

S is stoichiometric representation of metabolic networks corresponding to the reactions in the biochemical pathway. In an each column of the S is a biochemical reaction and in each row is a precise metabolite. There is a stoichiometric coefficient of zero, which means that metabolite not participate in that distinct reaction. The coefficient also can be positive when the appropriate metabolite is produced, or negative for every metabolite depleted [1].

```

ReactionFormulas = {'glc-D[e] -> glc-D[c]',...
  'glc-D[c] + atp[c] -> H[c] + adp[c] + g6p[c]',...
  'g6p[c] <=> f6p[c]',...
  'atp[c] + f6p[c] -> H[c] + adp[c] + fdp[c]',...
  'fdp[c] + h2o[c] -> f6p[c] + pi[c]',...
  'fdp[c] -> g3p[c] + dhap[c]',...
  'dhap[c] -> g3p[c]'};
ReactionNames = {'GLCt1', 'HEX1', 'PGI', 'PFK', 'FBP', 'FBA', 'TPI'};
lowerbounds = [-20, 0, -20, 0, 0, -20, -20];
upperbounds = [20, 20, 20, 20, 20, 20, 20];
model = createModel(ReactionNames, ReactionNames, ReactionFormulas, 'lowerBoundList', lowerbounds

```

We can now have a look at the different model fields created. The stoichiometry is stored in the S field of the model, which was described above. Since this is commonly a sparse matrix (i.e. it does contain a lot of zeros, to display it it is useful to use the full representation)

```
full(model.S)
```

Some descriptive fields always present are model.mets and model.rxns which represent the metabolites and the reactions respectively.

```

model.mets
model.rxns

```

Fields in the COBRA model are commonly column vectors, which can be an important detail when writing functions manipulating these fields.

There are a few more fields present in each COBRA model:

model.lb, indicating the lower bounds of each reaction, and model.ub indicating the upper bound of a reaction.

```

[{'Reaction ID', 'Lower Bound', 'Upper Bound'}; model.rxns, num2cell(model.lb), num2cell(model.ub)
printFluxBounds(model); %This is a convenience function which does pretty much the same as the

```

Before we start to modify the model, it might be useful to store some of the current properties of the model

```

mets_length = length(model.mets);
rxns_length = length(model.rxns);

```

Creating, adding and handling reactions

If we want to add a reaction to the model or modify an existing reaction we are using function addReaction.

We will add some more reactions from glycolysis.

- The formula approach

```

model = addReaction(model, 'GAPDH', 'reactionFormula', 'g3p[c] + NAD[c] + 2 pi[c] -> NADH[c] +
model = addReaction(model, 'PGK', 'reactionFormula', '13bpg[c] + adp[c] -> atp[c] + 3pg[c]');

```

```
model = addReaction(model, 'PGM', 'reactionFormula', '3pg[c] <=> 2pg[c] ');
```

Display of stoichiometric matrix after adding reactions. Note the enlarge link when you move your mouse over the output to display the full matrix:

```
full(model.S)  
% one extra column is added(for added reaction) and 4 new rows(for A, B, C, D metabolites)
```

The following functions are used when we want to search reactions sequence in the model and change the order of the selected reaction.

```
rxnID = findRxnIDs(model, model.rxns);  
model = moveRxn(model, 8, 1);
```

While the latter does not modify the structure as such it can help in keeping a model tidy.

- [The list approach](#)

The addReaction function has ability to recognize duplicate reactions when an order of metabolites and an abbreviation of the reaction are different.

```
model = addReaction(model, 'GAPDH2', 'metaboliteList', {'g3p[c]', 'NAD[c]', 'pi[c]', '13bpg[c]'
```

Since the second call should not have added anything we will check if the number of the reaction increased by the three reactions we added (and not by the one duplicated) and the number of metabolites was incremented by five (13bpg, NAD, NADH, 23bpg and 2pg).

```
assert(length(model.rxns) == rxns_length + 3);  
assert(length(model.mets) == mets_length + 5);
```

Adding Exchange, Sink and Demand reactions

Specific type of reactions in the constraint-based models are reactions that are using and recycling accumulated metabolites, or producing required metabolites in the model.

1. *Exchange reactions* - Reactions added to the model to move metabolites across the created *in silico* compartments. Those compartments represent intra- and intercellular membranes.
2. *Sink reactions* - The metabolites, produced in reactions that are outside of an ambit of the system or in unknown reactions, are supplied to the network with reversible sink reactions.
3. *Demand reactions* - Irreversible reactions added to the model to consume metabolites that are deposited in the system.

There are two ways to implement that kind of reactions:

1. **Use addReaction with the documented function call:**

```
model = addReaction(model, 'EX_glc-D[e]', 'metaboliteList', {'glc-D[e]'}, 'stoichCoeffList', [-
```

In the bigger networks we can find our exchange reactions with the following functions:

```
[selExc,selUpt] = findExcRxns(model, 0, 1) % determines whether a reaction is a general exchange
```

2. Use a utility function to create a particular reaction type: addExchangeRxn, addSinkReactions, addDemandReaction.

```
model = addExchangeRxn(model, {'glc-D[e]', 'glc-D[c]'})
```

```
model = addSinkReactions(model, {'13bpg[c]', 'NAD[c]'})
```

```
model = addDemandReaction(model, {'dhap[c]', 'g3p[c]'})
```

Setting ratio between the reactions and changing reactions boundary

It is important to emphasize that previous knowledge base informations should be taken into account. Most of them could disrupt future analysis of the model.

For instance, if it is familiar that flux through one reaction is X times the flux through another reaction, it is recommended to specify that in your model.

E.g. $1 \nu \text{EX_glc} - D[c] = 2 \nu \text{EX_glc} - D[e]$

```
model = addRatioReaction (model, {'EX_glc-D[c]', 'EX_glc-D[e]'}, [1; 2]);
```

Altering Reaction bounds

In order to respect the transport and exchange potential of a particular metabolite, or to resemble the different conditions in the model, we frequently need to set appropriate limits of the reactions.

```
model = changeRxnBounds(model, 'EX_glc-D[e]', -18.5, 'l');
```

Modifying Reactions

The addReaction function also is a good choice when modifying reactions. By supplying a new stoichiometry, the old will be overwritten. For example further up, we added awrong stoichiometry for the GAP-Dehydrogenase with a phosphate coefficient of 2. (easily visiple by printing the reaction)

```
printRxnFormula(model, 'rxnAbbrList', 'GAPDH');
```

We can correct this by simply calling ddReaction again with the corrected stoichiometry. In essence parts which are not provided are taken from the old reaction, and only the new ones overwrite the existing data

```
model = addReaction(model, 'GAPDH', 'metaboliteList', {'g3p[c]', 'NAD[c]', 'pi[c]', '13bpg[c]'}
```

We might also want to add a gene rule to the reaction. this can either be done using

```
model = changeGeneAssociation(model, 'GAPDH', 'G1 and G2');
```

```
printRxnFormula(model, 'rxnAbbrList', {'GAPDH'}, 'gprFlag', true);
```

The other option to achieve this is to use `addReaction` and the `geneRule` parameter

```
model = addReaction(model, 'PGK', 'geneRule', 'G2 or G3', 'printLevel', 0);  
printRxnFormula(model, 'gprFlag', true);
```

Remove reactions and metabolites

In order to detach reactions from the model, the following function has been used:

```
model = removeRxns(model, {'EX_glc-D[c]', 'EX_glc-D[e]', 'sink_13bpg[c]', 'sink_NAD[c]', 'DM_13bpg[c]'});  
assert(rxns_length + 3 == length(model.rxns));  
% The reaction length has been reevaluated
```

Remove metabolites

```
model = removeMetabolites(model, {'3pg[c]', '2pg[c]'}, false);
```

For instance, in previous code the many metabolites from 'GAPDH' were deleted, but the reaction is still present in the model (since there are more metabolites left). The false indicates, that empty reactions should not be removed.

To delete metabolites and reactions with zero rows and columns, the following function can be used:

```
model = removeTrivialStoichiometry(model)  
model = removeRxns(model, {'GAPDH', 'PGK'});
```

Search for duplicates and comparison of two models

Since genome-scale metabolic models are expanding every day [2], the need for comparison and merge of them is also spreading.

The elementary functions for the model manipulation, besides main actions, simultaneously perform the structural analysis and comparison (e.g. `addReaction`). Likewise, there are additional functions that are only dealing with analysing similarities and differences within and between the models.

- Checking for reaction duplicates by reaction abbreviation, by using method 'S' that will not detect reverse reactions, and method 'FR' that will neglect reactions direction:

```
[model, removedRxn, rxnRelationship] = checkDuplicateRxn(model, 'S', 1, 1);
```

Adding duplicate reaction to the model:

```
model = addReaction(model, 'GLCt1_duplicate_reverse', 'metaboliteList', {'glc-D[e]', 'glc-D[c]'},  
    'stoichCoeffList', [1 -1], 'lowerBound', 0, 'upperBound', 20, 'checkDuplicate', 0);  
  
fprintf('>> Detecting duplicates using S method\n');  
method = 'S';  
%will not be removed as does not detect reverse reaction
```

```
[model, removedRxn, rxnRelationship] = checkDuplicateRxn(model, method, 1, 1);
assert(rxns_length + 1 == length(model.rxns));
% The reaction length has been reevaluated

fprintf('>> Detecting duplicates with using FR method\n');
method = 'FR';
%will be removed as detects reverse reaction
[model, removedRxn, rxnRelationship] = checkDuplicateRxn(model, method, 1, 1);
assert(rxns_length == length(model.rxns));
```

- Function checkCobraModelUnique marks the reactions and metabolites that are not unique in the model.

```
model = checkCobraModelUnique(model, false)
```

Changing the model's objective

Simulating different conditions in the model is often necessary in favor of performing calculations that investigate a specific objective. One of the elementary objectives is optimal growth [3]. Model can be modified to get different conditions with changing the model objective:

```
modelNew = changeObjective(model, 'GLCt1', 0.5);

% multiple rxns, default coefficient (1)
modelNew = changeObjective(model, {'PGI'; 'PFK'; 'FBP'});
```

The direction of reactions

When reaction is reversible and we want to remove it, it is necessary to first change the reverse sign and then remove reaction afterwards.

```
fprintf('>> Converting to Irreversible\n');
load('testModelManipulation.mat', 'model', 'modelIrrev');
[testModelIrrev, matchRev, rev2irrev, irrev2rev] = convertToIrreversible(model);
```

Following function is comparing the differences and similarities between two models:

```
assert(isSameCobraModel(modelIrrev, testModelIrrev));
```

Converting to reversible

```
fprintf('>> Convert to Reversible\n');
testModelRev = convertToReversible(testModelIrrev);
load('testModelManipulation.mat', 'modelRev');

assert(isSameCobraModel(modelRev, testModelRev));
```

References

[1] Orth, J. D., Thiele I., and Palsson, B. Ø. (2010). What is flux balance analysis? *Nat. Biotechnol.*, 28(3), 245–248.

[2] Feist, A. M., Palsson, B. (2008). The Growing Scope of Applications of Genome-scale Metabolic Reconstructions: the case of *E. coli*. *Nature Biotechnology*, 26(6), 659–667.

[3] Feist, A. M., Palsson, B. O. (2010). The Biomass Objective Function. *Current Opinion in Microbiology*, 13(3), 344–349.