Creating a Model

Author(s): Thomas Pfau, University of Luxembourg

Reviewer(s):

INTRODUCTION

This tutorial explains the most basic functions provided by The COBRA Toolbox to create a model from scratch (i.e. define all relevant reactions and build a model from them).

MATERIALS

In this tutorial, two models are created: a small toy model and a simple model of glycolysis. The latter is also used in the model manipulation tutorial.

PROCEDURE

1. Create a simple toy model

To create a new model, there is a simple function createModel:

```
emptymodel = createModel()

emptymodel =
    rxns: {0x1 cell}
    S: []
    lb: [0x1 double]
    ub: [0x1 double]
```

genes: {0×1 cell}
osense: -1
csense: ''
rxnGeneMat: []

c: [0×1 double]
mets: {0×1 cell}
b: [0×1 double]
rules: {0×1 cell}

Calling it as above, yields an empty model struct with all required fields defined. To add reactions or metabolites please have a look at the tutorial for model manipulation.

There is also a possibility to immediately create a model with multiple reactions using <code>createModel</code>. To do so, a list of reaction identifiers, reaction names and reaction formulas has to be supplied.

Let's consider the following toy model:

The reactions are:

```
R1 = '1 Met1 + Met2 -> Met3'

R1 = 
1 Met1 + Met2 -> Met3

R2 = 'Met3 <=> 2 Met4'

R2 = 
Met3 <=> 2 Met4
```

Reaction formulas are given as metabolites and their stoichiometric coefficient concatenated by +. Products and substrates are separated by a reversibility indicator, with -> indicating an irreversible reaction and <=> indicating a reversible reaction.

To be able to use <code>createModel</code> to build this model, we also have to define the reaction Identifiers and the reaction names:

```
reactionIdentifiers = {'R1', 'R2'}

reactionIdentifiers =
    'R1'    'R2'

reactionNames = {'Reaction 1', 'Reaction 2'}

reactionNames =
    'Reaction 1'    'Reaction 2'
```

And we have to combine the reactions:

```
reactionFormulas = {R1, R2}

reactionFormulas =
   '1 Met1 + Met2 -> Met3'   'Met3 <=> 2 Met4'
```

now we can call

to create the model including the two reactions.

2. Explanation of options for the createModel function

createModel offers a couple of additional optional parameters. Those include:

• revFlagList - a double array of indications whether the reaction is reversible or not this will overwrite the indicator from the formula. (default: 1 for reversible formulas, 0 for irreversible formulas)

- lowerBoundList a double array indicating the lower bounds of the providing reactions (again, this will overwrite both revFlagList and the indication from the formula). E.g. if a revFlagList entry indicates a reversible reaction, but the lower bound is >0, the reaction will be considered as irreversible. (The default is to assume 0 for irreversible and 1000 for reversible reactions)
- upperBoundList a double array indicating the upper bounds of the reactions. (default: 1000)
- subSystemList a cell array indicating the subSystems of the reactions
- grRuleList a cell array indicating the GPR rules for a formula (in textual format e.g. Gene1 and Gene2)
- geneNameList a List of genes present in the grRuleList array
- systNameList a List (of equal size as geneNameList), that is used to translate the genes from those used in the geneNameList to those used in this list.

3. Creating a model with Gene-Protein-Reaction Association (GPR) rules

Let's assume, our network has the following GPR associations:

i.e. a complex of G1 and G2 catalyses R2 and either G3 or G4 catalyse R1. We further assume, that the flux maximum through R1 is 10 and 30 through R2

```
upperBounds = [10, 30];
grRuleR1 = 'G3 or G4';
grRuleR2 = 'G1 and G2';
grRuleList = {grRuleR1, grRuleR2};
```

The model creation call would then be:

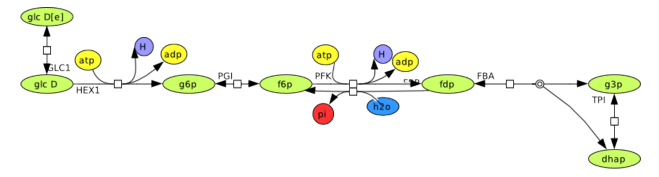
If we now compare the reactions, printing the GPR rules in both models

```
printRxnFormula(model1, 'gprFlag', 1);
R1
     Met1[c] + Met2[c]
                         <=>
                                 Met3[c]
R2
     Met3[c]
                <=>
                       2 Met4[c]
printRxnFormula(model2, 'gprFlag', 1);
R1
     Met1[c] + Met2[c]
                          <=>
                                 Met3[c]
                                            G3 or G4
R2
     Met3[c]
                <=>
                       2 Met4[c]
                                    G1 and G2
```

we see, that model2 has assigned GPR rules, while model1 does not have those.

4. Create a model of the upper part of glycolysis

We will now create a slightly more complex model (essentially, the upper part of the glycolysis) which will be used in other tutorials (e.g. ModelManipulation)



To create this model, we have to define the reactions:

```
Warning: Metabolite glc-D[e] not in model - added to the model
Warning: Metabolite glc-D[c] not in model - added to the model
        glc-D[e]
                    <=>
                           glc-D[c]
Warning: Metabolite atp[c] not in model - added to the model
Warning: Metabolite H[c] not in model - added to the model
Warning: Metabolite adp[c] not in model - added to the model
Warning: Metabolite g6p[c] not in model - added to the model
HEX1 glc-D[c] + atp[c]
                          -> H[c] + adp[c] + g6p[c]
Warning: Metabolite f6p[c] not in model - added to the model
                       f6p[c]
      g6p[c]
                <=>
Warning: Metabolite fdp[c] not in model - added to the model
     atp[c] + f6p[c]
                       ->
                               H[c] + adp[c] + fdp[c]
Warning: Metabolite h2o[c] not in model - added to the model
Warning: Metabolite pi[c] not in model - added to the model
```

```
FBP fdp[c] + h2o[c] -> f6p[c] + pi[c]
Warning: Metabolite g3p[c] not in model - added to the model
Warning: Metabolite dhap[c] not in model - added to the model
FBA fdp[c] <=> g3p[c] + dhap[c]
TPI dhap[c] <=> g3p[c]
```