## E.coli Core Model for Beginners (PART 2)

(please run PART 1 of this tutorial first)

### 3. Flux Balance Analysis

Flux balance analysis (FBA) is used to calculate the flow of metabolites through a metabolic network making it possible to predict an organism's growth-rate or the production-rate of a bioproduct. Combining the stoichiometric matrix and the objective function can create a system of linear equations that can be used to calculate the fluxes through all the reactions in the network. In flux balance analysis, these equations are solved using linear programming algorithms that can quickly identify optimal solutions to large systems of equations.

Once the external conditions have been set, which include 1) defining the allowed carbon sources, 2) defining the oxygen uptake level, and 3) setting the objective function, then the simulation conditions are setup to perform FBA. This is accomplished through the use of the "optimizeCbModel(model,osenseStr)", a COBRA toolbox function where the first argument is the model name and the second argument determines if the optimization algorithm maximizes ('max') or minimizes ('min') the objective function. Below is an example for an aerobic environment with glucose as the carbon source optimizing for maximum growth-rate. [Timing: Seconds]

```
model = e_coli_core; % Starting with the original model
model = changeRxnBounds(model,'EX_glc(e)',-10,'l'); % Set maximum glucose uptake
model = changeRxnBounds(model,'EX_o2(e)',-30,'l'); % Set maximum oxygen uptake
model = changeObjective(model,'Biomass_Ecoli_core_w_GAM'); % Set the objective function
FBAsolution = optimizeCbModel(model,'max') % FBA analysis
```

```
FBAsolution =
         full: [95×1 double]
          obj: 0.8739
        rcost: [95×1 double]
         dual: [72×1 double]
       solver: 'gurobi'
    algorithm: 'default'
         stat: 1
     origStat: 'OPTIMAL'
         time: 0.7348
        basis: [1x1 struct]
            x: [95 \times 1 \text{ double}]
            f: 0.8739
            y: [72×1 double]
            w: [95×1 double]
            v: [95×1 double]
```

"FBAsolution" is a Matlab structure that contains the following outputs. "FBAsolution.f" is the value of objective function as calculated by FBA, thus if the biomass reaction is the objective function then "FBAsolution.f" corresponds to the growth-rate of the cell. In the example above, it can be seen that the growth-rate "FBAsolution.f" is listed as  $0.8739~\rm hr^{-1}$ . "FBAsolution.x" is a vector listing the calculated fluxes flowing through the network. "FBAsolution.y" and "FBAsolution.w" contain vectors representing the shadow prices and reduced costs for each metabolite or reaction, respectively.

The flux values found in the structure "FBAsolution.x" can be printed out using the "printFluxVector(model,fluxData,nonZeroFlag,excFlag)" where the second argument is a vector of the flux values, the nonZeroFlag only prints nonzero rows (Default = false), and excFlag only prints exchange reaction

# fluxes (Default = false). Examples of printing non-zero fluxes and exchange reaction only fluxes are shown below. [Timing: Seconds]

```
printFluxVector(model,FBAsolution.x,true) % only prints nonzero rows
ACONTa
       6.00725
ACONTb
       6.00725
AKGDH 5.06438
      8.39
ATPM
ATPS4r
       45.514
Biomass_Ecoli_core_w_GAM 0.873922
CO2t -22.8098
CS 6.00725
CYTBD 43.599
ENO 14.7161
EX_co2(e) 22.8098
EX_glc(e) -10
EX_h(e) 17.5309
EX_h2o(e) 29.1758
EX_nh4(e) -4.76532
EX_02(e) -21.7995
EX_pi(e) -3.2149
FBA 7.47738
FRD7 994.936
FUM 5.06438
G6PDH2r 4.95998
GAPD 16.0235
GLCpts
      10
GLNS
      0.223462
GLUDy
      -4.54186
     4.95998
GND
H2Ot
      -29.1758
ICDHyr 6.00725
MDH 5.06438
NADH16 38.5346
NH4t 4.76532
O2t 21.7995
PDH 9.28253
PFK 7.47738
PGI 4.86086
PGK -16.0235
PGL 4.95998
PGM
     -14.7161
PIt2r
      3.2149
PPC
     2.50431
    1.75818
PYK
    2.67848
RPE
     -2.2815
RPI
SUCDi
       1000
SUCOAS
        -5.06438
      1.49698
TALA
TKT1
      1.49698
TKT2
      1.1815
      7.47738
TPI
```

```
printFluxVector(model,FBAsolution.x,true,true) % only print exchange reaction fluxes
```

```
Biomass_Ecoli_core_w_GAM 0.873922

EX_co2(e) 22.8098

EX_glc(e) -10

EX_h(e) 17.5309

EX_h2o(e) 29.1758

EX_nh4(e) -4.76532
```

```
EX_o2(e) -21.7995
EX_pi(e) -3.2149
```

Printing all the zero and nonzero fluxes can be achieved using "printFluxVector(model,FBAsolution.x)."

These fluxes can also be overlayed on a map of the model as shown below, [Timing: Seconds]

```
map=readCbMap('ecoli_core_map');
options.zeroFluxWidth = 0.1;
options.rxnDirMultiplier = 10;
drawFlux(map, model, FBAsolution.x, options); % Draw the flux values on the map "target"
```

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This overlayed map will be written to a file named "target.svg" that should be located in your working directory. Figure 7 is a screenshot of that map.

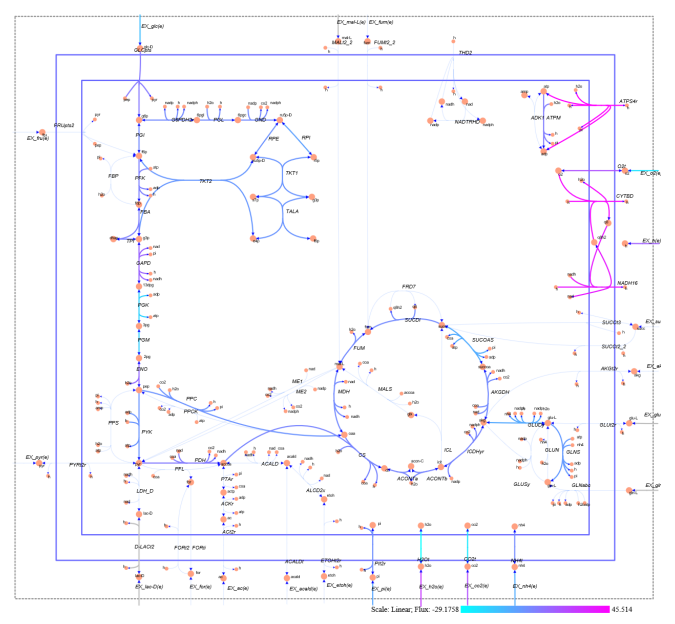
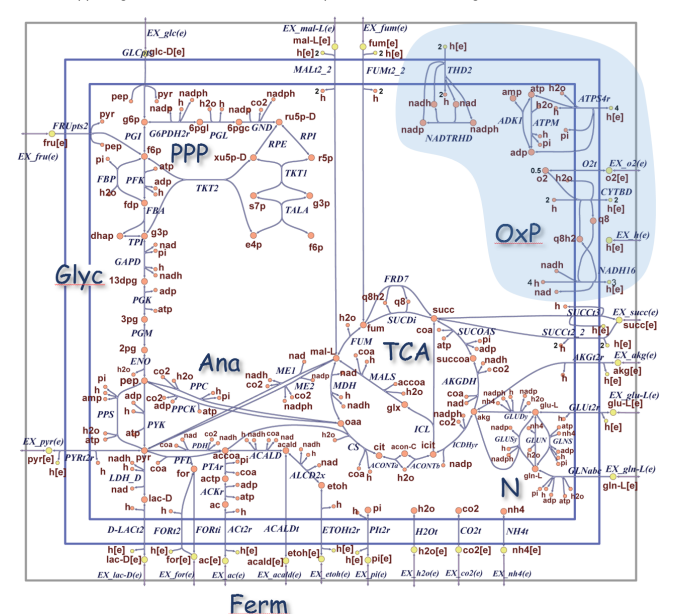


Figure 7. Screenshot of the network map of the *E.coli* core model with  $EX_glc(e) \ge -10 \text{ } \text{mmol} \cdot gDW^{-1} \cdot hr^{-1}$  and  $EX_o2(e) \ge -30 \text{ } \text{mmol} \cdot gDW^{-1} \cdot hr^{-1}$ .

As a cautionary note, the default condition for the E.coli core model sets the carbon source as glucose with an uptake rate of -10  $\text{mmol} \cdot \text{gDW}^{-1} \cdot \text{hr}^{-1}$ , the oxygen uptake is -1000  $\text{mmol} \cdot \text{gDW}^{-1} \cdot \text{hr}^{-1}$  which implies an aerobic environment with the objective function defined as 'Biomass\_Ecoli\_core\_w\_GAM'. It is a good practice to define the conditions of your simulation explicity to avoid unexpected results and long troubleshooting times.

## 4. The Subsystems of the *E.coli* Core Model

Now with these basic Matlab and COBRA toolbox skills behind us, it is time to start exploring the subsystems that make up the *E.coli* core model. We will start by looking at the "energy production and management" section of the model that is referred to as the "oxidative phosphorylation" subsystem in this core model. This subsystem is located in the upper right corner of the *E.coli* core map as shown below in Figure 8.



**Figure 8.** The location of the energy management subsystem and it's reactions highlighted in blue on the *E.coli* core map [3].

As you will see in this section, this subsystem not only includes the reactions for oxydative phosphorylation, it also includes reactions that are required for managing the reducing power needed in the cell. This subsystem will be followed by an exploration of the glycolysis pathway, the pentose phosphate pathway, the tricarboxylic acid cycle, the glycoxylate cycle, gluconeogenesis, and anapleurotic reactions, fermentation pathways, and the nitrogen metabolism.

#### 4.A. Energy Production & Management

Perhaps the most important requirement of an operational cell is the production and management of energy and reducing power. There are two main mechanisms available within the *E.coli* core model for the production of ATP (atp[c]) energy: 1) substrate level phosphorylation, and 2) oxidative phosphorylation through the use of the electron transport chain. Substrate level phosphorylation occurs when specific metabolic pathways within the cell are net producers of energy. In these cases, atp[c] is formed by a reaction between ADP (adp[c]) and a phosphorylated intermediate within the pathway. In the core model this occurs in the glycolysis pathway with both phosphoglycerate kinase (PGK), and pyruvate kinase (PYK), and in the tricarboxylic acid cycle with succinyl-CoA synthetase (SUCOAS). Through these substrate level phosphorylation enzymes each molecule of glucose can potentially add four molecules to the total cellular flux of atp[c].

The second mechanism for energy generation is oxidative phosphorylation through the electron transport chain, which under aerobic conditions, produces the bulk of the cell's atp[c]. In the simple core model, the electron transport chain is used to transport protons (h[c]) from the cytoplasm across the cytoplasmic membrane into the extracellular space (periplasmic space in actual cells) to create a proton-motive force which drives ATP synthase (ATPS4r) to produce atp[c].

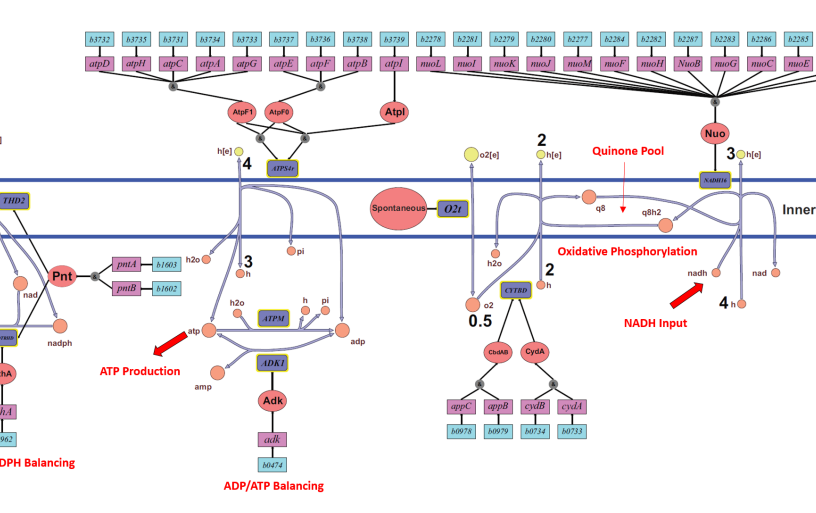


Figure 9. Oxidative Phosphorylation and Transfer of Reducing Equivalents [3].

#### Aerobic Respiration

For aerobic respiration, the primary source of atp[c] is produced through oxidative phosphorylation. This is illustrated in Figure 9 where NADH (nadh[c]), acting as a substrate for NADH dehydrogenase (NADH16), provides the reducing power necessary to trigger the electron transport chain. The *E. coli* core model combines the electron transport chain into two reactions. In the first of these two reactions, NADH16 catalyzes the oxidation of nadh[c] to form NAD+ (nad[c]) while extracting four protons (h[c]) from the cytoplasm. It then transports three protons to the extracellular space while combining the fourth proton with a proton and two electrons from NADH to transform ubiquinone-8) (q8[c]) to its reduced form ubiquinol-8 (q8h2[c]). Both q8[c] and q8h2[c] are oil soluble coenzymes that can diffuse freely within the lipid environment of the cytoplasmic membrane allows q8h2[c] to eventually transfer its two electrons and two protons to cytochrome oxidase (CYTBD). The two protons (h[e]) are then transferred into the extracellular space where they add to the proton-motive force. The two electrons from q8h2[c] are then combined with two cytoplasmic protons and an oxygen atom, the terminal electron acceptor, to form water. In this model, oxygen (o2[c]) spontaneously diffuses from the environment into the cell through the spontaneous 02t reaction.

With a proton-motive force now created by the pumping of protons from the cytoplasm to the extracellular space, the reaction ATPS4r can synthesize atp[c] from adp[c]. For this simple model the *P*/0 ratio is stoichiometrically set to 1.25. Another reaction included in the energy management suite is adenylate kinase (ADK1), a

phosphotransferase enzyme that catalyzes the interconversion of adenine nucleotides, and plays an important role in the adp[c]/atp[c] balance or cellular energy homeostasis.

Finally, the ATP maintenance function (ATPM), which is set at  $8.39 \, \text{mmol} \cdot \text{gDW}^{-1} \cdot \text{hr}^{-1}$  accounts for the energy (in form of atp[c]) necessary to replicate a cell, including for macromolecular synthesis (e.g., proteins, DNA, and RNA). Thus, for growth to occur in the *E.coli* model, the flux rate through ATPM must be greater than  $8.39 \, \text{mmol} \cdot \text{gDW}^{-1} \cdot \text{hr}^{-1}$ . If the model detects that ATPM has not reached its minimum value it will not produce FBA results.

Another part of the energy management of a cell is the reducing power that is required for both cellular catabolism and anabolism. Catabolism refers to a set of metabolic pathways that break down molecules into smaller units and release energy. For this core model, nadh[c] provides the reducing power necessary for the catabolic activities of the cell.

Anabolism, on the other hand, is the set of metabolic pathways that construct molecules from smaller units. These anabolic reactions are endergonic and therefore require an input of energy. In this case, NADPH (nadph[c]) is the reducing power required for biosynthesis using the cell's precursor metabolites.

Maintaining the proper balance between anabolic reduction charge, nadph[c]/ nadp[c], and catabolic reduction charge, nadh[c]/ nadpc], is achieved by reactions catalyzed by transhydrogenase enzymes, as shown in Figure 9. Using the proton-motive force, NAD(P) transhydrogenase (THD2) catalyzes the transfer of a hydride ion, a negative ion of hydrogen, from nadh[c] to create nadph[c]. The opposite transfer, of a hydride ion from nadph[c], to create nadh[c], is catalyzed by another enzyme, NAD+ transhydrogenase (NADTRHD), but it is not coupled to the translocation of protons. These pair of reactions effectively allow transfer of reducing equivalents between anabolic and catabolic reduction charge.

Now let's use the COBRA Toolbox to explore the details of the energy managing elements of the *E.coli* core model. In this tutorial, we will focus on exploring the role of cofactors in a core model that is optimized for growth-rate. There is a good discussion of how to find the maximum cofactor fluxes possible in a COBRA-based model in Chapter 19 of Palsson's book [1]. To start with let's print out a table that includes all the reaction abbreviations, names, and their formulas for the reactions invovled in oxidative phosphorylation and the cell's energy and reducing power management (see Figure 9). [Timing: Seconds]

```
model = e_coli_core; % Starting this section with the original model
energySubSystems = {'Oxidative Phosphorylation'};
energyReactions = model.rxns(ismember(model.subSystems,energySubSystems));
[~,energy_rxnID] = ismember(energyReactions,model.rxns);
reactionNames = model.rxnNames(energy_rxnID);
reactionFormulas = printRxnFormula(model,energyReactions,0);
T = table(reactionNames,reactionFormulas,'RowNames',energyReactions)
```

T = reactionNames reactionFormulas

```
'amp[c] + atp[c] <=> 2 adp[c] '
ADK1
           'adenylate kinase'
ATPM
           'ATP maintenance requirement'
                                                               'atp[c] + h2o[c] -> adp[c] + h[c] + pi
           'ATP synthase (four protons for one ATP)'
ATPS4r
                                                               'adp[c] + 4 h[e] + pi[c] <=> atp[c] +
CYTBD
           'cytochrome oxidase bd (ubiquinol-8: 2 protons)'
                                                               '2 h[c] + 0.5 o2[c] + q8h2[c] -> h2o[c]
                                                               fum[c] + q8h2[c] -> q8[c] + succ[c]'
FRD7
           'fumarate reductase'
NADH16
           'NADH dehydrogenase (ubiquinone-8 & 3 protons)'
                                                               '4 h[c] + nadh[c] + q8[c] -> 3 h[e] +
```

```
NADTRHD 'NAD transhydrogenase' 'nad[c] + nadph[c] -> nadh[c] + nadp[c] 
SUCDi 'succinate dehydrogenase (irreversible)' 'q8[c] + succ[c] -> fum[c] + q8h2[c] 'THD2 'NAD(P) transhydrogenase' '2 h[e] + nadh[c] + nadp[c] -> 2 h[c]
```

Although this is a specifc table for the reactions associated with energy management, it illustrates how you can pull up the full reaction (enzyme) name and formula for any subsystem in the core model. It should be pointed out that although the reactions succinate dehydrogenase (SUCDi) and fumarate reductase (FRD7) are included in the oxidative phosphorylation subsystem because they are membrane-bound enzymes that interact with the quinone pool, they are a better fit functionally in the TCA cycle, as will be seen later.

Now lets explore the flux through these reactions in aerobic conditons with the glucose uptake set at -10 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup> and the oxygen uptake at -30 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup>. [Timing: Seconds]

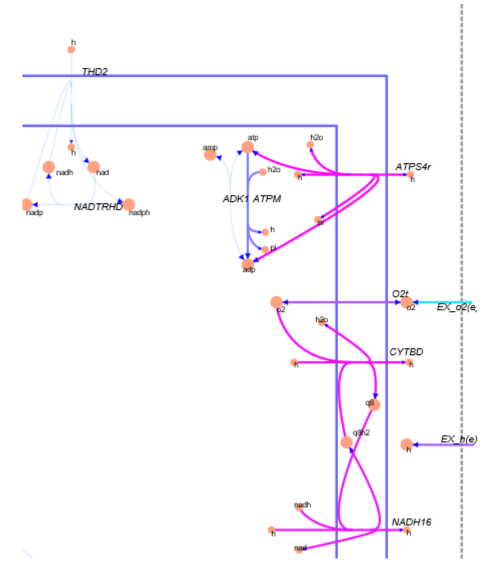
NADTRHD SUCDi

THD2

1000

```
model = changeRxnBounds(model,'EX_glc(e)',-10,'l'); % Set maximum glucose uptake
model = changeRxnBounds(model,'EX_o2(e)',-30,'1'); % Set oxygen uptake
model = changeObjective(model, 'Biomass_Ecoli_core_w_GAM'); % Set the objective function
FBAsolution = optimizeCbModel(model, 'max'); % Perform FBA
printLabeledData(energyReactions,FBAsolution.x(energy_rxnID))
ADK1
ATPM
      8.39
       45.514
ATPS4r
       43.599
CYTBD
FRD7
      994.936
NADH16
        38.5346
```

Below in Figure 10 is screenshot showing these fluxes flowing through the oxidative phosphorylation section of the core map (upper right corner). In this figure we can see the electrons from nadh[c] entering the electron transport change at NADH16, flowing through the quinone pool, and then finding their way to reduce oxygen through CYTBD and O2t. With the proton-motive force in place, ATPS4r can now use that energy to convert adp[c] to atp[c]. We can also see the flux flowing through the dummy reaction ATPM that is used to model the atp[c] load required for cell growth. Finally, THD2, NADTRHD or ADK1 are not required to recycle any of the key energy cofactors.



**Figure 10:** Close-up of the oxidative phosphorylation section of the *E.coli* core map in aerobic conditions with glucose as the sole carbon source (see Figure 7).

#### **ATP Production**

Now let's explore in more detail the production and consumption of atp[c] in the core model. The atp[c] produced by ATPS4r is added to the total cellular atp[c] flux that provides the cell's energy. Remember that in aerobic conditions, atp[c] is produced by both substrate phosphorylation and oxidative phosphorylation. All of the reactions that either produce or consume atp[c] can be found using the "surfNet" COBRA toolbox function. [Timing: Seconds]

```
surfNet(model, 'atp[c]', 0,FBAsolution.x,1,1)

Met #17 atp[c], ATP, C10H12N5013P3
Consuming reactions with non-zero fluxes:
    #11 ATPM (8.39), Bd: 8.39 / 1000, ATP maintenance requirement
    atp[c] + h2o[c] -> adp[c] + h[c] + pi[c]
    #13 Biomass_Ecoli_core_w_GAM (0.87392), Bd: 0 / 1000, Biomass Objective Function with GAM
    1.496 3pg[c] + 3.7478 accoa[c] + 59.81 atp[c] + 0.361 e4p[c] + 0.0709 f6p[c] + 0.129 g3p[c] + 0.205 g6p[
```

```
#51 GLNS (0.22346), Bd: 0 / 1000, glutamine synthetase atp[c] + glu-L[c] + nh4[c] -> adp[c] + gln-L[c] + h[c] + pi[c] #72 PFK (7.47738), Bd: 0 / 1000, phosphofructokinase atp[c] + f6p[c] -> adp[c] + fdp[c] + h[c] Producing reactions with non-zero fluxes:

#12 ATPS4r (45.514), Bd: -1000 / 1000, ATP synthase (four protons for one ATP) adp[c] + 4 h[e] + pi[c] <=> atp[c] + h2o[c] + 3 h[c] #75 PGK (-16.0235), Bd: -1000 / 1000, phosphoglycerate kinase 3pg[c] + atp[c] <=> 13dpg[c] + adp[c] #83 PYK (1.75818), Bd: 0 / 1000, pyruvate kinase adp[c] + h[c] + pep[c] -> atp[c] + pyr[c] #90 SUCOAS (-5.06438), Bd: -1000 / 1000, succinyl-CoA synthetase (ADP-forming) atp[c] + coa[c] + succ[c] <=> adp[c] + pi[c] + succoa[c]
```

These results show that under aerobic conditions with glucose as the sole carbon source there are four producers of atp[c] within the core model. These include ATPS4r (oxidative phosphorylation) as the primary contributor and PGK, PYK, and SOCAS (substrate phosphoylation) as secondary sources. This also shows the consumers to be GLNS, PFK, ATPM and the biomass function. As we will see later, the atp[c] associated with PFK is required by the glycolysis pathway. The atp[c] used by ATPM must be greater than or equal to 8.39 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup> to allow the cell to grow. Finally the biomass function shows that 52.27 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup> (0.873922 x 59.81) is used for the cell's biosynthesis needs.

One of the important concepts associated with these constraint-based steady state models is that the total cell fluxes for key cofactors like atp[c] and adp[c] must be equal. This means that for every atp[c] metabolite that is produced, one adp[c] metabolite will be consumed, but to maintain the mass balance throughout the cell somewhere else in the cell an adp[c] molecule will be created from another atp[c] molecule. Thus, the total cellular atp[c] flux must equal the total cellular adp[c] flux. This can be observed using the COBRA Toolbox function called "computeFluxSplits" as shown below. [Timing: Seconds]

```
[P, C, vP, vC] = computeFluxSplits(model, {'adp[c]'}, FBAsolution.x);
total_adp_flux = sum(vP)

total_adp_flux = 68.3601

[P, C, vP, vC] = computeFluxSplits(model, {'adp[c]'}, FBAsolution.x);
total_adp_flux = sum(vP)

total_adp_flux = 68.3601
```

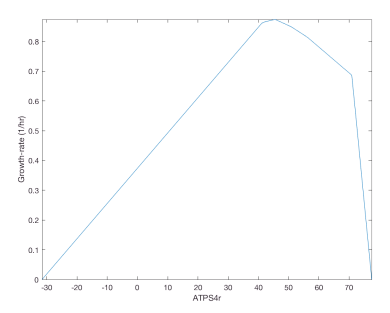
These results show that the amount of atp[c] flux in the cell equals the amount of adp[c] flux. Thus, the adp[c]/ atp[c] flux ratio is 1. This is also true for nadp[c]/nadph[c] and the nad[c]/nadh[c] flux ratios.

Another way to explore the ATPS4r's ability to produce atp[c] is through the use of robustness analysis [12]. Assuming that the objective function is the biomass function (growth-rate), then the following simulation illustrates that the maximum atp[c] flux that can be supported by ATPS4r under aerobic conditions with glucose as the sole carbon source. [Timing: Minutes]

```
model = changeRxnBounds(model,'EX_glc(e)',-10,'l'); % Set maximum glusose uptake
model = changeRxnBounds(model,'EX_o2(e)',-30,'l'); % Set oxygen uptake
[controlFlux, objFlux] = robustnessAnalysis(model,'ATPS4r',100);
```

]3%

```
ylabel('Growth-rate (1/hr)');
```



This graph shows the entire capability of ATPS4r when the carbon source glucose has a maximum uptake rate greater than or equal to  $-10~\text{mmol} \cdot \text{gDW}^{-1} \cdot \text{hr}^{-1}$ . If we start at the left of this of this figure, it can be seen that ATPS4r takes on negative values which implies that instead of producing atp[c] through the proton-motive force, it has become an energy-dependent proton pump removing protons from the cytoplasm and transporting them to the extracellular space. Note that the growth-rate under these anaerobic conditions is small. As the flux through ATPS4r becomes positive it starts producing atp[c] providing the majority of the atp[c] required for aerobic operation. At the beginning of aerobic operation there is a nice linear relationship between the produced atp[c] and the growth-rate. Eventually the growth-rate reaches a maximum of 0.8738 hr<sup>-1</sup> when the ATPS4r flux level reaches 45.54 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup>. After the maximum growth-rate has been achieved the cell then needs to find ways to recycle the extra ATP. This can be seen below by fixing the flux through ATPS4r to a value greater than 45.54 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup>. [Timing: Seconds]

```
model = e_coli_core; % Starting the original model
model = changeRxnBounds(model,'EX_glc(e)',-10,'l'); % Set maximum glucose uptake
model = changeRxnBounds(model,'ATPS4r',60,'b'); % Fix ATPS4r flux rate
FBAsolution = optimizeCbModel(model,'max'); % Perform FBA
surfNet(model, 'atp[c]', 0,FBAsolution.x,1,1)
```

```
Met #17 atp[c], ATP, C10H12N5013P3
Consuming reactions with non-zero fluxes :
    #7 ADK1 (4.01673), Bd: -1000 / 1000, adenylate kinase
    amp[c] + atp[c] <=> 2 adp[c]
    #11 ATPM (13.7463), Bd: 8.39 / 1000, ATP maintenance requirement
    atp[c] + h2o[c] -> adp[c] + h[c] + pi[c]
    #13 Biomass_Ecoli_core_w_GAM (0.78116), Bd: 0 / 1000, Biomass Objective Function with GAM
    1.496 3pg[c] + 3.7478 accoa[c] + 59.81 atp[c] + 0.361 e4p[c] + 0.0709 f6p[c] + 0.129 g3p[c] + 0.205 g6p[
    #51 GLNS (0.19974), Bd: 0 / 1000, glutamine synthetase
    atp[c] + glu-L[c] + nh4[c] -> adp[c] + gln-L[c] + h[c] + pi[c]
```

```
atp[c] + f6p[c] -> adp[c] + fdp[c] + h[c]
#81    PPS (4.01673),    Bd: 0 / 1000, phosphoenolpyruvate synthase
atp[c] + h2o[c] + pyr[c] -> amp[c] + 2 h[c] + pep[c] + pi[c]

Producing reactions with non-zero fluxes:
    #12    ATPS4r (60),    Bd: 60 / 60,    ATP synthase (four protons for one ATP)
    adp[c] + 4 h[e] + pi[c] -> atp[c] + h2o[c] + 3 h[c]
    #75    PGK (-9.79587),    Bd: -1000 / 1000, phosphoglycerate kinase
    3pg[c] + atp[c] <=> 13dpg[c] + adp[c]

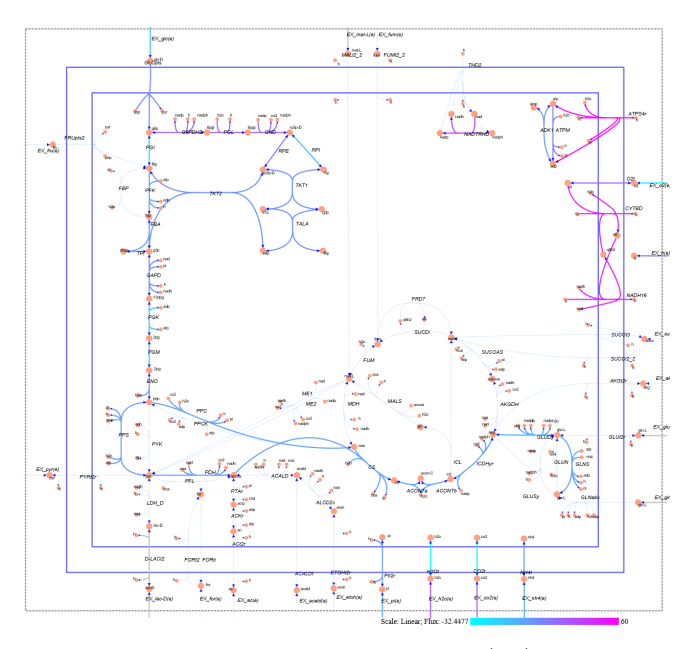
Show previous steps...

map=readCbMap('ecoli_core_map');
    options.zeroFluxWidth = 0.1;
    options.rxnDirMultiplier = 10;
    drawFlux(map, model, FBAsolution.x, options); % Draw the flux values on the map "target"
```

#72 PFK (1.0954), Bd: 0 / 1000, phosphofructokinase

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If we compare these results with the previous fluxes calculated for the optimized cell performance under aerobic conditions with a similar glucose carbon source uptake, we can see the differences in atp[c] flux distribution. To start with it can be seen that the flux through ATPM increases (13.74 > 8.39). Notice that ADK1 has been activated to recycle atp[c] to adp[c]. Since the growth-rate decreases, we would also expect the flux used by the biomass function to decrease along with other parts of the cell by selecting alternate pathways to help absorb the extra atp[c]. This is illustrated in the core metabolic map shown below.



**Figure 11.** A screenshot of the core map with ATPS4r fixed at  $60 \, \mathrm{mmol} \cdot \mathrm{gDW^{-1}} \cdot \mathrm{hr^{-1}}$ .

#### **NADH Production**

Now that we have explored the production and consumption of atp[c], let's look at the producers and consumers of nadh[c]. [Timing: Seconds]

```
model = e_coli_core; % Starting with the original model
model = changeRxnBounds(model,'EX_glc(e)',-10,'l'); % Set maximum glucose uptake
model = changeRxnBounds(model,'EX_o2(e)',-30,'l'); % Set oxygen uptake
FBAsolution = optimizeCbModel(model,'max'); % Perform FBA
surfNet(model,'nadh[c]',0,FBAsolution.x,1,1)
Met #51 nadh[c], Nicotinamide-adenine-dinucleotide-reduced, C21H27N7014P2
Consuming reactions with non-zero fluxes:
```

#67 NADH16 (38.5346), Bd: 0 / 1000, NADH dehydrogenase (ubiquinone-8 & 3 protons)

4 h[c] + nadh[c] + q8[c] -> 3 h[e] + nad[c] + q8h2[c]

Note that in this case, the only consumer of nadh[c] is NAD16 which is the beginning of the electron transport chain. The producing reactions, as we will discuss later, are primarily located in the glycolysis and TCA pathways. Note that for this core model, the biomass function is also listed as a producer. Since the biomass function represents all the functionality not included in the core model (e.g. biosynthesis pathways), this implies that NADH would be produced in other parts of the cell that are not included in this simple core model. The flux supplied through the biomass function is calculated by multiplying the total biomass flux (0.873922) times the nadh[c] biomass function coefficient (3.547) to yielding a total nadh[c] biomass flux of 3.0998 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup>. This can also be calculated using the COBRA Toolbox function "computeFluxSplits" as follows. [Timing: Seconds]

```
[nadh_P, nadh_C, nadh_vP, nadh_vC] = computeFluxSplits(model, {'nadh[c]'}, FBAsolution
[~,nadh_rxnID] = ismember('Biomass_Ecoli_core_w_GAM',model.rxns);
nadhBiomassFlux = nadh_vP(nadh_rxnID)
```

nadhBiomassFlux = 3.0998

#### **NADPH** production

Finally, we can also obtain this same information for nadph[c], the reducing power for cellular biosynthesis. [Timing: Seconds]

```
surfNet(model, 'nadph[c]', 0, FBAsolution.x,1,1)

Met #53    nadph[c], Nicotinamide-adenine-dinucleotide-phosphate-reduced, C21H26N7017P3
Consuming reactions with non-zero fluxes:
    #13    Biomass_Ecoli_core_w_GAM (0.87392), Bd: 0 / 1000, Biomass Objective Function with GAM
    1.496    3pg[c] + 3.7478    accoa[c] + 59.81    atp[c] + 0.361    e4p[c] + 0.0709    f6p[c] + 0.129    g3p[c] + 0.205    g6p[
    #53    GLUDy (-4.54186), Bd: -1000 / 1000, glutamate dehydrogenase (NADP)
    glu-L[c] + h2o[c] + nadp[c] <=> akg[c] + h[c] + nadph[c] + nh4[c]

Producing reactions with non-zero fluxes:
    #48    G6PDH2r (4.95998), Bd: -1000 / 1000, glucose 6-phosphate dehydrogenase
    g6p[c] + nadp[c] <=> 6pg1[c] + h[c] + nadph[c]
    #57    GND (4.95998), Bd: 0 / 1000, phosphogluconate dehydrogenase
    6pgc[c] + nadp[c] -> co2[c] + nadph[c] + ru5p-D[c]
    #59    ICDHyr (6.00725), Bd: -1000 / 1000, isocitrate dehydrogenase (NADP)
    icit[c] + nadp[c] <=> akg[c] + co2[c] + nadph[c]
Show previous steps...
```

Due to the simplicity of the E.coli core model, most of the nadph[c] is consumed by the biomass function (0.873922 x 13.0279 = 11.385) to support the cell's biosynthesis needs. The other consumer is the nitrogen metabolism (GLUDy). On the other hand, nadph[c] is produced by reactions in the oxidative phosphorylation

pathways, pentose phosphate pathway, and the TCA cycle. It is worth pointing out that in the larger models, that incorporate most of the cells biosynthesis pathways, the number of reactions consuming nadph[c] could be very large. [Timing: Seconds]

#### Anaerobic Respiration

Now let's turn our attention to anaerobic cell operation. During aerobic respiration, oxygen is the terminal electron acceptor for the electron transport chain, which yields the bulk of atp[c] required for biosynthesis. Anaerobic respiration refers to respiration without molecular oxygen. For anaerobic respiration, *E. coli* only generates atp[c] by substrate level phosphorylation. Glycolysis results in the net production of two atp[c] per glucose by substrate level phosphorylation, but this is low compared to the total atp[c] production of 17.5 atp[c] per glucose for aerobic respiration [1].

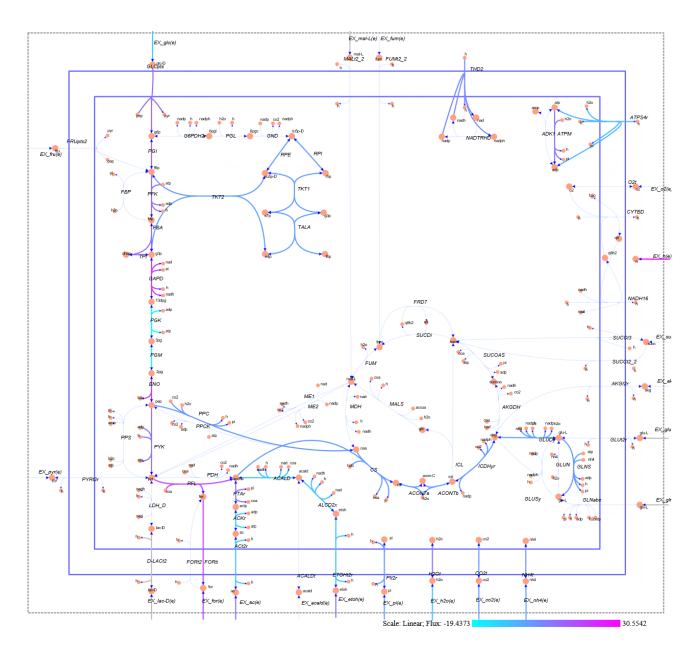
The substrates of fermentation are typically sugars, so during fermentative growth, it is necessary for each cell to support large flux values through glycolysis to generate sufficient atp[c] to drive cell growth. Glycolysis also produces two molecules of nadh[c] for each molecule of glucose [1]. As a result, nadh[c] must be reoxidized by fermentation in order to regenerate nad[c] necessary to maintain the oxidation-reduction balance of the cell.

Figure 12 is a map of anaerobic operation using glucose as the only carbon source.

```
model = e_coli_core; % Starting with the original model
model = changeRxnBounds(model,'EX_glc(e)',-10,'l'); % Set maximum glusose uptake
model = changeRxnBounds(model,'EX_o2(e)',-0,'l'); % Set maximum oxygen uptake
model = changeObjective(model,'Biomass_Ecoli_core_w_GAM'); % Set the objective function
FBAsolution = optimizeCbModel(model,'max'); % Perform FBA
map=readCbMap('ecoli_core_map');
options.zeroFluxWidth = 0.1;
options.rxnDirMultiplier = 10;
drawFlux(map, model, FBAsolution.x, options); % Draw the flux values on the map "target"
```

Document Written

A screenshot of the produced map of anaerobic operation is shown below.



**Figure 12.** Network map of the *E.coli* core model with glucose as the carbon source (EX\_glc(e)  $\geq$  -10 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup>) in an anaerobic environment (EX\_o2(e)  $\geq$  0 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup>).

Note that for anaerobic operation the flux through oxidative phosphorylation pathways (electron transport chain) is zero. Let's look at the nonzero fluxes associated with anaerobic operation to understand the role of THD2 and ATPS4r. [Timing: Seconds]

```
Reactions = transpose({'ATPS4r','THD2'});
[~,rxnID] = ismember(Reactions,model.rxns);
printLabeledData(Reactions,FBAsolution.x(rxnID))
ATPS4r -5.45205
THD2 3.62919
```

Now let's look at the fomulas for these reactions to understand what is happening in this condition. [Timing: Seconds]

#### printRxnFormula(model,Reactions)

```
ATPS4r adp[c] + 4 h[e] + pi[c] <=> atp[c] + h2o[c] + 3 h[c]
THD2 2 h[e] + nadh[c] + nadp[c] -> 2 h[c] + nad[c] + nadph[c]
ans =
    'adp[c] + 4 h[e] + pi[c] <=> atp[c] + h2o[c] + 3 h[c] '
    '2 h[e] + nadh[c] + nadp[c] -> 2 h[c] + nad[c] + nadph[c] '
```

Since the flux for ATPS4r is negative, we can assume that ATPS4r is operating in reverse and pumping protons from the cytoplasm into the extracellular space. Some of these protons can now be used by THD2 to convert nadh[c], which is not needed for the electron transport chain, into nadph[c] where they can be used for cellular biosynthesis.

All the nonzero fluxes for this anaerobic example are printed below. [Timing: Seconds]

printFluxVector(model,FBAsolution.x,true) % only print nonzero reaction fluxes

```
ACALD
      -8.27946
ACKr -8.50359
ACONTa 0.228363
ACONTb 0.228363
      -8.50359
ACt2r
ALCD2x -8.27946
ATPM 8.39
ATPS4r -5.45205
Biomass_Ecoli_core_w_GAM 0.211663
CO2t 0.378178
    0.228363
CS
    19.1207
ENO
ETOHt2r
         -8.27946
         8.50359
EX_ac(e)
EX_co2(e)
           -0.378178
EX_etoh(e) 8.27946
EX_for(e) 17.8047
EX_glc(e)
           -10
EX_h(e) 30.5542
EX_h2o(e) -7.1158
EX_nh4(e) -1.15416
EX_pi(e)
         -0.778644
FBA 9.78946
FORti
      17.8047
GAPD 19.4373
GLCpts 10
GLNS 0.0541222
GLUDy -1.10003
      7.1158
H2Ot.
ICDHyr 0.228363
      1.15416
NH4t
    9.78946
PFK
    17.8047
PFL
PGI
     9.95661
      -19.4373
      -19.1207
PGM
       0.778644
PIt2r
      0.606541
PPC
      8.50359
PTAr
PYK
     8.40427
RPE -0.152143
     -0.152143
RPI
TALA -0.0378665
THD2
      3.62919
```

```
TKT1 -0.0378665
TKT2 -0.114277
TPI 9.78946
```

So one question that could be asked is this anaerobic environment is, where is the nadh[c] produced and where is it consumed. Using "surfNet" we can find out. [Timing: Seconds]

```
surfNet(model, 'nadh[c]',0,FBAsolution.x,1,1)

Met #51 nadh[c], Nicotinamide-adenine-dinucleotide-reduced, C21H27N7014P2
Consuming reactions with non-zero fluxes:
    #1 ACALD (-8.27946), Bd: -1000 / 1000, acetaldehyde dehydrogenase (acetylating)
    acald[c] + coa[c] + nad[c] <=> accoa[c] + h[c] + nadh[c]
    #10 ALCD2x (-8.27946), Bd: -1000 / 1000, alcohol dehydrogenase (ethanol)
    etoh[c] + nad[c] <=> acald[c] + h[c] + nadh[c]
    #92 THD2 (3.62919), Bd: 0 / 1000, NAD(P) transhydrogenase
    2 h[e] + nadh[c] + nadp[c] -> 2 h[c] + nad[c] + nadph[c]

Producing reactions with non-zero fluxes:
    #13 Biomass_Ecoli_core_w_GAM (0.21166), Bd: 0 / 1000, Biomass Objective Function with GAM
    1.496 3pg[c] + 3.7478 accoa[c] + 59.81 atp[c] + 0.361 e4p[c] + 0.0709 f6p[c] + 0.129 g3p[c] + 0.205 g6p[
    #49 GAPD (19.4373), Bd: -1000 / 1000, glyceraldehyde-3-phosphate dehydrogenase
    g3p[c] + nad[c] + pi[c] <=> 13dpg[c] + h[c] + nadh[c]
Show previous steps...
```

In this case, the nadh[c] is primarily used to support mixed fermentation through the ethanol pathway. This will be described in the fermentation section.

Now let's explore the production of atp[c] in an anaerobic environment. [Timing: Seconds]

```
surfNet(model, 'atp[c]',0,FBAsolution.x,1,1)
Met #17 atp[c], ATP, C10H12N5O13P3
Consuming reactions with non-zero fluxes :
      #11 ATPM (8.39), Bd: 8.39 / 1000, ATP maintenance requirement
      atp[c] + h2o[c] -> adp[c] + h[c] + pi[c]
      #12 ATPS4r (-5.45205), Bd: -1000 / 1000, ATP synthase (four protons for one ATP)
      adp[c] + 4 h[e] + pi[c] <=> atp[c] + h2o[c] + 3 h[c]
      #13 Biomass_Ecoli_core_w_GAM (0.21166), Bd: 0 / 1000, Biomass Objective Function with GAM
      1.496 \ 3pg[c] + 3.7478 \ accoa[c] + 59.81 \ atp[c] + 0.361 \ e4p[c] + 0.0709 \ f6p[c] + 0.129 \ g3p[c] + 0.205 \ g6p[c] + 
      #51 GLNS (0.05412), Bd: 0 / 1000, glutamine synthetase
      atp[c] + glu-L[c] + nh4[c] -> adp[c] + gln-L[c] + h[c] + pi[c]
      #72 PFK (9.78946), Bd: 0 / 1000, phosphofructokinase
      atp[c] + f6p[c] -> adp[c] + fdp[c] + h[c]
Producing reactions with non-zero fluxes :
      #3 ACKr (-8.50359), Bd: -1000 / 1000, acetate kinase
      ac[c] + atp[c] <=> actp[c] + adp[c]
      #75 PGK (-19.4373), Bd: -1000 / 1000, phosphoglycerate kinase
      3pg[c] + atp[c] <=> 13dpg[c] + adp[c]
      #83 PYK (8.40427), Bd: 0 / 1000, pyruvate kinase
      adp[c] + h[c] + pep[c] -> atp[c] + pyr[c]
Show previous steps...
```

As can be seen above, the production of atp[c] is exclusively through substrate phosphorylation (ACKr, PGK, PYK).

Finally, the nadph[c] producers and consumers are shown below. [Timing: Seconds]

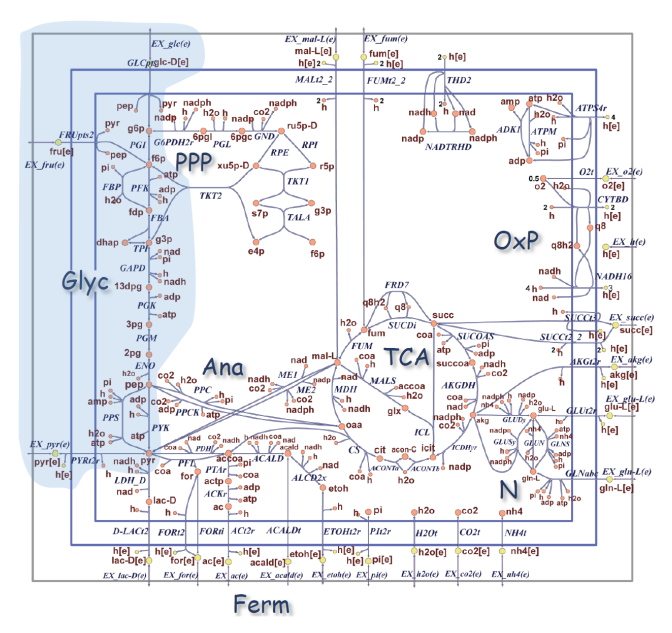
```
surfNet(model, 'nadph[c]',0,FBAsolution.x,1,1)
```

```
Met #53     nadph[c], Nicotinamide-adenine-dinucleotide-phosphate-reduced, C21H26N7017P3
Consuming reactions with non-zero fluxes :
    #13     Biomass_Ecoli_core_w_GAM (0.21166), Bd: 0 / 1000, Biomass Objective Function with GAM
    1.496 3pg[c] + 3.7478 accoa[c] + 59.81 atp[c] + 0.361 e4p[c] + 0.0709 f6p[c] + 0.129 g3p[c] + 0.205 g6p[
#53     GLUDy (-1.10003), Bd: -1000 / 1000, glutamate dehydrogenase (NADP)
    glu-L[c] + h2o[c] + nadp[c] <=> akg[c] + h[c] + nadph[c] + nh4[c]
Producing reactions with non-zero fluxes :
    #59     ICDHyr (0.22836), Bd: -1000 / 1000, isocitrate dehydrogenase (NADP)
    icit[c] + nadp[c] <=> akg[c] + co2[c] + nadph[c]
    #92     THD2 (3.62919), Bd: 0 / 1000, NAD(P) transhydrogenase
    2     h[e] + nadh[c] + nadp[c] -> 2     h[c] + nadph[c]
Show previous steps...
```

Note that the primary producer of nadph[c] in this anaerobic environment is THD2, which converts the surplus nadh[c] to nadph[c].

## 4.B. Glycolysis Pathway

Now that we have completed the exploration of the energy management subsystem of the core model, it is time to start looking at the other included subsytems. Glycolysis is the metabolic pathway in the *E.coli* core model that converts glucose and fructose into pyruvate. The free energy released in this process is used to form the high-energy compounds of atp[c] and nadh[c]. The location of the glycolysis pathway on the *E.coli* core map is highlighted in the Figure 13.



**Figure 13.** The location of the glycolysis pathway subsystem reactions are highlighted in blue on the *E.coli* core map [3].

A table showing the reactions associated with the glycolysis pathway can be extracted from the core model as follows: [Timing: Seconds]

```
model = e_coli_core; % Starting with the original model
model = changeRxnBounds(model,'EX_glc(e)',-10,'l');
model = changeRxnBounds(model,'EX_o2(e)',-30,'l');
model = changeObjective(model,'Biomass_Ecoli_core_w_GAM');
glycolysisSubystem = {'Glycolysis/Gluconeogenesis'};
glycolysisReactions = model.rxns(ismember(model.subSystems,glycolysisSubystem));
[~,glycolysis_rxnID] = ismember(glycolysisReactions,model.rxns);
Reaction_Names = model.rxnNames(glycolysis_rxnID);
Reaction_Formulas = printRxnFormula(model,glycolysisReactions,0);
T = table(Reaction_Names,Reaction_Formulas,'RowNames',glycolysisReactions)
```

T =

```
<=> h2o[c] + pep[c] '
ENO
                                                       '2pg[c]
                                                               <=> dhap[c] + g3p[c] '
FBA
        'fructose-bisphosphate aldolase'
                                                       'fdp[c]
FBP
        'fructose-bisphosphatase'
                                                       'fdp[c] + h2o[c] -> f6p[c] + pi[c] '
                                                       'g3p[c] + nad[c] + pi[c] <=> 13dpg[c] + h[c] +
        'glyceraldehyde-3-phosphate dehydrogenase'
GAPD
PDH
        'pyruvate dehydrogenase'
                                                       'coa[c] + nad[c] + pyr[c] -> accoa[c] + co2[c]
        'phosphofructokinase'
PFK
                                                       'atp[c] + f6p[c] -> adp[c] + fdp[c] + h[c] '
        'glucose-6-phosphate isomerase'
                                                       'g6p[c] <=> f6p[c] '
PGI
PGK
        'phosphoglycerate kinase'
                                                       '3pg[c] + atp[c] <=> 13dpg[c] + adp[c] '
PGM
        'phosphoglycerate mutase'
                                                       '2pg[c] <=> 3pg[c] '
PPS
        'phosphoenolpyruvate synthase'
                                                       'atp[c] + h2o[c] + pyr[c] -> amp[c] + 2 h[c] +
                                                       'adp[c] + h[c] + pep[c] -> atp[c] + pyr[c] '
PYK
        'pyruvate kinase'
TPI
        'triose-phosphate isomerase'
                                                       'dhap[c] <=> q3p[c] '
```

It should be pointed out that although the reaction pyrvate dehydrogenase (PDH) is included in the glycolysis subsystem it is functionally a better fit in the "Glycoxylate Cycle, Gluconeogenesis, and Anapleurotic Reactions" subsystem, as described in section 4.E.

In addition to providing some atp[c] through substrate phosphorylation (PGK and PYK), the glycolysis pathway also proves a major source of nadh[c] (GAPD) that is used to power the electron transport chain. It also supplies several key precursors needed for the biosynthesis pathways. These precursors include: D-Glucose 6-phosphate (g6p[c]) a precursor for sugar nucleotides, D-Fructose 6-phosphate (f6p[c]) a precursor for amino sugars, glyceraldehyde 3-phosphate (g3p[c]) a precursor for phospholipids, 3-Phospho-D-glycerate (3pg[c]) a precursor for cysteine, glycine, and serine, phosphoenolpyruvate (pep[c]) a precursor for tyrosine, tryptophan and phenylalanine, and finally pyruvate (pyr[c]) the precursor for alanine, leucine, and valine [5]. These precursors and their location on the glycolysis pathway are illustrated in Figure 14.

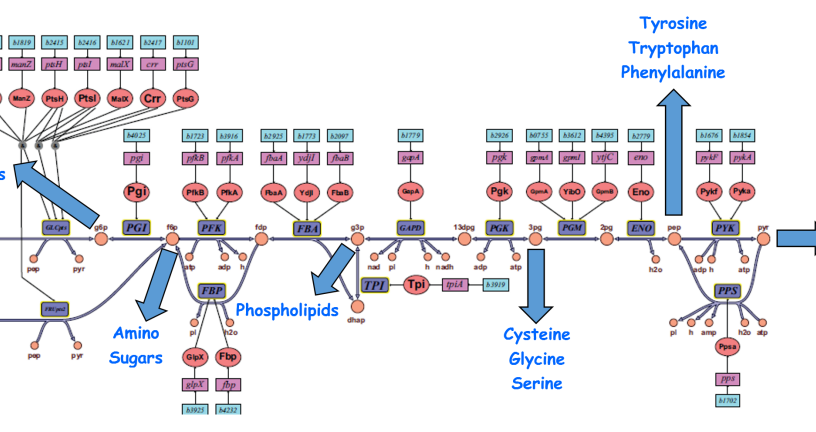


Figure 14. Precursors produced in the glycolysis pathway [3].

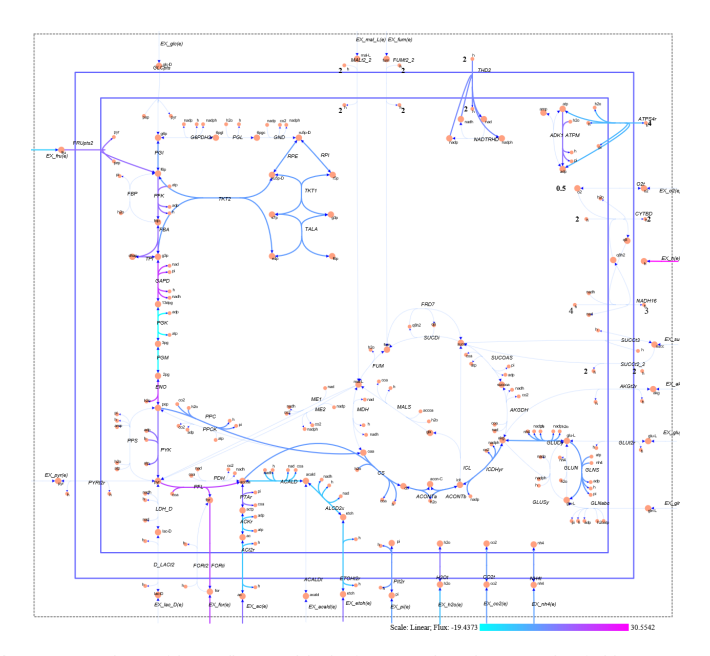
Visualizing the flux though the glycolysis pathways can be seen by using the draw package available with COBRA Toolbox. This is illustrated in the Matlab and COBRA toolbox code listed below for the case of anaerobic operation with fructose as the carbon source. [Timing: Seconds]

```
model = e_coli_core; % Starting with the original model
model = changeRxnBounds(model,'EX_glc(e)',0,'l');
model = changeRxnBounds(model,'EX_fru(e)',-10,'l');
model = changeRxnBounds(model,'EX_o2(e)',-0,'l');
model = changeObjective(model,'Biomass_Ecoli_core_w_GAM');
FBAsolution = optimizeCbModel(model,'max',0,0);

% Import E.coli core map and adjust parameters
map=readCbMap('ecoli_Textbook_ExportMap');
options.zeroFluxWidth = 0.1;
options.rxnDirMultiplier = 10;
drawFlux(map, model, FBAsolution.x, options);
```

Document Written

A screenshot of the saved "target.svg" file is shown in the Figure 15.



**Figure 15.** Network map of the *E.coli* core model using fructose as the carbon source (EX\_fru(e)  $\geq$  -10 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup>) in an anaerobic environment (EX\_o2(e)  $\geq$  0 mmol  $\cdot$  gDW<sup>-1</sup>  $\cdot$  hr<sup>-1</sup>).

Note that the fructose enters the network on the top left of the map. The detailed flux values for all the active reactions are shown below. [Timing: Seconds]

```
% Print the non-zero flux values
printFluxVector(model, FBAsolution.x, true)

ACALD -8.27946
ACKr -8.50359
ACONTa 0.228363
ACONTb 0.228363
ACt2r -8.50359
```

ATPM 8.39 ATPS4r -5.45205 Biomass\_Ecoli\_core\_w\_GAM 0.211663 CO2t 0.378178

-8.27946

ALCD2x

```
CS
     0.228363
ENO
    19.1207
ETOHt2r
         -8.27946
EX_ac(e)
          8.50359
           -0.378178
EX_co2(e)
EX_etoh(e)
            8.27946
         17.8047
EX_for(e)
EX_fru(e)
           -10
         30.5542
EX_h(e)
EX_h2o(e) -7.1158
EX_nh4(e) -1.15416
EX_pi(e)
          -0.778644
    9.78946
FBA
FORti 17.8047
FRUpts2 10
     19.4373
GAPD
GLNS
     0.0541222
       -1.10003
GLUDV
H2Ot.
      7.1158
       0.228363
ICDHyr
NH4t
       1.15416
      9.78946
PFK
     17.8047
PFL
PGI
      -0.0433909
PGK
      -19.4373
PGM
      -19.1207
PIt2r
       0.778644
PPC
      0.606541
PTAr
      8.50359
      8.40427
PYK
      -0.152143
RPE
RPI
      -0.152143
       -0.0378665
TALA
THD2
      3.62919
TKT1
      -0.0378665
TKT2
       -0.114277
TPI
      9.78946
```

The consumers of precursors formed in the glycolysis pathways can be found using the "surfNet" COBRA Toolbox function. An example looking for both the producers and consumers of "f6p[c]," a precursor for amino sugars is shown below. [Timing: Seconds]

surfNet(model, 'f6p[c]',0,FBAsolution.x,1,1)

Producing reactions with non-zero fluxes :

fru[e] + pep[c] -> f6p[c] + pyr[c]

```
Met #26     f6p[c], D-Fructose-6-phosphate, C6H1109P
Consuming reactions with non-zero fluxes :
    #13     Biomass_Ecoli_core_w_GAM (0.21166), Bd: 0 / 1000, Biomass Objective Function with GAM
    1.496     3pg[c] + 3.7478     accoa[c] + 59.81     atp[c] + 0.361     e4p[c] + 0.0709     f6p[c] + 0.129     g3p[c] + 0.205     g6p[
    #72     PFK (9.78946), Bd: 0 / 1000, phosphofructokinase
    atp[c] + f6p[c] -> adp[c] + fdp[c] + h[c]
    #74     PGI (-0.04339), Bd: -1000 / 1000, glucose-6-phosphate isomerase
    g6p[c] <=> f6p[c]
    #91     TALA (-0.03787), Bd: -1000 / 1000, transaldolase
    g3p[c] + s7p[c] <=> e4p[c] + f6p[c]
    #94     TKT2 (-0.11428), Bd: -1000 / 1000, transketolase
    e4p[c] + xu5p-D[c] <=> f6p[c] + g3p[c]
```

Show previous steps...

#45 FRUpts2 (10), Bd: 0 / 1000, Fructose transport via PEP:Pyr PTS (f6p generating)

Note that the majority of the f6p[c] flux is directed down the glycolysis pathway (PFK), a modest amount is directed to the pentose phosphate pathway (PGI, TALA, TKT2), with a small amount directed to the biomass function  $(0.211663 \times 0.0709 = 0.015)$  which represents the biosynthesis load of the precursors. A similar approach can be used to understand the producer/consumer relationships with the other glycolytic precursors.

Using the COBRA Toolbox, it is possible to create a table of reactions and their flux values for both glycolysis supported carbon sources, glucose and fructose. This is illustrated below. [Timing: Seconds]

```
% Starting with the original model
model = e_coli_core;
% Obtain the rxnIDs for the glycolysis pathway reactions
[tmp,glycolysis_rxnID] = ismember(glycolysisReactions,model.rxns);
% Glucose aerobic flux
FBAsolution = optimizeCbModel(model, 'max', 0, 0);
Glucose_Aerobic_Flux = FBAsolution.x(glycolysis_rxnID);
% Fructose aerobic flux
model = changeRxnBounds(model, 'EX_glc(e)',-0,'l');
model = changeRxnBounds(model, 'EX_fru(e)',-10,'1');
FBAsolution = optimizeCbModel(model, 'max', 0, 0);
Fructcose_Aerobic_Flux = FBAsolution.x(glycolysis_rxnID);
% Set anaerobic conditions
model = changeRxnBounds(model, 'EX_o2(e)',-0,'1');
% Glucose anaerobic flux
model = changeRxnBounds(model, 'EX_glc(e)',-10,'1');
FBAsolution = optimizeCbModel(model, 'max', 0, 0);
Glucose_Anaerobic_Flux = FBAsolution.x(glycolysis_rxnID);
% Fructose anaerobic flux
model = changeRxnBounds(model, 'EX_glc(e)',-0,'1');
model = changeRxnBounds(model, 'EX_fru(e)',-10,'1');
FBAsolution = optimizeCbModel(model, 'max', 0, 0);
Fructose_Anaerobic_Flux = FBAsolution.x(glycolysis_rxnID);
T = table(Glucose_Aerobic_Flux,Fructcose_Aerobic_Flux,Glucose_Anaerobic_Flux,...
    Fructose_Anaerobic_Flux,'RowNames',glycolysisReactions)
```

T =				
	Glucose_Aerobic_Flux	Fructcose_Aerobic_Flux	Glucose_Anaerobic_Flux	Fructose_Anaerobic
ENO	14.716	14.716	37.855	19.121
FBA	7.4774	7.4774	19.486	9.7895
FBP	0	0	0	0
GAPD	16.024	16.024	38.628	19.437
PDH	9.2825	9.2825	0	0
PFK	7.4774	7.4774	19.486	9.7895
PGI	4.8609	-5.1391	9.8942	-0.043391
PGK	-16.024	-16.024	-38.628	-19.437
PGM	-14.716	-14.716	-37.855	-19.121

PPS	0	0	0	0
PYK	1.7582	1.7582	16.108	8.4043
TPI	7.4774	7.4774	19.486	9.7895

From this table, it can be seen that in all four situations, the flux flows from the carbon source at the top left of the metabolic maps down the glycolysis pathway to form pyruvate in the lower right. In aerobic conditions, part of the flux is diverted to the G6PDH2r entrance to the pentose phosphate pathways. For the anaerobic case, the flux is only diverted to the lower half of the pentose phosphate pathway (TKT2) to produce the pentose phosphate pathway precursors. Also note that the flux through GAPD has almost doubled since the number of g3p[c] metabolites leaving the FBA and TPI reaction are double the number of fdp[c] metabolites entering FBA. This is possible since the output of FBA provides both a molecule of g3p[c] and a molecule of dhap[c]. The dhap[c] is rapidly converted to g3p[c] thus creating the effect of doubling the g3p[c] entering GAPD. A more detailed understanding of the fluxes through glycolysis using the COBRA toolbox is left as an exploration opportunity for the reader.