

# Package ‘abcdeFBA’

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**Type** Package

**Title** ABCDE\_FBA: A-Biologist-Can-Do-Everything of Flux Balance  
Analysis with this package.

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**Author** Abhilash Gangadharan, Neha Rohatgi

**Maintainer** Abhilash Gangadharan <gangutalk@gmail.com>

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**Suggests** LIM,sybil

**Description** Functions for Constraint Based Simulation using Flux  
Balance Analysis and informative analysis of the data generated  
during simulation.

**License** GPL-2

**Lazyload** yes

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BFD_Processor	<i>BFD_Processor, a Bi-Flux-Del-Processor function</i>
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### Description

BFD\_Processor reassembles the "result(n)" files created by Exhaustive\_double\_deletion, the result files should be a complete set from 1-n and put in a folder BKO in pwd before execution. The output is a tab delimited spreadsheet Fatal\_Double\_knockouts\_unique.xls

### Usage

```
BFD_Processor(fba_object, EXSDR)
```

### Arguments

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
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EXSDR            A list of the type generated by the Exhaustive\_single\_deletion function, this list may be passed to this function or the function will generate the list on its own. It is used for removing the results of lethal single knockouts which will form pairs with all other knockouts leading to false positives of double synthetic lethality.

### Examples

```
#Function to process the results of the double knockout
#data(Ecoli_core)
#BFD_Processor(Ecoli_core,EXSDR=Exhaustive_single_deletion_results)
# A prompt will appear asking for the number of simulation pieces
#and also to make sure you have your results filled in a folder
#called BKO in the present working directory.
```

---

BYPASS\_REACTIONS\_SUBSTRATE

*BYPASS\_REACTIONS\_SUBSTRATE, to find the influx and efflux reactions available to a particular metabolite/substrate*

---

### Description

this function computes the production and consumption fluxes available to a particular metabolite from the fba\_object and returns a list of with elements Production and Consumption

### Usage

```
BYPASS_REACTIONS_SUBSTRATE(substrate_number, fba_object, verbose)
```

### Arguments

fba\_object        Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all\_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite\_name contains list of all the metabolites, reaction\_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp\_name.

substrate\_number        this is the metabolite number of interest which may be found by using the SEARCH\_metabolite function

verbose            a boolean indicating if the name of the reactions should be printed to the terminal output, default is TRUE

**Examples**

```
#To find the Bypass Reactions for a Substrate
data(Ecoli_core)
ATP_prod_consump<-BYPASS_REACTIONS_SUBSTRATE(17,Ecoli_core)
```

---

CHANGE\_OBJ\_FUNCTION    *CHANGE\_OBJ\_FUNCTION, a function to change the objective for optimization*

---

**Description**

a function akin to the COBRA function to change the objective function for FBA

**Usage**

```
CHANGE_OBJ_FUNCTION(obj_reaction, fba_object, new_wt, old_wt)
```

**Arguments**

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
obj_reaction	a reaction number which is to be made the new objective function; retrieved using the SEARCH reaction function
new_wt	the weight of the new objective, defaults to 1 but can be any number from 0~1
old_wt	the weight of the old objective, defaults to 0 but can be any number from 0~(1-new_obj_weight) or any other co-efficient if you wish for a customized objective function.

**Examples**

```
#To change the objective function of the model.
data(Ecoli_core)
ec_new_obj<-CHANGE_OBJ_FUNCTION(11,Ecoli_core,0.5,0.5)
#ec_new_obj will be identical to the Ecoli_core model except that
#the objective function would change
FBA_solve(ec_new_obj)
```

---

CHANGE\_RXN\_BOUNDS      *CHANGE\_RXN\_BOUNDS, Change Reaction Bounds*

---

### Description

this function helps to modify the bounds on the fba\_object, it returns an object of the same type as the model, inspired by the COBRA-function of the same name

### Usage

```
CHANGE_RXN_BOUNDS(reaction_number, fba_object, lb, ub)
```

### Arguments

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
lb	the new value of the lower bound
ub	the new value of the upper bound
reaction_number	the reaction number of the reaction the bounds of which have to be changed

### Examples

```
#Changing Reaction Bounds, to simulate a reaction deletion.  
data(Ecoli_core)  
Ec_mutant<-CHANGE_RXN_BOUNDS(reaction_number=36, fba_object=Ecoli_core,  
lb=0, ub=0)
```

---

DEGREE_MEASURE	<i>DEGREE_MEASURE</i> , measures the in-degree and out-degree of the metabolites in the network.
----------------	--

---

### Description

this is a simple function to determine the degree measure, it uses the fba\_object and computes and writes the measures to disk as a tab-separated spreadsheet

### Usage

```
DEGREE_MEASURE(fba_object, file="Degree_measure")
```

### Arguments

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
file	a filename for the tab delimited output file which is generated by this function

### Examples

```
#Determining the Degree Measure of the Core E.coli Metabolic Network
data(Ecoli_core)
#DEGREE_MEASURE(fba_object=Ecoli_core)
```

---

Ecoli_core.rda	<i>E.coli core model 72 reactions and 95 metabolites</i>
----------------	--

---

### Description

This "list" form of the E.coli core model was created from the S4 object created by readSBMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

### Usage

```
Ecoli_core
```

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database

**References**

Systems Biology Properties of Reconstructed Networks

---

*Ec\_iAF1260\_flux1.rda* *E.coli model 1260 ORF's*

---

**Description**

This "list" form of the E.coli model iAF1260 was created from the S4 object created by readSBMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

**Usage**

`Ec_iAF1260_flux1`

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database

**References**

Feist AM, Henry CS, Reed JL, Krummenacker M, Joyce AR, Karp PD, Broadbelt LJ, Hatzimanikatis V and Palsson BO, A genome-scale metabolic reconstruction for Escherichia coli K-12 MG1655 that accounts for 1260 ORFs and thermodynamic information, Molecular Systems Biology.

---

Ec\_iAF1260\_flux2      *E.coli model 1260 ORF's*

---

**Description**

This "list" form of the E.coli model iAF1260 was created from the S4 object created by readSBMLmod function from the package Sybil using the Sybil\_2\_FBA\_obj function.

**Usage**

Ec\_iAF1260\_flux2

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database

**References**

Feist AM, Henry CS, Reed JL, Krummenacker M, Joyce AR, Karp PD, Broadbelt LJ, Hatzimanikatis V and Palsson BO, A genome-scale metabolic reconstruction for Escherichia coli K-12 MG1655 that accounts for 1260 ORFs and thermodynamic information, Molecular Systems Biology.

---

Ec\_iJR904      *E.coli model 904 ORF's*

---

**Description**

This "list" form of the E.coli model iJR904 was created from the S4 object created by readSBMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

**Usage**

Ec\_iJR904

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database



## References

Reed JL, Vo TD, Schilling CH and Palsson BO, An expanded genome-scale metabolic reconstruction for Escherichia coli K-12 (iJR904 GSM/GPR) Genome Biology 2003

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Exhaustive\_double\_deletion

*Exhaustive\_double\_deletion, a function for computing synthetic double knockouts.*

---

## Description

Exhaustive\_double\_deletion enables parallel computing of double knockouts by splitting and running the simulation as different instances on the same multi-core machine. The results of the simulation are stored into files named result1, result2 etc. which may be reassembled using the BFDProcessor function

## Usage

Exhaustive\_double\_deletion(fba\_object, thread\_no, core\_number)

## Arguments

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
thread_no	This specifies the thread number, it is useful to think of the thread number as a chunk of the complete combination of all pairs of reactions that can be formed. This depends on the number of the cores you are employing which must obviously be a fixed number. For instance if you have a 40 core computer you can divide the double-knockout simulation into 40 chunks. In that case the thread number would refer to the chunks of simulation as 0-39, with thread 0 performing the first 1/40th part of the simulation
core_number	core_number specifies the number of cores that are available to you for performing double knockout simulations. It should be a fixed number for a particular instance of simulations on one model

**Examples**

```
#Performing double knockouts in a 2 core computer
data(Ecoli_core)
#Exhaustive_double_deletion(Ecoli_core,0,2)
#New instance of R
data(Ecoli_core)
#Exhaustive_double_deletion(Ecoli_core,1,2)
```

---

**Exhaustive\_single\_deletion**

*Exhaustive\_single\_deletion, a function that deletes each reaction in the network one at a time and returns results describing reaction lethality*

---

**Description**

This function takes the argument of type `fba_object` which would be a FBA object and performs an exhaustive deletion of all the reactions in the reaction network and returns a list of biomass generated for all the deletions, the sub-optimal deletions, the super-optimal deletions and non-lethal deletions, also generates a pdf containing a distribution of the fatal reactions according to their sub-systems and a histogram of the biomass distribution for each deletion.

**Usage**

```
Exhaustive_single_deletion(fba_object, reactions, plot_to_file)
```

**Arguments**

<code>fba_object</code>	Is a list containing the data required to perform flux balance analysis. The elements of the list are <code>mat</code> which is the stoichiometric matrix, <code>dir</code> which gives the direction of the equality constraints, <code>obj</code> specifies the objective function for the simulation, <code>bounds</code> specifies the lower and upper inequality constraints, <code>rhs</code> is the right hand side of the steady state expression, <code>types</code> refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", <code>max</code> is a Boolean specifying the type of optimization, "Maximization" by default, <code>all_genes</code> is all the genes present in the model, <code>gpr</code> contains boolean expressions of gene essentiality for the corresponding reactions in the model, <code>metabolite_name</code> contains list of all the metabolites, <code>reaction_list</code> contains all the reactions present in the model, <code>compartment</code> is a numeric identifier for each reaction the key for which is in <code>comp_name</code> .
<code>reactions</code>	is a vector containing the reaction numbers to be deleted, if none are supplied all the reactions in the model are deleted iteratively.
<code>plot_to_file</code>	is a boolean indicating if a PDF output of the results of the single deletion should be made.

**Examples**

```
#Performing an exhaustive single reaction deletion
data(Ecoli_core)
#Results<-Exhaustive_single_deletion(Ecoli_core)
```

---

FBA\_solve

*FBA\_solve, a function to solve CBM problems*


---

**Description**

This function sugar-coats Rglpk\_solve\_LP which is a function provided by the Rglpk package, FBA\_solve solves FBA problems using Rglpk\_solve\_LP and gives the solver output in a CBM context including solver error messages and graceful degradation.

**Usage**

```
FBA_solve(fba_object, precision, verbosity, maximize)
```

**Arguments**

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
precision	A number indicating the precision of the flux solution after the decimal point, defaults to 6
verbosity	is a Boolean indicating if the verbose output of the LP solver should be displayed during simulation, defaults to FALSE
maximize	is a Boolean that can over-ride the default mode of optimization (maximization) and minimize if FALSE

**Examples**

```
# Flux Balance Analysis performed on a core-metabolism model of E.coli
data(Ecoli_core)
FBA_solve(fba_object=Ecoli_core, precision=6)
```

---

 flux\_difference\_plotter

*flux\_difference\_plotter, a function to plot two pre-existing flux distributions obtained using FBA\_solve*

---

## Description

To analyse the effect of reaction deletions on the fluxome requires a contextual visualization, simple plots give little insight on what the results of the simulation mean. This function uses the annotations inherent to SBML models and generates comparative overlapping fluxome bar graphs depicting the overlap/change in flux based on Sub-system wise classification and generates PDF's for the same. The color scheme is green for wild-type and red for mutant. Overlaps of red and green generate brown while overshoot mutant fluxes show up as magenta-pink, also separate PDF's are generated for increased and decreased fluxes.

## Usage

```
flux_difference_plotter(wt_flux, mut_flux, fba_object, graph_fname)
```

## Arguments

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
wt_flux	A list containing the solution to an FBA problem which is returned by the FBA_solve function. By convention the fluxes in this list represent the Wild-type strain and will appear as green bar plots.
mut_flux	A list containing the solution to an FBA problem which is returned by the FBA_solve function. By convention the fluxes in this list represent the Mutant-type strain and will appear as red bar plots.
graph_fname	A string to name the output files

## Examples

```
# A comparison of two flux distributions generated by FBA_solve
data(Ecoli_core)
```

```

#The reaction number for O2 exchange is 36 by setting
#the corresponding bounds to zero we make a aerobic
#respiration deficient mutant

Ec_Mutant<-CHANGE_RXN_BOUNDS(36,Ecoli_core,0,0)

mut_flux<-FBA_solve(Ec_Mutant)
wt_flux<-FBA_solve(Ecoli_core)

#flux_difference_plotter(wt_flux,mu_flux,Ecoli_core,
#graph_fname="Flux_comparison")

```

---

Flux\_Ranger

*Flux\_Ranger - a function to create a flux ramp.*


---

## Description

a function to create a linear ramp for the specified flux automatically/intelligently.

## Usage

```
Flux_Ranger(reaction_number, fba_object, divs, art_limit_range)
```

## Arguments

reaction_number	reaction of which the ramp is to be created
fba_object	list containing the necessary elements making up the flux balance model.
divs	the number of divisions for the ramp
art_limit_range	in case the "intelligent ramp" misbehaves you can force the ramp into a particular numeric range by concatenating the lower and upper numeric limits into this variable.

## Examples

```

#Creating a ramp for any reaction
data(Ecoli_core)
flux_range<-Flux_Ranger(reaction_number=12, fba_object=Ecoli_core, divs=10)

```

---

FLUX_VAR_ANALYSIS	<i>FLUX_VAR_ANALYSIS, a function to perform a flux variability analysis.</i>
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---

### Description

FVA can indicate the decrease in network robustness caused by non-lethal deletions.

### Usage

```
FLUX_VAR_ANALYSIS(fba_object, reactions, filename)
```

### Arguments

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
reactions	subset of reaction numbers on which to perform FVA, if left empty, all reactions of the network will be subject to FVA
filename	A string which will be the name of the file containing the output of the FVA simulation

### Examples

```
#Simple flux variability Analysis
data(Ecoli_core)
#FLUX_VAR_ANALYSIS(Ecoli_core, filename="Wt_FVA.xls")
```

---

FVA_robustness	<i>FVA_robustness, a function to determine the change in robustness of the network caused by a mutation.</i>
----------------	--

---

### Description

FVA\_robustness builds upon FLUX\_VAR\_ANALYSIS to give the user a convenient function to examine the change in network robustness caused by a user supplied mutation. Graphical results for absolute flux span comparisons may be expected in the working directory

**Usage**

```
FVA_robustness(fba_object, mutation)
```

**Arguments**

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
mutation	A reaction number obtained using the SEARCH_reaction, or in the non-lethal reaction "list" result of an Exhaustive_single_deletion run. This reaction will be deleted during the FVA analysis

**Examples**

```
#Mutant network robustness, removing Formate Exchange which has no effect
#on fluxes
data(Ecoli_core)
#FVA_robustness(Ecoli_core,25)
```

---

Gene\_del

*Gene\_del, a function to create Gene Deletion mutants*


---

**Description**

Gene\_del interprets the boolean rules inherent to the model and returns a mutant model when supplied with the Wild-type model and a vector of gene names to be deleted.

**Usage**

```
Gene_del(query_genes, fba_object, return_reactions)
```

**Arguments**

query_genes	query_genes should be a vector the elements of which are the genes that are to be deleted from the model.
fba_object	A flux balance model containing the necessary elements for simulation

```
return_reactions
```

a boolean specifying if the reaction numbers deleted as a consequence of the gene deletion should be returned or if the mutant model itself be returned by the function

## Examples

```
#To create a mutant model .  
{data(Ec_iAF1260_flux1)  
ho<-Gene_del(c("b3040"),Ec_iAF1260_flux1,return_reactions=TRUE)  
}
```

---

H\_pylori\_iIT341

*Helicobacter pylori* by Ines Thiele

---

## Description

This "list" form of the H. pylori model iIT341 was created from the S4 object created by readSBMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

## Usage

```
H_pylori_iIT341
```

## Format

A list containing the information required to perform FBA and annotations for intelligible output.

## Source

BiGG database

## References

Thiele I, Vo TD, Price ND, Palsson BO, Expanded Metabolic Reconstruction of *Helicobacter pylori* (iIT341 GSM/GPR): an In Silico Genome-Scale Characterization of Single- and Double-Deletion Mutants, *Journal of Bacteriology*.



---

H_sapien_Recon1	<i>H. sapien Recon1, human metabolic network reconstruction</i>
-----------------	---

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**Description**

This "list" form of the human model Recon1 was created from the S4 object created by readSBMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

**Usage**

```
H_sapien_Recon1
```

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database

**References**

Duarte NC, Becker SA, Jamshidi N, Thiele I, Mo ML, Vo TD, Srivas R and Palsson BO, Global reconstruction of the human metabolic network based on genomic and bibliomic data, PNAS.

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M_barkeri_iAF629	<i>Genome scale metabolic model for the archaeal methanogen M. Barkeri</i>
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---

**Description**

This "list" form of the M.Barkeri model was created from the S4 object created by readSBMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

**Usage**

```
M_barkeri_iAF629.rda
```

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database

**References**

Modeling methanogenesis with a genome scale metabolic reconstruction of *Methanosarcina barkeri* by Feist AM, Scholten JCM, Palsson BO

---

M\_tb\_iNJ661                      *M. tuberculosis model iNJ661*

---

**Description**

This "list" form of the M.tb model iNJ661 was created from the S4 object created by readSBMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

**Usage**

M\_tb\_iNJ661

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database

**References**

Bordbar A, Lewis NE, Schellenberger J, Palsson BO, Jamshidi N, Insight into human alveolar macrophage and *M. tuberculosis* interactions via metabolic reconstructions, *Molecular Systems Biology*.

---

PERTURBATION\_analysis    *PERTURBATION\_analysis, a function for robustness analysis.*

---

**Description**

Robustness analysis is described procedurally in the COBRA-ToolBox manual. This function encodes the basic principle of the procedure and returns the result as an X versus Y list and generates a plot on successful completion.

**Usage**

```
PERTURBATION_analysis(reaction_number, fba_object, y_axis_rxn=NULL,  
plot_to_file=FALSE, write_FLD_file=FALSE, ret_FLD_matrix=FALSE)
```

**Arguments**

<code>fba_object</code>	Is a list containing the data required to perform flux balance analysis. The elements of the list are <code>mat</code> which is the stoichiometric matrix, <code>dir</code> which gives the direction of the equality constraints, <code>obj</code> specifies the objective function for the simulation, <code>bounds</code> specifies the lower and upper inequality constraints, <code>rhs</code> is the right hand side of the steady state expression, <code>types</code> refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", <code>max</code> is a Boolean specifying the type of optimization, "Maximization" by default, <code>all_genes</code> is all the genes present in the model, <code>gpr</code> contains boolean expressions of gene essentiality for the corresponding reactions in the model, <code>metabolite_name</code> contains list of all the metabolites, <code>reaction_list</code> contains all the reactions present in the model, <code>compartment</code> is a numeric identifier for each reaction the key for which is in <code>comp_name</code> .
<code>reaction_number</code>	Reaction number of the reaction to be ramped down. This number may be obtained by using the function <code>SEARCH_reaction</code> for a text search through the <code>FBA_obj-reaction</code> list.
<code>y_axis_rxn</code>	Reaction to be plotted on the y axis in case it is not the objective function.
<code>plot_to_file</code>	Boolean indicating if the graph to be printed to a file.
<code>write_FLD_file</code>	The flux vector for each point of perturbation is stored into the columns of a matrix, this boolean indicates if that matrix should be written out to a tab-delimited file.
<code>ret_FLD_matrix</code>	The flux vector for each point of perturbation is stored into the columns of a matrix, this boolean indicates if that matrix should be added to the list returned by this function.

**Examples**

```
#Perturbation analysis of fluxes
data(Ecoli_core)
#Ec_xy_O2<-PERTURBATION_analysis(reaction_number=36,Ecoli_core)
```

---

 PHPP

---

*PHPP, phenotypic phase plane analysis*


---

**Description**

this function helps in performing a phenotypic phase plane analysis, a visualization of the effect of two input fluxes on the value of the objective function.

**Usage**

```
PHPP(reaction_number, fba_object, PCS, flux_range,
ret_OBJ_mat, surf_col, divs, dimension, animate, objective)
```

**Arguments**

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
PCS	a string to search for the primary carbon source of the model, for example PCS="D glucose" will display all the reactions with that string in them, you need to choose the appropriate reaction number of the Carbon source before continuing. In case you are performing a PhPP for alternate carbon sources then the primary carbon source is supposed to be shut-down to get a correct picture of the flux cone.
reaction_number	two reaction numbers that specify the two input fluxes that make the x and y axes of the PhPP
flux_range	the range between which the input fluxes are to be varied
ret_OBJ_mat	boolean indicating if the PHPP matrix should be returned
surf_col	character string for surface color
divs	number of divisions of ramp- note this increases computation time of PHPP by n-squared
dimension	characters- "2" or "3" specifying if the PHPP should be viewed in 2-D or 3-D
animate	boolean indicating if the 3-D plot should be animated; spins it around once on each axes
objective	reaction_number to be used as the objective parameter for PhPP

**Examples**

```
#Performing a phenotypic phase plane analysis of
#glucose and oxygen in Core E.coli Metabolism
data(Ecoli_core)
#PHPP(reaction_number=c(28,36),fba_object=Ecoli_core,
#PCS="glucose",flux_range=c(1,15),ret_OBJ_mat=FALSE,surf_col="red")
# a menu pops up asking to select the primary carbon source,
#select D glucose for Ecoli_core
```

---

SEARCH_metabolite	<i>SEARCH_metabolite, a function to search for metabolites in a model using a simple text query</i>
-------------------	---

---

### Description

this function helps to search for metabolites in a model using a simple text query, it returns a list of possible hits along with their metabolite numbers and locations in the compartments of the model

### Usage

```
SEARCH_metabolite(metabolite_name, fba_object)
```

### Arguments

**fba\_object** Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all\_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite\_name contains list of all the metabolites, reaction\_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp\_name.

**metabolite\_name** is a character string containing a part or the complete name of the metabolite to be searched for in the fba object. If you give a number it will retrieve the name.

### Examples

```
#To search for the metabolite in the model
data(Ecoli_core)
SEARCH_metabolite("ATP",Ecoli_core)
```

---

SEARCH_reaction	<i>SEARCH_reaction, a function to search for reactions in a model using a simple text query</i>
-----------------	---

---

### Description

this function helps to search for reactions in a model using a simple text query, it returns a list of possible hits along with their reaction numbers, these reaction numbers are to be used with other perturbation/knockout/ optimality/robustness analysis functions

**Usage**

```
SEARCH_reaction(react_name, fba_object)
```

**Arguments**

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name.
react_name	is a character string containing a part or the complete name of the reaction to be searched for in the fba object. If you give a number it will give the name of the reaction

**Examples**

```
#Search for reactions involving glucose
data(Ecoli_core)
SEARCH_reaction("glucose", Ecoli_core)
```

---

SINGLE_DEL_FLUXOME	<i>SINGLE_DEL_FLUXOME, a function to generate comparative fluxome graphs</i>
--------------------	--

---

**Description**

To analyse the effect of reaction deletions on the fluxome requires a contextual visualization, simple plots give little insight on what the results of the simulation mean. This function uses the annotations inherent to SBML models and generates comparative overlapping fluxome bar graphs depicting the overlap/change in flux based on Sub-system wise classification and generates PDF's for the same. The color scheme is green for wild-type and red for mutant. Overlaps of red and green generate brown while overshoot mutant fluxes show up as magenta-pink, also separate PDF's are generated for increased and decreased fluxes.

**Usage**

```
SINGLE_DEL_FLUXOME(fba_object, deletion_number)
```

**Arguments**

`fba_object` Is a list containing the data required to perform flux balance analysis. The elements of the list are `mat` which is the stoichiometric matrix, `dir` which gives the direction of the equality constraints, `obj` specifies the objective function for the simulation, `bounds` specifies the lower and upper inequality constraints, `rhs` is the right hand side of the steady state expression, `types` refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", `max` is a Boolean specifying the type of optimization, "Maximization" by default, `all_genes` is all the genes present in the model, `gpr` contains boolean expressions of gene essentiality for the corresponding reactions in the model, `metabolite_name` contains list of all the metabolites, `reaction_list` contains all the reactions present in the model, `compartment` is a numeric identifier for each reaction the key for which is in `comp_name`.

`deletion_number` Reaction number of the reaction to be deleted. This number may be obtained by using the function `SEARCH_reaction` for a text search through the `fba_object`-reaction list.

**Examples**

```
#Compare the flux distributions of the mutant with the wild-type
data(Ecoli_core)
#SINGLE_DEL_FLUXOME(Ecoli_core,36)
```

---

<code>Sybil_2_FBA_obj</code>	<i>Sybil_2_FBA_obj, a function to convert a model generated by Sybil into the one used by abcdeFBA.</i>
------------------------------	---

---

**Description**

Changes the S4 object read by Sybil which is available on CRAN into a list usable by abcdeFBA.

**Usage**

```
Sybil_2_FBA_obj(Sybil_S4_object)
```

**Arguments**

`Sybil_S4_object` Is an S4 object of class `modelorg`

**Examples**

```
#Model conversion
#data(Ecoli_core) - # the Ecoli_core model included in the Sybil package
#FBA_obj<-Sybil_2_FBA_obj(Ecoli_core)
```

---

S\_aureus\_iSB619

*Staphylococcus aureus* model iSB619 by S.Becker

---

**Description**

This "list" form of the S. aureus model iSB619 was created from the S4 object created by readS-BMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

**Usage**

S\_aureus\_iSB619

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database

**References**

Becker SA, Palsson BO, Genome-scale reconstruction of the metabolic network in Staphylococcus aureus N315: an initial draft to the two-dimensional annotation

---

S\_cerevisiae\_IND750

*Saccharomyces Cerevisiae* iND750

---

**Description**

This "list" form of the yeast model iND750 was created from the S4 object created by readS-BMLmod from the package Sybil using the Sybil\_2\_FBA\_obj function.

**Usage**

S\_cerevisiae\_IND750

**Format**

A list containing the information required to perform FBA and annotations for intelligible output.

**Source**

BiGG database



## References

Duarte NC ,Herrgard MJ , Palsson BO, Reconstruction and Validation of *Saccharomyces cerevisiae* iND750, a Fully Compartmentalized Genome-Scale Metabolic Model, Genome Research

---

View_objective	<i>View_objective, a function that shows the components of the objective function.</i>
----------------	--

---

## Description

This function takes the argument of type fba\_object and prints out the components of the objective function, i.e substrates and products involved and their corresponding pseudo-stoichiometry

## Usage

```
View_objective(fba_object)
```

## Arguments

fba_object	Is a list containing the data required to perform flux balance analysis. The elements of the list are mat which is the stoichiometric matrix, dir which gives the direction of the equality constraints, obj specifies the objective function for the simulation, bounds specifies the lower and upper inequality constraints, rhs is the right hand side of the steady state expression, types refers to the numeric nature of the variables which in case of FBA happens to be "Continuous", max is a Boolean specifying the type of optimization, "Maximization" by default, all_genes is all the genes present in the model, gpr contains boolean expressions of gene essentiality for the corresponding reactions in the model, metabolite_name contains list of all the metabolites, reaction_list contains all the reactions present in the model, compartment is a numeric identifier for each reaction the key for which is in comp_name
------------	---

## Examples

```
#Viewing the objective function of the E.coli core metabolism
data(Ecoli_core)
View_objective(Ecoli_core)
```

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