

Parameter estimation for iJO1366-based E. coli RBA model

November 2, 2018

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1 Introduction

Here we detail the procedure of parameterizing a Resource Balance Analysis (RBA)(2) model of model bacterium *Escherichia coli* based on the genome scale metabolic reconstruction *iJO1366* (7). We first explain how one can compute parameters related to compartment densities, those being the cytosolic density and total protein concentration, both in $[\frac{mmol.AA}{gCDW}]$. These parameters enable RBA models to give meaningful quantitative predictions of appropriate scale and unit. We then proceed to explain how one can compute the efficiencies of cellular process machineries, such as ribosomes, chaperones, secretion apparatus etc. We shortly explain the choice of some default values in the model. Finally, we elaborate an additional parameter modification of the model - the efficiency of the respiratory chain.

2 Compartment densities

RBA modeling paradigm incorporates the idea of limited cellular space in terms of inequality constraints putting an upper bound on the amount of macromolecules that can fit into each compartment. For the sake of simplicity in formulating these constraints, the size taken up by all the macromolecules in RBA is expressed in terms of the size of an average amino acid. Since the volume of each macromolecule is difficult to know or predict, we assume the size is proportional to their molecular weight.

Average amino acid molecular weight \bar{MW}_{AA} will depend on the percentage of each amino acid in the total amino acid pool. The amino acid composition data was taken from (6), and the weighter average amino acid molecular weight computed to be $\bar{MW}_{AA} = 108.3 \frac{g}{mol}$.

Average rRNA molecular weight \bar{MW}_{RNA} is computed from the composition of the ribosome, and is $\bar{MW}_{RNA} = 340 \frac{g}{mol}$.

Ribosome conversion factor $d_{r/AA}$ is a constant describing the molecular weight of all the nucleic acids in one ribosome, expressed in terms of number of average amino acids. Total number of nucleic acids in an *E. coli* is $N_{NA/r} = 4593$. This parameters is computed to be $d_{r/AA} = N_{NA/r} \times \frac{\bar{MW}_{RNA}}{\bar{MW}_{AA}} = 4593 \times 3.14 = 14422$.

2.1 Cytosolic density

Cytosolic density is a constant independent of the growth rate representing an idea that the concentration of macromolecules in the cytosol is kept constant to ensure proper diffusion of all cytosolic cellular components. It represents the space taken up by all the enzymes and ribosomes, thus taking into account the protein and rRNA content. To compute the cytosolic density, we consolidated three relevant datasets (1; 5; 6) and have thus chosen a growth rate which is present in all three datasets, that of $1[h^{-1}]$. Cytosolic density can be expressed as a sum of portion of protein that is assigned to cytosol and

of the RNA content of ribosomes converted to average amino acid:

$$D_{cyt} = p_{cyt}(\mu)P_{tot}(\mu) + d_{r/AA}R(\mu) \quad \left[\frac{mmol.AA}{gCDW}\right] \quad (1)$$

Fraction of protein assigned to cytosol $p_{cyt}(\mu)$ is estimated using the package `RBapy.estim` from proteomics data (8) as a linear function of the growth rate:

$$p_{cyt} = 0.04\mu + 0.73 \quad (2)$$

What remains to be computed are the total amino acid and ribosome concentration. To compute the total amino acid concentration in $\left[\frac{mmol}{gCDW}\right]$, we use the comprehensive dataset of (1).

$$C_{AA/CDW} = N_{AA/CDW} \left[\frac{\#AA}{\mu gCDW}\right] \times \frac{1}{N_a} [mol] \times 10^6 \times 10^3 \quad \left[\frac{mmol.AA}{gCDW}\right] \quad (3)$$

N_a is the Avogadro constant, and $N_{AA/CDW}$ is the number of amino acids in a μg of cell dry weight. For ease of comparison, the abbreviations used here to express the formula for total amino acid count are the same as used in the (1). We compute $N_{AA/CDW}$ in the following way:

$$N_{AA/CDW} = \frac{P_M}{\frac{M_C(\mu g)}{M_C}} \quad \left[\frac{\#AA}{\mu gCDW}\right] \quad (4)$$

We finally obtain a value for total concentration of amino acids:

$$C_{AA/CDW} = 4.98 \quad \left[\frac{mmol.AA}{gCDW}\right] \quad (5)$$

for the growth rate of $\mu = 1[h^{-1}]$.

We compute the ribosome concentration by assuming that the cell has as many ribosomes as needed to translate the flux of total protein at steady state:

$$R = \frac{\mu C_{AA/CDW}}{k_T \times p_{R_a}} \quad (6)$$

where k_T is the ribosome efficiency in $\left[\frac{AA}{h}\right]$ and p_{R_a} is the percentage of active ribosomes expressed as $e^{\mu\tau}$, τ being the maturation time of ribosomes, set to be $5min$ (4). The ribosome concentration thus computed is:

$$R = 7.24 \times 10^{-5} \quad \left[\frac{mmol}{gCDW}\right] \quad (7)$$

Now we can compute the cytosolic density for $\mu = 1[h^{-1}]$ from equation 1 to be:

$$D_{cyt} = 4.89 \quad \left[\frac{mmol.AA}{gCDW}\right] \quad (8)$$

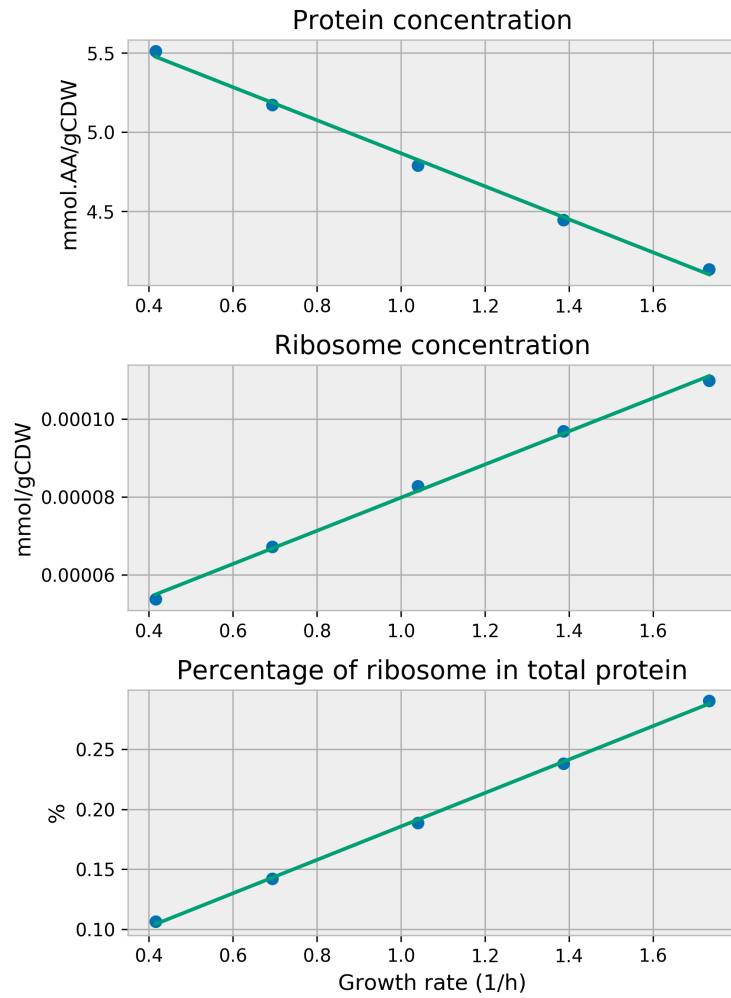


Figure 1: Computed values for total amino acid concentration, ribosome concentration and percentage of ribosome in total protein.

2.2 Total amino acid concentration

We compute total cytosolic concentration as described in the supplementary Text E1 in (3), by solving the following system of equations:

$$\mu C_{AA/CDW} = k_T e^{\mu\tau} R \quad (9)$$

$$p_{cyt} C_{AA/CDW} + d_{r/AA} R = D_{cyt} \quad (10)$$

for growth rates available in (1). We used the computed values to estimate the linear dependency of amino acid concentration to the growth rate:

$$C_{AA/CDW} = -1.04\mu + 5.91 \quad (11)$$

The results of the estimation can be seen on figure 1.

3 Machinery efficiencies

The efficiencies of process machineries were estimated using proteomics datasets from (8) for growth on 12 different carbon sources and 1 supplemented with 20 amino acids. We basically estimate the total amino acid concentration flux that needs to be processed by a particular machinery, and divide it by the abundance of the machinery, obtaining process machinery efficiency in $[\frac{\#AA}{h}]$.

3.1 Folding

We consider two chaperoning systems (GroEL/S and DnaJK) as one folding machinery that needs to fold 10% of all protein. We estimate the total concentration of amino acids that needs to be folded per unit time using the estimation obtained in the section 2.

$$\nu_{P_{fold}} = \mu \times 0.1 \times P_{tot}(\mu) = 0.31 \quad \left[\frac{mmol.AA}{gCDW}\right] \quad (12)$$

The total number of amino acids in the chaperone complex consisting of all subunits in their correct stoichiometries (tig, dnaJ, dnaK, groL, groS, grpE) is $N_{AA/ch} = 10829$. Number of measured amino acids of the same complex is $N_{AA/ch/mes} = 3.8 \times 10^7$. Efficiency of the chaperone complex becomes:

$$k_{CH} = \frac{\nu_{P_{fold}}}{\frac{N_{AA/ch}}{N_{AA/ch/mes}}} \quad (13)$$

The folding efficiency as a linear function of the growth rate is:

$$k_{CH}(\mu) = 7.2\mu + 1.59 \quad (14)$$

and can be seen on Figure 2a.

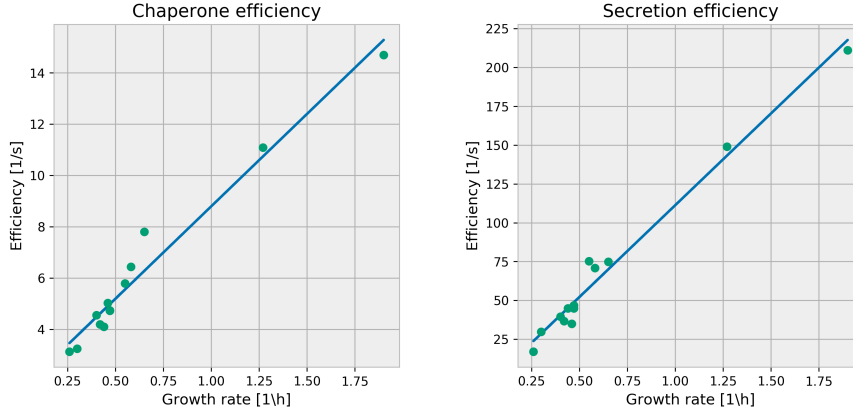


Figure 2: Efficiency of cellular process machineries (folding and secretion) estimated from proteomics data

3.2 Secretion

In the case of secretion, we say that all non-cytosolic protein needs to be secreted. Thus our concentration of amino acids to be secreted per unit time will be:

$$\nu_{P_{sec}} = \mu \times (1 - p_{cyt}(\mu)) \times P_{tot}(\mu) \quad (15)$$

The rest of the procedure is the same as in the case of folding, and the final linear relation between the growth rate and secretion efficiency is

$$k_{SEC}(\mu) = 118.23\mu - 6.94 \quad (16)$$

and can be seen on Figure 2b.

4 Default values

5 Further modifications

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