

Creating a Model

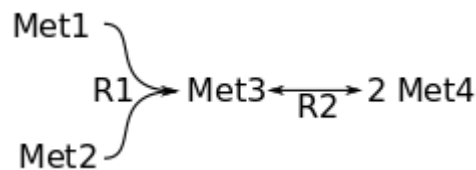
To create a new model, there is a simple function (`createModel`):

```
emptymodel = createModel()
```

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 `createModel`. To do so, a list of reaction identifiers, reaction names and reaction formulas has to be supplied.

Lets consider the following toy model:



The reactions are:

```
R1 = '1 Met1 + Met2 -> Met3'  
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 `createModel()` to build this model, we also have to define the reaction Identifiers and the reaction names:

```
reactionIdentifiers = {'R1','R2'}  
reactionNames = {'Reaction 1','Reaction 2'}
```

And we have to combine the reactions:

```
reactionFormulas = {R1,R2}
```

now we can call

```
modell = createModel(reactionIdentifiers,reactionNames,reactionFormulas);
```

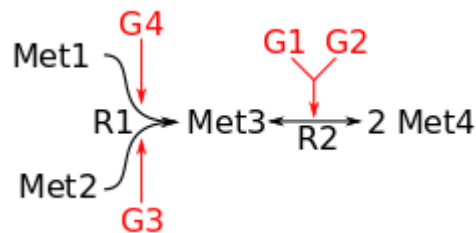
to create the model including the two reactions.

`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 irreversible 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.

Lets 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:

```

model2 = createModel(reactionIdentifiers, reactionNames, reactionFormulas, 'upperBoundList', upper
  
```

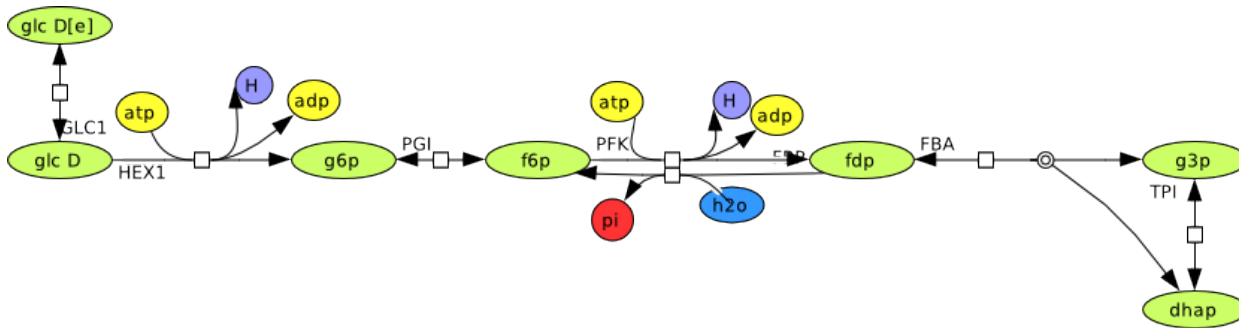
If we now compare the reactions, printing the GPR rules in both models

```

printRxnFormula(model1, 'gprFlag', 1);
printRxnFormula(model2, 'gprFlag', 1);
  
```

we see, that model 2 has assigned GPR rules, while model 1 does not have those.

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:

```
ReactionFormulas = {'glc-D[e]  -> glc-D',...
  'glc-D + atp  -> H + adp + g6p',...
  'g6p  <=> f6p',...
  'atp + f6p  -> H + adp + fdp',...
  'fdp + h2o  -> f6p + pi',...
  'fdp  -> g3p + dhap',...
  'dhap  -> g3p'};
ReactionNames = {'GLCt1', 'HEX1', 'PGI', 'PFK', 'FBP', 'FBA', 'TPI'};
lowerbounds = [-20, 0, -20, 0, 0, -20, -20];
upperbounds = [20, 20, 20, 20, 20, 20, 20];
glycolysisModel = createModel(ReactionNames, ReactionNames, ReactionFormulas, 'lowerBoundList', 1);
```