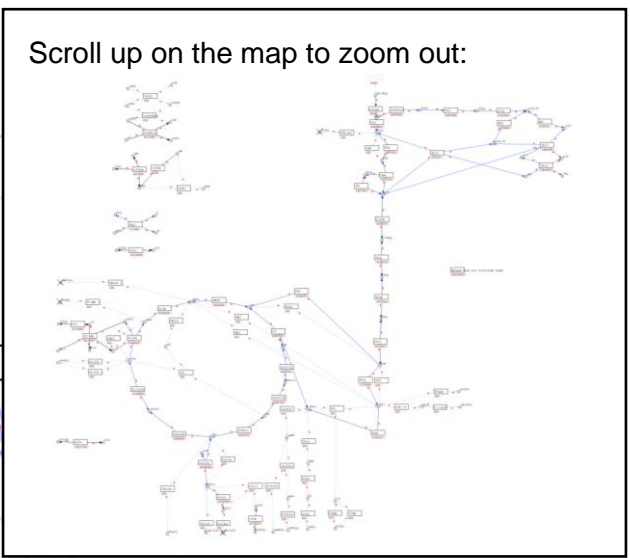
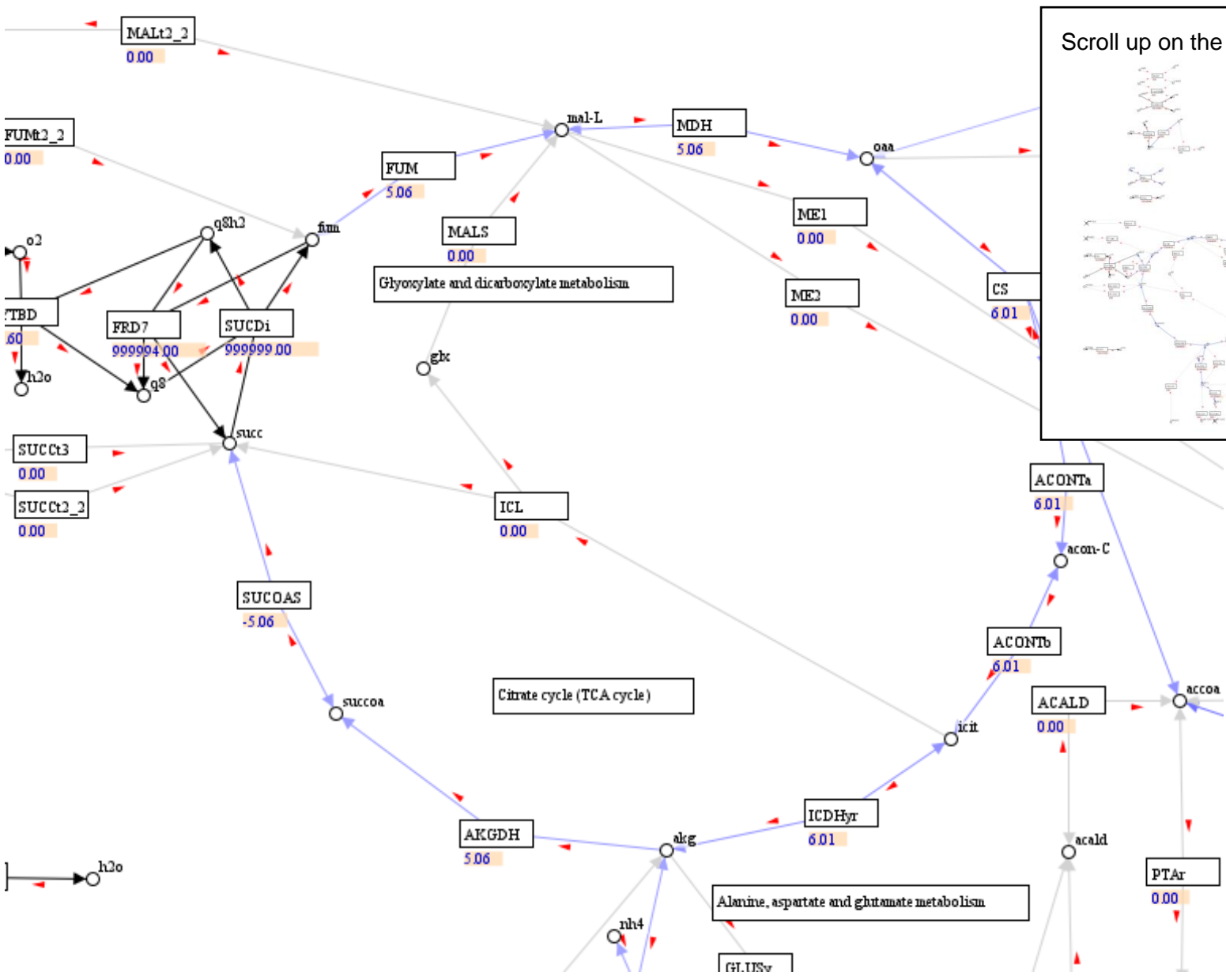
To ☒ maximize or ☐ minimize
the objective reaction:

Biomass_Ecoli_core_N(w/GAM)_Nmet2

☐ Output nonzero fluxes only

Execute Simulation Close

A visualization map with reaction fluxes:

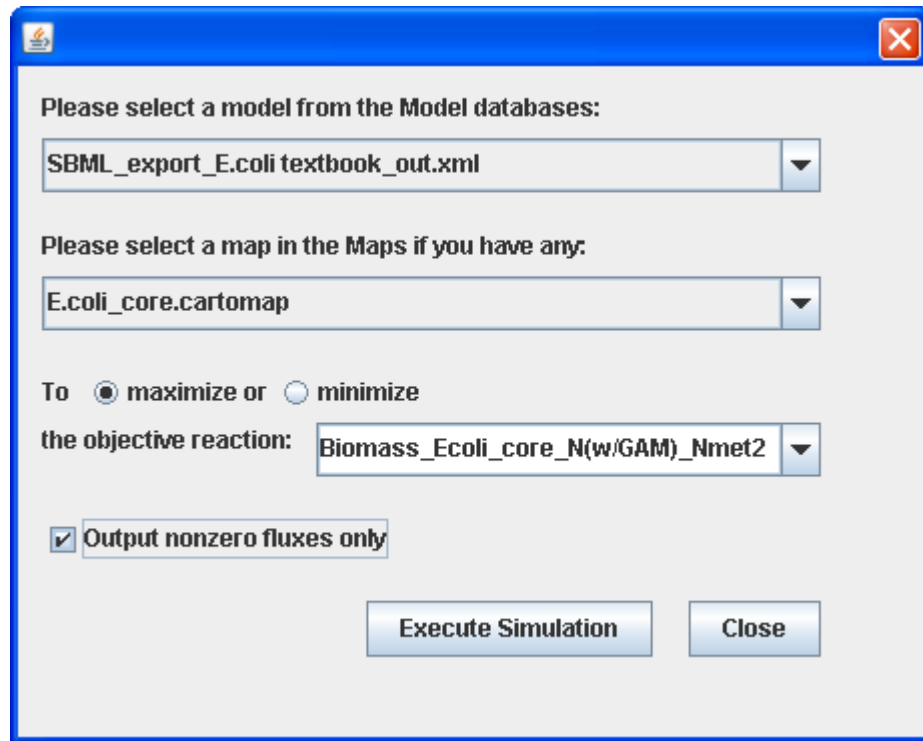


Flux result:



E.coli_core.cartomap	
Flux result	
save	
GLNabc	0.0
GLUDy	-4.54186
GLUN	0.0
GLUSy	0.0
GLUt2r	0.0
GND	4.95998
H2Ot	-29.1758
ICDHyr	6.00725
ICL	0.0
LDH_D	0.0
MAL8	0.0
MALt2_2	0.0
MALt	5.06422

To check the checkbox for outputting nonzero fluxes only.



Please select a model from the Model databases:

SBML_export_E.coli_textbook_out.xml ▼

Please select a map in the Maps if you have any:

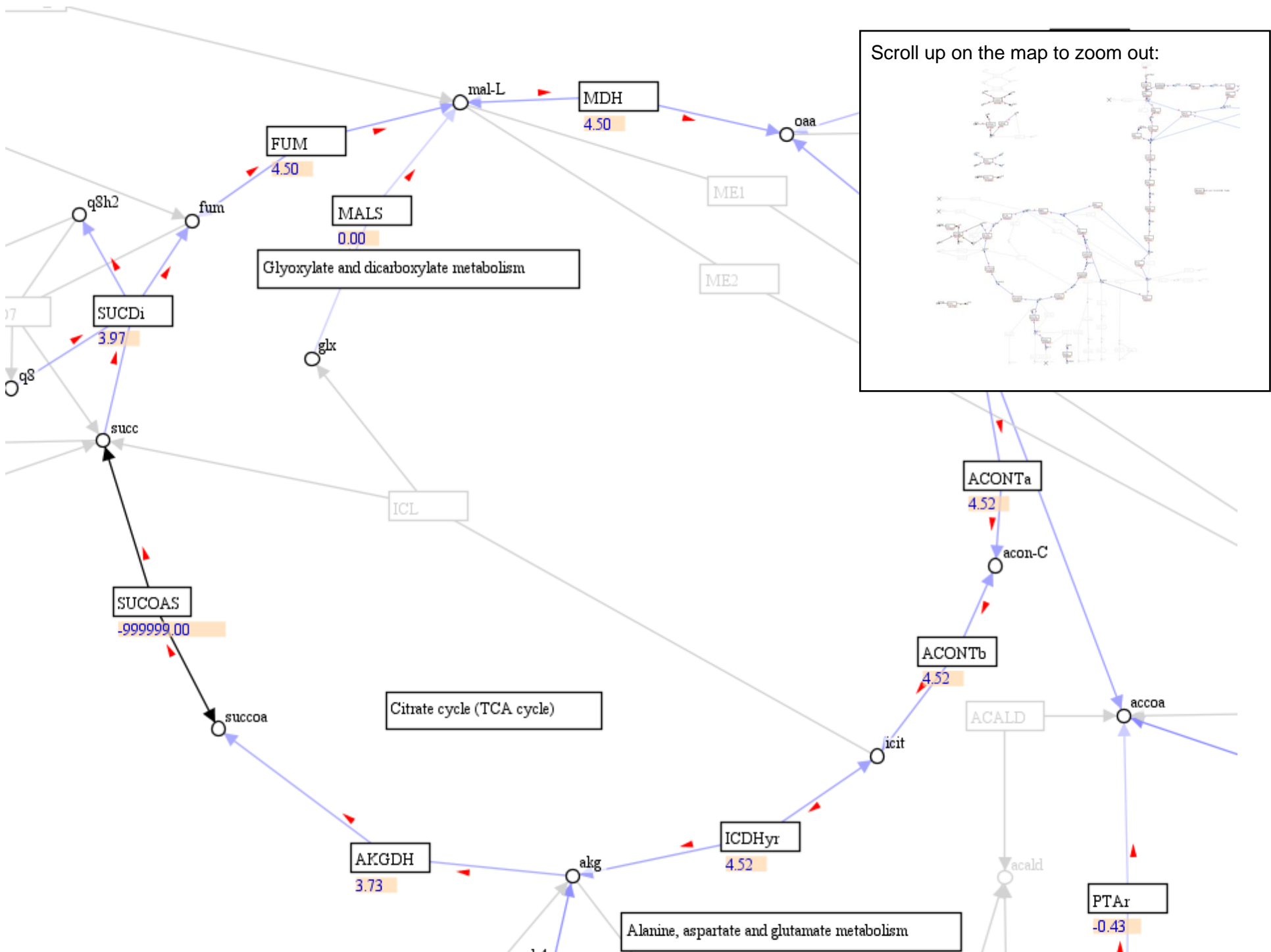
E.coli_core.cartomap ▼

To ☒ maximize or ☐ minimize
the objective reaction:

Biomass_Ecoli_core_N(w/GAM)_Nmet2 ▼

☒ Output nonzero fluxes only

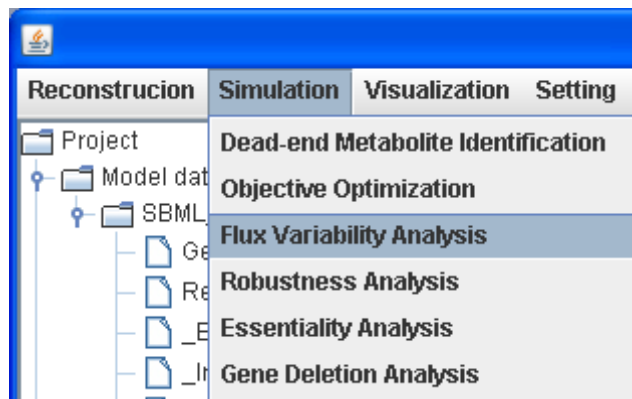
Execute Simulation Close



Flux variability analysis

Flux variability analysis can be used to study the redundancy of reactions in a network. GEMSiRV can determine the minimum and maximum flux values for each reaction in the model and thus identify the blocked reactions which carry zero fluxes for the both conditions and tag them with crosses in a map as well.

You can select a model and a map (if you have) for flux variability analysis. The min and max fluxes of reaction are plotted in the map and the blocked reaction are tagged with crosses.



Please select a model from the Model databases:

SBML_export_E.coli_textbook_out.xml ▼

Please select a map in the Maps if you have any:

E.coli_core.cartomap ▼

☒ To plot the result in the map

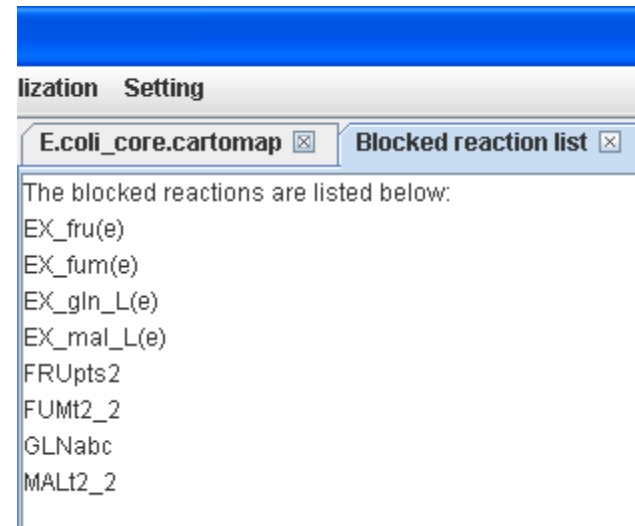
☒ To identify the blocked reactions

You can define the Min/Max flux cutoff on map.

Min: Max:

Please assign a reaction list file, if you won't run FVA for all reactions:

A list for the blocked reactions:

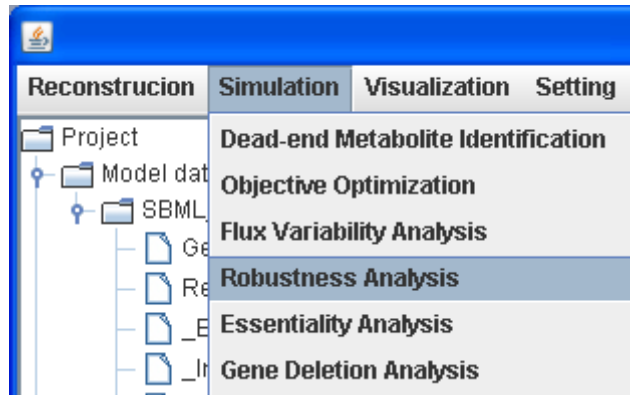


Flux variability result:

E.coli_core.cartomap <input type="button" value="x"/>			
Blocked reaction list <input type="button" value="x"/>			
Flux variability result <input type="button" value="x"/>			
<input type="button" value="save"/>			
#Reaction	Min	Max	
ACALD	-20.0	-1.169719405E-29	
ACALDt	-20.0	0.0	
ACKr	-20.0	-7.316790248E-29	
ACONTa	1.243854342E-28	20.0	
ACONTb	1.242966364E-28	20.0	
Act2r	-20.0	0.0	
ADK1	0.0	166.61	
AKGDH	0.0	20.0	
AKGt2r	-10.0	0.0	
ALCD2x	-20.0	-1.169719405E-29	
ATPM	8.39	8.39	
ATPS4r	-31.61	150.0	
Biomass_Ecoli_core_N(w/GAM)_Nmet2	0.0	0.873921507	
CO2t	-60.0	11.10424242	
CS	1.241908217E-28	20.0	
CYTBD	0.0	120.0	
D_LACT	20.0	0.0	

Robustness analysis

Robustness analysis can be used to study the effect of changing a reaction flux on the other reaction flux, especially on the objective of interest (e.g. growth rate). Therefore, you can select the reactions of interest in the model for robustness analysis.



To see how sensitive of the objective reaction (Biomass) is to the glucose uptake rate in the range of -20 to 0 mmol/gDW/h.

Please select a model from the Model databases:

SBML_export_E.coli_textbook_out.xml ▼

To ☒ maximize or ☐ minimize
the objective reaction: Biomass_Ecoli_core_N(w/GAM)_Nmet2 ▼

To see the robustness to reaction:

EX_glc(e) ▼ in ☐ Min and Max or ☒ -20 and 0

and reaction (optional):

▼ in ☒ Min and Max or ☐ -1000 and 1000

Number of points spaced in the flux range 20

Execute Simulation Close

Robustness analysis for growth rate maximization while changing glucose uptake rate (uptaking 0-20 mmol/gDW/h) with oxygen uptake fixed at 17 mmol/gDW/h (set LB to -17)



To see how sensitive of the objective reaction (Biomass) is to the oxygen uptake rate in the range of -25 to 0 mmol/gDW/h.

Please select a model from the Model databases:

SBML_export_E.coli_textbook_out.xml ▼

To ☒ maximize or ☐ minimize
the objective reaction: Biomass_Ecoli_core_N(w/GAM)_Nmet2 ▼

To see the robustness to reaction:

EX_o2(e) ▼ in ☐ Min and Max or ☒ -25 and 0

and reaction (optional):

▼ in ☒ Min and Max or ☐ -1000 and 1000

Number of points spaced in the flux range 20 ▼

Execute Simulation Close

Robustness analysis for growth rate maximization while changing oxygen uptake rate (uptaking 0-25 mmol/gDW/h) with glucose uptake fixed at 10 mmol/gDW/h (set LB to -10).



To change two reactions simultaneously. GEMSiRV can plot the results as a phenotypic phase plane.

Please select a model from the Model databases:

SBML_export_E.coli_textbook_out.xml ▼

To ☒ maximize or ☐ minimize
the objective reaction: Biomass_Ecoli_core_N(w/GAM)_Nmet2 ▼

To see the robustness to reaction:

EX_o2(e) ▼ in ☐ Min and Max or ☒ -20 and 0

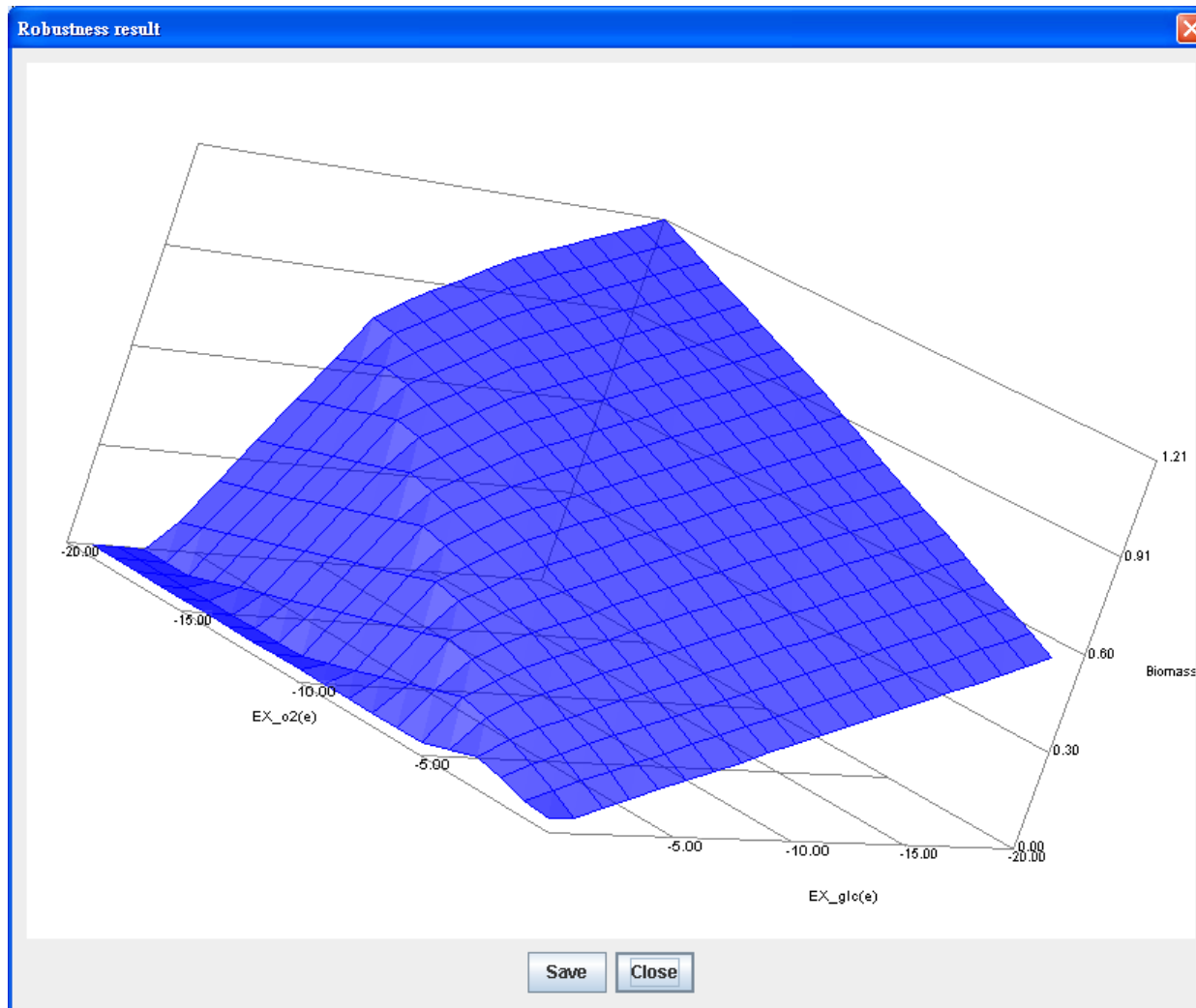
and reaction (optional):

EX_glc(e) ▼ in ☐ Min and Max or ☒ -20 and 0

Number of points spaced in the flux range 20 ▼

Execute Simulation Close

The phenotypic phase plane for growth rate maximization while changing glucose and oxygen uptake rates in the range of -20 to 0 mmol/gDW/h.

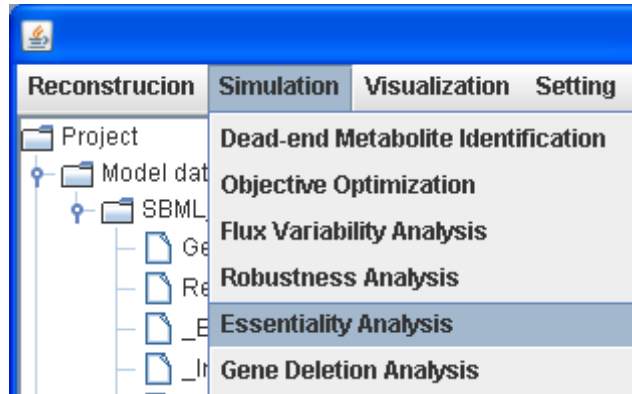


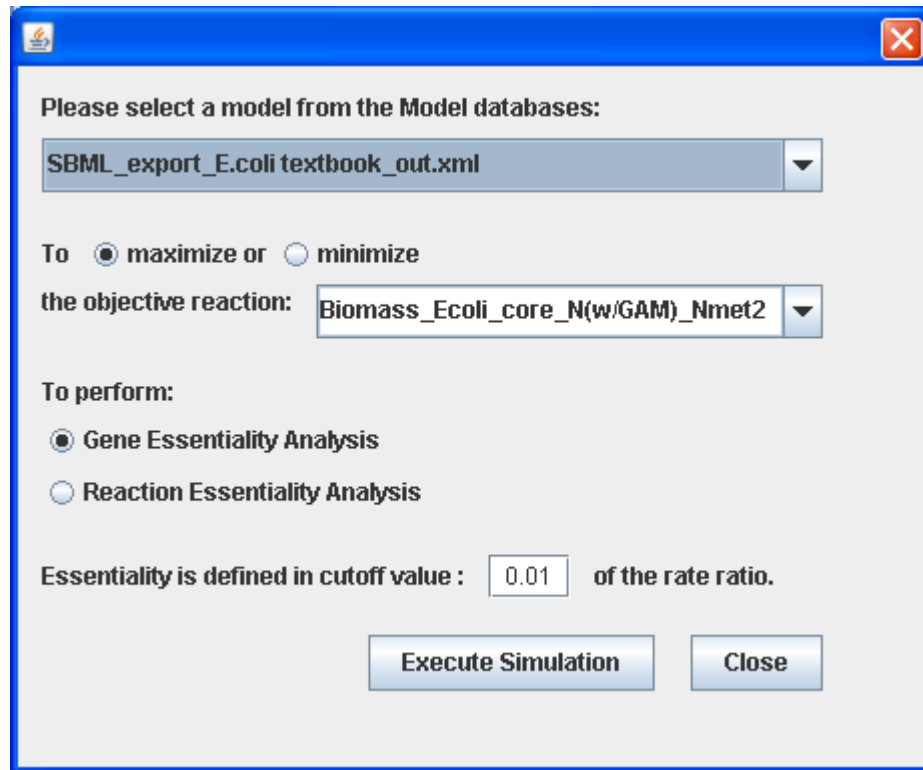
Essentiality analysis

To constrain a reaction in a zero flux can simulate the reaction deletion. Likewise, to constrain the reaction corresponding to a deleted gene can simulate the gene deletion. GEMSiRV performs essentiality analysis for gene and reaction separately and determines the rate

ratio (objective flux of deleted model to that of wild-type model) for every single-knockout condition.

You can select a metabolic model for essentiality analysis, the computational essential genes or reactions can be identified.





Please select a model from the Model databases:

SBML_export_E.coli textbook_out.xml ▼

To ☒ maximize or ☐ minimize
the objective reaction: Biomass_Ecoli_core_N(w/GAM)_Nmet2 ▼

To perform:

☒ Gene Essentiality Analysis
☐ Reaction Essentiality Analysis

Essentiality is defined in cutoff value : 0.01 of the rate ratio.

Execute Simulation Close

Results of gene essentiality analysis:

E.coli_core.cartomap		Essential gene result
Computationally essential genes:		
b0451		
b0720		
b1136		
b1779		
b2415		
b2416		
b2779		
b2926		
KO gene	Reaction	Rate ratio
b0114	[R_PDH]	0.9116332748634368
b0115	[R_PDH]	0.9116332748634368
b0116	[R_AKGDH, R_PDH]	0.895216
b0474	[R_ADK1]	1.0
b0721	[R_SUCDi]	0.9317741936519247
b0722	[R_SUCDi]	0.9317741936519247
b0723	[R_SUCDi]	0.9317741936519247
b0724	[R_SUCDi]	0.9317741936519247
b0726	[R_AKGDH]	0.9821332935796487
b0727	[R_AKGDH]	0.9821332935796487
b0728	[R_SUCOAS]	0.9821332935796487
b0729	[R_SUCOAS]	0.9821332935796487
b0767	[R_PGL]	0.9884335178628347
b0809	[R_GLNabc]	1.0
b0810	[R_GLNabc]	1.0

Results of reaction essentiality analysis:

E.coli_core.cartomap

Essential reaction result

Computationally essential reactions:

ACONTa

ACONTb

Biomass_Ecoli_core_N(w/GAM)_Nmet2

CS

ENO

EX_glc(e)

EX_h(e)

EX_nh4(e)

EX_pi(e)

GAPD

GLCpts

GLNS

ICDHyr

NH4t

PGK

PGM

Pit2r

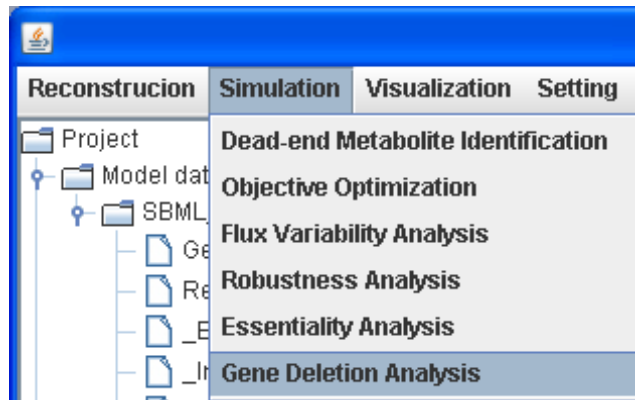
RPI

KO reaction	Rate ratio
ACALD	1.0
ACALDt	1.0
ACKr	1.0
Act2r	1.0
ADK1	1.0
AKGDH	0.9821332935796487

Gene deletion analysis

Gene deletion analysis is carried out by imposing a single-gene deletion or a set of gene deletions at a time, which simulates biological knockout mutant or transcriptional regulatory constraints. GEMSiRV performs the gene deletion analysis to generate the flux result and a

SBML model for the specified condition.



To delete a single gene:

Please select a model from the Model databases:

SBML_export_E.coli textbook_out.xml

To ☒ maximize or ☐ minimize
the objective reaction: Biomass_Ecoli_core_N(w/GAM)_Nmet2

To perform:

☒ Single Gene Deletion: b

☐ Gene Deletion:

☐ To export the model in

b0008
b0114
b0115
b0116
b0118
b0351
b0356
b0451

Browse...

Close

Or you can upload a list of genes for multiple-gene deletion.

To export the SBML file with single gene deletion by checking the checkbox.

Please select a model from the Model databases:

SBML_export_E.coli textbook_out.xml ▼

To ☒ maximize or ☐ minimize
the objective reaction: Biomass_Ecoli_core_N(w/GAM)_Nmet2 ▼

To perform:

☒ Single Gene Deletion: b0114 ▼

☐ Gene Deletion: Browse...

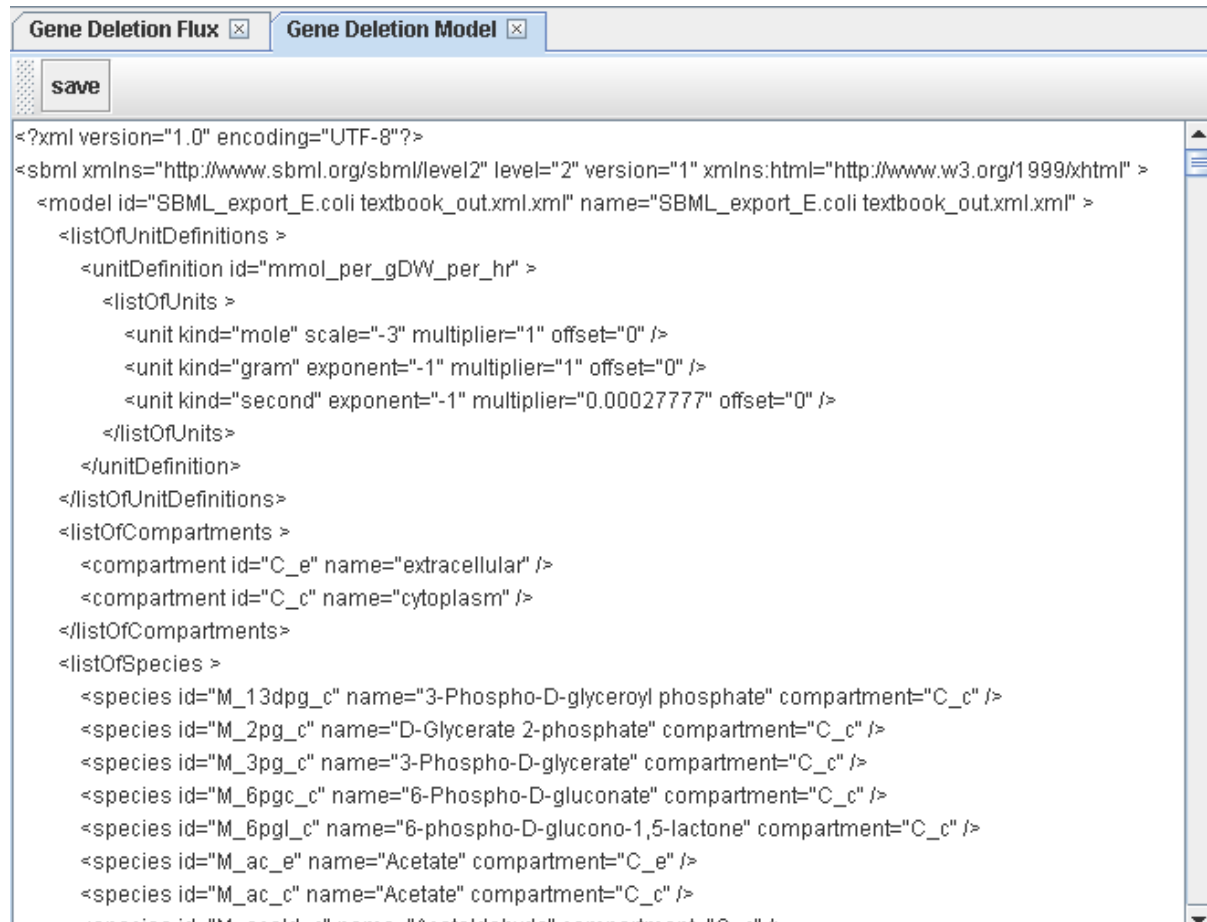
☒ To export the model in SBML format

Execute Simulation Close

The flux result for the single gene deletion model:

Gene Deletion Flux		Gene Deletion Model
save		
#Reaction	Flux	
ACALD	0.0	
ACALDt	0.0	
ACKr	0.0	
ACONTa	4.75726	
ACONTb	4.75726	
ACT2r	0.0	
ADK1	0.0	
AKGDH	3.89771	
AKGt2r	0.0	
ALCD2x	0.0	
ATPM	8.39	
ATPS4r	43.6718	
Biomass_Ecoli_core_N(w/GAM)_Nmet2	0.796696	
CO2t	-18.3531	
CS	4.75726	
CYTBD	42.6072	
D_LACT2	0.0	
ENO	12.6966	
ETOht2r	0.0	
EX_ac(e)	0.0	
EX_acald(e)	0.0	
EX_akg(e)	0.0	
EX_co2(e)	18.3531	
EX_etoh(e)	0.0	

The gene-deletion model in SBML format:



```

<?xml version="1.0" encoding="UTF-8"?>
<sbml xmlns="http://www.sbml.org/sbml/level2" level="2" version="1" xmlns:html="http://www.w3.org/1999/xhtml" >
  <model id="SBML_export_E.coli textbook_out.xml.xml" name="SBML_export_E.coli textbook_out.xml.xml" >
    <listOfUnitDefinitions >
      <unitDefinition id="mmol_per_gDW_per_hr" >
        <listOfUnits >
          <unit kind="mole" scale="-3" multiplier="1" offset="0" />
          <unit kind="gram" exponent="-1" multiplier="1" offset="0" />
          <unit kind="second" exponent="-1" multiplier="0.00027777" offset="0" />
        </listOfUnits>
      </unitDefinition>
    </listOfUnitDefinitions>
    <listOfCompartments >
      <compartment id="C_e" name="extracellular" />
      <compartment id="C_c" name="cytoplasm" />
    </listOfCompartments>
    <listOfSpecies >
      <species id="M_13dpg_c" name="3-Phospho-D-glyceroyl phosphate" compartment="C_c" />
      <species id="M_2pg_c" name="D-Glycerate 2-phosphate" compartment="C_c" />
      <species id="M_3pg_c" name="3-Phospho-D-glycerate" compartment="C_c" />
      <species id="M_6pgc_c" name="6-Phospho-D-gluconate" compartment="C_c" />
      <species id="M_6pgl_c" name="6-phospho-D-glucono-1,5-lactone" compartment="C_c" />
      <species id="M_ac_e" name="Acetate" compartment="C_e" />
      <species id="M_ac_c" name="Acetate" compartment="C_c" />
    </listOfSpecies>
  </model>
</sbml>

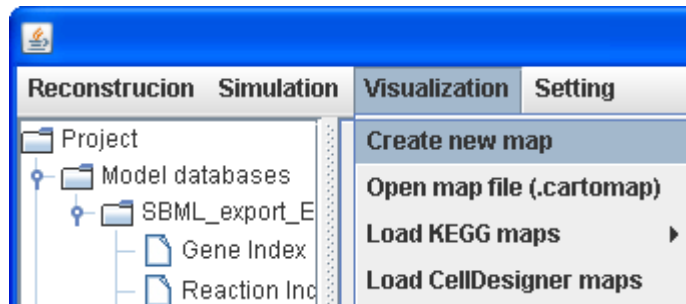
```

Such a model can be further imported into GEMSiRV for the other network evaluations as described early.

Visualization

Metabolic map creation

Click on Visualization in the menu bar to Create new map. You can create a metabolic map by clicking and moving network objects from the toolbar onto the main network view window.



The toolbar for creating/editing a map:



Add a map

Add a reaction

Add a metabolite

Add a label

Add a dotted line

Add an undirected line

Add a directed line

Add a bidirected line

Delete the selected item(s)

Search

Pan: You can pan a map by dragging and dropping left-click button over an empty point.

Zoom: You can zoom out or zoom in a map by scrolling up or down, respectively.

Select: You can click on an object to select it or you can hold right-click button to drag the mouse to select groups of objects.

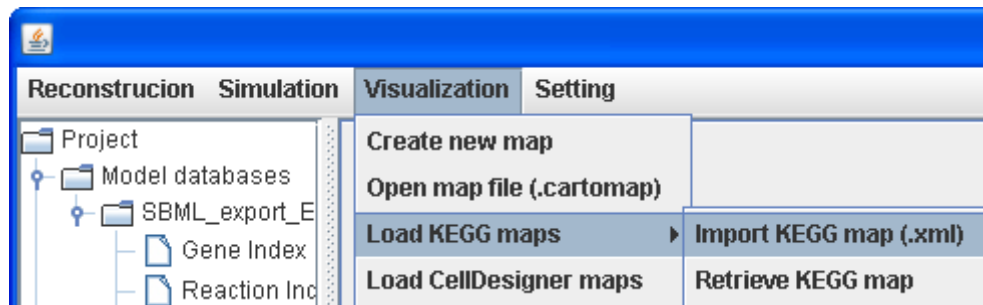
Move: You can move any selected object by dragging and dropping it.

Delete: After selecting objects, you can right click over the selected items or click the Delete button in the toolbar for deletion.

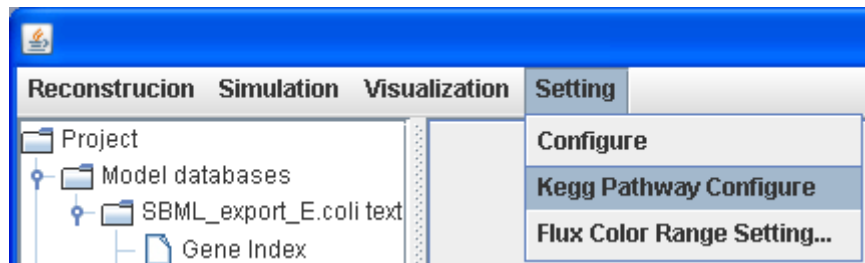
Merge: After selecting identical objects, you can right click over the selected items to merge.

KEGG map loading

Click on Visualization in the menu bar to Load KEGG maps by either Import KEGG map (.xml) or Retrieve KEGG map depending on whether you have KEGG maps in hand.



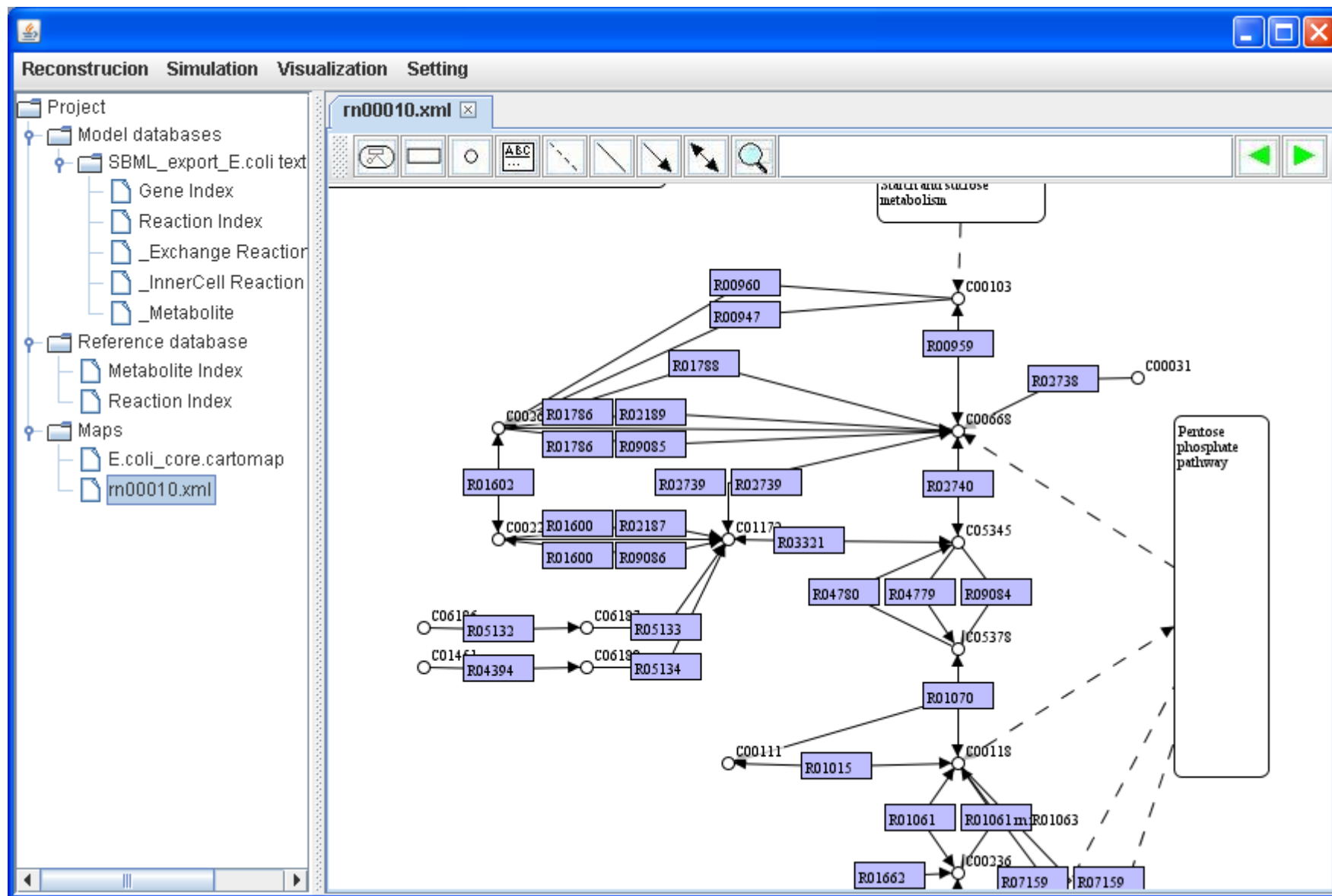
If not, you can click on Setting in the menu bar to KEGG pathway Configure and set the link to where the KEGG pathway maps can be retrieved, e.g. <http://www.genome.jp/kegg-bin/download>.



Then you can retrieve KEGG pathway by choosing from the KEGG Pathway List.

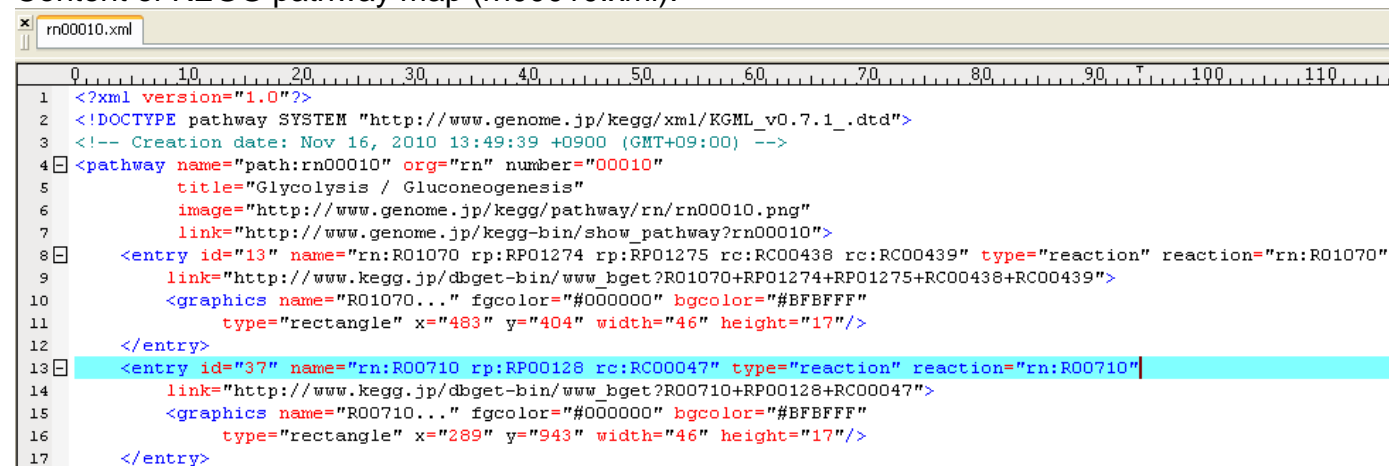


The KEGG pathway map of rn00010:



We set rectangular nodes to represent reactions and define node name and node caption for each reaction. We directly use the entry name and reaction in KEGG maps as the node name and node caption respectively. Therefore, you can decide to Show node name or Show node caption by right clicking on a map.

Content of KEGG pathway map (rn00010.xml):

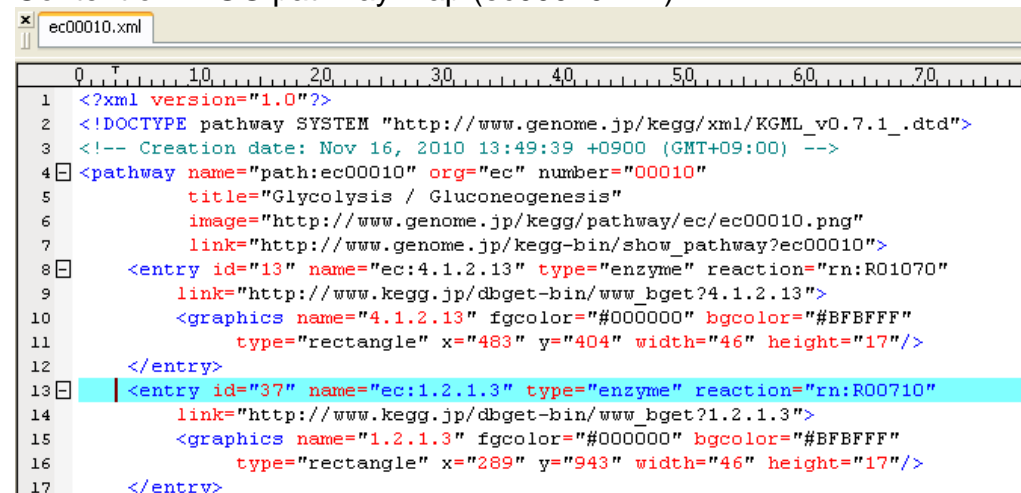


```

1  <?xml version="1.0"?>
2  <!DOCTYPE pathway SYSTEM "http://www.genome.jp/kegg/xml/KGML_v0.7.1_.dtd">
3  <!-- Creation date: Nov 16, 2010 13:49:39 +0900 (GMT+09:00) -->
4  <pathway name="path:rn00010" org="rn" number="00010"
5      title="Glycolysis / Gluconeogenesis"
6      image="http://www.genome.jp/kegg/pathway/rn/rn00010.png"
7      link="http://www.genome.jp/kegg-bin/show_pathway?rn00010">
8  <entry id="13" name="rn:R01070 rp:RP01274 rp:RP01275 rc:RC00438 rc:RC00439" type="reaction" reaction="rn:R01070"
9      link="http://www.kegg.jp/dbget-bin/www_bget?R01070+RP01274+RP01275+RC00438+RC00439">
10     <graphics name="R01070..." fgcolor="#000000" bgcolor="#BFBFFF"
11         type="rectangle" x="483" y="404" width="46" height="17"/>
12   </entry>
13  <entry id="37" name="rn:R00710 rp:RP00128 rc:RC00047" type="reaction" reaction="rn:R00710"
14      link="http://www.kegg.jp/dbget-bin/www_bget?R00710+RP00128+RC00047">
15     <graphics name="R00710..." fgcolor="#000000" bgcolor="#BFBFFF"
16         type="rectangle" x="289" y="943" width="46" height="17"/>
17   </entry>

```

Content of KEGG pathway map (ec00010.xml):

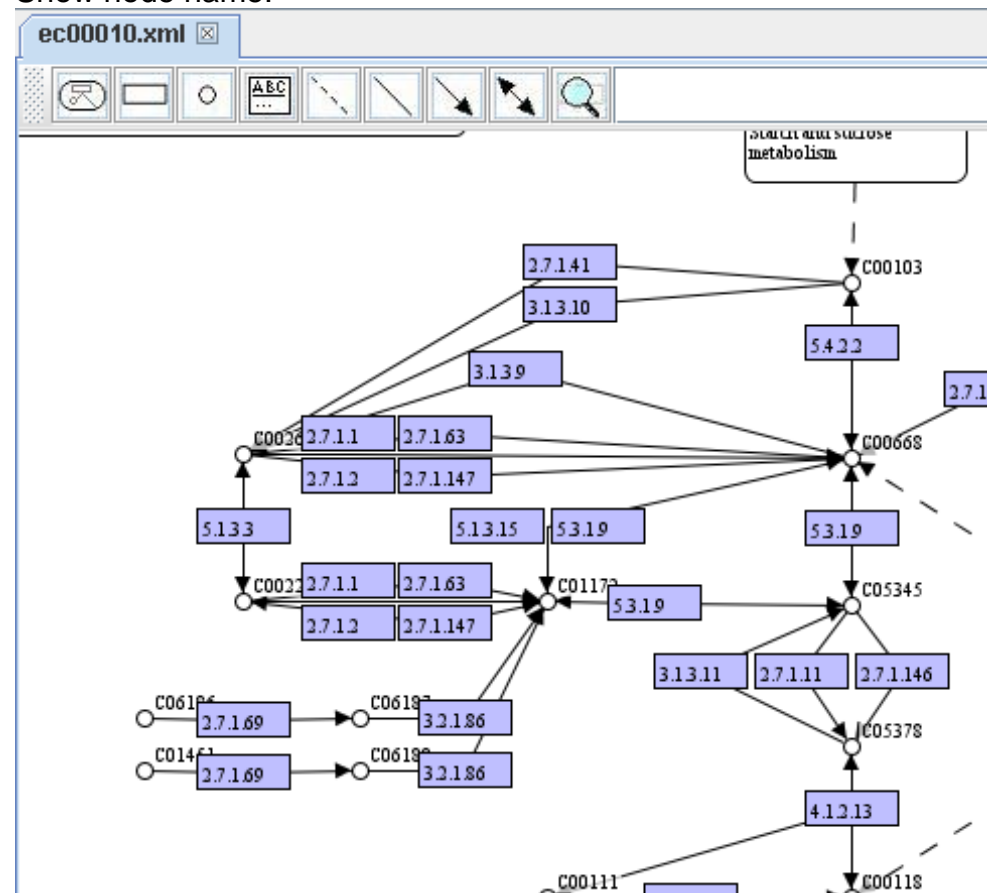


```

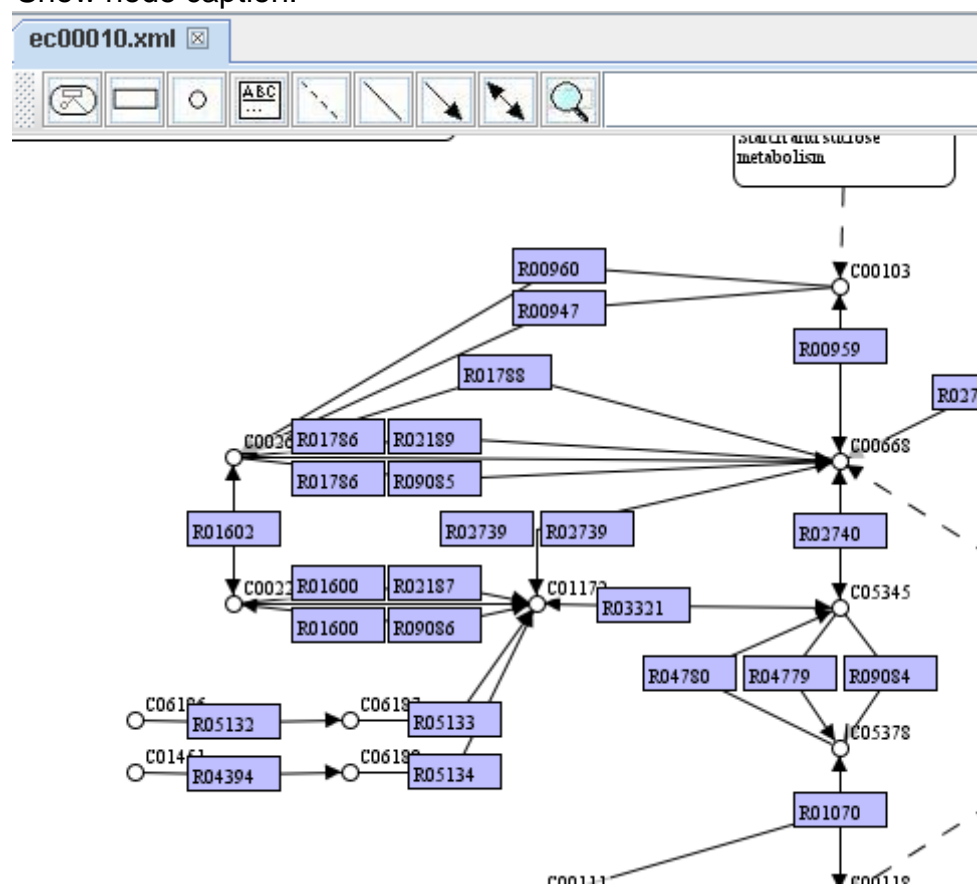
1  <?xml version="1.0"?>
2  <!DOCTYPE pathway SYSTEM "http://www.genome.jp/kegg/xml/KGML_v0.7.1_.dtd">
3  <!-- Creation date: Nov 16, 2010 13:49:39 +0900 (GMT+09:00) -->
4  <pathway name="path:ec00010" org="ec" number="00010"
5      title="Glycolysis / Gluconeogenesis"
6      image="http://www.genome.jp/kegg/pathway/ec/ec00010.png"
7      link="http://www.genome.jp/kegg-bin/show_pathway?ec00010">
8  <entry id="13" name="ec:4.1.2.13" type="enzyme" reaction="rn:R01070"
9      link="http://www.kegg.jp/dbget-bin/www_bget?4.1.2.13">
10     <graphics name="4.1.2.13" fgcolor="#000000" bgcolor="#BFBFFF"
11         type="rectangle" x="483" y="404" width="46" height="17"/>
12   </entry>
13  <entry id="37" name="ec:1.2.1.3" type="enzyme" reaction="rn:R00710"
14      link="http://www.kegg.jp/dbget-bin/www_bget?1.2.1.3">
15     <graphics name="1.2.1.3" fgcolor="#000000" bgcolor="#BFBFFF"
16         type="rectangle" x="289" y="943" width="46" height="17"/>
17   </entry>

```

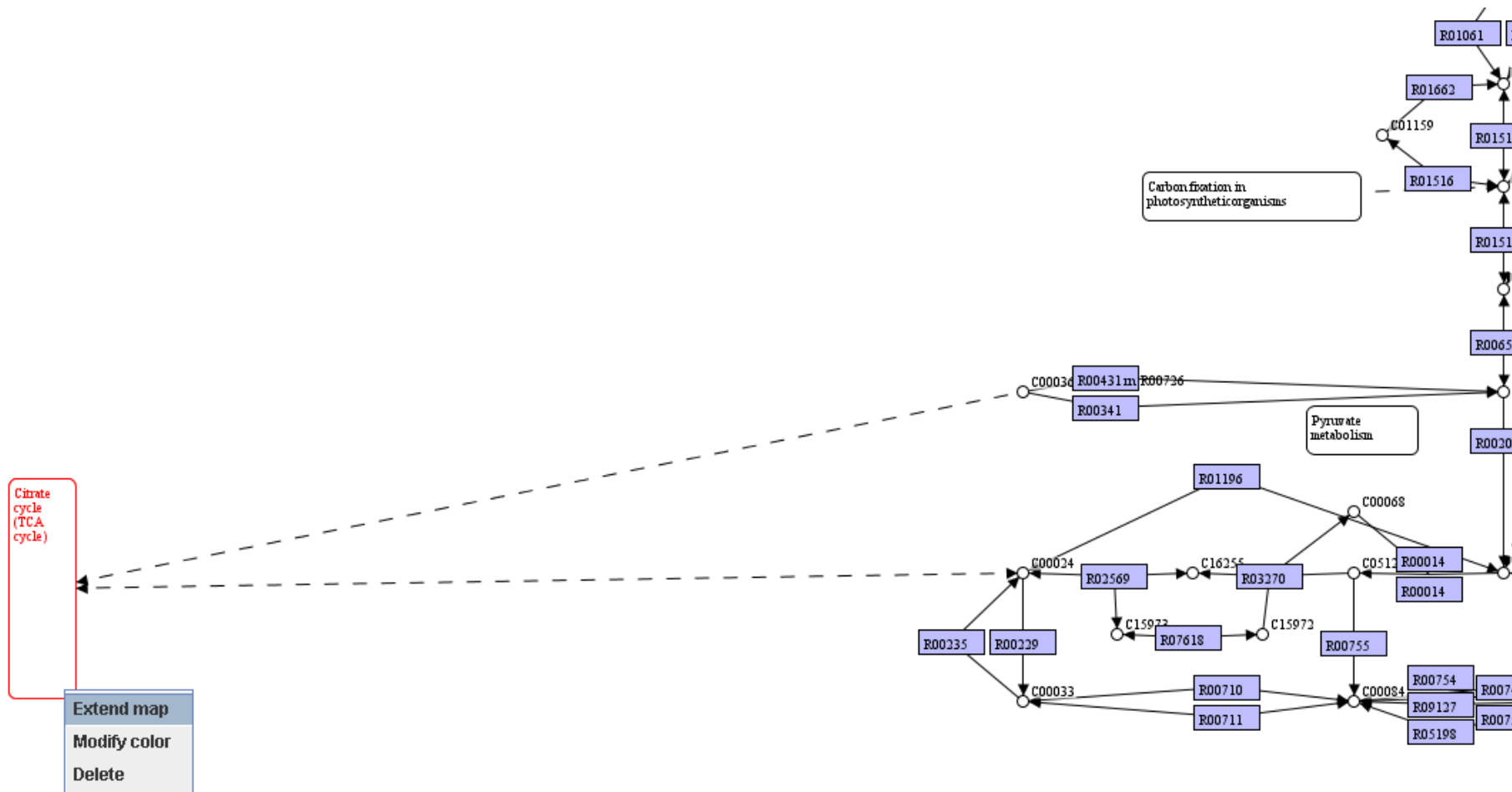
Show node name:



Show node caption:

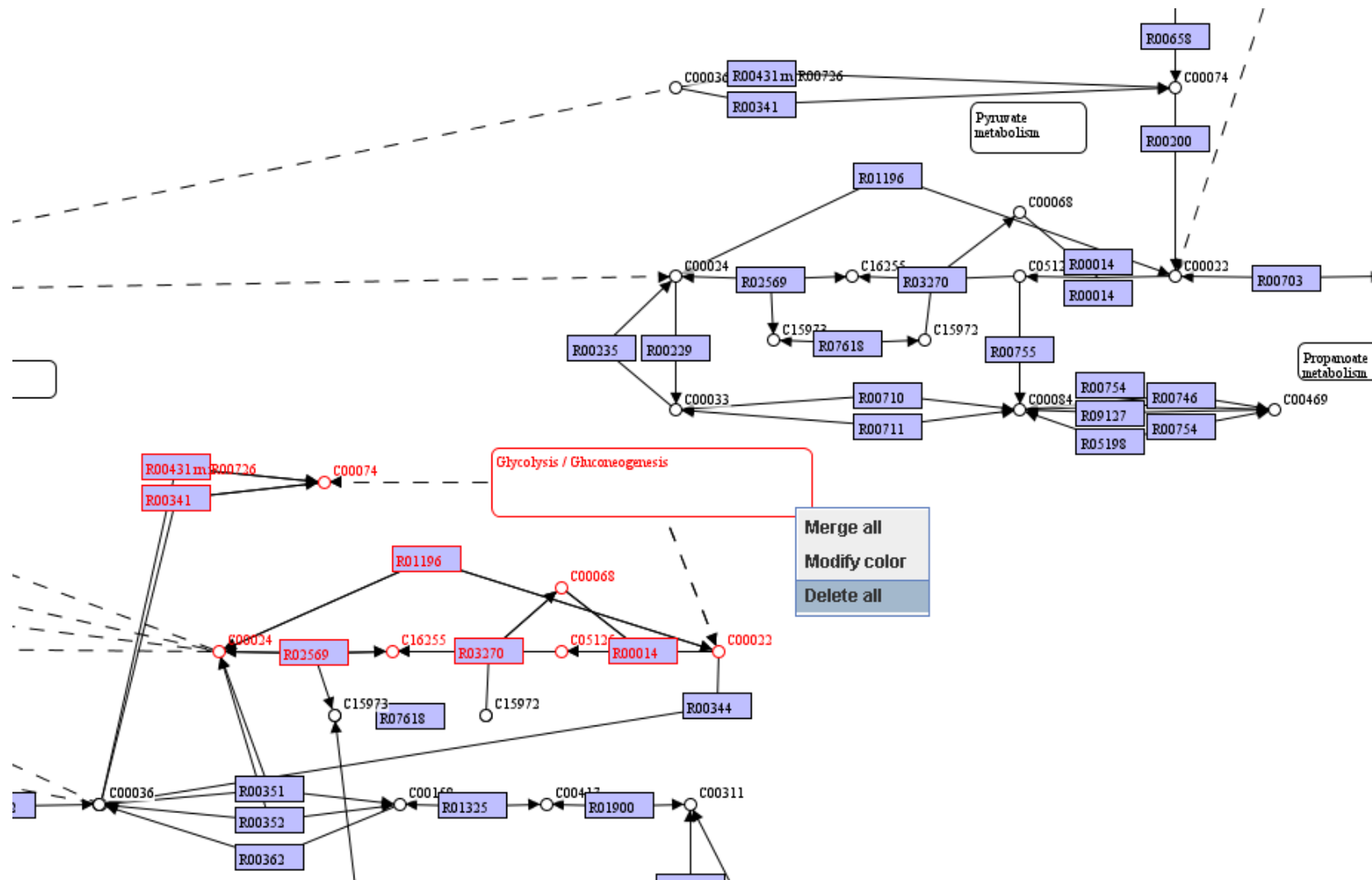


You can extend other pathway maps in the map you have in the main network view window. A pathway map is represented in a rounded rectangle. We can move the pathway that you would like to extend to an empty region and right click on it to Extend map.

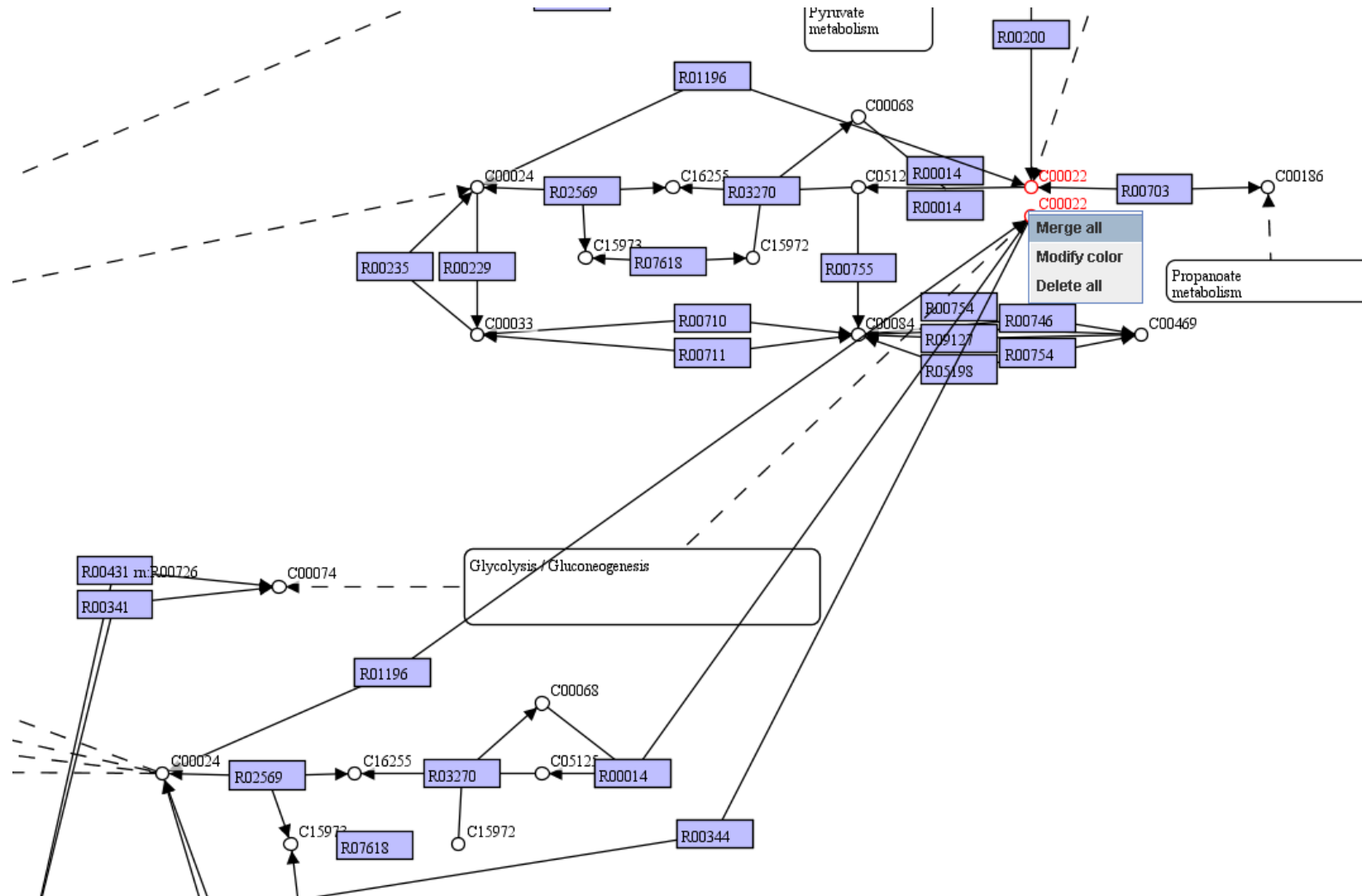


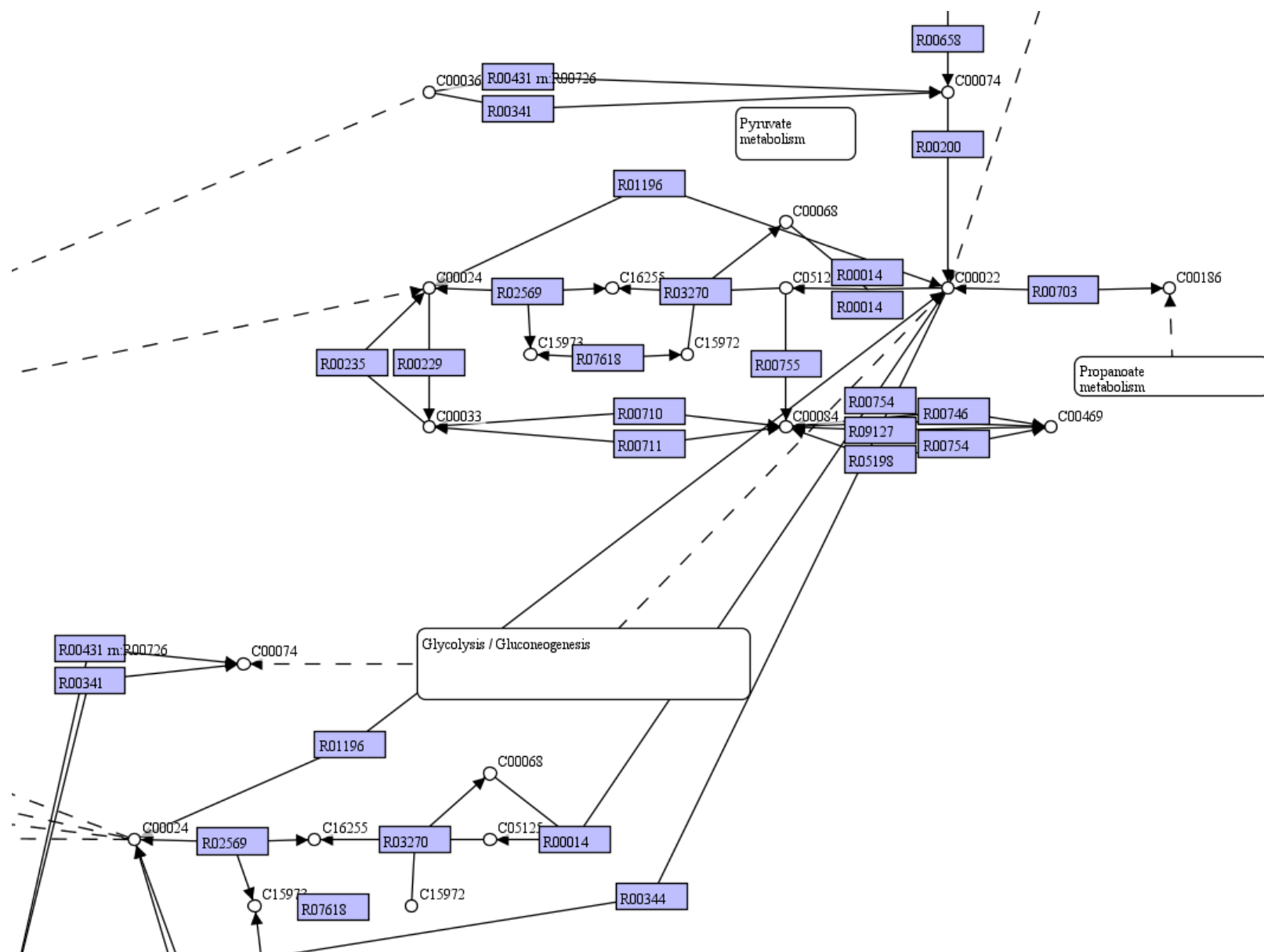
Map of Citrate cycle (TCA cycle) is extended in the map:

You can hold right-click button on the map and drag a rectangular region for selecting groups of objects, then right click over the selected objects to delete them all.

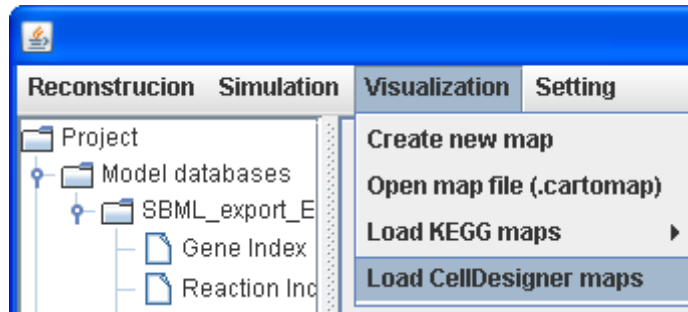


You can move identical objects close to each other. Select the identical objects and then right click on them to Merge all.

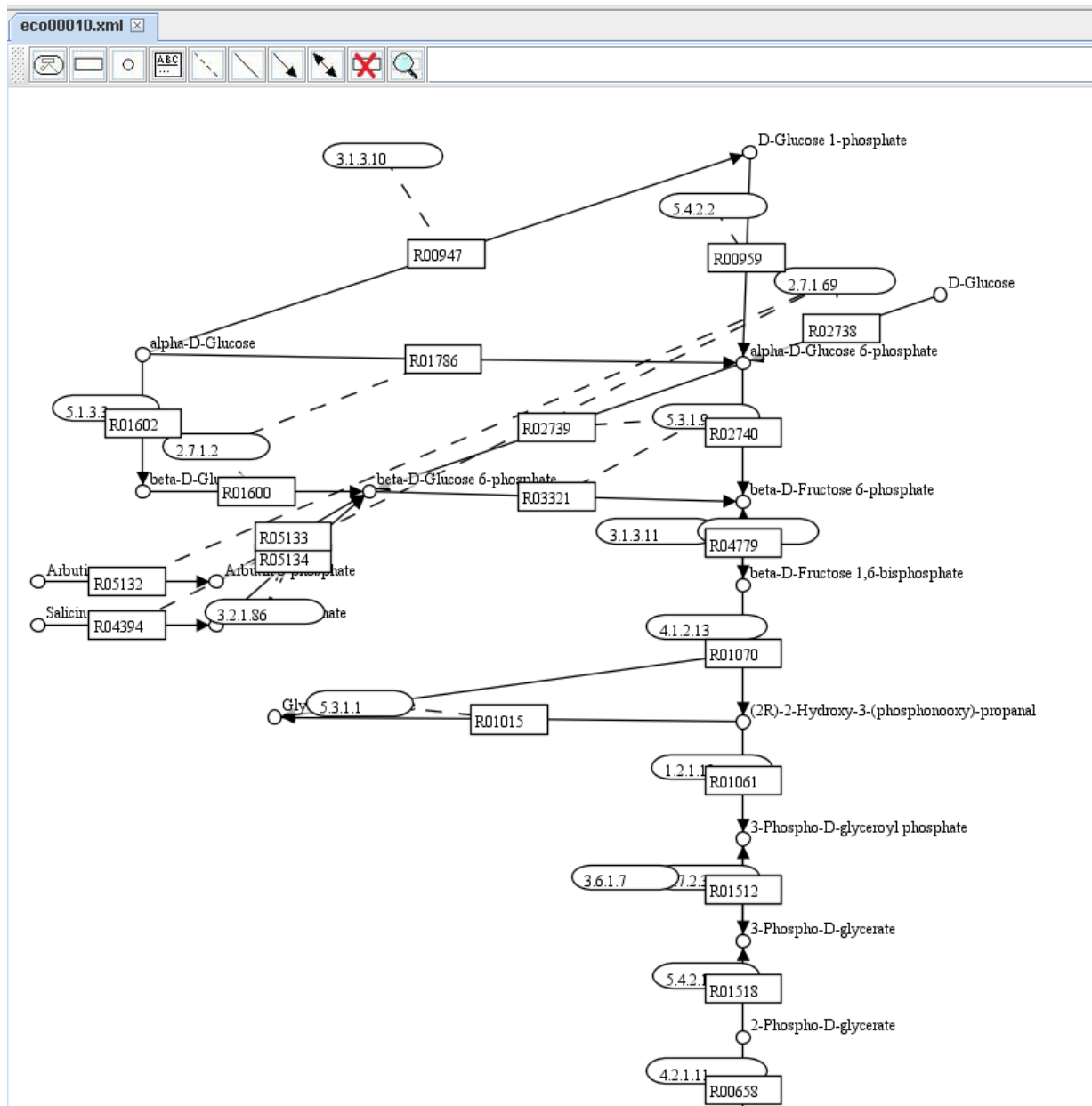




You can also load SBML models compatible to CellDesigner (<http://www.celldesigner.org/index.html>) to GEMSiRV. The SBML models for KEGG can be found and downloaded in <http://www.systems-biology.org/001/001.html>. You can click on Visualization in the menu bar to Load CellDesigner maps.

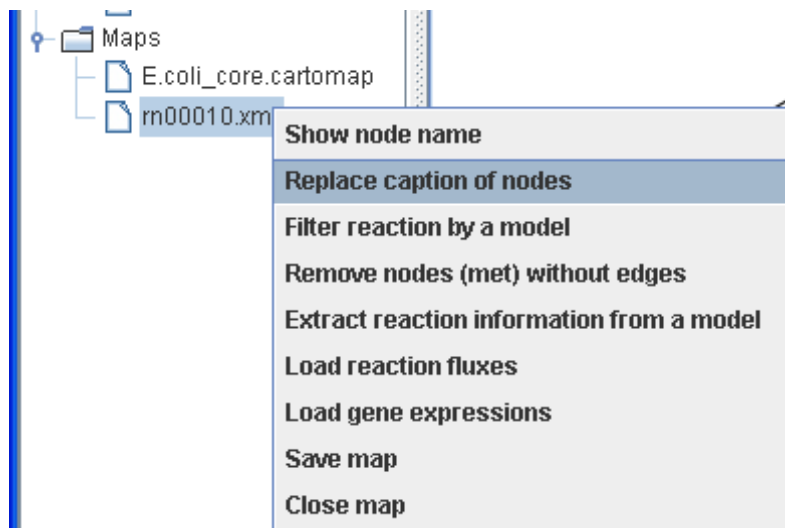


A SBML file eco00010.xml provided in http://sb.nhri.org.tw/GEMSiRV/en/Metabolic_Maps can be downloaded for demonstration.



Map replacement

In order to ease the creation of customized maps, GEMSiRV provide a function in map replacement. You can right click on a map to Replace caption of nodes to convert the map to a customized map.

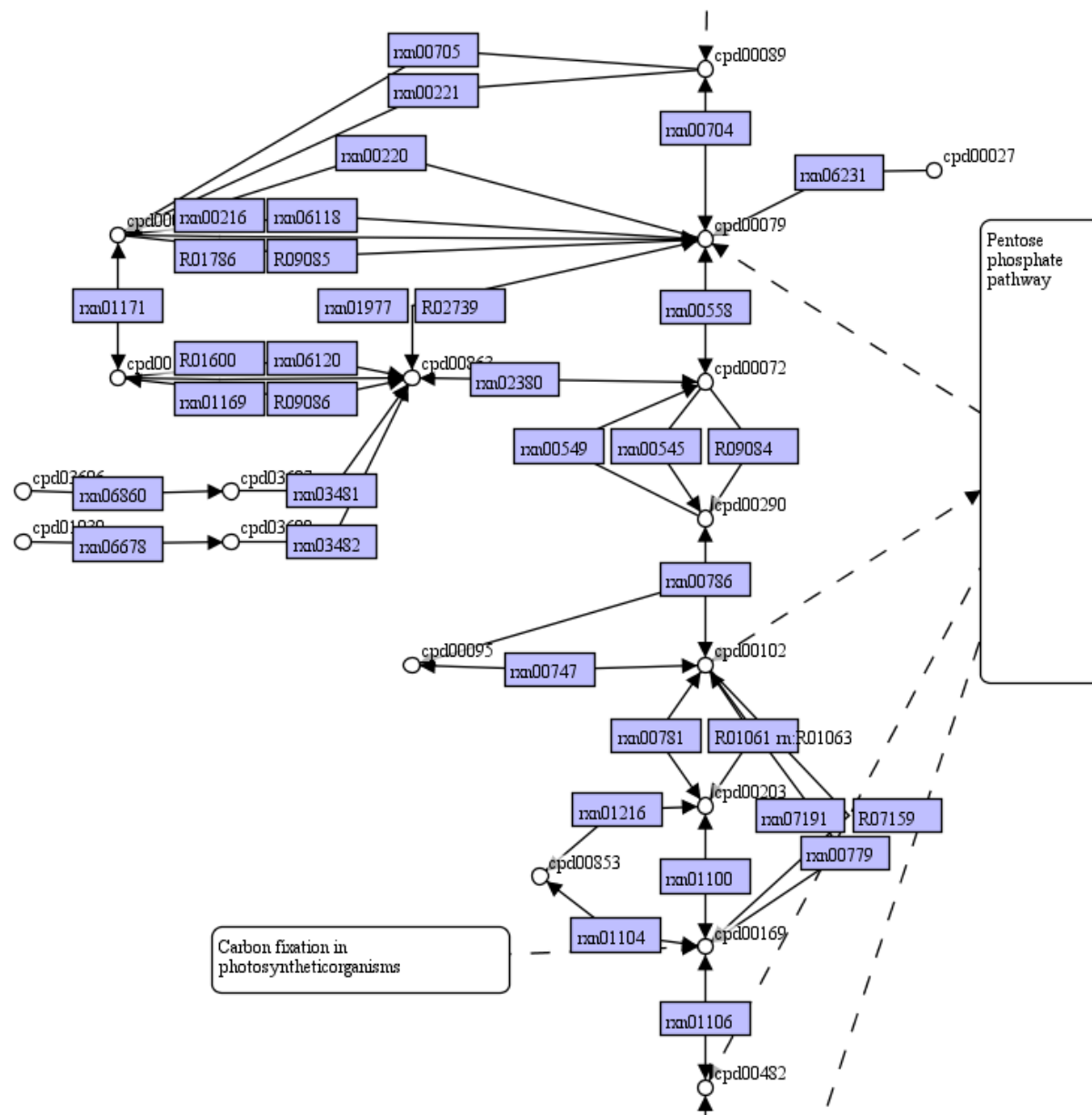


For example, we replace a KEGG map (e.g. rn00010.xml) to a Model SEED-based map by providing two separate lists for metabolite and reaction mapping. The KEEG to Model SEED mapping lists can be found and downloaded in <http://sb.nhri.org.tw/GEMSiRV/en/Manual>.

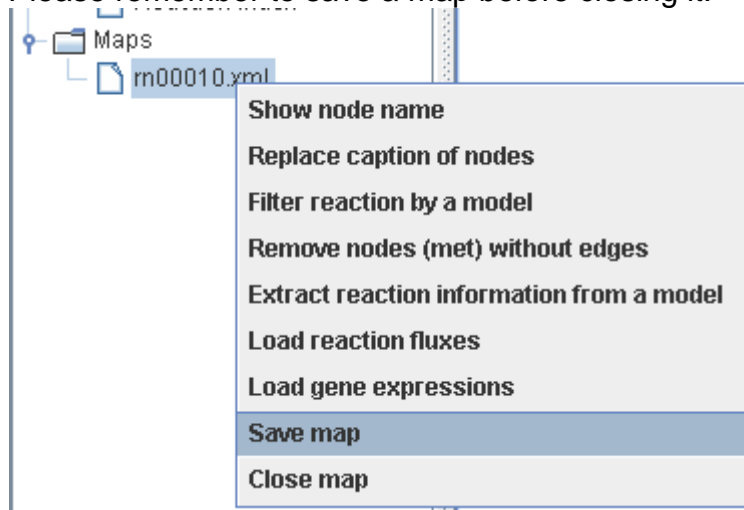
met_KEGGtoSEED.TXT		
	0	1.0
1	C16254	cpd00860
2	C00149	cpd00130
3	C00002	cpd00002
4	C00003	cpd00003
5	C00004	cpd00004
6	C00005	cpd00005
7	C00006	cpd00006
8	C00007	cpd00007
9	C00008	cpd00008
10	C00009	cpd00009

rxn_KEGGtoSEED.TXT		
	0	1.0
1	R00432	rxn00306
2	R07618	rxn01241
3	R00001	rxn05757
4	R00002	rxn11947
5	R00004	rxn00001
6	R00005	rxn00002
7	R00008	rxn00004
8	R00009	rxn00006
9	R00010	rxn00007
10	R00011	rxn00008

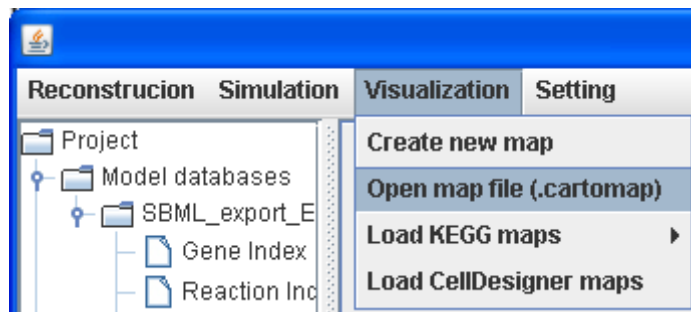
Therefore, some nodes of metabolite and reaction can be replaced to form a Model SEED-based map.



Please remember to save a map before closing it.

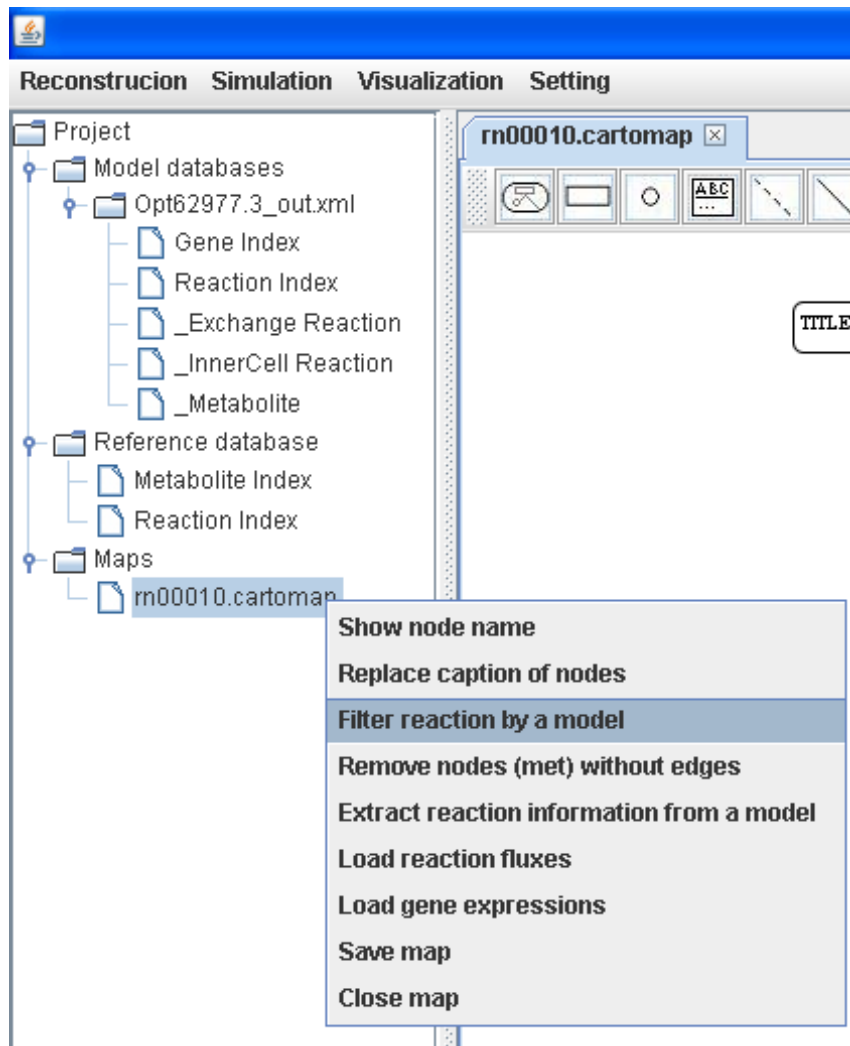


You can open a map saved in cartomap format.

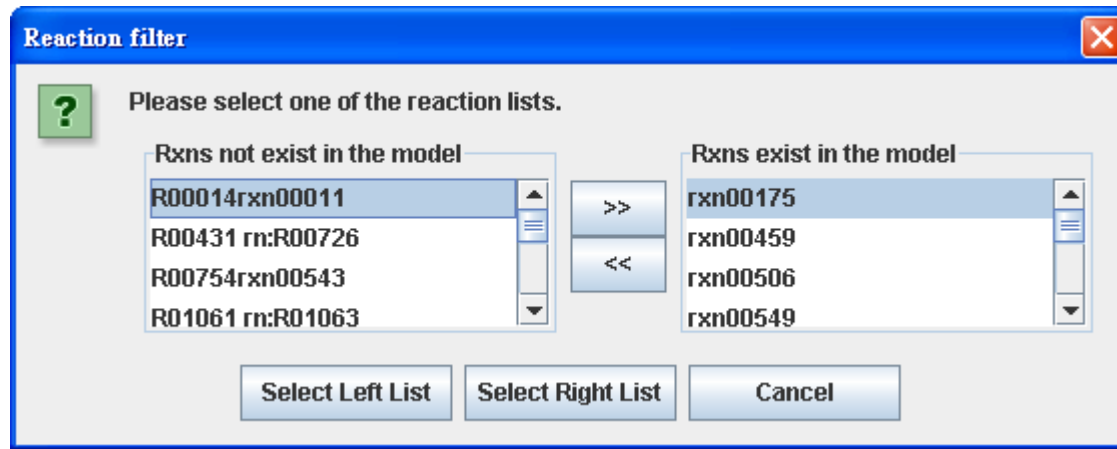


In order to create a useful map for visualization, an interactive function between model reconstruction and map visualization is implemented in GEMSiRV.

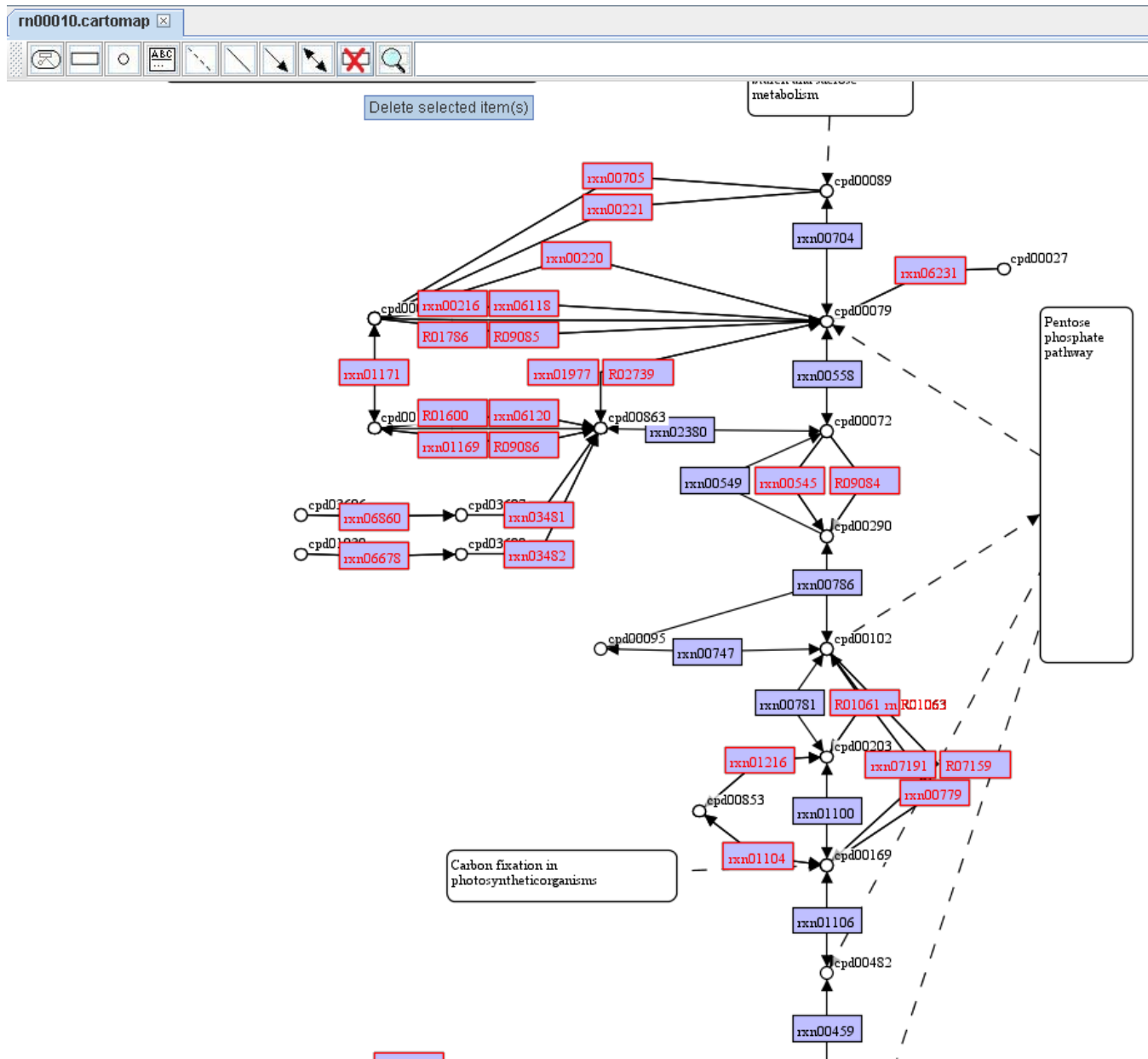
For demonstration, we import a Model SEED model *Acinetobacter* sp. ADP1 (Opt 62977.3.xml).

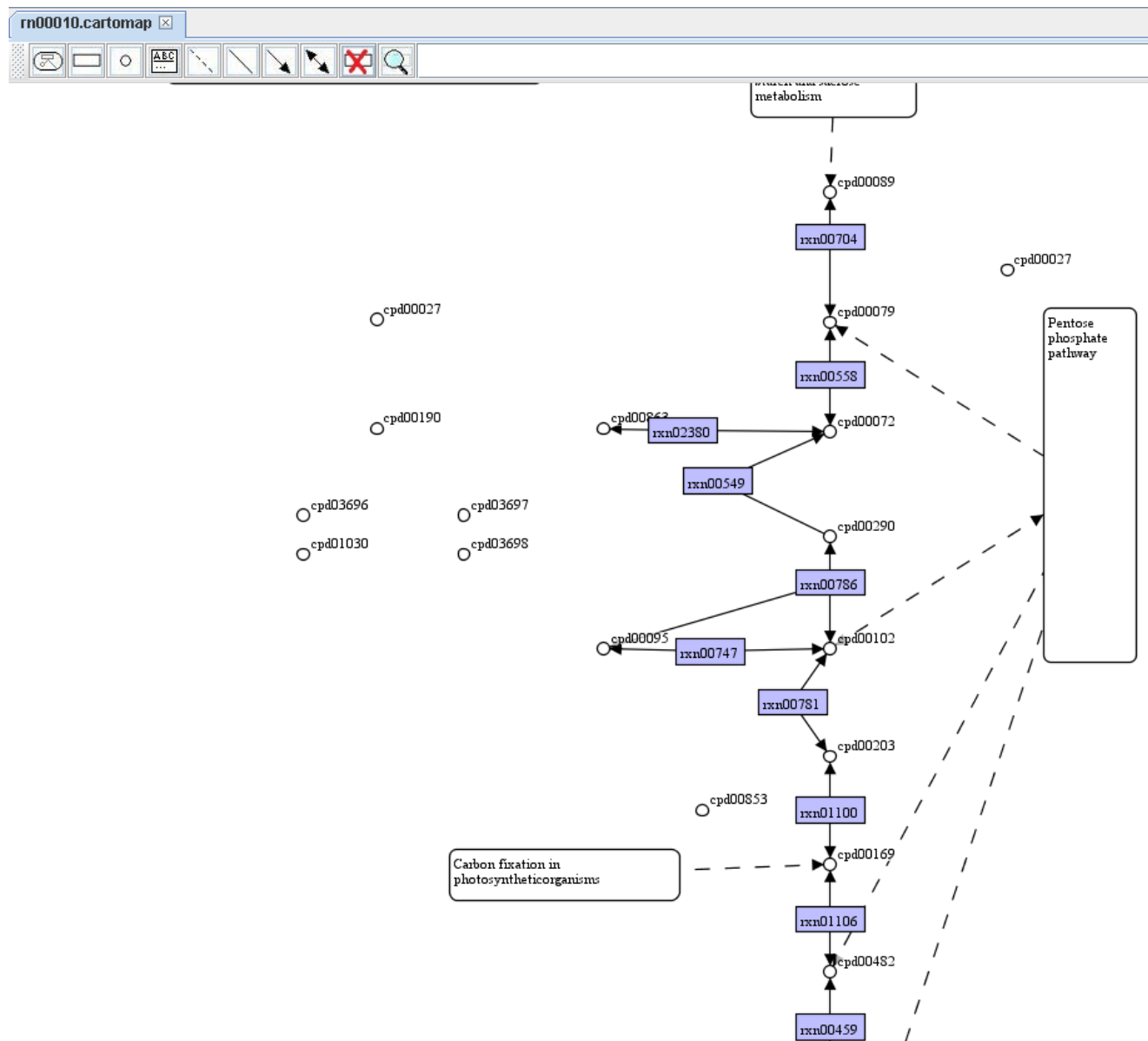


You can filter reactions by comparing with the metabolic model you select and you can get the reaction lists for reactions not existing or existing in the model.

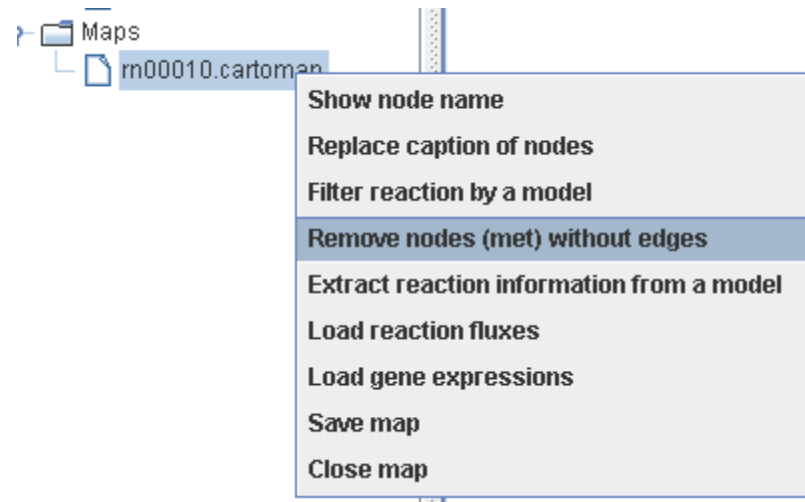


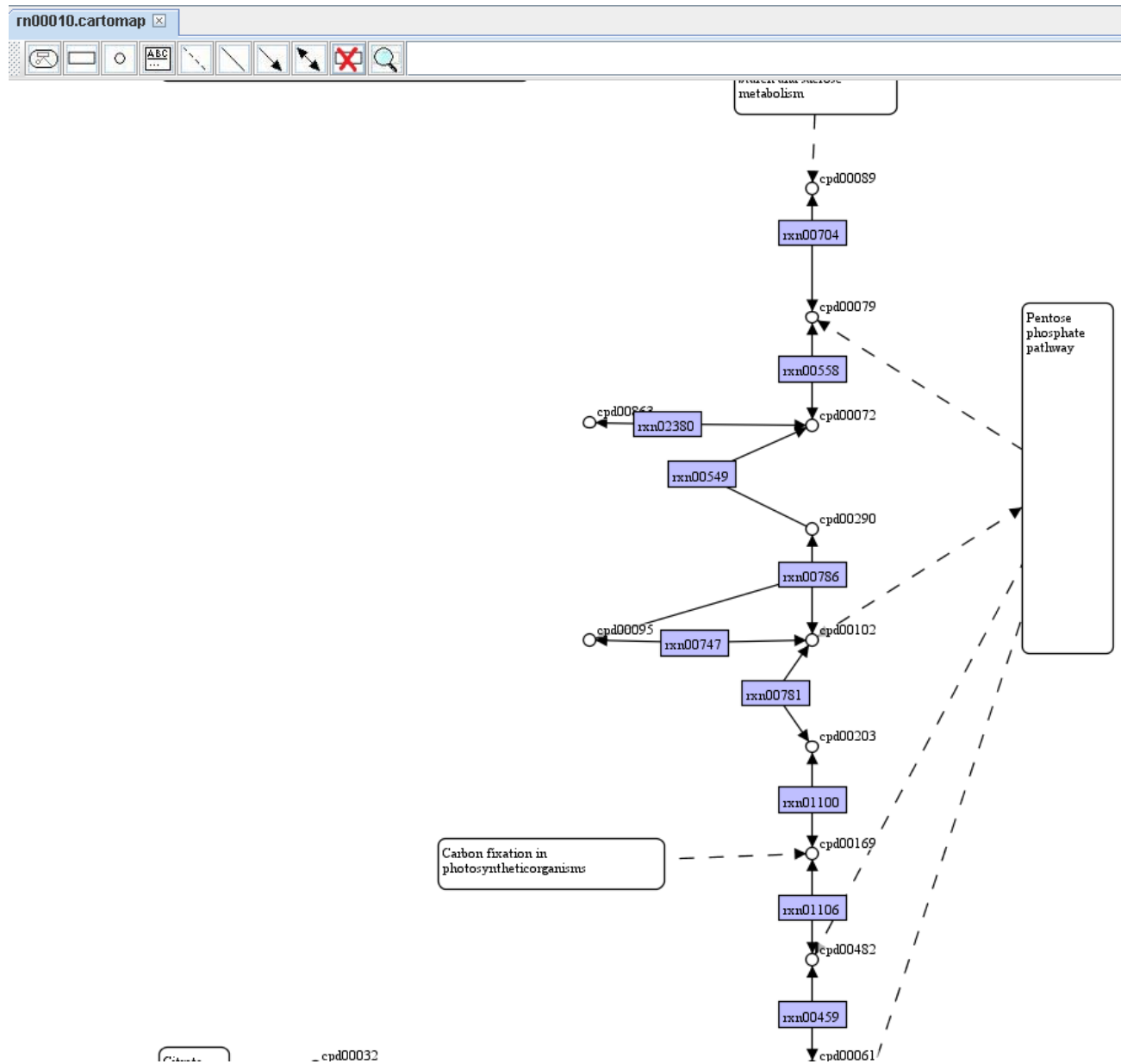
We select and delete the left list of reactions for creating a model-specific map.



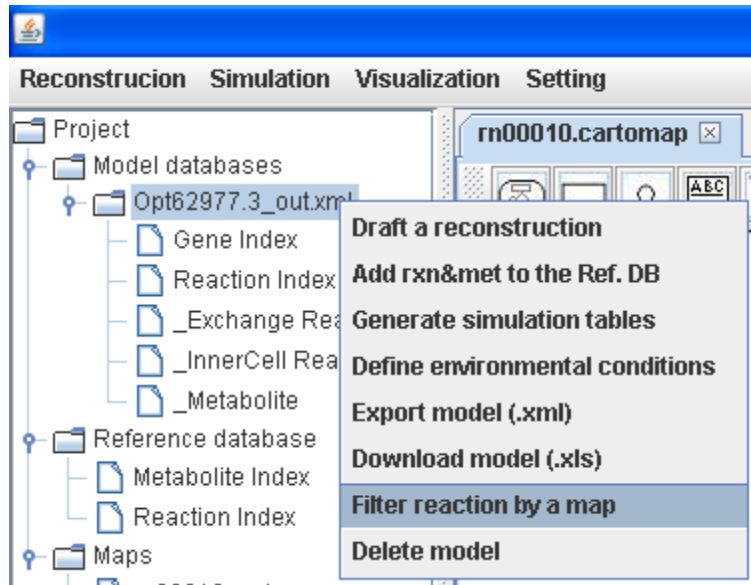


You can remove those nodes of metabolite without linking to reaction by right clicking on the map to Remove nodes (met) without edges.





You can also filter reactions in a metabolic model by comparing with a map. Right click on the model to Filter reaction by a map and choose a map you want to compare with. Then you can get a comparison report as well as a temporary map including those reactions not present in the map you chosen.



A comparison report showing what reactions are present in the model only, in the map only, and in the both.

Reconstrucion Simulation Visualization Setting

Project

- Model databases
 - Opt62977.3_out.xml
 - Gene Index
 - Reaction Index
 - _Exchange Reaction
 - _InnerCell Reaction
 - _Metabolite
- Reference database
 - Metabolite Index
 - Reaction Index
- Maps
 - rn00010.cartomap
 - Temporary Map

rn00010.cartomap Comparison Report Temporary Map

The reactions matched in both table and map:

rxn02342	rxn00781	rxn00459	rxn00175	rxn00558	rxn00786	rxn00549
rxn02380	rxn01100	rxn01241	rxn00747	rxn00704		

The reactions listed in the table only:

EX_cpd11416_c		bio00025	rxn00001	rxn00002	rxn00006	rxn00011
rxn00062	rxn00077	rxn00083	rxn00085	rxn00095	rxn00097	rxn00100
rxn00117	rxn00119	rxn00122	rxn00126	rxn00127	rxn00132	rxn00138
rxn00146	rxn00147	rxn00159	rxn00161	rxn00166	rxn00173	rxn00178
rxn00184	rxn00187	rxn00189	rxn00190	rxn00192	rxn00193	rxn00199
rxn00209	rxn00211	rxn00213	rxn00214	rxn00224	rxn00225	rxn00231
rxn00248	rxn00251	rxn00256	rxn00260	rxn00262	rxn00269	rxn00283
rxn00291	rxn00292	rxn00293	rxn00295	rxn00297	rxn00299	rxn00300
rxn00305	rxn00313	rxn00321	rxn00322	rxn00324	rxn00327	rxn00328
rxn00338	rxn00340	rxn00342	rxn00346	rxn00347	rxn00350	rxn00351
rxn00379	rxn00392	rxn00405	rxn00407	rxn00409	rxn00410	rxn00412
rxn00433	rxn00438	rxn00441	rxn00461	rxn00463	rxn00469	rxn00470
rxn00503	rxn00508	rxn00509	rxn00512	rxn00514	rxn00515	rxn00527
rxn00568	rxn00569	rxn00584	rxn00588	rxn00589	rxn00598	rxn00601
rxn00615	rxn00616	rxn00623	rxn00640	rxn00646	rxn00649	rxn00650
rxn00675	rxn00676	rxn00679	rxn00686	rxn00689	rxn00691	rxn00692
rxn00710	rxn00711	rxn00714	rxn00726	rxn00727	rxn00737	rxn00758
rxn00775	rxn00776	rxn00777	rxn00782	rxn00785	rxn00789	rxn00790
rxn00800	rxn00802	rxn00806	rxn00831	rxn00832	rxn00834	rxn00836
rxn00858	rxn00863	rxn00867	rxn00868	rxn00874	rxn00875	rxn00881
rxn00902	rxn00903	rxn00907	rxn00910	rxn00912	rxn00913	rxn00915

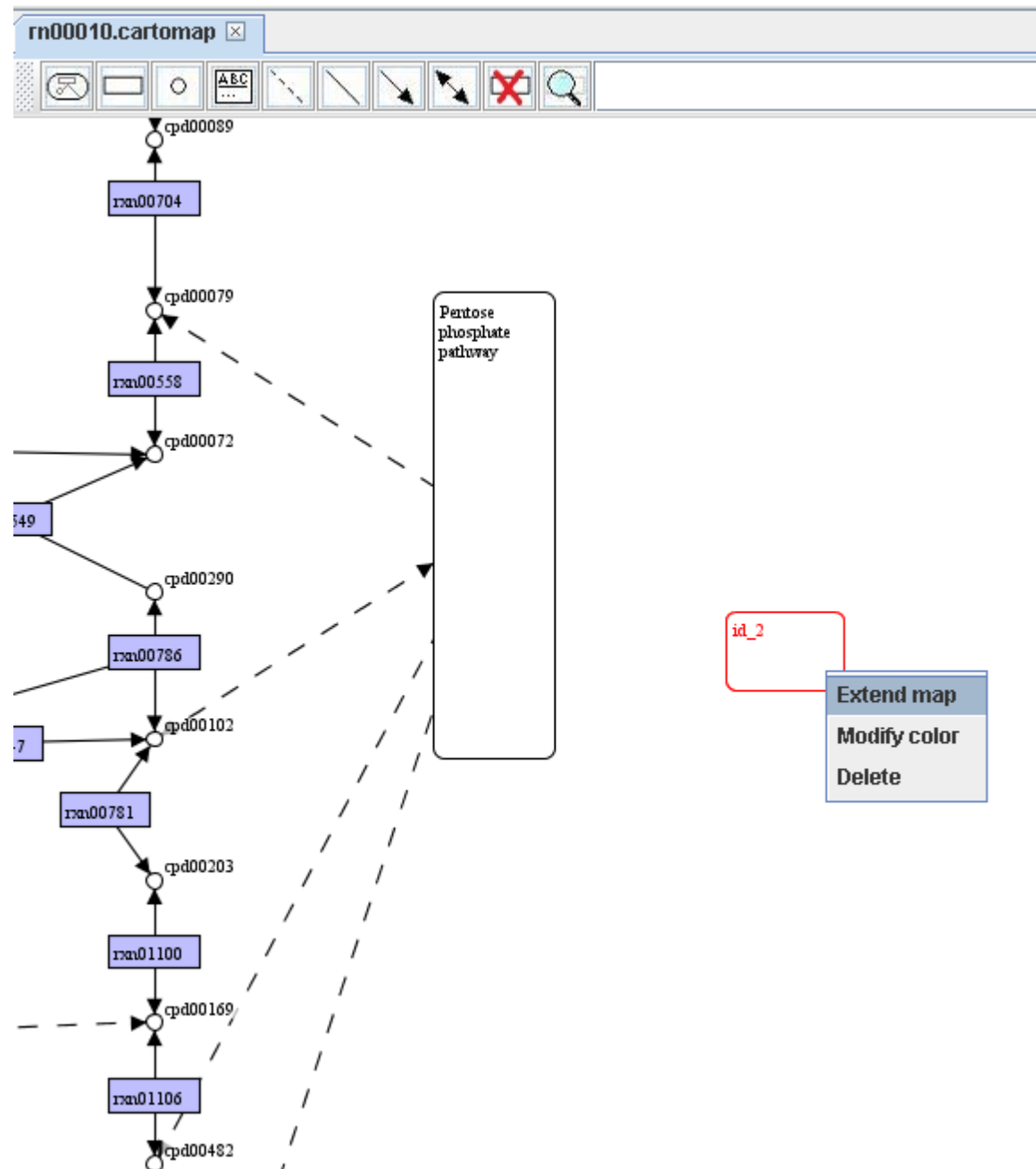
A temporary map including those reactions in the model but not in the map

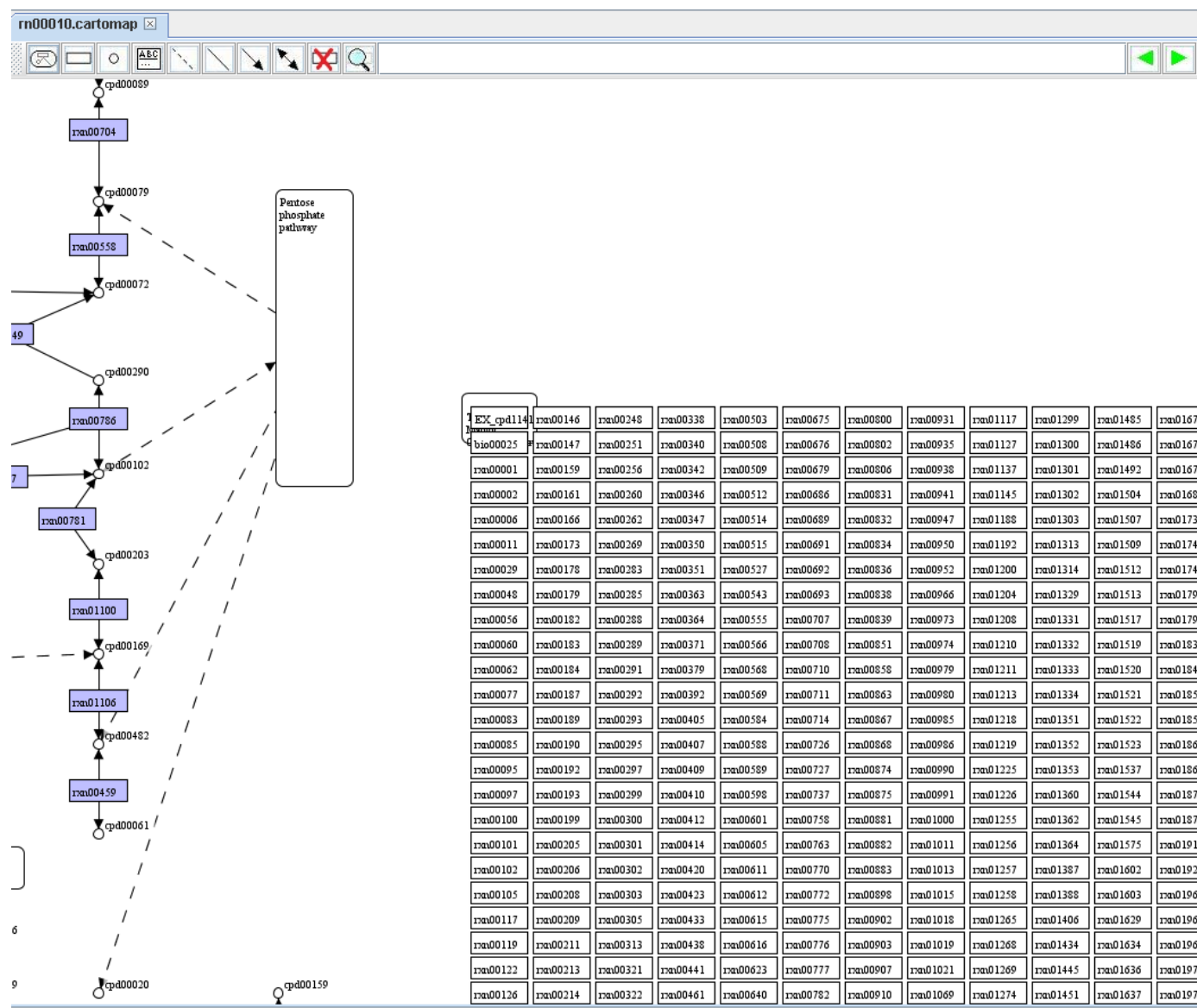
The screenshot shows a software interface with a menu bar (Reconstruction, Simulation, Visualization, Setting) and a sidebar. The sidebar contains a tree view under 'Project' with folders for 'Model databases' and 'Reference database'. The 'Model databases' folder is expanded, showing 'Opt62977.3_out.xml' and its sub-items: 'Gene Index', 'Reaction Index', '_Exchange Reaction', '_InnerCell Reaction', and '_Metabolite'. The 'Reference database' folder contains 'Metabolite Index' and 'Reaction Index'. The 'Maps' folder contains 'rn00010.cartomap' and 'Temporary Map'. The main window has three tabs: 'rn00010.cartomap', 'Comparison Report', and 'Temporary Map'. The 'Temporary Map' tab is active, showing a grid of reaction IDs. The grid has 12 columns and 20 rows. The first column contains reaction IDs starting with 'rn02666', and the subsequent columns contain reaction IDs starting with 'rn03013', 'rn03242', 'rn03540', 'rn03990', 'rn05153', 'rn05256', 'rn05333', 'rn05373', 'rn05413', 'rn05448', 'rn05516', and 'rn05594'. The grid is titled 'Temporary Map from reaction filter'.

rn02666	rn03013	rn03242	rn03540	rn03990	rn05153	rn05256	rn05333	rn05373	rn05413	rn05448	rn05516	rn05594
rn02680	rn03030	rn03243	rn03638	rn03991	rn05171	rn05287	rn05334	rn05375	rn05414	rn05449	rn05517	rn05596
rn02774	rn03031	rn03244	rn03839	rn04070	rn05172	rn05289	rn05335	rn05376	rn05415	rn05450	rn05526	rn05599
rn02782	rn03052	rn03245	rn03841	rn04113	rn05176	rn05293	rn05336	rn05377	rn05417	rn05451	rn05527	rn05602
rn02789	rn03060	rn03246	rn03843	rn04132	rn05183	rn05294	rn05337	rn05379	rn05418	rn05452	rn05528	rn05603
rn02791	rn03061	rn03247	rn03869	rn04133	rn05195	rn05295	rn05338	rn05380	rn05419	rn05453	rn05533	rn05605
rn02792	rn03062	rn03248	rn03884	rn04139	rn05197	rn05296	rn05339	rn05381	rn05421	rn05454	rn05534	rn05613
rn02804	rn03068	rn03249	rn03887	rn04308	rn05198	rn05297	rn05340	rn05383	rn05422	rn05455	rn05535	rn05614
rn02811	rn03075	rn03250	rn03891	rn04456	rn05199	rn05298	rn05341	rn05384	rn05423	rn05456	rn05536	rn05616
rn02834	rn03080	rn03384	rn03892	rn04457	rn05200	rn05299	rn05342	rn05385	rn05425	rn05457	rn05537	rn05619
rn02835	rn03084	rn03393	rn03893	rn04674	rn05201	rn05301	rn05343	rn05386	rn05426	rn05458	rn05538	rn05620
rn02853	rn03087	rn03394	rn03897	rn04713	rn05202	rn05305	rn05344	rn05388	rn05427	rn05459	rn05539	rn05621
rn02866	rn03108	rn03395	rn03898	rn04724	rn05205	rn05306	rn05345	rn05389	rn05429	rn05460	rn05540	rn05625
rn02889	rn03130	rn03397	rn03901	rn04745	rn05211	rn05312	rn05346	rn05390	rn05430	rn05461	rn05541	rn05627
rn02895	rn03135	rn03406	rn03903	rn04748	rn05215	rn05315	rn05347	rn05392	rn05431	rn05462	rn05542	rn05634
rn02911	rn03136	rn03407	rn03904	rn04750	rn05217	rn05316	rn05348	rn05393	rn05433	rn05464	rn05543	rn05638
rn02914	rn03137	rn03408	rn03906	rn04783	rn05219	rn05317	rn05349	rn05394	rn05434	rn05465	rn05544	rn05649
rn02916	rn03146	rn03409	rn03907	rn04794	rn05221	rn05318	rn05350	rn05396	rn05435	rn05466	rn05545	rn05651
rn02927	rn03147	rn03419	rn03908	rn04954	rn05229	rn05319	rn05358	rn05397	rn05436	rn05467	rn05546	rn05654
rn02929	rn03159	rn03421	rn03909	rn04996	rn05231	rn05322	rn05359	rn05398	rn05437	rn05468	rn05547	rn05656
rn02933	rn03164	rn03422	rn03910	rn05029	rn05233	rn05323	rn05360	rn05400	rn05438	rn05469	rn05552	rn05663
rn02934	rn03167	rn03433	rn03916	rn05039	rn05239	rn05324	rn05361	rn05401	rn05439	rn05481	rn05555	rn05667
rn02937	rn03174	rn03435	rn03917	rn05040	rn05243	rn05325	rn05363	rn05402	rn05440	rn05484	rn05560	rn05669

You can save the temporary map and add it into the map you are working with.

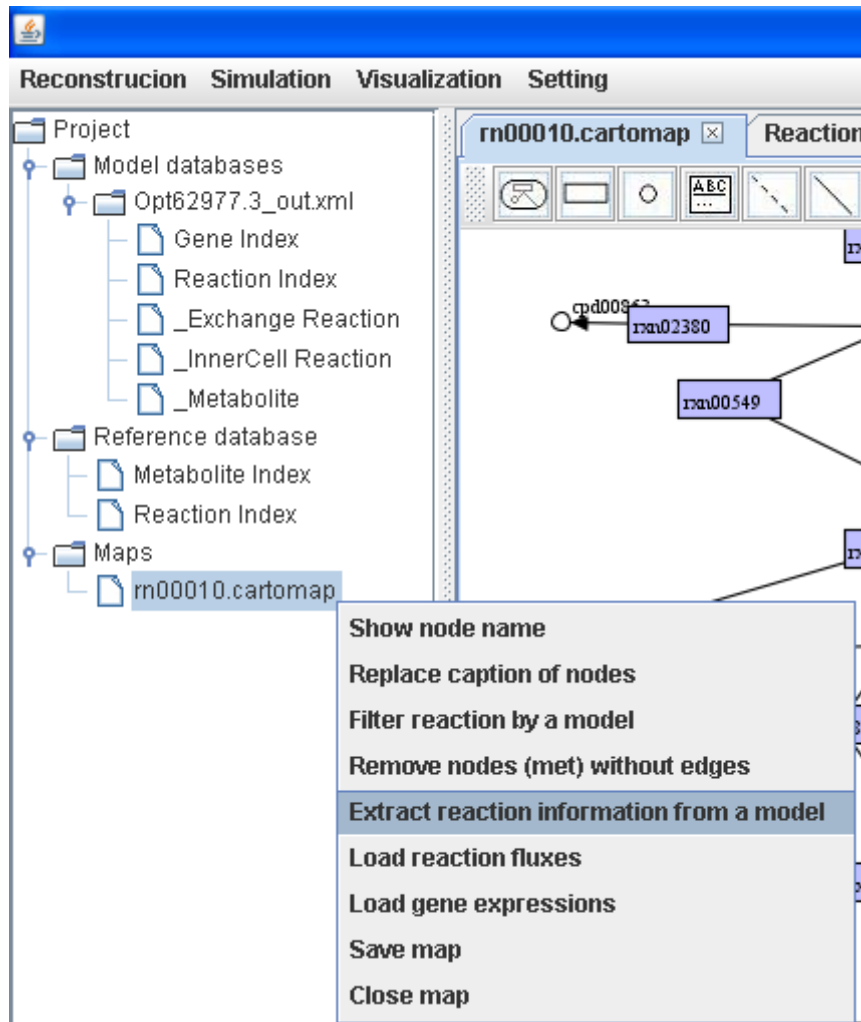
Add a map by clicking Add a map in the toolbar and dropping in an empty region of the map and extend the map by right clicking on the added map to Extend map.

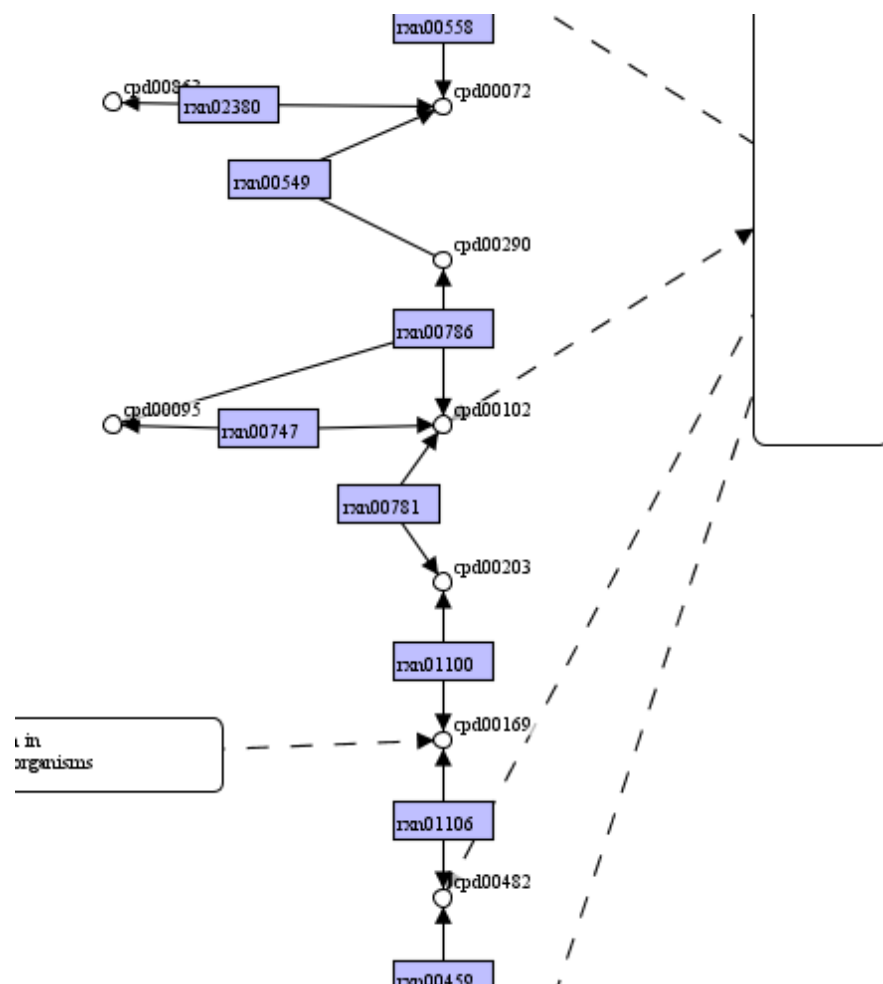




Information extraction

Right click on a map to Extract reaction information from a model and choose a model you want to extract information from. Then you can show the extra information of reaction in the map by right clicking a reaction to Show extra info.





EX_cpd114	rxn00146	rxn00248	rxn00338	rxn00428
bio00025	rxn00147	Show extra info.		rxn00429
rxn00001	rxn00159	Update caption		rxn00430
rxn00002	rxn00161	Modify color		rxn00431
rxn00006	rxn00166	Delete		rxn00432
rxn00011	rxn00173	rxn00269	rxn00350	rxn00433
rxn00029	rxn00178	rxn00283	rxn00351	rxn00434
rxn00048	rxn00179	rxn00285	rxn00363	rxn00435
rxn00056	rxn00182	rxn00288	rxn00364	rxn00436
rxn00060	rxn00183	rxn00289	rxn00371	rxn00437
rxn00062	rxn00184	rxn00291	rxn00379	rxn00438
rxn00077	rxn00187	rxn00292	rxn00392	rxn00439
rxn00083	rxn00189	rxn00293	rxn00405	rxn00440
rxn00085	rxn00190	rxn00295	rxn00407	rxn00441
rxn00095	rxn00192	rxn00297	rxn00409	rxn00442
rxn00097	rxn00193	rxn00299	rxn00410	rxn00443

The gene and equation information for the reaction can be seen on the map.

Gene: ACIAD0109 Equation: [c] : (2.0) cpd00109 + cpd00221 <==> cpd00020 + (2.0) cpd00067 + (2.0) cpd00110									
EX_cpd114	rxn00146	rxn00248	rxn00338	rxn00503	rxn00675	rxn00800	rxn00931	rxn01117	17
bio00025	rxn00147	rxn00251	rxn00340	rxn00508	rxn00676	rxn00802	rxn00935	rxn01127	17
rxn00001	rxn00159	rxn00256	rxn00342	rxn00509	rxn00679	rxn00806	rxn00938	rxn01137	17
rxn00002	rxn00161	rxn00260	rxn00346	rxn00512	rxn00686	rxn00831	rxn00941	rxn01145	17
rxn00006	rxn00166	rxn00267	rxn00347	rxn00514	rxn00689	rxn00837	rxn00947	rxn01188	17

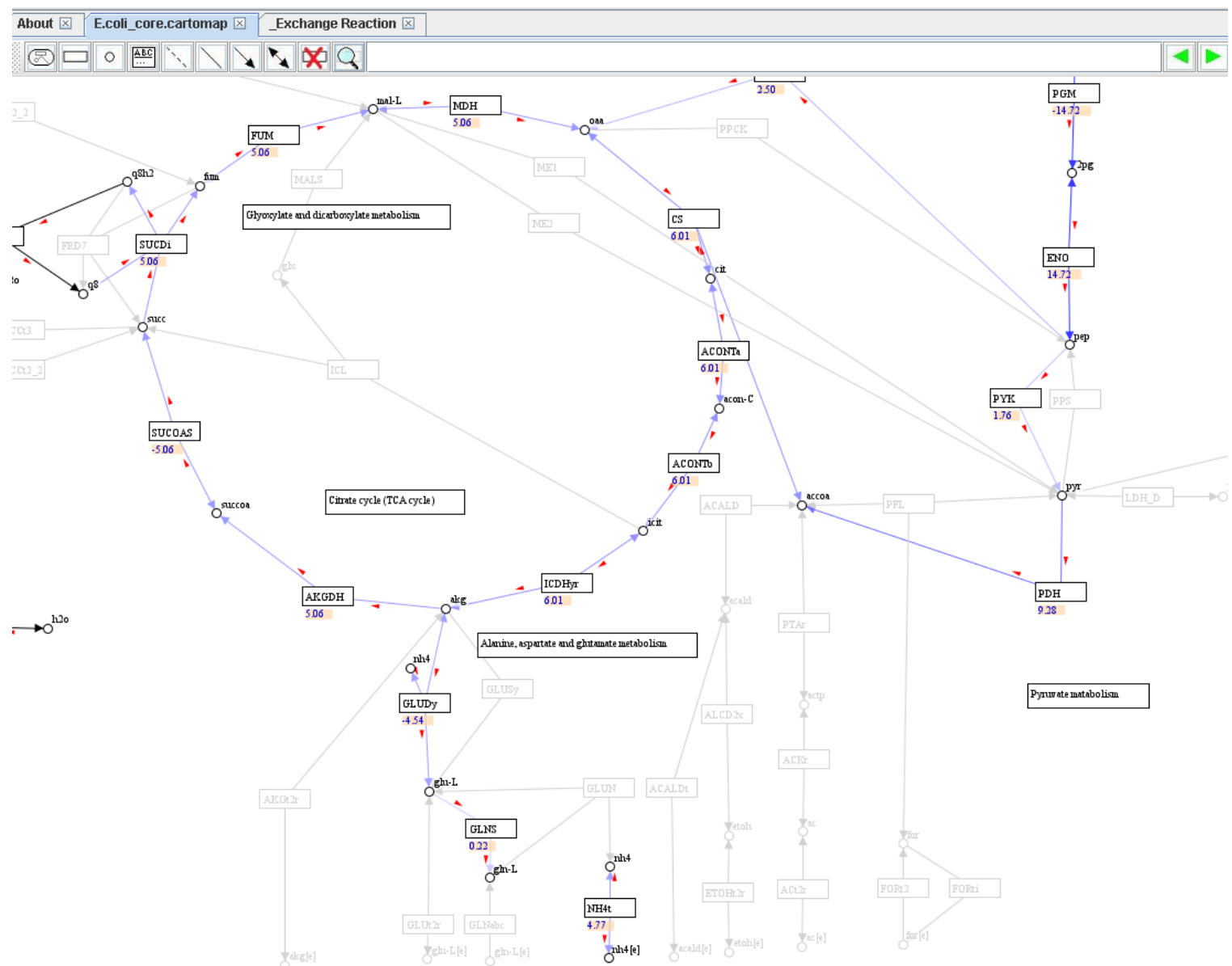
The function in information extraction can aid in metabolic map creation and providing associated genes for later loading gene expression.

Flux visualization

As mentioned early in Simulation, reaction fluxes can be loaded into a map for visualization. GEMSiRV provides a function in loading reaction fluxes by right clicking on a map to Load reaction fluxes.

A single run of simulation: (A header line beginning with “#” is optional)

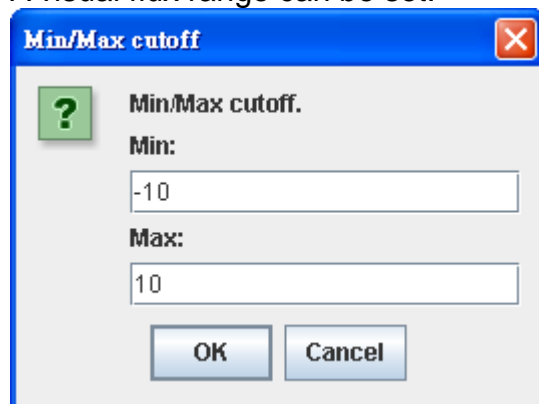
```
#Reaction Flux
ACONTa 6.00725
ACONTb 6.00725
AKGDH 5.06438
ATPM 8.39
ATPS4r 45.514
Biomass_Ecoli_core_N(w/GAM)_Nmet2 0.873922
CO2t -22.8098
CS 6.00725
CYTBD 43.599
ENO 14.7161
EX_co2(e) 22.8098
EX_glc(e) -10.0
```



Multiple runs of simulation: (A header line beginning with “#” is preferred for labeling in legend)

```
#Reaction F1 F2 F3 F4
ACALD 0 0 5 10
ACALDt 0 0 5 10
ACKr 0 0 5 10
ACONTa 6.00725 6.00725 5 10
ACONTb 6.00725 6.00725 5 10
ACt2r 0 10 5 10
ADK1 0 10 5 10
AKGDH 5.06438 5.06438 5 10
AKGt2r 0 0 5 10
ALCD2x 0 0 5 10
ATPM 8.39 8.39 5 10
ATPS4r 45.514 45.514 5 10
Biomass_Ecoli_core_N(w/GAM)_Nmet2 0.873922 0.873922 5 10
CO2t -22.8098 -22.8098 5 10
CS 6.00725 6.00725 5 10
CYTBD 43.599 43.599 5 10
D_LACt2 0 0 5 10
ENO 14.7161 14.7161 5 10
ETOht2r 0 0 5 10
EX_ac(e) 0 0 5 10
EX_acald(e) 0 0 5 10
```

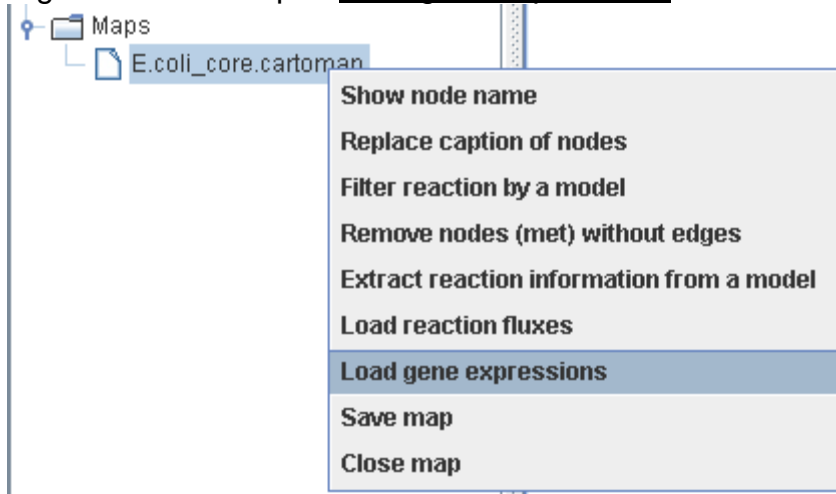
A visual flux range can be set:



Gene expression visualization

Because GEMSiRV allows users to extract information from a model to a map, in addition to reaction fluxes, gene expressions can also be loaded into a map for visualization. In this circumstance, we can simultaneously compare the differences of reaction fluxes with that of gene expressions in two conditions (e.g. aerobic and anaerobic conditions).

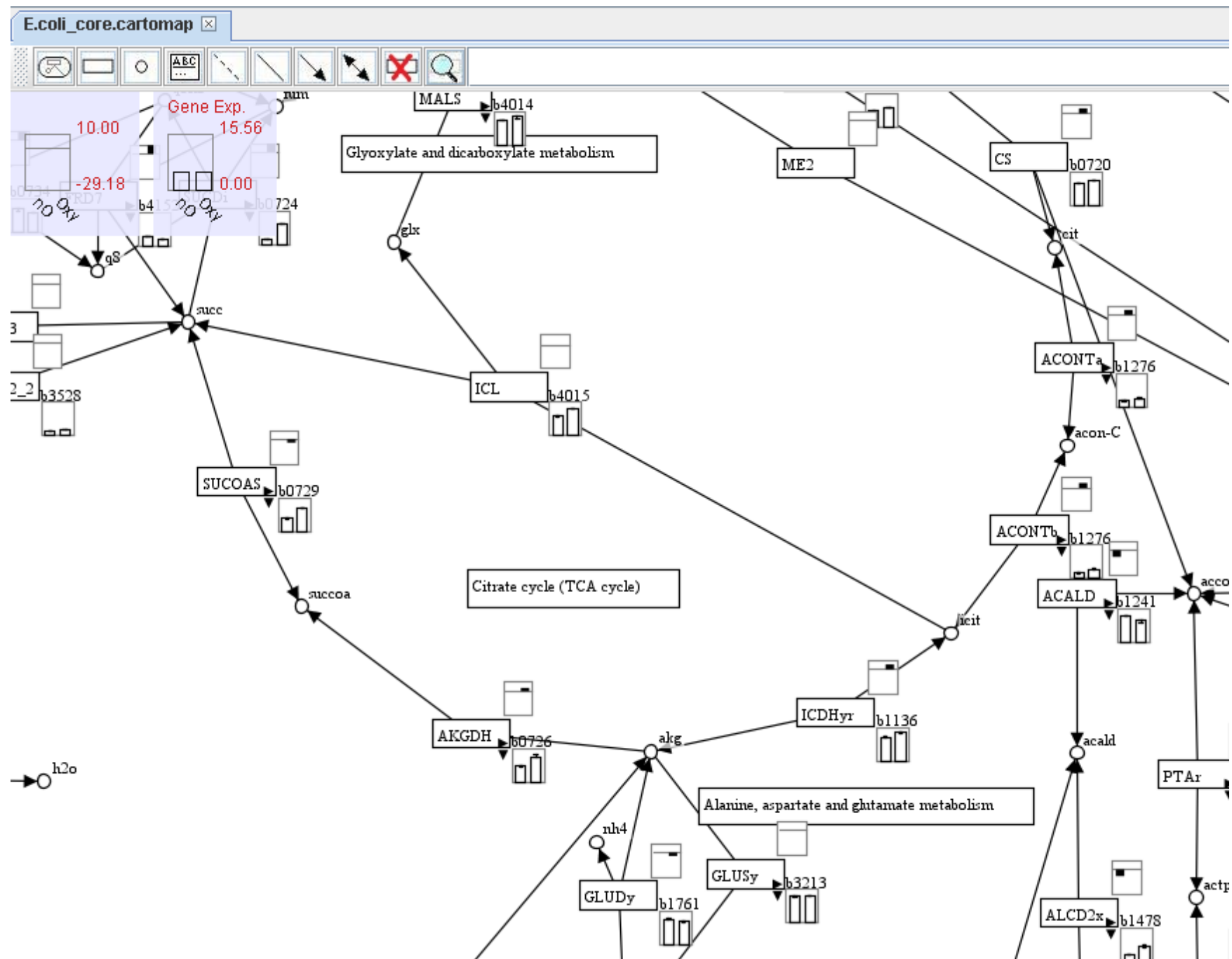
Right click on a map to Load gene expressions.



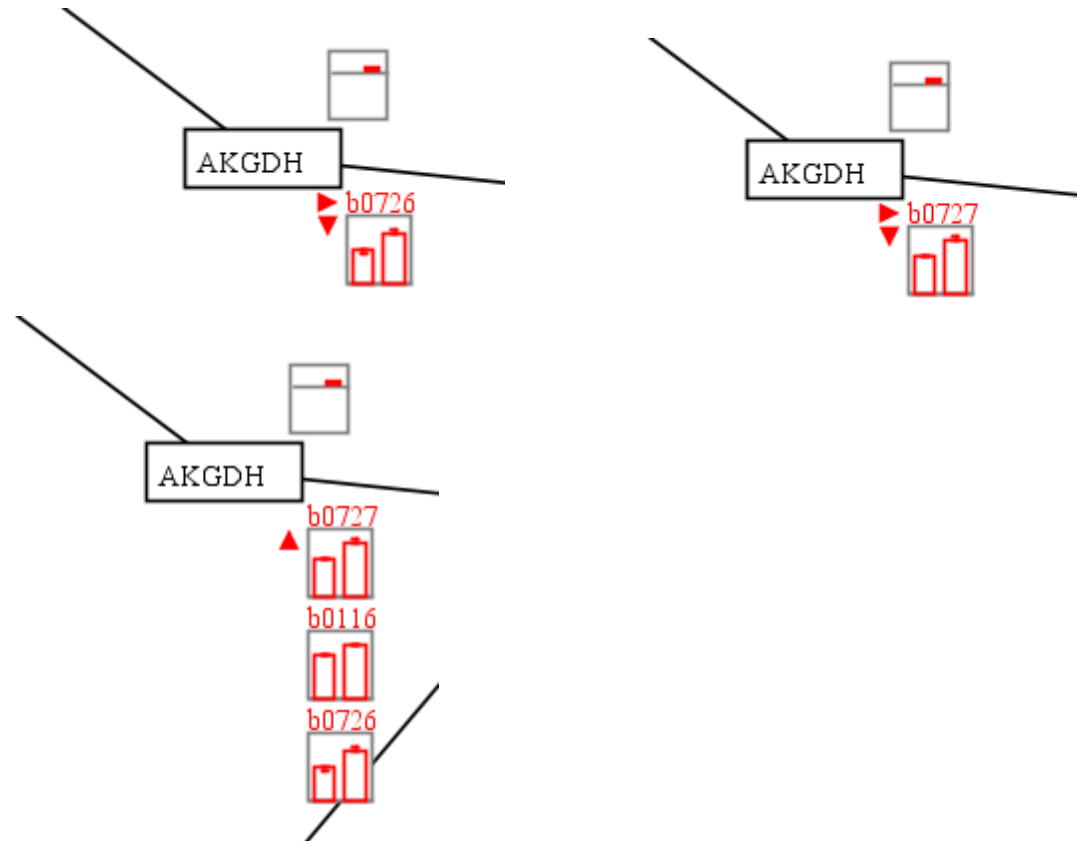
You can use identical header to represent the replicates of condition. Then the mean and standard deviation of gene expression for a specific gene will be shown in the map. Here, we used the expression data (array number 42-48) available in http://systemsbiology.ucsd.edu/In_Silico_Organisms/E_coli/E_coli_expression2.

Expr_o2 effect on Ecoli_rep.TXT									
#Gene	nO	nO	nO	nO	Oxy	Oxy	Oxy	Oxy	Oxy
1	b2836	5.769096757	5.879973845	5.889739712	5.867358643	5.760884085	5.873523604	5.957984412	
2	b0885	3.535727661	3.469050784	4.049806246	3.645329898	3.879638515	3.597923384	3.593089884	
3	b0199	9.804808142	10.40118062	10.57760507	10.25695657	9.301827015	9.898372638	10.09632395	
4	b0715	2.565546658	2.567760685	2.536834823	2.513905138	2.533385516	2.555976111	2.552486193	
5	b0185	6.661911901	6.094619316	7.696563672	6.963963632	7.476477903	7.20705541	7.645863323	
6	b3255	12.24120191	12.17555935	12.10939238	12.14068385	11.88957437	11.98381957	11.53553843	
7	b3256	8.626216539	8.9348643	8.754524155	9.051725652	8.586089409	8.749223826	8.216142737	
8	b2316	10.30507868	10.30964272	10.70406165	10.96516547	10.13253158	10.50413996	9.648748283	
9	b4015	9.463208813	9.972113349	8.9044422	9.819722455	12.72892109	12.51954356	12.59735882	
10	b4014	11.97537857	12.07528508	11.84044276	11.72425199	13.23488622	13.29639349	13.19317442	
11	b0114	11.76711824	11.58164936	11.88583332	11.57104461	12.49915551	12.55704041	12.55320524	
12	b0115	11.02053116	10.81358943	11.14457251	11.36213807	11.755529	12.04086088	12.18448931	
13	b4016	2.963904777	2.966189525	2.93471414	2.895148444	4.296326343	3.846552601	3.597761521	

Because we want to visualize reaction fluxes and gene expressions on a map, we firstly load the reaction fluxes which were simulated by setting the LB and UB of EX_o2(e) to close and open bound for anaerobic and aerobic conditions (nO and Oxy), respectively. The right upper panel of reaction shows the reaction fluxes for the two conditions.



The right lower panel of reaction shows the expressions of associated genes. In default, the gene with the largest expression among the associated genes will be present. You can click on the small right arrow to present other gene expressions of associated genes or you can click on the small down arrow to show all gene expressions of associated genes.



In this example, we can see that the AKGDH reaction-associated genes b0116, b0726 and b0727 were up-regulated in aerobic condition and the corresponding reaction flux was increased.