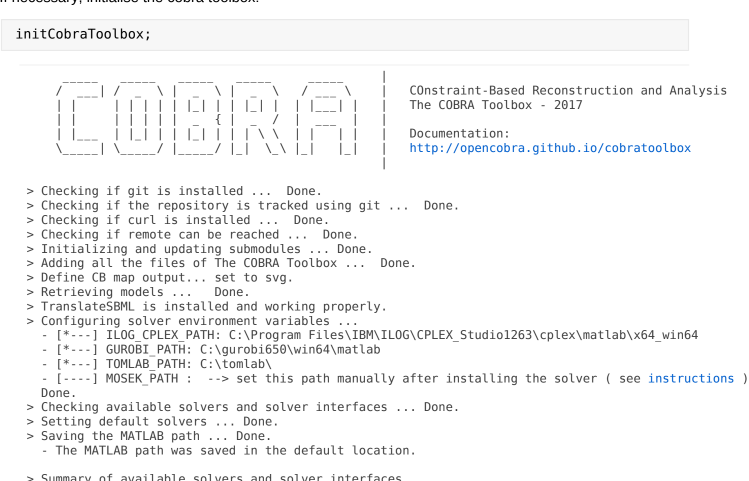
# Varying Parameters analysis

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In this tutorial, we show how computations are performed by varying one or two parameters over a fixed range of numerical values.

#### **EQUIPMENT SETUP**

If necessary, initialise the cobra toolbox:



>	Summarv	οf	available	solvers	and	solver	interfaces

Support		LP	MILP		QP	MIQP	NLP			
cplex_direct	full			0		0	0	0	-	
dqqMinos	full			0		-	-	-	-	
glpk	full			1		1	-	-	-	
gurobi	full			1		1	1	1	-	
ibm cplex	full			0		0	0	-	-	
matlab	full			1		-	-	-	1	
mosek	full			0		0	0	-	-	
pdco	full			1		-	1	-	-	

```
quadMinos full
                                             0
                                                                                   0
                                            1
tomlab cplex full
                                                                1
                                                                         1
           experimental
                                                                1
qpng
                                         - -
0 0
0 -
tomlab_snopt experimental
                                                                                   1
gurobi_mex legacy
lindo_old legacy
                                                                0
                                                                         0
lindo_legacy legacy
                                          0
                                           1
lp solve
                  legacy
                   legacy
opti
Total
                                                                                   2
+ Legend: - = not applicable, 0 = solver not compatible or not installed, 1 = solver installed.
> You can solve LP problems using: 'glpk' - 'gurobi' - 'matlab' - 'pdco' - 'tomlab_cplex' - 'lp_solve'
> You can solve MILP problems using: 'glpk' - 'gurobi' - 'tomlab_cplex'
> You can solve QP problems using: 'gurobi' - 'pdco' - 'tomlab_cplex' - 'qpng'
> You can solve MIQP problems using: 'gurobi' - 'tomlab_cplex'
> You can solve NLP problems using: 'matlab' - 'tomlab_snopt'
> Checking for available updates ...
--> You cannot update your fork using updateCobraToolbox(). [535a88 @ develop].
      Please use the MATLAB.devTools (https://github.com/opencobra/MATLAB.devTools).
```

For solving linear programming problems in the analysis, certain solvers are required:

```
changeCobraSolver ('gurobi', 'all', 1);
%changeCobraSolver ('glpk', 'all', 1);

> Solver for LPproblems has been set to glpk.
> Solver for MILPproblems has been set to glpk.
> Solver glpk not supported for problems of type MIQP. Currently used: tomlab_cplex
> Solver glpk not supported for problems of type NLP. Currently used: matlab
> Solver glpk not supported for problems of type QP. Currently used: qpng
```

The present tutorial can run with 'glpk' package, which does not require additional installation and configuration. Although, for the analysis of large models is recommended to use the 'gurobi' package. For detail information, refer to the solver installation guide: https://github.com/opencobra/cobratoolbox/blob/master/docs/source/installation/solvers.md

### **PROCEDURE**

Before proceeding with the simulations, the path for the model needs to be set up. In this tutorial, the used model is the generic model of human metabolism, Recon 3 [1]. Therefore, we assume, that the cellular objectives include energy production or optimisation of uptake rates and by-product secretion for various physiological functions of the human body. If Recon 3 is not available, please use Recon 2.

```
if exist('2017_04_28_Recon3dForCurrentDistribution.mat','file')==2
    filename = '2017_04_28_Recon3dForCurrentDistribution.mat';
    load(filename);
    model=modelRecon3model;
    clear modelRecon3model;
    model.csense(1:size(model.S,1),1)='E';
else
    filename2='Recon2.0model.mat';
    if exist('Recon2.0model.mat','file')==2
        load(filename2);
        model=Recon2model;
```

```
clear Recon2model;
    model.csense(1:size(model.S,1),1)='E';
end
end
```

If Recon2 is used, the reaction nomenclature needs to be adjusted.

```
model.rxns(find(ismember(model.rxns, 'EX_glc(e)')))={'EX_glc_D[e]'};
model.rxns(find(ismember(model.rxns, 'EX_o2(e)')))={'EX_o2[e]'};
```

#### TROUBLESHOOTING

If there are multiple energy sources available in the model; Specifying more constraints is necessary. If we do not do that, we will have additional carbon and oxygen energy sources available in the cell and the maximal ATP production.

To avoid this issue, all external carbon sources need to be closed.

```
%Closing the uptake of all energy and oxygen sources
for i=1:length(model.rxns)
    if strncmp(model.rxns{i}, 'EX ',3)
        model.subSystems{i}='Exchange/demand reaction';
    end
end
idx=strmatch('Exchange/demand reaction', model.subSystems);
for i=1:length(idx)
    if model.lb(idx(i))~=0
        c=c+1:
        uptakes{c}=model.rxns{idx(i)};
    end
end
modelalter = model;
modelalter = changeRxnBounds(modelalter, uptakes, 0, 'l');
% The alternative way to do that, in case you were using another large model,
% that does not contain defined Subsystem is
% to find uptake exchange reactions with following codes:
% [selExc, selUpt] = findExcRxns(model);
% uptakes = model.rxns(selUpt);
% Selecting from the exchange uptake reactions those
% which contain at least 1 carbon in the metabolites included in the reaction:
% subuptakeModel = extractSubNetwork(model, uptakes);
% hiCarbonRxns = findCarbonRxns(subuptakeModel,1);
% Closing the uptake of all the carbon sources
% modelalter = model;
% modelalter = changeRxnBounds(modelalter, hiCarbonRxns, 0, 'l');
```

### Robustness analysis

Robustness analysis is applied to estimate and visualise how changes in the concentration of an environmental parameter (exchange rate) or internal reaction effect on the objective [2]. If we are interested in varying  $v_i$  between two values, i.e.,  $v_{i,min}$  and  $v_{i,max}$ , we can solve l optimisation problems:

```
\max Z_k = c^T v s.t. k = 1, ..., l, Sv = 0, fixing v_j = v_{j,min} + \frac{(k-1)}{(l-1)} * (v_{j,max} - v_{j,min}) constraints v_{i,min} \le v_i \le v_{i,max}, i = 1, ..., n, i \ne j
```

The function robustnessAnalysis() is used for this analysis:

```
% [controlFlux, objFlux] = robustnessAnalysis(model, controlRxn, nPoints,...
% plotResFlag, objRxn,objType)
```

where inputs are a COBRA model, a reaction that has been analysed and optional inputs:

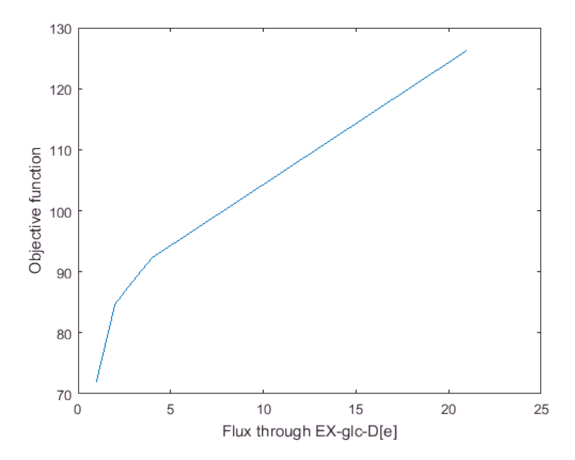
```
% INPUTS
% model
                COBRA model structure
% controlRxn
                Reaction of interest whose value is to be controlled
% OPTIONAL INPUTS
               Number of points to show on plot (Default = 20)
% nPoints
% plotResFlag
                Plot results (Default true)
% objRxn
                Objective reaction to be maximized
                (Default = whatever is defined in model)
               Maximize ('max') or minimize ('min') objective
% objType
                (Default = 'max')
%
% OUTPUTS
                Flux values within the range of the maximum and minimum for
% controlFlux
                a reaction of interest
                Optimal values of objective reaction at each control
% objFlux
                reaction flux value
```

Here, we will investigate how robust the maximal ATP production of the network (i.e., the maximal flux through 'DM atp c ') is with respect to varying glucose uptake rates and fixed oxygen uptake.

```
modelrobust = modelalter;
modelrobust = changeRxnBounds(modelrobust, 'EX_o2[e]', -17, 'b');
AtpRates = zeros(21, 1);
for i = 0:20
    modelrobust = changeRxnBounds(modelrobust, 'EX_glc_D[e]', -i, 'b');
    modelrobust = changeObjective(modelrobust, 'DM_atp_c_');
    FBArobust = optimizeCbModel(modelrobust, 'max');
    AtpRates(i+1) = FBArobust.f;
end
plot (1:21, AtpRates)
```

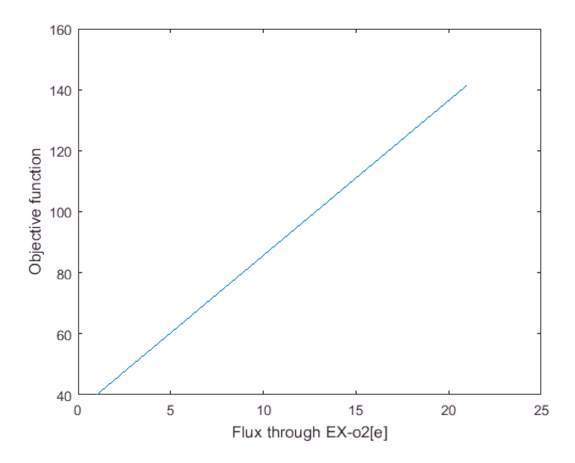
Warning: MATLAB has disabled some advanced graphics rendering features by switching to software OpenGL. For more information, click <a href="here">here</a>.

```
xlabel('Flux through EX-glc-D[e]')
ylabel('Objective function')
```



We can also investigate the robustness of the maximal ATP production when the available glucose amount is fixed, while different levels of oxygen are available.

```
modelrobustoxy = modelalter;
modelrobustoxy = changeRxnBounds(modelrobustoxy, 'EX_glc_D[e]', -20, 'b');
AtpRatesoxy = zeros(21, 1);
for i = 0:20
    modelrobustoxy = changeRxnBounds(modelrobustoxy, 'EX_o2[e]', -i, 'b');
    modelrobustoxy = changeObjective(modelrobustoxy, 'DM_atp_c_');
    FBArobustoxy = optimizeCbModel(modelrobustoxy, 'max');
    AtpRatesoxy(i+1) = FBArobustoxy.f;
end
plot (1:21, AtpRatesoxy)
xlabel('Flux through EX-o2[e]')
ylabel('Objective function')
```



### Double robust analysis

Performs robustness analysis for a pair of reactions of interest and an objective of interest. The double robust analysis is implemented with the function doubleRobustnessAnalysis().

```
% [controlFlux1, controlFlux2, objFlux] = doubleRobustnessAnalysis(model,...
% controlRxn1, controlRxn2, nPoints, plotResFlag, objRxn, objType)
```

The inputs are a COBRA model, two reactions for the analysis and optional inputs:

```
%INPUTS
% model
                COBRA model to analyse,
% controlRxn1
                The first reaction for the analysis,
% controlRxn2
                The second reaction for the analysis;
%OPTIONAL INPUTS
% nPoints
                The number of flux values per dimension (Default = 20)
                Indicates whether the result should be plotted (Default = true)
% plotResFlag
% objRxn
                is objective to be used in the analysis (Default = whatever
%
                is defined in model)
                Direction of the objective (min or max)
 objType
                (Default = 'max')
```

```
modeldrobustoxy = changeRxnBounds(modeldrobustoxy, 'EX_o2[e]', -17, 'l');
[controlFlux1, controlFlux2, objFlux] = doubleRobustnessAnalysis(modeldrobustoxy,...
    'EX glc D[e]', 'EX o2[e]', 10, 1, 'DM atp c ', 'max')
Double robustness analysis in progress ...
                                                           [
                                                  ]2%
controlFlux1 =
  -20.0000
  -17.7225
  -15.4451
  -13.1676
  -10.8902
   -8.6127
   -6.3353
   -4.0578
   -1.7804
    0.4971
controlFlux2 =
  -17.0000
  -15.1111
  -13.2222
  -11.3333
   -9.4444
   -7.5556
   -5.6667
   -3.7778
   -1.8889
    0.0000
objFlux =
  126.2944 116.7061
                                 97.5296
                                           87.9413
                                                      78.3531
                                                                68.7648
                      107.1179
                                                                          59.1765
  121.7395
            112.1512
                      102.5630
                                  92.9747
                                            83.3864
                                                      73.7982
                                                                64.2099
                                                                          54.6216
  117.1846
            107.5963
                       98.0081
                                  88.4198
                                            78.8315
                                                      69.2433
                                                                59.6550
                                                                          50.0667
  112.6297
            103.0414
                       93.4532
                                  83.8649
                                           74.2766
                                                      64.6884
                                                                55.1001
                                                                          45.5118
  108.0748
             98.4865
                       88.8983
                                 79.3100
                                           69.7217
                                                      60.1335
                                                                50.5452
                                                                          40.9569
  103.5199
             93.9316
                       84.3434
                                  74.7551
                                           65.1668
                                                      55.5785
                                                                45.9903
                                                                          36.4020
   98.9650
                                  70.2002
                                                                41.4354
             89.3767
                       79.7884
                                            60.6119
                                                      51.0236
                                                                          31.8471
   94.4101
             84.8218
                       75.2335
                                  65.6453
                                            56.0570
                                                      46.4687
                                                                36.8805
                                                                          27.2922
```

51.5021

41.9138

32.3256

0

22.7373

69.7997

78.7732

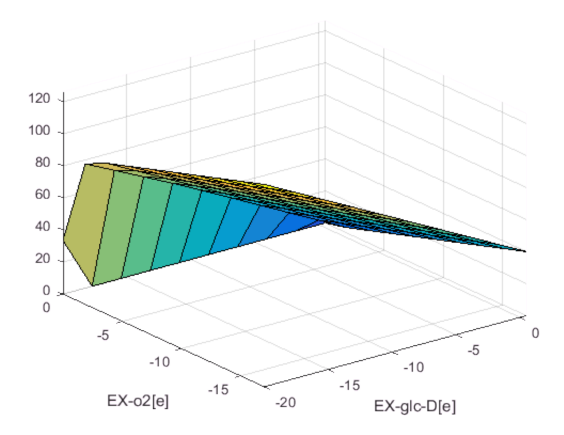
87.7466

33.5029

60.8263

]3%

modeldrobustoxy = changeRxnBounds(modeldrobustoxy, 'EX\_glc\_D[e]', -20, 'l');



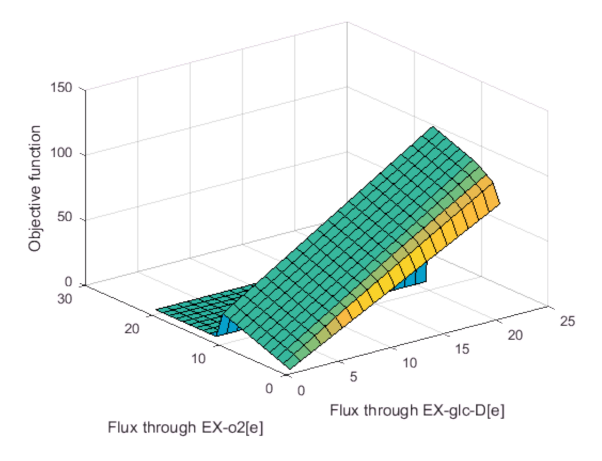
### Phenotypic phase plane analysis (PhPP)

The PhPP is a method for describing in two or three dimensions, how the objective function would change if additional metabolites were given to the model [3].

Essentially PhPP performs a doubleRobustnessAnalysis(), with the difference that shadow prices are retained. The code is as follows-

```
modelphpp = modelalter;
ATPphppRates = zeros(21);
for i = 0:10
    for j = 0:20
        modelphpp = changeRxnBounds(modelphpp, 'EX_glc_D[e]', -i, 'b');
        modelphpp = changeRxnBounds(modelphpp, 'EX_o2[e]', -j, 'b');
        modelphpp = changeObjective(modelphpp, 'DM_atp_c_');
        FBAphpp = optimizeCbModel(modelphpp, 'max');
        ATPphppRates(i+1,j+1) = FBAphpp.f;
    end
end

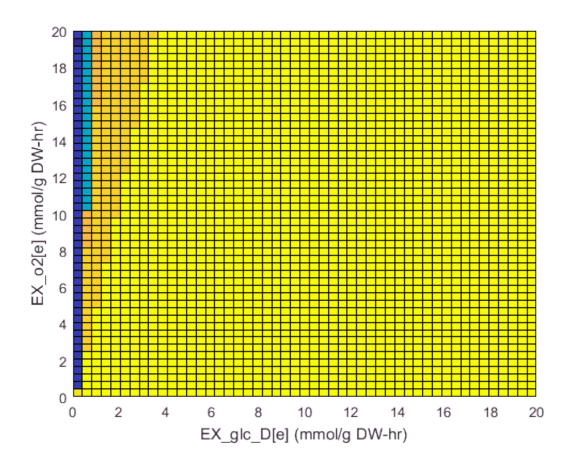
surfl(ATPphppRates) % 3d plot
xlabel('Flux through EX-glc-D[e]')
ylabel('Flux through EX-o2[e]')
zlabel('Objective function')
```

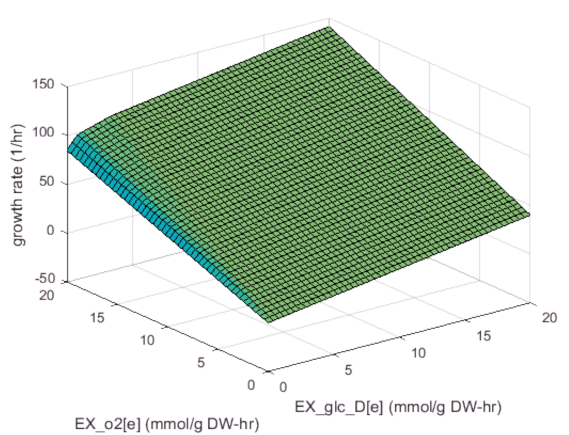


To generate a 2D plot: pcolor (ATPphppRates)

Alternatively, use the function <code>phenotypePhasePlane()</code>. This function also draws the line of optimality, as well as the shadow prices of the metabolites from the two control reactions. In this case, control reactions are  $\texttt{Ex\_glc\_D[e]'}$  and  $\texttt{Ex\_o2[e]'}$ . The line of optimality signifies the state wherein, the objective function is optimal. In this case it is  $\texttt{'DM\_atp\_c\_'}$ .

generating PhPP





## **REFERENCES**

- [1] Noronha A., et al. (2017). ReconMap: an interactive visualization of human metabolism. *Bioinformatics.*, 33 (4): 605-607.
- [2] Edwards, J.S. and and Palsson, B. Ø. (2000). Robustness analysis of the Escherichia coli metabolic network. *Biotechnology Progress*, 16(6):927-39.
- [3] Edwards, J.S., Ramakrishna, R. and and Palsson, B. Ø. (2002). Characterizing the metabolic phenotype: A phenotype phase plane analysis. *Biotechnology and Bioengineering*, 77:27-36.