Browse Networks in the Matlab Command Window Using surfNet

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Reviewer(s):

INTRODUCTION

In this tutorial, we will demonstrate how to browse a COBRA model in verbal format in the Matlab command window through an initial call and interactive mouse clicking.

MATERIALS

EQUIPMENT SETUP

cplex direct full dqqMinos full

gurobi full
ibm_cplex full
matlab full

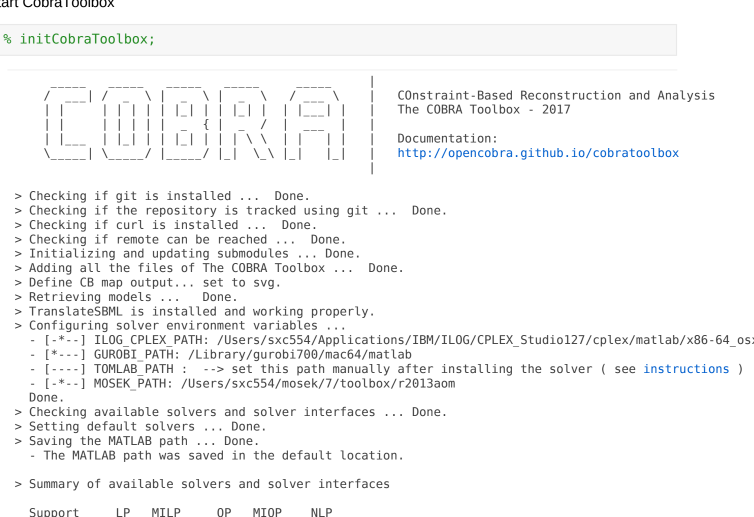
full

full

glpk

matlab

Start CobraToolbox



1

1 1 0 0

1 0

```
full
mosek
                                   1
                                                  1
                                   1
               full
                                                  1
pdco
                                   1
quadMinos
              full
                                                                  1
tomlab_cplex full
                                   0
                                           0
                                                  Θ
                                                          0
gpng
             experimental
                                                  1
tomlab_snopt experimental
gurobi_mex legacy
lindo_old legacy
                                   0
                                                  0
                                                          0
                                   0
lindo legacy legacy
                                   0
                                   1
lp solve
              legacy
                                   0
                                           0
                                                   0
                                                          0
                                                                  0
opti
               legacy
Total
                                   8
                                           3
                                                          1
                                                                  2
+ Legend: - = not applicable, 0 = solver not compatible or not installed, 1 = solver installed.
> You can solve LP problems using: 'dqqMinos' - 'glpk' - 'gurobi' - 'matlab' - 'mosek' - 'pdco' - 'quad' > You can solve MILP problems using: 'glpk' - 'gurobi' - 'mosek'
> You can solve QP problems using: 'gurobi' - 'mosek' - 'pdco' - 'qpng'
> You can solve MIQP problems using: 'gurobi'
> You can solve NLP problems using: 'matlab' - 'quadMinos'
> Checking for available updates ...
--> You cannot update your fork using updateCobraToolbox(). [blda0e @ add-tutorial-browseNetwork].
    Please use the MATLAB.devTools (https://github.com/opencobra/MATLAB.devTools).
```

PROCEDURE

Load the E. coli iJO1366 model as an example model.

```
load('iJ01366.mat')
```

Browse a network

Browse the network by starting from an initial metabolite, e.g., D-glucose in the extracellular compartment.

```
surfNet(iJ01366, 'glc__D_e')

Met #1195    glc__D_e, D-Glucose, C6H1206
Consuming reactions:
    #164    EX_glc__D_e, Bd: -10 / 1000, D-Glucose exchange
    glc__D_e <=>
    #1355    GLCtex_copy1, Bd: -1000 / 1000, Glucose transport via diffusion (extracellular to periplasm)
    glc__D_e <=>    glc__D_p
    #1356    GLCtex_copy2, Bd: 0 / 1000, Glucose transport via diffusion (extracellular to periplasm)
    glc__D_e ->    glc__D_p
    Producing reactions: none

Show previous steps...
```

All reactions producing or consuming 'glc__D_e' will have their reaction indices (#xxx), ids (.rxns), bounds (.lb/.ub), names (.rxnNames) and formulae printed on the command window. All reactions and the participating metabolites are hyperlinked. For example, **click** on the reaction 'GLCtex_copy1'. (This is equivalent to run the following command.)

```
% called by clicking 'GLCtex_copy1' surfNet([], 'GLCtex_copy1', 0, 'none', 0, 1, [], 0)
```

Details for the metabolites will appear, e.g., indeices, ids, stoichiometric coefficients, names and chemical formulae. By iteratively clicking on the reactions and metabolites that you are interested in, you can browse through the metabolic network.

Now, say you have gone through a series of metabolites and reactions (glc__D_e, GLCtex_copy1, glc__D_p, GLCptspp, g6p_c):

Click glc__D_p:

```
% called by clicking 'glc__D_p' surfNet([], 'glc__D_p', 0, 'none', 0, 1, [], 0)
```

```
Met #1587 glc D p, D-Glucose, C6H12O6
Consuming reactions:
 #1336 GLCDpp, Bd: 0 / 1000, Glucose dehydrogenase (ubiquinone-8 as acceptor) (periplasm)
  q8_c + glc_D_p + h2o_p -> q8h2_c + glcn_p + h_p
 #1352 GLCabcpp, Bd: 0 / 1000, D-glucose transport via ABC system (periplasm)
  atp c + h2o c + glc D p \rightarrow adp c + glc D c + h c + pi c
 #1353 GLCptspp, Bd: 0 / 1000, D-glucose transport via PEP:Pyr PTS (periplasm)
  pep_c + glc__D_p -> g6p_c + pyr_c
  #1354 GLCt2pp, Bd: 0 / 1000, D-glucose transport in via proton symport (periplasm)
  glc _Dp + h_p -> glc_Dc + h_c
Producing reactions:
  #1252 G1PPpp, Bd: 0 / 1000, Glucose-1-phosphatase
  g1p_p + h2o_p \rightarrow glc_D_p + pi_p
 #1355 GLCtex copy1, Bd: -1000 / 1000, Glucose transport via diffusion (extracellular to periplasm)
  glc D_e <=> glc__D_p
 #1356 GLCtex copy2, Bd: 0 / 1000, Glucose transport via diffusion (extracellular to periplasm)
  glc De->glc Dp
 #1607 LACZpp, Bd: 0 / 1000, B-galactosidase
 h2o p + lcts p \rightarrow qal p + qlc D p
 #2463 TREHpp, Bd: 0 / 1000, Alpha, alpha-trehalase (periplasm)
 h2o p + tre p \rightarrow 2 qlc D p
Show previous steps...
```

Click GLCptspp:

#853

pyr_c 1

```
% called by clicking 'GLCptspp'
surfNet([], 'GLCptspp', 0, 'none', 0, 1, [], 0)
Rxn #1353 GLCptspp, Bd: 0 / 1000, D-glucose transport via PEP:Pyr PTS (periplasm)
pep c + qlc D p \rightarrow q6p c + pyr c
  id
                   Stoich
                              metNames, metFormulas
Reactant:
            pep_c -1
 #784
                              Phosphoenolpyruvate, C3H2O6P
 #1587
         glc Dp -1
                              D-Glucose, C6H12O6
Product:
 #508
            g6p c 1
                              D-Glucose 6-phosphate, C6H1109P
```

Pyruvate, C3H3O3

Show previous steps...

Click g6p_c:

```
% called by clicking 'g6p_c' surfNet([], 'g6p_c', 0, 'none', 0, 1, [], 0)
```

```
Met #508 g6p c, D-Glucose 6-phosphate, C6H1109P
Consuming reactions:
 #1283 G6PDH2r, Bd: -1000 / 1000, Glucose 6-phosphate dehydrogenase
 g6p_c + nadp_c <=> 6pgl_c + h_c + nadph_c
 #1284 GGPP, Bd: 0 / 1000, Glucose-6-phosphate phosphatase
 g6p_c + h2o_c -> glc_D_c + pi_c
 #2077 PGI, Bd: -1000 / 1000, Glucose-6-phosphate isomerase
 g6p c <=> f6p c
 #2461 TRE6PS, Bd: 0 / 1000, Alpha, alpha-trehalose-phosphate synthase (UDP-forming)
 g6p c + udpg c -> h c + tre6p c + udp c
Producing reactions:
 #477 AB6PGH, Bd: 0 / 1000, Arbutin 6-phosphate glucohydrolase
 arbt6p c + h2o c -> q6p c + hqn c
 #1214 FFSD, Bd: 0 / 1000, Beta-fructofuranosidase
 h2o c + suc6p c \rightarrow fru c + g6p c
 #1231 FRULYSDG, Bd: -1000 / 1000, Fructoselysine phosphate deglycase
 frulysp c + h2o c <=> g6p c + lys L c
 #1285 GGPt6 2pp, Bd: 0 / 1000, Glucose-6-phosphate transport via phosphate antiport (periplasm)
 2 pi c + g6p p -> g6p c + 2 pi p
 #1353 GLCptspp, Bd: 0 / 1000, D-glucose transport via PEP:Pyr PTS (periplasm)
 pep_c + glc__D_p -> g6p_c + pyr_c
 #1500 HEX1, Bd: 0 / 1000, Hexokinase (D-glucose:ATP)
 atp_c + glc_
              D c \rightarrow adp c + g6p c + h c
 #2082 PGMT, Bd: -1000 / 1000, Phosphoglucomutase
 glp c <=> g6p c
 #2459 TRE6PH, Bd: 0 / 1000, Trehalose-6-phosphate hydrolase
 h2o_c + tre6p_c -> g6p_c + glc__D_c
Show previous steps...
```

In each click, there is also a button '**Show previous steps...**' at the bottom. Clicking on it will show the metabolites and reactions that you have visited in order. This is equivalent to calling:

```
% called by clicking 'Show previous steps...'
surfNet([], [], 0, 'none', 0, 1, [], 0, struct('showPrev', true))
```

```
glc D e>>GLCtex copy1>>glc D p>>GLCptspp>>g6p c>>
```

You can go back to any of the intermediate metabolites/reactions by clicking the hyperlinked mets/rxns shown.

Call options

Shown below are various call options for including flux vectors and customizing display. All call options are preserved during the interactive browsing by mouse clicking.

Show objective reactions

Omit the 'metrxn' (2nd) argument to print objective reactions:

```
surfNet(iJ01366)
```

```
Rxn #8 BIOMASS Ec iJ01366 core 53p95M, Bd: 0 / 1000, E. coli biomass objective function (iJ01366) - col
0.000223 10fthf c + 2.6e-05 2fe2s c + 0.000223 2ohph c + 0.00026 4fe4s c + 0.51369 ala L c + 0.000223
     0.29579 arg L c + 0.24105 asn L c + 0.24105 asp L c + 54.1248 atp c + 0.000122 bmocogdp c + 2e-06 l 0.005205 ca2 c + 0.005205 cl c + 0.000576 coa c + 2.5e-05 cobalt2 c + 0.13351 ctp c + 0.000709 cu2 c -
                                         L c + 0.026166 datp c + 0.027017 dctp c + 0.027017 dgtp c + 0.026166 dttp c + 0.000223 fa
     0.09158 cys
     0.006715 \ \text{fe2\_c} + 0.007808 \ \text{fe3\_c} + 0.26316 \ \text{gln\_L\_c} + 0.26316 \ \text{glu\_L\_c} + 0.61264 \ \text{gly\_c} + 0.2151 \ \text{gtp\_c} + 0.2151 \ \text{gt
     48.6015 h2o_c + 0.094738 his_L_c + 0.29053 ile_L_c + 0.19519 k_c + 0.45053 leu_L_c + 0.34316 lys_I
     0.15369 \ \text{met} \ \_L\_c \ + \ 0.008675 \ \text{mg} \ \_c \ + \ 0.000223 \ \text{mlthf} \ \_c \ + \ 0.000691 \ \text{mn} \ \_c \ + \ 7e-06 \ \text{mobd} \ \_c \ + \ 0.001831 \ \text{nad} \ \_c
     0.000447 \text{ nadp_c} + 0.013013 \text{ nh4_c} + 0.000323 \text{ ni2_c} + 0.017868 \text{ pe160_c} + 0.054154 \text{ pe161_c} + 0.18527 \text{ phe}
     0.000223 pheme_c + 0.22106 pro_L_c + 0.000223 pydx5p_c + 0.000223 ribflv_c + 0.21579 ser_L_c + 0.000
     + 0.004338 \text{ so4} \text{ c} + 0.000223 \text{ thf} \text{ c} + 0.000223 \text{ thmpp} \text{ c} + 0.25369 \text{ thr} \text{ L} \text{ c} + 0.056843 \text{ trp} \text{ L} \text{ c} + 0.1379 \text{ thr}
     5.5e-05 \ udcpdp_c + 0.1441 \ utp_c + 0.42316 \ val__L_c + 0.000341 \ zn2_c + 0.019456 \ kdo2lipid4_e + 0.000341 \ zn2_c + 0.019456 \ kdo2lipid4_e + 0.000341 \ zn2_c + 0.019456 \ kdo2lipid4_e + 0.000341 \ zn2_c + 0.0003456 \ kdo2lipid4_e + 0.000341 \ zn2_c + 0.000341 \ zn2_c + 0.0003456 \ kdo2lipid4_e + 0.000341 \ zn2_c + 0.000341 \ zn2_c + 0.0003456 \ kdo2lipid4_e + 0.000341 \ zn2_c + 0.000341 \ zn2_c + 0.0003456 \ kdo2lipid4_e + 0.000341 \ zn2_c + 0.0003456 \ kdo2lipid4_e + 0.000341 \ zn2_c + 0.0003456 \ kdo2lipid4_e + 0.000346 \ k
     0.013894 \text{ murein5px4p p} + 0.045946 \text{ pe160 p} + 0.02106 \text{ pe161 p} -> 53.95 \text{ adp c} + 53.95 \text{ h} \text{ c} + 53.9457 \text{ pi c}
     0.7739 ppi c
     id
                         Met
                                                                     Stoich
                                                                                                     metNames, metFormulas
Reactant:
                                        10fthf c -0.000223 10-Formyltetrahydrofolate, C20H21N707
  #1
  #69
                                           2fe2s c
                                                                   -0.000026 [2Fe-2S] iron-sulfur cluster, S2Fe2
                                                                   -0.000223 2-Octaprenyl-6-hydroxyphenol, C46H7002
  #82
                                           2ohph c
                                                                                                     [4Fe-4S] iron-sulfur cluster, S4Fe4
  #167
                                           4fe4s_c
                                                                   -0.00026
  #255
                                        ala L c
                                                                   -0.513689 L-Alanine, C3H7N02
  #265
                                              amet c
                                                                   -0.000223 S-Adenosyl-L-methionine, C15H23N605S
                                                                   -0.295792 L-Arginine, C6H15N402
  #294
                                        arg L c
                                        asn__L_c
asp__L_c
  #298
                                                                    -0.241055
                                                                                                   L-Asparagine, C4H8N2O3
                                                                    -0.241055 L-Aspartate, C4H6N04
  #302
  #307
                                                                    -54.124831 ATP, C10H12N5013P3
                                                 atp c
  #314
                                                                    -0.000122 Bis-molybdopterin quanine dinucleotide, C40H44N20027P4S4Mo
                                  bmocogdp c
  #317
                                                 btn c
                                                                   -0.000002 Biotin, C10H15N2O3S
                                                                   -0.005205 Calcium, Ca
  #326
                                                 ca2 c
  #355
                                                   cl_c -0.005205 Chloride, Cl
  #358
                                                 coa c -0.000576 Coenzyme A, C21H32N7016P3S
  #359
                                     cobalt2 c -0.000025 Co2+, Co
  #377
                                                 ctp c -0.133508 CTP, C9H12N3014P3
                                                 cu2_c -0.000709 Cu2+, Cu
  #379
                                        cys__L_c -0.09158
                                                                                                     L-Cysteine, C3H7N02S
  #383
  #392
                                              datp c -0.026166 DATP, C10H12N5012P3
  #401
                                              dctp_c -0.027017 DCTP, C9H12N3013P3
  #412
                                              dgtp_c -0.027017 DGTP, C10H12N5013P3
                                              dttp c -0.026166 DTTP, C10H13N2014P3
  #451
                                                fad_c -0.000223 Flavin adenine dinucleotide oxidized, C27H31N9015P2
  #468
  #474
                                                fe2 c -0.006715 Fe2+, Fe
                                                 fe3 c -0.007808 Fe3+, Fe
  #475
```

Call with a list of mets/rxns

The 'metrxn' arguement can be a string of id for a metabolite or reaction. It can also be a cell array of ids, e.g.,

```
Met #1587 glc_D_p, D-Glucose, C6H1206
Consuming reactions:
    #1336 GLCDpp, Bd: 0 / 1000, Glucose dehydrogenase (ubiquinone-8 as acceptor) (periplasm)
    q8_c + glc_D_p + h2o_p -> q8h2_c + glcn_p + h_p
    #1352 GLCabcpp, Bd: 0 / 1000, D-glucose transport via ABC system (periplasm)
    atp_c + h2o_c + glc_D_p -> adp_c + glc_D_c + h_c + pi_c
    #1353 GLCptspp, Bd: 0 / 1000, D-glucose transport via PEP:Pyr PTS (periplasm)
    pep_c + glc_D_p -> g6p_c + pyr_c
    #1354 GLCt2pp, Bd: 0 / 1000, D-glucose transport in via proton symport (periplasm)
    glc_D_p + h_p -> glc_D_c + h_c
Producing reactions:
    #1252 G1PPpp, Bd: 0 / 1000, Glucose-1-phosphatase
    glp_p + h2o_p -> glc_D_p + pi_p
```

```
#1355 GLCtex_copy1, Bd: -1000 / 1000, Glucose transport via diffusion (extracellular to periplasm)
 glc De <=> glc Dp
 #1356 GLCtex copy2, Bd: 0 / 1000, Glucose transport via diffusion (extracellular to periplasm)
 glc De->glc Dp
 #1607 LACZpp, Bd: 0 / 1000, B-galactosidase
 h2o p + lcts p -> gal p + glc D p
 #2463 TREHpp, Bd: 0 / 1000, Alpha, alpha-trehalase (periplasm)
 h2o p + tre p \rightarrow 2 qlc D p
Rxn #1353 GLCptspp, Bd: 0 / 1000, D-glucose transport via PEP:Pyr PTS (periplasm)
pep_c + glc__D_p -> g6p_c + pyr_c
                  Stoich
                             metNames, metFormulas
 id
        Met
Reactant:
#784
                              Phosphoenolpyruvate, C3H2O6P
           pep c -1
#1587
        glc Dp -1
                             D-Glucose, C6H12O6
Product:
#508
           g6p c 1
                              D-Glucose 6-phosphate, C6H1109P
#853
           pyr c 1
                             Pyruvate, C3H3O3
Met #508 g6p c, D-Glucose 6-phosphate, C6H1109P
Consuming reactions:
 #1283 G6PDH2r, Bd: -1000 / 1000, Glucose 6-phosphate dehydrogenase
 g6p_c + nadp_c <=> 6pgl_c + h_c + nadph_c
 #1284 GGPP, Bd: 0 / 1000, Glucose-6-phosphate phosphatase
 g6p_c + h2o_c \rightarrow glc_D_c + pi_c
 #2077 PGI, Bd: -1000 / 1000, Glucose-6-phosphate isomerase
 g6p c <=> f6p c
 #2461 TRE6PS, Bd: 0 / 1000, Alpha,alpha-trehalose-phosphate synthase (UDP-forming)
 g6p c + udpg c \rightarrow h c + tre6p c + udp c
Producing reactions:
 #477 AB6PGH, Bd: 0 / 1000, Arbutin 6-phosphate glucohydrolase
 arbt6p c + h2o c -> g6p c + hqn c
 #1214 FFSD, Bd: 0 / 1000, Beta-fructofuranosidase
 h2o_c + suc6p_c -> fru_c + g6p_c
 #1231 FRULYSDG, Bd: -1000 / 1000, Fructoselysine phosphate deglycase
 frulysp c + h2o c <=> g6p c + lys L c
 #1285 GGPt6 2pp, Bd: 0 / 1000, Glucose-6-phosphate transport via phosphate antiport (periplasm)
 2 pi c + g6p p -> g6p c + 2 pi p
 #1353 GLCptspp, Bd: 0 / 1000, D-glucose transport via PEP:Pyr PTS (periplasm)
 pep_c + glc__D_p -> g6p_c + pyr_c
 #1500 HEX1, Bd: 0 / 1000, Hexokinase (D-glucose:ATP)
 atp c + glc D c -> adp c + g6p c + h c
```

Show metabolite names in reaction formulae

Some models may use generic ids for mets/rxns. In this case, call surfNet() with the 'metNameFlag' (3rd) argument turned on to show the names for metabolites (.metNames) in the reaction formulae, e.g.,

```
Met #484 fgam_c, N2-Formyl-N1-(5-phospho-D-ribosyl)glycinamide, C8H13N2O9P
Consuming reactions:
    #2207 PRFGS, Bd: 0 / 1000, Phosphoribosylformylglycinamidine synthase
    ATP + N2-Formyl-N1-(5-phospho-D-ribosyl)glycinamide + L-Glutamine + H2O -> ADP + 2-(Formamido)-N1-(5-phospho-D-ribosyl)glycinamide + L-Glutamine + H2O -> ADP + 2-(Formamido)-N1-(5-phospho-Indicated + H2O -> ADP + Indicated + H2O -> ADP + Indicated + Indicated
```

Hide reaction detials

Turn off the 'showMets' (6th) arguement to suppress details for reactions, e.g.,

```
surfNet(iJ01366, iJ01366.rxns(1001:1010), [], [], [], 0)
```

```
Rxn #1001 DHPPDA2, Bd: 0 / 1000, Diaminohydroxyphosphoribosylaminopryrimidine deaminase (25drapp)
25drapp c + h c + h2o c -> 5apru c + nh4 c
Rxn #1002 DHPS2, Bd: 0 / 1000, Dihydropteroate synthase
4abz c + 6hmhptpp c -> dhpt c + ppi c
Rxn #1003 DHPTDCs2, Bd: 0 / 1000, 4,5-dihydroxy-2,3-pentanedione cyclization (spontaneous)
dhptd c -> mdhdhf c
Rxn #1004 DHPTDNR, Bd: 0 / 0, Dihydropteridine reductase
dhptdn c + 3 h c + nadph c -> nadp c + thptdn c
Rxn #1005 DHPTDNRN, Bd: 0 / 0, Dihydropteridine reductase (NADH)
dhptdn c + 3 h c + nadh c -> nad c + thptdn c
Rxn #1006 DHPTPE, Bd: -1000 / 1000, Dihydroneopterin triphosphate 2'-epimerase
ahdt c <=> dhmptp c
Rxn #1007 DHQS, Bd: 0 / 1000, 3-dehydroquinate synthase
2dda7p_c -> 3dhq_c + pi_c
Rxn #1008 DHQTi, Bd: 0 / 1000, 3-dehydroquinate dehydratase, irreversible
3dhq c \rightarrow 3dhsk c + h2o c
Rxn #1009 DIMPtex, Bd: -1000 / 1000, DIMP transport via diffusion (extracellular to periplasm)
dimp e <=> dimp_p
Rxn #1010 DINSt2pp, Bd: 0 / 1000, Deoxyinosine transport in via proton symport (periplasm)
din p + h p \rightarrow din c + h c
Show previous steps...
```

Look at one or more flux distributions

First, get a flux distribution by optimizing the biomass production of the model (the standard flux balance analysis ¹). Then call surfNet with the flux distribution (4th argument) to look at how the flux through pyruvate is distributed:

```
s = optimizeCbModel(iJ01366, 'max', 'one');
surfNet(iJ01366, 'pyr_c', [], s.x)
```

```
Met #853 pyr_c, Pyruvate, C3H3O3
Consuming reactions with non-zero fluxes:

#511 ACHBS (0.28541), Bd: 0 / 1000, 2-aceto-2-hydroxybutanoate synthase
20but_c + h_c + pyr_c -> 2ahbut_c + co2_c

#513 ACLS (0.85886), Bd: 0 / 1000, Acetolactate synthase
h_c + 2 pyr_c -> alac_S_c + co2_c

#618 ALATA_L (-0.57111), Bd: -1000 / 1000, L-alanine transaminase
akg_c + ala_L_c <=> glu_L_c + pyr_c

#987 DHDPS (0.36441), Bd: 0 / 1000, Dihydrodipicolinate synthase
aspsa_c + pyr_c -> 23dhdp_c + h_c + 2 h2o_c

#1053 DXPS (0.00279), Bd: 0 / 1000, 1-deoxy-D-xylulose 5-phosphate synthase
g3p_c + h_c + pyr_c -> co2_c + dxyl5p_c

#2047 PDH (7.96919), Bd: 0 / 1000, Pyruvate dehydrogenase
```

```
coa_c + nad_c + pyr_c -> accoa_c + co2_c + nadh_c
 #2171 POR5 (0.10684), Bd: -1000 / 1000, Pyruvate synthase
 coa c + 2 flxso c + pyr c <=> accoa c + co2 c + 2 flxr c + h c
 #2466 TRPAS2 (-0.05584), Bd: -1000 / 1000, Tryptophanase (L-tryptophan)
 h2o c + trp L c <=> indole c + nh4 c + pyr c
Producing reactions with non-zero fluxes:
 #554 ADCL (0.00066), Bd: 0 / 1000, 4-aminobenzoate synthase
 4adcho c \rightarrow 4abz c + h c + pyr c
 #666 ANS (0.05584), Bd: 0 / 1000, Anthranilate synthase
 chor c + gln L c \rightarrow anth c + glu L c + h c + pyr c
 #813 CHRPL (0.00022), Bd: 0 / 1000, Chorismate pyruvate lyase
 chor c -> 4hbz c + pyr c
 #908 CYSTL (0.1512), Bd: 0 / 1000, Cystathionine b-lyase
 cyst L c + h2o c -> hcys L c + nh4 c + pyr c
 #978 DHAPT (0.86538), Bd: 0 / 1000, Dihydroxyacetone phosphotransferase
 dha c + pep c -> dhap c + pyr c
 #1353 GLCptspp (10), Bd: 0 / 1000, D-qlucose transport via PEP:Pyr PTS (periplasm)
 pep c + glc D p -> g6p c + pyr c
Show previous steps...
```

All reactions involving pyruvate with non-zero fluxes are printed. The flux values are in the parentheses following the reaction ids. Note that reactions stated as consuming or producing the metabolite have taken the directions of the fluxes into account. Therefore, supplying a different flux distribution or not supplying may give different display. By default, only reactions with non-zero fluxes are printed if a flux distribution is supplied. Turn the 'nonzerofluxflag' (5th) argument off to show all reactions:

```
surfNet(iJ01366, 'pyr_c', [], s.x, 0)
```

```
Met #853 pyr c, Pyruvate, C3H3O3
Consuming reactions:
 #511 ACHBS (0.28541), Bd: 0 / 1000, 2-aceto-2-hydroxybutanoate synthase
  2obut_c + h_c + pyr_c -> 2ahbut c + co2 c
 #513 ACLS (0.85886), Bd: 0 / 1000, Acetolactate synthase
 h c + 2 pyr c \rightarrow alac S c + co2 c
  #618 ALATA L (-0.57111), Bd: -1000 / 1000, L-alanine transaminase
              _L_c <=> glu__L_c + pyr_c
  akg c + ala
  #987 DHDPS (0.36441), Bd: 0 / 1000, Dihydrodipicolinate synthase
  aspsa_c + pyr_c \rightarrow 23dhdp_c + h_c + 2 h2o_c
 #1053 DXPS (0.00279), Bd: 0 / 1000, 1-deoxy-D-xylulose 5-phosphate synthase
  g3p c + h c + pyr c \rightarrow co2 c + dxyl5p c
 #2047 PDH (7.96919), Bd: 0 / 1000, Pyruvate dehydrogenase
  coa c + nad c + pyr c -> accoa c + co2 c + nadh c
 #2067 PFL (0), Bd: 0 / 1000, Pyruvate formate lyase
  coa c + pyr c -> accoa c + for c
 #2171 POR5 (0.10684), Bd: -1000 / 1000, Pyruvate synthase
  coa c + 2 flxso c + pyr c \iff accoa c + co2 c + 2 flxr c + h c
 #2172 POX (0), Bd: 0 / 1000, Pyruvate oxidase
 h2o_c + pyr_c + q8_c -> ac_c + co2_c + q8h2_c
  #2198 PPS (0), Bd: 0 / 1000, Phosphoenolpyruvate synthase
 atp_c + h2o_c + pyr_c \rightarrow amp_c + 2 h_c + pep_c + pi_c
 #2466 TRPAS2 (-0.05584), Bd: -1000 / 1000, Tryptophanase (L-tryptophan)
  h2o c + trp L c <=> indole c + nh4 c + pyr c
Producing reactions:
  #507 ACGAptspp (0), Bd: 0 / 1000, N-Acetyl-D-glucosamine transport via PEP:Pyr PTS (periplasm)
  pep c + acgam p -> acgam6p c + pyr c
 #516 ACMANAptspp (0), Bd: 0 / 1000, N-acetyl-D-mannosamine transport via PTS (periplasm)
  pep c + acmana p -> acmanap c + pyr c
 #518 ACMUMptspp (0), Bd: 0 / 1000, N-acetylmuramate transport via PEP:Pyr PTS (periplasm)
  pep_c + acmum_p -> acmum6p_c + pyr_c
 #522 ACNML (0), Bd: 0 / 1000, N-Acetylneuraminate lyase
  acnam c -> acmana_c + pyr_c
 #554 ADCL (0.00066), Bd: 0 / 1000, 4-aminobenzoate synthase
  4adcho c -> 4abz_c + h_c + pyr_c
 #617 ALATA_D2 (0), Bd: 0 / 1000, D-alanine transaminase
```

```
ala__D_c + pydx5p_c -> pyam5p_c + pyr_c
#619 ALATA_L2 (0), Bd: 0 / 1000, Alanine transaminase
ala__L_c + pydx5p_c -> pyam5p_c + pyr_c
#666 ANS (0.05584), Bd: 0 / 1000, Anthranilate synthase
chor_c + gln__L_c -> anth_c + glu__L_c + h_c + pyr_c
#698 ARBTptspp (0), Bd: 0 / 1000, Arbutin transport via PEP:Pyr PTS (periplasm)
pep_c + arbt_p -> arbt6p_c + pyr_c
#716 ASCBptspp (0), Bd: 0 / 1000, L-ascorbate transport via PEP:Pyr PTS (periplasm)
```

You can also compare multiple flux distributions by supplying them in a matrix format, each column being a flux distribution. For example, get another flux distribution maximizing the biomass production using D-fructose instead of glucose as substrate. Then call surfNet to look at reactions with different fluxes.

Original uptake rates:

```
printUptakeBound(iJ01366);
```

```
EX ca2 e -1000
EX cbl1 e -0.01
EX cl e -1000
EX_co2_e -1000
EX cobalt2 e -1000
EX_cu2_e -1000
EX fe2 e -1000
EX fe3 e -1000
EX_glc__D_e -10
EX_h_e -1000
EX h2o e -1000
EX k e -1000
EX mq2 e -1000
EX mn2 e -1000
EX mobd e -1000
EX na1 e -1000
EX nh4 e -1000
EX ni2 e -1000
EX o2 e -1000
EX pi e -1000
EX sel e -1000
EX slnt e -1000
EX so4 e -1000
EX tungs e -1000
EX zn2 e -1000
```

Use fructose instead of glucose as substrate:

```
iJ01366 = changeRxnBounds(iJ01366, {'EX_glc__D_e'; 'EX_fru_e'},...
[0; -10], {'L'; 'L'});
printUptakeBound(iJ01366);
```

```
EX_ca2_e -1000

EX_cbl1_e -0.01

EX_cl_e -1000

EX_co2_e -1000

EX_cobalt2_e -1000

EX_cu2_e -1000

EX_fe2_e -1000

EX_fe3_e -1000

EX_fru_e -10

EX_h_e -1000

EX_h_e -1000
```

```
EX_k_e -1000
EX_mg2_e -1000
EX_mn2_e -1000
EX_mobd_e -1000
EX_nal_e -1000
EX_nh4_e -1000
EX_ni2_e -1000
EX_o2_e -1000
EX_pi_e -1000
EX_sel_e -1000
EX_sel_e -1000
EX_sel_e -1000
EX_sunt_e -1000
EX_so4_e -1000
EX_tungs_e -1000
EX_zn2_e -1000
```

Run FBA again to get a flux distribution using fructose as substrate. Then look at reactions with different fluxes in the glucose and fructose cases using surfNet.

```
sFru = optimizeCbModel(iJ01366, 'max', 'one'); % FBA
fluxMatrix = [s.x, sFru.x]; % put two flux vectors in a matrix
% reactions with different fluxes
rxnDiff = abs(fluxMatrix(:, 1) - fluxMatrix(:, 2)) > 1e-6;
surfNet(iJ01366, iJ01366.rxns(rxnDiff), [], fluxMatrix, [], 0)
Rxn #139 EX_fru_e (0, -10), Bd: -10 / 1000, D-Fructose exchange
fru e <=>
Rxn #164 EX glc D e (-10, 0), Bd: 0 / 1000, D-Glucose exchange
glc De->
Rxn #623 ALAt2pp copy2 (-0.00511, 0), Bd: -1000 / 1000, L-alanine transport in via proton symport (per
ala Lp + hp \ll ala Lc + hc
Rxn #624 ALAt4pp (0.00511, 0), Bd: 0 / 1000, L-alanine transport in via sodium symport (periplasm)
ala_L_p + na1_p -> ala_L_c + na1_c
Rxn #1230 FRUK (0, 5.75203), Bd: 0 / 1000, Fructose-1-phosphate kinase
atp c + flp c \rightarrow adp c + fdp c + h c
Rxn #1238 FRUpts2pp (0, 4.24797), Bd: 0 / 1000, Fructose transport via PEP:Pyr PTS (f6p generating) (pe
pep_c + fru_p -> f6p_c + pyr_c
Rxn #1239 FRUptspp (0, 5.75203), Bd: 0 / 1000, D-fructose transport via PEP:Pyr PTS (periplasm)
pep_c + fru_p -> f1p_c + pyr_c
Rxn #1240 FRUtex (-0, 10), Bd: -1000 / 1000, D-fructose transport via diffusion (extracellular to peri
fru_e <=> fru_p
Rxn #1353 GLCptspp (10, 0), Bd: 0 / 1000, D-glucose transport via PEP:Pyr PTS (periplasm)
pep_c + glc__D_p -> g6p_c + pyr_c
Rxn #1356 GLCtex copy2 (10, 0), Bd: 0 / 1000, Glucose transport via diffusion (extracellular to peripla
glc De->glc Dp
Rxn #1377 GLUt2rpp (0, -0.00511), Bd: -1000 / 1000, L-glutamate transport via proton symport, reversib
glu Lp + hp <=> glu Lc + hc
Rxn #1378 GLUt4pp (0, 0.00511), Bd: 0 / 1000, Na+/qlutamate symport (periplasm)
glu L p + na1 p \rightarrow glu L c + na1 c
Rxn #1758 MDH (4.82506, 4.82528), Bd: -1000 / 1000, Malate dehydrogenase
mal L c + nad c \iff h c + nadh c + oaa c
```

Rxn #1837 MOX (0.0016, 0.00138), Bd: -1000 / 1000, Malate oxidase

 $mal L_c + o2_c \iff h2o2_c + oaa_c$

```
Rxn #2048 PDX5P02 (0.00022, 0), Bd: 0 / 1000, Pyridoxine 5'-phosphate oxidase (anaerboic nad_c + pdx5p_c -> h_c + nadh_c + pydx5p_c

Rxn #2049 PDX5P0i (0, 0.00022), Bd: 0 / 1000, Pyridoxine 5'-phosphate oxidase o2_c + pdx5p_c -> h2o2_c + pydx5p_c

Rxn #2064 PFK (5.75203, 0), Bd: 0 / 1000, Phosphofructokinase atp_c + f6p_c -> adp_c + fdp_c + h_c

Rxn #2077 PGI (5.91807, -4.08193), Bd: -1000 / 1000, Glucose-6-phosphate isomerase g6p_c <=> f6p_c
Show previous steps...
```

Customize model data to be displayed

Customize the fields for metabolites and reactions to be printed by supplying the 'field2print' (7th) argument. It is defaulted to be:

```
{{ 'metNames', 'metFormulas'}, { 'rxnNames', 'lb', 'ub'}}
```

The first cell contains the metabolite-related fields to be printed and the second cell contains the reaction-related fields to be printed. It can also be inputted as a single cell array of strings, as long as from the size (equal to #mets or #rxns) or from the name of the field (starting with 'met' or 'rxn'), the fields are recognizable to be met- or rxn-related. For example, show the grRules for rxns but omit the bounds and show the constraint sense (csense) associated with each metabolite. Note the difference from the original call:

```
surfNet(iJ01366, 'fdp c', [], [], [], ...
Met #473 fdp c, D-Fructose 1,6-bisphosphate, C6H10012P2, csense: E
Consuming reactions with non-zero fluxes :
  #1151 FBA (5.75203, 5.75203), Fructose-bisphosphate aldolase, grRules: b2097 or b1773 or b2925
  fdp c \ll  dhap c + q3p c
Producing reactions with non-zero fluxes:
  #1230 FRUK (0, 5.75203), Fructose-1-phosphate kinase, grRules: b2168
  atp_c + f1p_c \rightarrow adp_c + fdp_c + h_c
  #2064 PFK (5.75203, 0), Phosphofructokinase, grRules: b3916 or b1723
  atp c + f6p c \rightarrow adp c + fdp c + h c
Show previous steps...
    {'metNames', 'metFormulas', 'rxnNames', 'grRules', 'csense'})
surfNet(iJ01366, 'fdp_c')
Met #473 fdp c, D-Fructose 1,6-bisphosphate, C6H10012P2
Consuming reactions:
  #1151 FBA, Bd: -1000 / 1000, Fructose-bisphosphate aldolase
  fdp c \iff dhap c + g3p c
  #1153 FBP, Bd: 0 / 1000, Fructose-bisphosphatase
  fdp c + h2o_c -> f6p_c + pi_c
Producing reactions:
  #1230 FRUK, Bd: 0 / 1000, Fructose-1-phosphate kinase
  atp_c + f1p_c \rightarrow adp_c + fdp_c + h_c
  #2064 PFK, Bd: 0 / 1000, Phosphofructokinase
  atp c + f6p c \rightarrow adp c + fdp c + h c
Show previous steps...
```

The last argument (8th) 'nCharBreak' sets the number of characters printed per line. By default, it is equal to the width of the Matlab command window. Note the difference:

Characters per line = width of the command window (default):

```
surfNet(iJ01366, [], [], [], 0)
```

```
Rxn #8 BIOMASS_Ec_iJ01366_core_53p95M, Bd: 0 / 1000, E. coli biomass objective function (iJ01366) - colono223 10fthf_c + 2.6e-05 2fe2s_c + 0.000223 2ohph_c + 0.00026 4fe4s_c + 0.51369 ala__L_c + 0.000223 ala__S_c + 0.29579 arg__L_c + 0.24105 asn__L_c + 0.24105 asp_L_c + 54.1248 atp_c + 0.000122 bmocogdp_c + 2e-06 lag. 0.005205 ca2_c + 0.005205 cl_c + 0.000576 coa_c + 2.5e-05 cobalt2_c + 0.13351 ctp_c + 0.000709 cu2_c - 0.09158 cys__L_c + 0.026166 datp_c + 0.027017 dctp_c + 0.027017 dgtp_c + 0.026166 dttp_c + 0.000223 fallono6715 fe2_c + 0.007808 fe3_c + 0.26316 gln_L_c + 0.26316 glu_L_c + 0.61264 gly_c + 0.2151 gtp_c - 48.6015 h2o_c + 0.094738 his__L_c + 0.29053 ile_L_c + 0.19519 k_c + 0.45053 leu_L_c + 0.34316 lys__l 0.15369 met__L_c + 0.008675 mg2_c + 0.000223 mlthf_c + 0.000691 mn2_c + 7e-06 mobd_c + 0.001831 nad_c 0.000447 nadp_c + 0.013013 nh4_c + 0.000323 ni2_c + 0.017868 pe160_c + 0.054154 pe161_c + 0.18527 phe 0.000223 pheme_c + 0.22106 pro__L_c + 0.000223 pydx5p_c + 0.000223 ribflv_c + 0.21579 ser_L_c + 0.00044338 so4_c + 0.000223 thf_c + 0.000223 thmpp_c + 0.25369 thr_L_c + 0.056843 trp_L_c + 0.1379 5.5e-05 udcpdp_c + 0.1441 utp_c + 0.42316 val_L_c + 0.000341 zn2_c + 0.019456 kdo2lipid4_e + 0.013894 murein5px4p_p + 0.045946 pe160_p + 0.02106 pe161_p -> 53.95 adp_c + 53.95 h_c + 53.9457 pi_c 0.7739 ppi_c
```

40 characters per line:

Show previous steps...

 $0.013894 \text{ murein5px4p_p} +$

```
surfNet(iJ01366, [], [], [], 0, [], 40)
```

```
Rxn #8 BIOMASS_Ec_iJ01366_core_53p95M, Bd: 0 / 1000, E. coli biomass objective function (iJ01366) - col
0.000223 10fthf_c + 2.6e-05 2fe2s_c +
  0.000223 20hph_c + 0.00026 4fe4s_c +
  0.51369 ala_L_c + 0.000223 amet_c +
  0.29579 \text{ arg}_Lc + 0.24105 \text{ asn}_Lc +
  0.24105 asp L c + 54.1248 atp c +
  0.000122 \text{ bmocogdp c} + 2e-06 \text{ btn c} +
  0.005205 ca2 c + 0.005205 cl c +
  0.000576 coa c + 2.5e-05 cobalt2 c +
  0.13351 \text{ ctp c} + 0.000709 \text{ cu2 c} +
  0.09158 cys L c + 0.026166 datp c +
  0.027017 \ dctp_c + 0.027017 \ dgtp_c +
  0.026166 \text{ dttp_c} + 0.000223 \text{ fad_c} +
  0.006715 \text{ fe2 c} + 0.007808 \text{ fe3 c} +
  0.26316 gln__L_c + 0.26316 glu__L_c +
  0.61264 gly c + 0.2151 gtp c +
  48.6015 h2o_c + 0.094738 his_
  0.29053 ile_L_c + 0.19519 k_c + 0.45053 leu_L_c + 0.34316 lys_L_c + 0.15369 met_L_c + 0.008675 mg2_c +
  0.000223 \ mlthf_c + 0.000691 \ mn2_c +
  7e-06 \ mobd_c + 0.001831 \ nad \ c +
  0.000447 \text{ nadp\_c} + 0.013013 \text{ nh4\_c} +
  0.000323 \text{ ni2 c} + 0.017868 \text{ pe160 c} +
  0.054154 pe161_c + 0.18527 phe_L_c +
  0.000223 pheme c + 0.22106 pro L c +
  0.000223 \text{ pydx5p c} + 0.000223 \text{ ribflv c}
  + 0.21579 ser_L_c + 0.000223 sheme_c
  + 0.004338 \text{ so4 c} + 0.000223 \text{ thf c} +
  0.000223 thmpp c + 0.25369 thr L c +
  0.056843 \text{ trp } L c + 0.1379 \text{ tyr } L c +
  5.5e-05 \ udcpdp_c + 0.1441 \ utp_c +
  0.42316 \text{ val}\_\_L\_c + 0.000341 \text{ zn2}\_c +
  0.019456 kdo2lipid4 e +
```

```
0.045946 pe160_p + 0.02106 pe161_p -> 53.95 adp_c + 53.95 h_c + 53.9457 pi_c + 0.7739 ppi_c
```

80 characters per line:

Show previous steps...

```
surfNet(iJ01366, [], [], [], [], 0, [], 80)
```

```
Rxn #8 BIOMASS_Ec_iJ01366_core_53p95M, Bd: 0 / 1000, E. coli biomass objective function (iJ01366) - col
0.000223 10fthf_c + 2.6e-05 2fe2s_c + 0.000223 2ohph_c + 0.00026 4fe4s_c +
  0.51369 \text{ ala\_L\_c} + 0.000223 \text{ amet\_c} + 0.29579 \text{ arg\_L\_c} + 0.24105 \text{ asn\_L\_c} +
  0.24105 \text{ asp\_L_c} + 54.1248 \text{ atp\_c} + 0.000122 \text{ bmocogdp\_c} + 2e-06 \text{ btn\_c} +
  0.005205 ca2_c + 0.005205 cl_c + 0.000576 coa_c + 2.5e-05 cobalt2_c +
  0.13351 ctp_c + 0.000709 cu2_c + 0.09158 cys_L_c + 0.026166 datp_c +
  0.027017 \text{ dctp } c + 0.027017 \text{ dgtp } c + 0.026166 \text{ dttp } c + 0.000223 \text{ fad } c +
  0.006715 fe2 c + 0.007808 fe3 c + 0.26316 gln L c + 0.26316 glu L c +
  0.61264 \text{ gly c} + 0.2151 \text{ gtp c} + 48.6015 \text{ h2o c} + 0.094738 \text{ his } \text{L c} +
  0.29053 ile L c + 0.19519 k c + 0.45053 leu L c + 0.34316 lys L c +
  0.15369 \text{ met} \quad L \text{ c} + 0.008675 \text{ mg2 c} + 0.000223 \text{ mlthf c} + 0.000691 \text{ mn2 c} +
  7e-06 \mod c + 0.001831 \mod c + 0.000447 \mod c + 0.013013 \mod c +
  0.000323 \text{ ni2 c} + 0.017868 \text{ pe} 160 \text{ c} + 0.054154 \text{ pe} 161 \text{ c} + 0.18527 \text{ phe} \text{ L c} +
  0.000223 pheme c + 0.22106 pro L c + 0.000223 pydx5p c + 0.000223 ribfly c +
  0.21579 \text{ ser} \text{L}_{c} + 0.000223 \text{ sheme}_{c} + 0.004338 \text{ so4 } c + 0.000223 \text{ thf } c +
  0.000223 \text{ thmpp c} + 0.25369 \text{ thr } L \text{ c} + 0.056843 \text{ trp } L \text{ c} + 0.1379 \text{ tyr } L \text{ c} +
  5.5e-05 udcpdp c + 0.1441 utp c + 0.42316 val L c + 0.000341 zn2 c +
  0.019456 kdo2lipid4_e + 0.013894 murein5px4p_p + 0.045946 pe160_p +
  0.02106 \text{ pe}161_p -> 53.95 \text{ adp_c} + 53.95 \text{ h_c} + 53.9457 \text{ pi_c} + 0.7739 \text{ ppi_c}
Show previous steps...
```

REFERENCES

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