# Toward a Genome Scale Dynamic Model of Cell-Free Protein Synthesis in *Escherichia coli*

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# **Abstract**

Cell-free protein expression systems have become widely used in systems and synthetic biology. In this study, we developed an ensemble of dynamic E. coli cell-free protein synthesis (CFPS) models. Model parameters were estimated from a training dataset for the cell-free production of a protein product, chloramphenicol acetyltransferase (CAT). The dataset consisted of measurements of glucose, organic acids, energy species, amino acids, and CAT. The ensemble accurately predicted these measurements, especially those of the central carbon metabolism. We then used the trained model to evaluate the optimality of protein production. CAT was produced with a carbon yield of 7% and an energy efficiency of 5%, suggesting that the process could be further optimized. Reaction group knockouts showed that protein productivity and the metabolism as a whole depend most on oxidative phosphorylation and glycolysis and gluconeogenesis. Amino acid biosynthesis is also important for productivity, while the overflow metabolism and TCA cycle affect the overall system state. In addition, CAT production was robust to allosteric control, as was most of the network, with the exception of the organic acids in central carbon metabolism. This study is the first to use kinetic modeling to predict dynamic protein production in a cell-free *E. coli* system, and should provide a foundation for genome scale, dynamic modeling of cell-free *E. coli* protein synthesis.

**Keywords:** Biochemical engineering, systems biology, *E. coli*, cell-free protein synthesis, kinetic modeling

# Introduction

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Cell-free systems offer many advantages for the study, manipulation and modeling of metabolism compared to *in vivo* processes. Central amongst these is direct access to metabolites and the biosynthetic machinery without the interference of a cell wall, or complications associated with cell growth. This allows us to interrogate the chemical environment while the biosynthetic machinery is operating, potentially at a fine time resolution. Cell-free protein synthesis (CFPS) systems are arguably the most prominent examples of cell-free systems used today [1]. However, CFPS is not new; CFPS in crude *E. coli* extracts has been used since the 1960s to explore fundamentally important biological mechanisms [2, 3]. Today, cell-free systems are used in a variety of applications ranging from therapeutic protein production [4] to synthetic biology [5, 6]. However, if CFPS is to become a mainstream technology for applications such as point of care manufacturing, we must first understand the performance limits of these systems. One tool we can use to achieve this understanding is mathematical modeling.

Mathematical modeling has long contributed to our understanding of metabolism. Dec-15 ades before the genomics revolution, mechanistically structured metabolic models arose 16 from the desire to predict microbial phenotypes resulting from changes in intracellular or 17 extracellular states [7]. The single cell E. coli models of Shuler and coworkers pioneered 18 the construction of large-scale, dynamic metabolic models that incorporated multiple regulated catabolic and anabolic pathways constrained by experimentally determined kinetic 20 parameters [8]. Shuler and coworkers generated many single cell kinetic models, in-21 cluding single cell models of eukaryotes [9, 10], minimal cell architectures [11], as well as DNA sequence based whole-cell models of E. coli [12]. However, cell-free genome scale kinetic models of industrially important organisms such as E. coli have yet to be constructed.

In this study, we developed an ensemble of kinetic cell-free protein synthesis (CFPS)

models using dynamic metabolite measurements in an E. coli cell-free extract. Model parameters were estimated from measurements of glucose, organic acids, energy species, amino acids, and the protein product, chloramphenicol acetyltransferase (CAT). The parameter estimation problem was constrained by characteristic values for model param-30 eters from literature. The ensemble of parameter sets described the training data with 31 a median cost that was more than two orders of magnitude smaller than random sets 32 constructed using the literature parameter constraints. We then used the ensemble of 33 kinetic models to analyze the optimality of the CFPS system, and the pathways most im-34 portant to CAT production. We calculated that CAT was produced with a carbon yield of 35 7% and an energy efficiency of 5%, suggesting that much of the resources for protein synthesis were diverted to non-productive pathways. By knocking out metabolic enzymes 37 in groups, we showed that the system metabolism and protein production depend most 38 on oxidative phosphorylation and glycolysis/gluconeogenesis. Taken together, we have integrated traditional kinetics with a logical rule-based description of allosteric control to simulate a comprehensive CFPS dataset. This study provides a foundation for genome scale, dynamic modeling of cell-free *E. coli* protein synthesis.

# 43 Results

The cell-free E. coli metabolic network was constructed by removing growth associated reactions from the iAF1260 reconstruction of K-12 MG1655 E. coli [13], and by adding reactions describing chloramphenicol acetyltransferase (CAT) biosynthesis, a model pro-46 tein for which there exists a comprehensive training dataset [14]. In addition, reactions 47 that were knocked out from the cell extract preparation were removed from the network ( $\triangle$ speA,  $\triangle$ tnaA,  $\triangle$ sdaA,  $\triangle$ sdaB,  $\triangle$ gshA,  $\triangle$ tonA,  $\triangle$ endA). The CFPS model equations were formulated using the hybrid cell-free modeling framework of Wayman et al. [15]. An initial ensemble of model parameter sets (N > 30,000) was estimated from measurements of glucose, CAT, organic acids (pyruvate, lactate, acetate, succinate, malate), energy species (A(x)P, G(x)P, C(x)P, U(x)P), and 18 of the 20 proteinogenic amino acids 53 using a constrained Markov Chain Monte Carlo (MCMC) approach. A final ensemble of parameter sets (N = 100) was constructed by selecting the sets with the lowest errors, the 55 lowest of which was defined as the best-fit set. Parameter sets in the final ensemble had a mean Pearson correlation coefficient of 0.77; thus, an accurate yet diverse ensemble 57 was created. 58

The ensemble of kinetic CFPS models captured the time evolution of CAT biosynthe-59 sis. Central carbon metabolites (Fig. 1, top), energy species (Fig. 2), and amino acids 60 (Fig. 3) were captured by the ensemble and the best-fit set. The constrained MCMC 61 approach estimated parameter sets with a median error more than two orders of mag-62 nitude less than random parameter sets generated within the same parameter bounds 63 (Fig. 4); thus, we have confidence in the predictive capability of the estimated param-64 eters. The model captured the biphasic CAT production: during the first hour glucose 65 powers production, and CAT is produced at 10  $\mu$ M/h; subsequently, pyruvate and lactate reserves are consumed to power metabolism, and CAT is produced less efficiently at 3  $\mu$ M/h. Allosteric control was important to central carbon metabolism, especially pyruvate, acetate, and succinate (Fig. 1, bottom). The difference between the allosteric control and no-control cases is mostly seen in the second phase of CAT production, after glucose is exhausted. Specifically, there is an increase in the consumption of pyruvate, succinate, and malate, as well as the accumulation of acetate. Taken together, we produced an ensemble of kinetic models that was consistent with time series measurements of the production of a model protein. Although the ensemble described the experimental data, it was unclear which kinetic parameters most influenced CAT production, and whether the performance of the CFPS reaction was optimal.

To better understand the effect of network reactions on system performance we con-77 ducted a group knockout analysis (Fig. 5). The network was divided into 19 groups of 78 reactions, spanning central carbon metabolism, energetics, and amino acid biosynthesis. 79 The enzymes in each of these groups were knocked out, and the resulting change in 80 productivity and system state were recorded. Then each pair of groups was knocked out 81 to determine pairwise effects. These were summed with the first-order effect to obtain a total-order coefficient for each group for the change in productivity and system state. Gly-83 colysis/gluconeogenesis and oxidative phosphorylation were seen to have the greatest effect on both productivity and system state. This supports previous findings that oxidative phosphorylation does occur in a cell-free system [1]. Jewett and coworkers observed a decrease in CAT yield, ranging from 1.5-fold to 4-fold, when knocking out oxidative phosphorylation reactions in cell-free (the Cytomim platform) with both pyruvate and glutamate as substrates. CAT productivity was also affected by two sectors of amino acid biosynthesis: alanine/aspartate/asparagine, and glutamate/glutamine. This is likely be-90 cause aspartate, glutamate, and glutamine are key reactants in the biosynthesis of many 91 other amino acids, all of which are required for CAT synthesis. Meanwhile, the TCA cycle and overflow metabolism (which includes acetyl-coA/acetate reactions and the intercon-93 version of pyruvate and lactate) have a significant effect on the system state, defined as all species for which ata exist. These reactions directly impact key species in the system state: succinate and malate in the TCA cycle, and acetate, pyruvate, and lactate in the overflow metabolism.

To understand whether the CFPS performance was optimal, we calculated the carbon 98 yield and energy efficiency of the two phases of CAT production for the best-fit set (Fig. 6). 99 The equivalent ATP numbers for glucose, amino acids, and organic acids were calculated 100 from the network stoichiometry; for example, 21 molecules of ATP shold be generated 101 from one glucose molecule if the optimal path through glycolysis, the TCA cycle, and ox-102 idative phosphorylation is taken. The ATP number for CAT was simply equal to the cost 103 of transcription and translation. During the first phase, with glucose as the substrate, CAT 104 was produced with a carbon yield of 5% and an energy efficiency of 3%. Of the remain-105 ing carbon, 4% is accounted for by the accumulation of amino acids (alanine, isoleucine, 106 glutamine, proline, and tyrosine), 39% by organic acid accumulation (pyruvate, lactate, 107 acetate, succinate, and malate), and 52% by the accumulation of other byproducts, pri-108 marily glycolytic intermediates and carbon dioxide. The breakdown is very similar with 109 the energy efficiency: 4% amino acids, 41% organic acids, and 52% other byproducts. 110 This suggests that for glucose-driven production, the best ways to improve efficiency are by ensuring that the flux through glycolysis and the TCA cycle is constant throughout, preventing any accumulation of intermediates. However, the accumulated organic acids (except acetate) were then utilized as substrates in the second phase once glucose was 114 depleted. Although succinate and malate were consumed in the second phase, they only 115 account for 14% of the substrate consumption; thus, it may be reasonable to consider 116 this as pyruvate-driven production. Interestingly, this mode of protein production showed 117 higher carbon yield and energy efficiency: 6% in each case. Of the remaining carbon, 118 11% went to amino acids (alanine, glutamine, proline, and serine), 31% went to organic 119 acids (only acetate in this case), and 52% went to other byproducts. The remainder of the 120

energy was accounted by 21% in amino acids, 49% in organic acids, and 24% in other byproducts. While the efficiency of production is higher for the pyruvate-driven phase, it is still relatively low, and could be improved by utilizing more of the resources that are generated (amino aicds and glycoltyic/TCA intermediates).

# Discussion

In this study we present an ensemble of *E. coli* cell-free protein synthesis (CFPS) models that accurately predict a comprehensive CFPS dataset of glucose, CAT, central carbon metabolites, energy species, and amino acid measurements. We used the hybrid cell-free modeling approach of Wayman and coworkers, which integrates traditional kinetic modeling with a logic-based description of allosteric regulation. Our ensemble of models accurately predicts dynamic experimental measurements of central carbon metabolism, energy species, and amino acids over 100 times better than random sets in the same region of parameter space. CFPS was seen to be biphasic, relying on glucose during the first hour and pyruvate and lactate afterward. Allosteric control was essential to the maintenance of the network, specifically central carbon metabolism. Without it, pyruvate, succinate, and malate were consumed more quickly following glucose exhaustion to power downstream reactions and ultimately CAT synthesis. Interestingly, CAT production is virtually unaffected; this is because the amino acids and energy species that are reactants for CAT synthesis were also not affected by allosteric control.

Having captured the experimental data, we investigated if CAT yield and CFPS performance could be further improved. We showed that the model predicts CAT production with a carbon yield of 5% and energy efficiency of 3% under glucose, and a carbon yield and energy efficiency of 6% under pyruvate. The accumulation of glycolytic intermediates and byproducts such as acetate and carbon dioxide were responsible for this sub-optimal performance. If fluxes could be balanced such that intermediates were fully utilized, CAT production would increase. Knocking out sections of network metabolism revealed that glycolysis/gluconeogenesis and oxidative phosphorylation were the most important to CAT production and the system as a whole. Productivity was also heavily dependent on the synthesis reactions of alanine, aspartate, asparagine, glutamate, and glutamine, while TCA cycle and overflow reactions affected the system state. Taken together, these

findings represent the first dynamic model of E. coli cell-free protein synthesis, and an important step toward a functional genome scale description. 152

We present an ensemble of models that quantitatively described the system behavior 153 of cell-free metabolism and production of CAT. Experimental observations of the metabo-154 lites validate the structure of the model and the estimation of kinetic parameters. This is 155 important in applying metabolic engineering principles to rationally design cell-free pro-156 duction processes and predicting the redirection of carbon fluxes to product-forming path-157 ways. In analyzing the effect of reaction groups on CAT production and the system state, 158 the regions of metabolism associated with substrate utilization and energy generation 159 are the most important. Oxidative phosphorylation is vital, since it provides most of the 160 energetic needs of CFPS. While it is unknown how active oxidative phosphorylation is compared to that of in vivo systems, our modeling approach suggests its importance to 162 CFPS performance and protein yield. However, the biphasic operation of CFPS high-163 lights the ability of the system to respond to an absence of glucose. During the first 164 phase, there is an accumulation of central carbon metabolites with the majority of flux 165 going toward acetate and some toward pyruvate, lactate, succinate and malate. While acetate continued to accumulate as a byproduct, the other organic acids were consumed as secondary substrates after glucose is no longer available. Glutamate also served as a substrate throughout both phases, powering amino acid synthesis. These results confirm experimental findings that CAT production can be sustained by other substrates in the ab-170 sence of glucose, providing alternative strategies to optimize CFPS performance. While CAT synthesis can be powered by other substrates, the productivity is significantly lower 172 (3  $\mu$ M/h, as opposed to 10  $\mu$ M/h). This is in accordance with literature, where pyruvate 173 provided a relatively slow but continuous supply of ATP [16]. Taken together, this shows 174 CFPS can be designed towards a specified application, either requiring a slow stable 175 energy source or faster production.

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This work represents the first dynamic model of *E. coli* cell-free protein synthesis. 177 We apply a hybrid modeling framework to capture an experimental dataset for production 178 of a test protein, and identify system limitations and areas of improvement for produc-179 tion efficiency. This work could be extended through further experimentation to gain a 180 deeper understanding of model performance under a variety of conditions. Specifically, 181 CAT production performed in the absence of amino acids could inform the system's ability 182 to manufacture them, while experimentation in the absence of glucose or oxygen could 183 shed light on the importance of those substrates. In addition, the approach should be ex-184 tended to other protein products. CAT is only a test protein used for model identification; 185 the modeling framework, and to some extent the parameter values, should be protein ag-186 nostic. An important extension of this study would be to apply its insights to other protein 187 applications, where possible.

# Materials and Methods

Formulation and solution of the model equations. We used ordinary differential equations (ODEs) to model the time evolution of metabolite  $(x_i)$  and scaled enzyme abundance  $(\epsilon_i)$  in hypothetical cell-free metabolic networks:

$$\frac{dx_i}{dt} = \sum_{j=1}^{\mathcal{R}} \sigma_{ij} r_j(\mathbf{x}, \epsilon, \mathbf{k}) \qquad i = 1, 2, \dots, \mathcal{M}$$
 (1)

$$\frac{d\epsilon_i}{dt} = -\lambda_i \epsilon_i \qquad i = 1, 2, \dots, \mathcal{E}$$
 (2)

where  $\mathcal{R}$  denotes the number of reactions,  $\mathcal{M}$  denotes the number of metabolites and 193  $\mathcal{E}$  denotes the number of enzymes in the model. The quantity  $r_i(\mathbf{x}, \epsilon, \mathbf{k})$  denotes the 194 rate of reaction j. Typically, reaction j is a non-linear function of metabolite and enzyme 195 abundance, as well as unknown kinetic parameters  $\mathbf{k}$  ( $\mathcal{K} \times 1$ ). The quantity  $\sigma_{ij}$  denotes 196 the stoichiometric coefficient for species i in reaction j. If  $\sigma_{ij} > 0$ , metabolite i is produced 197 by reaction j. Conversely, if  $\sigma_{ij} < 0$ , metabolite i is consumed by reaction j, while  $\sigma_{ij} = 0$ 198 indicates metabolite i is not connected with reaction j. Lastly,  $\lambda_i$  denotes the scaled 199 enzyme activity decay constant. The system material balances were subject to the initial 200 conditions  $\mathbf{x}(t_o) = \mathbf{x}_o$  and  $\epsilon(t_o) = 1$  (initially we have 100% cell-free enzyme abundance). 201 The reaction rate was written as the product of a kinetic term  $(\bar{r}_j)$  and a control term 202  $(v_i)$ ,  $r_i(\mathbf{x}, \mathbf{k}) = \bar{r}_i v_i$ . We used multiple saturation kinetics to model the reaction term  $\bar{r}_i$ :

$$\bar{r}_j = V_j^{max} \epsilon_i \prod_{s \in m_j^-} \frac{x_s}{K_{js} + x_s} \tag{3}$$

where  $V_j^{max}$  denotes the maximum rate for reaction j,  $\epsilon_i$  denotes the scaled enzyme activity which catalyzes reaction j,  $K_{js}$  denotes the saturation constant for species s, in reaction j and  $m_j^-$  denotes the set of *reactants* for reaction j. Transcription was modeled

as first-order in RNA polymerase, since it acts like an enzyme:

$$\bar{r}^{TX} = k_{cat}^{TX} \cdot \text{RNAP}\left(\frac{\text{GENE}}{K_{GENE}^{TX} + \text{GENE}}\right) \prod_{s \in m_{TX}^{-}} \frac{x_s}{K_s^{TX} + x_s}$$
 (4)

where  $k_{cat}^{TX}$  denotes the maximum transcription rate, RNAP denotes the RNA polymerase concentration, GENE denotes the gene concentration,  $K_{GENE}^{TX}$  denotes the gene saturation constant,  $K_s^{TX}$  denotes the saturation constant for species s, and  $m_{TX}^{-}$  denotes the set of *reactants* for transcription: ATP, GTP, CTP, UTP, and water. While transcription is modeled as saturating with respect to gene concentration, the gene is not considered a reactant in the stoichiometry as it is not consumed. Translation was modeled as first-order in ribosome, since it acts like an enzyme:

$$\bar{r}^{TL} = k_{cat}^{TL} \cdot \text{RIBO}\left(\frac{\text{mRNA}}{K_{mRNA}^{TL} + \text{mRNA}}\right) \prod_{s \in m_{TL}^{-}} \frac{x_s}{K_s^{TL} + x_s}$$
 (5)

where  $k_{cat}^{TL}$  denotes the maximum translation rate, RIBO denotes the ribosome concentration, mRNA denotes the transcript concentration,  $K_{mRNA}^{TL}$  denotes the transcript saturation constant,  $K_s^{TL}$  denotes the saturation constant for species s, and  $m_{TL}^-$  denotes the set of reactants for translation: GTP, water, and the 20 species representing tRNA charged with amino acids. While translation is modeled as saturating with respect to transcript concentration, the transcript is not considered a reactant in the stoichiometry as it is not consumed. Transcript degradation was modeled as first-order in transcript:

$$\bar{r}_{deg}^{mRNA} = k_{deg}^{mRNA} \cdot \text{mRNA}$$
 (6)

where  $k_{deg}^{mRNA}$  denotes the transcript degradation rate constant.

The control term  $0 \leq v_j \leq 1$  depended upon the combination of factors which in-

fluenced rate process j. For each rate, we used a rule-based approach to select from competing control factors. If rate j was influenced by  $1,\ldots,m$  factors, we modeled this relationship as  $v_j = \mathcal{I}_j\left(f_{1j}\left(\cdot\right),\ldots,f_{mj}\left(\cdot\right)\right)$  where  $0 \leq f_{ij}\left(\cdot\right) \leq 1$  denotes a transfer function quantifying the influence of factor i on rate j. The function  $\mathcal{I}_j\left(\cdot\right)$  is an integration rule which maps the output of regulatory transfer functions into a control variable. We used hill-like transfer functions and  $\mathcal{I}_j \in \{min, max\}$  in this study [15].

We included 17 allosteric regulation terms, taken from literature, in the CFPS model. PEP was modeled as an inhibitor for phosphofructokinase [17, 18], PEP carboxykinase [17], PEP synthetase [17, 19], isocitrate dehydrogenase [17, 20], and isocitrate lyase/malate synthase [17, 20, 21], and as an activator for fructose-biphosphatase [17, 22–24]. AKG was modeled as an inhibitor for citrate synthase [17, 25, 26] and isocitrate lyase/malate synthase [17, 21]. 3PG was modeled as an inhibitor for isocitrate lyase/malate synthase [17, 21]. FDP was modeled as an activator for pyruvate kinase [17, 27] and PEP carboxylase [17, 28]. Pyruvate was modeled as an inhibitor for pyruvate dehydrogenase [17, 29, 30] and as an activator for lactate dehydrogenase [31]. Acetyl CoA was modeled as an inhibitor for malate dehydrogenase [17].

Estimation of kinetic model parameters. We estimated an ensemble of diverse parameter sets using a constrained Markov Chain Monte Carlo (MCMC) random walk strategy. Starting from a single best-fit parameter set estimated by inspection and literature, we calculated the cost function, equal to the sum-squared-error between experimental data and model predictions:

$$cost = \sum_{i=1}^{\mathcal{D}} \left[ \frac{w_i}{\mathcal{Y}_i^2} \sum_{j=1}^{\mathcal{T}_i} \left( y_{ij} - x_i |_{t(j)} \right)^2 \right] \tag{7}$$

where  $\mathcal{D}$  denotes the number of datasets ( $\mathcal{D}=$  37),  $w_i$  denotes the weight of the  $i^{th}$  dataset,  $\mathcal{T}_i$  denotes the number of timepoints in the  $i^{th}$  dataset, t(j) denotes the  $j^{th}$  time-

point,  $y_{ij}$  denotes the measurement value of the  $i^{th}$  dataset at the  $j^{th}$  timepoint, and  $x_i|_{t(j)}$  denotes the simulated value of the metabolite corresponding to the  $i^{th}$  dataset, interpolated to the  $j^{th}$  timepoint. Lastly, the cost calculation was scaled by the maximum experimental value in the  $i^{th}$  dataset,  $\mathcal{Y}_i = \max_j{(y_{ij})}$ . We then perturbed each model parameter between an upper and lower bound that varied by parameter type:

$$k_i^{new} = \min\left(\max\left(k_i \cdot \exp(a \cdot r_i), l_i\right), u_i\right) \qquad i = 1, 2, \dots, \mathcal{P}$$
(8)

where  $\mathcal{P}$  denotes the number of parameters ( $\mathcal{P}=815$ ), which includes 163 maximum reaction rates ( $V^{max}$ ), 163 enzyme activity decay constants, 455 saturation constants ( $K_{js}$ ), and 34 control parameters,  $k_i^{new}$  denotes the new value of the  $i^{th}$  parameter,  $k_i$  denotes the current value of the  $i^{th}$  parameter, a denotes a distribution variance,  $r_i$  denotes a random sample from the normal distribution,  $l_i$  denotes the lower bound for that parameter type, and  $u_i$  denotes the upper bound for that parameter type.

Model parameters were constrained by literature [32]. The rate maximum for CAT transcription was calculated as  $123 \, h^{-1}$ :

$$k_{cat}^{TX} = \left(\frac{k_{TX}}{l_{mRNA}}\right) \cdot P \tag{9}$$

where  $k_{TX}$  denotes the average mRNA elongation rate (25 nt/s, [33]),  $l_{mRNA}$  denotes the CAT mRNA length (660 nt), and P denotes the level of promoter activity (estimated at 0.9). The rate maximum for CAT translation was calculated as 247 h<sup>-1</sup>:

$$k_{cat}^{TL} = \left(\frac{k_{TL}}{l_{protein}}\right) \cdot K_P \tag{10}$$

where  $k_{TL}$  denotes the peptide chain elongation rate (1.5 aa/s, [33]),  $l_{protein}$  denotes the CAT protein length (219 aa), and  $K_P$  denotes the polysome amplification constant (esti-

mated at 10). The transcript degradation rate constant was calculated as 5.2  $h^{-1}$ :

$$k_{deg}^{mRNA} = \frac{log(2)}{t_{1/2}} \tag{11}$$

where  $t_{1/2}$  denotes the mRNA degradation time (8 min, BNID 106253). Transcription, translation, mRNA degradation, and tRNA charging were bounded within two orders of magnitude of their calculated values. A characteristic cell-free enzyme concentration of 167 nM was calculated by diluting the one-tenth maximal amount lacZ (5  $\mu$ M, BNID 100735) by a cell-free dilution factor of 30. This enzyme level was then used to calculate rate maxima from turnover numbers for various enzymes from literature. Rate maxima were bounded within one order of magnitude of the calculated value where available; all other rate maxima were bounded within two orders of magnitude of the geometric mean of the available values.

The median maximum reaction rate was 7.8 mM/h; assuming a total cell-free enzyme concentration of 167 nM, this corresponds to a median catalytic rate of  $0.08 \, \text{s}^{-1}$  across the ensemble. Enzyme activity decay constants were bounded between 0 and 1 h<sup>-1</sup>, corresponding to half lives of 42 minutes and infinity; median = 156 h. Saturation constants were bounded between 0.0001 and 10 mM; median = 1.0 mM. Control parameters (gains and orders) were bounded between 0.1 and 10 (dimensionless); median = 0.74. For each newly generated parameter set, we re-solved the balance equations and calculated the cost function. All sets with a lower cost (and some with higher cost) were accepted into the ensemble. After generating over 30,000 sets, N = 100 sets with minimal error were selected for the final ensemble. The final ensemble had a mean Pearson correlation coefficient of 0.77.

Comparison against random ensemble. A random ensemble of 100 parameter sets was generated from within the same parameter bounds as the trained ensemble. Sets

were sampled using a Monte Carlo approach: each parameter was taken from a uniform distribution constructed between its upper and lower bounds. The model equations were then solved and the cost function was calculated in terms of the 37 separate experimental datasets. The random ensemble had a log median error of 0.80 across the datasets, as compared with a log median error of -1.43 for the trained ensemble. Thus, the trained ensemble fits the dataset over one hundred times better than a random ensemble generated within the same bounds.

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**Group knockouts.** The network was divided into 19 groups: glycolysis/gluconeogenesis, 295 pentose phosphate, Entner-Doudoroff, TCA cycle, oxidative phosphorylation, cofactor re-296 actions, anaplerotic/glyoxylate reactions, overflow metabolism, folate synthesis, purine/ 297 pyrimidine reactions, alanine/aspartate/asparagine synthesis, glutamate/glutamine synthesis, arginine/proline synthesis, glycine/serine synthesis, cysteine/methionine synthesis, threonine/lysine synthesis, histidine synthesis, tyrosine/tryptophan/phenylalanine syn-300 thesis, and valine/leucine/isoleucine synthesis. Each group of reactions was turned off 301 individually, and then in pairs, and the model equations were re-solved. The CAT pro-302 ductivity was calculated and compared to that of the best-fit set. The absolute difference 303 in productivity was recorded for each first-order knockout (diagonal elements) and each 304 pairwise knockout, and a total-order coefficient was calculated by summing the first-order 305 effect with all pairwise effects. Total-order coefficients were then normalized to fit within 306 the same colorbar range as the first-order and pairwise effects. The system state was 307 also calculated for each simulation, defined as the model predictions for all species for 308 which data exist. 309

Calculation of carbon yield. The CAT carbon yield  $(Y_C^{CAT})$  was calculated as the ratio of carbon produced as CAT divided by the carbon consumed as reactants:

$$Y_C^{CAT} = \frac{\Delta \text{CAT} \cdot C_{CAT}}{\sum_{i=1}^{R} \Delta m_i \cdot C_i}$$
(12)

where  $\Delta \mathtt{CAT}$  denotes the abundance of CAT produced,  $C_{CAT}$  denotes carbon number of 312 CAT,  ${\cal R}$  denotes the number of reactants,  $\Delta m_i$  denotes the amount of the  $i^{th}$  reactant 313 consumed, and  $C_i$  denotes the carbon number of the  $i^{th}$  reactant. This analysis was ex-314 tended to the accumulation of amino acids, organic acids, and other byproducts, to create 315 a complete carbon balance through the network. The first phase of CAT production was 316 defined as t = 0 h to t = 1.11 h, the time at which glucose concentration falls below 0.1 nM. 317 In the first phase, amino acid accumulation consisted of alanine, isoleucine, glutamine, 318 proline, and tyrosine, while organic acid accumulation consisted of pyruvate, lactate, ac-319 etate, succinate, and malate. Glucose and the amino acids that did not accumulate were 320 considered reactants. In the second phase, amino acid accumulation consisted of ala-321 nine, glutamine, proline, and serine, while organic acid accumulation consisted of acetate 322 only. Pyruvate, lactate, succinate, malate, and the amino acids that did not accumulate 323 were considered reactants. 324

Calculation of energy efficiency. Energy efficiency was calculated as the ratio of CAT production to substrate consumption, both in terms of equivalent ATP molecules:

$$\text{Efficiency} = \frac{\Delta \text{CAT} \cdot (2 \cdot (\text{ATP}_{\text{TX}} + \text{CTP}_{\text{TX}} + \text{GTP}_{\text{TX}} + \text{UTP}_{\text{TX}}) + 2 \cdot \text{ATP}_{\text{TL}} + \text{GTP}_{\text{TL}})}{\sum_{i=1}^{\mathcal{R}} \Delta m_i \cdot \text{ATP}_i }$$
 (13)

where ATP<sub>TX</sub>, CTP<sub>TX</sub>, GTP<sub>TX</sub>, UTP<sub>TX</sub> denote the stoichiometric coefficients of each energy species for CAT transcription, ATP<sub>TL</sub>, GTP<sub>TL</sub> denote the stoichiometric coefficients 328 of ATP and GTP for CAT translation,  $\Delta m_i$  denotes the amount of the  $i^{th}$  substrate con-329 sumed, and  $ATP_i$  denotes the equivalent ATP number of the  $i^{th}$  substrate.  $ATP_{TX}$  = 176, 330  $CTP_{TX} = 144$ ,  $GTP_{TX} = 151$ ,  $UTP_{TX} = 189$ ,  $ATP_{TL} = 219$ ,  $GTP_{TL} = 438$ ,  $ATP_{GLC} = 189$ 331 21,  $\mathrm{ATP_{PYR}}$  = 8,  $\mathrm{ATP_{LAC}}$  = 9.5,  $\mathrm{ATP_{SUCC}}$  = 11.5,  $\mathrm{ATP_{MAL}}$  = 10.5. This analysis was 332 also extended to the accumulation of amino acids, organic acids, and other byproducts 333 to create a complete energy balance through the network. These terms were calculated 334 as accumulation times equivalent ATP number over total energy consumption. Equiva-335 lent ATP numbers for glucose, amino acids, and organic acids were calculated from the 336 network stoichiometry. In the first phase, amino acid accumulation consisted of alanine, 337 isoleucine, glutamine, proline, and tyrosine, while organic acid accumulation consisted of 338 pyruvate, lactate, acetate, succinate, and malate. Glucose was considered the substrate. 339 In the second phase, amino acid accumulation consisted of alanine, glutamine, proline, and serine, while organic acid accumulation consisted of acetate only. Pyruvate, lactate, 341 succinate, and malate were considered the substrates. Equivalent ATP numbers for glu-342 cose, amino acids, and organic acids were calculated from the network stoichiometry.

# **Competing interests**

The authors declare that they have no competing interests.

# 346 Author's contributions

J.V directed the modeling study. K.C and J.S conducted the cell-free protein synthesis experiments. J.V, J.W, and N.H developed the cell-free protein synthesis mathematical model, and parameter ensemble. The manuscript was prepared and edited for publication by J.S, N.H, M.V, J.W and J.V.

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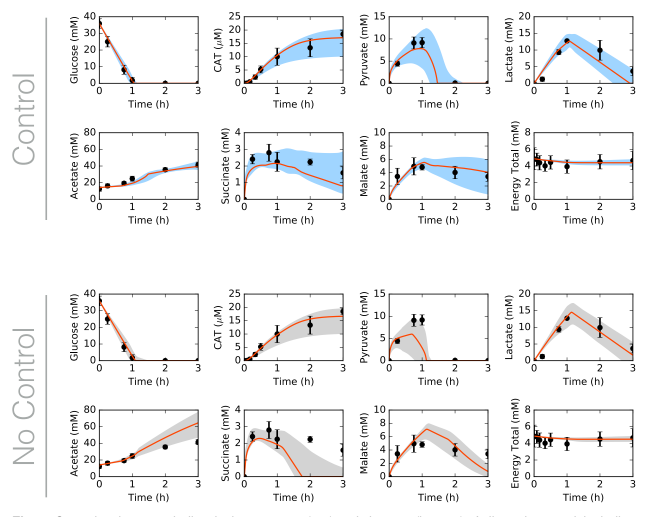
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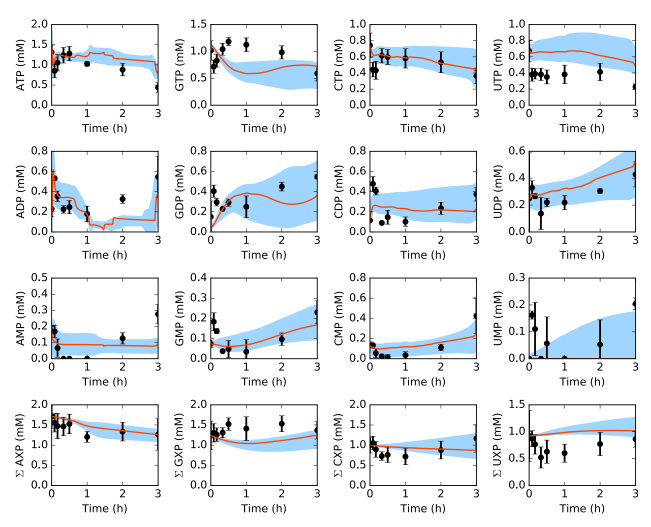
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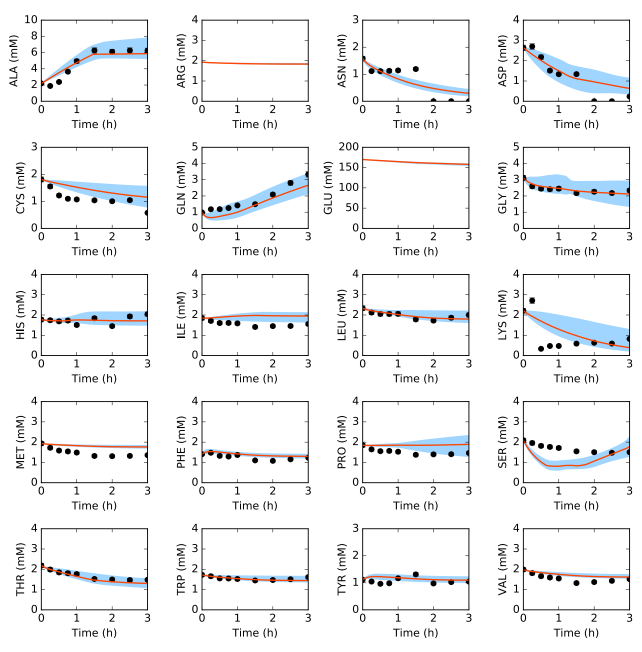
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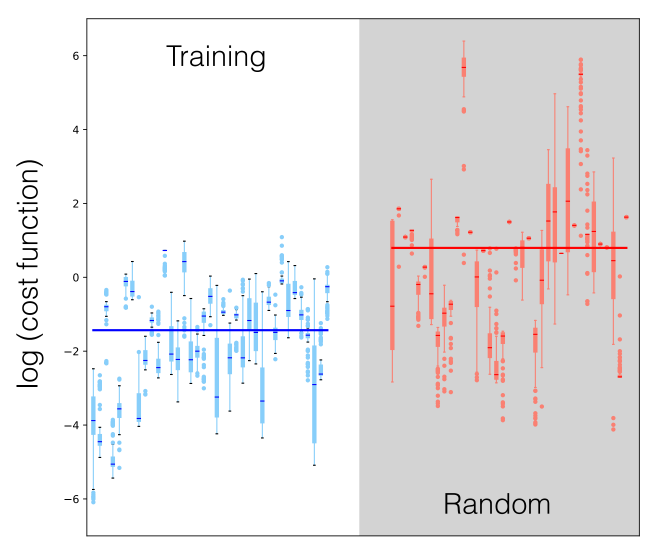
**Fig. 1:** Central carbon metabolism in the presence (top) and absence (bottom) of allosteric control, including glucose (substrate), CAT (product), and intermediates, as well as total concentration of energy species. Best-fit parameter set (orange line) versus experimental data (points). 95% confidence interval (blue or gray shaded region) over the ensemble of 100 sets.



**Fig. 2:** Energy species and energy totals by base in the presence of allosteric control. Best-fit parameter set (orange line) versus experimental data (points). 95% confidence interval (blue shaded region) over the ensemble of 100 sets.



**Fig. 3:** Amino acids in the presence of allosteric control. Best-fit parameter set (orange line) versus experimental data (points). 95% confidence interval (blue shaded region) over the ensemble of 100 sets.



# Measured species

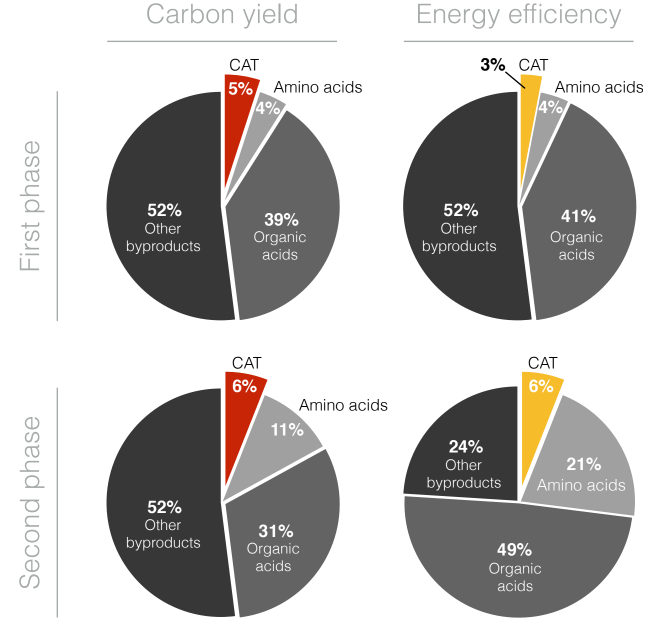
**Fig. 4:** Log of cost function across 37 datasets for data-trained ensemble (blue) and randomly generated ensemble (red, gray background). Median (bars), interquartile range (boxes), range excluding outliers (dashed lines), and outliers (circles) for each dataset. Median across all datasets (large bar overlaid).



**Fig. 5:** Effect of group knockouts on system. A. Change in CAT productivity when one (diagonal) or two (off-diagonal) reaction groups are turned off. B. Change in system state (only species for which data exist) when one (diagonal) or two (off-diagonal) reaction groups are turned off. Total-order effect for each group calculated as the sum of first-order effect and all pairwise effects. Larger and darker circles represent greater effects.

**Table 1:** Reference values for reaction rate maxima  $(V_{max})$  from literature.  $V_{max}$  values calculated from turnover numbers  $(K_{cat})$  from literature, and a characteristic enzyme concentration of 167 nM. Characteristic rate maximum for all other reactions calculated as geometric mean of calculated rate maxima.

Enzyme	Reaction	$K_{cat}$ (min $^{-1}$ )	$V_{max}$ (mM/h)	BNID#
Serine dehydrase	R_ser_deg	10400	104	101119
Isocitrate dehydrogenase	R₋icd	11900	119	101152
Lactate dehydrogenase	R₋ldh	5800	58	101036
Aspartate transaminase	R₋aspC, R₋tyr, R₋phe	25800	258	101108
Enolase	R_eno	13200	132	101028
Pyruvate kinase	R₋pyk	25000	250	101029 101030
Malic enzyme	R₋maeA, R₋maeB	35400	354	101167
Phosphofructokinase	R_pfk	554400	5544	104955
Malate dehydrogenase	$R_{-}mdh$	33000	330	101163
Citrate Synthase	R_gltA	42000	420	101149
6PG dehydrogenase	R_zwf, R_pgl, R_gnd	3200	32	101048
Succinate dehydrogenase	R_sdh	121	1.21	101162
Succinyl-coA synthetase	R₋sucCD	4700	47	101158
3PGA dehydrogenase	R₋gpm	1100	11	101135
PEP carboxylase	R₋ppc	35400	354	101139
3PGA kinase	R₋pgk	4300	43	101016
Characteristic rate maximum			110	



**Fig. 6:** Carbon and energy balances during the first and second phases of protein production for the best-fit set. Top left: Carbon moles produced as CAT, amino acids (alanine, isoleucine, glutamine, proline, and tyrosine), organic acids (pyruvate, lactate, acetate, succinate, and malate), and all other byproducts, as percentages of total carbon consumption (glucose and all other amino acids). Bottom left: Carbon moles produced as CAT, amino acids (alanine, glutamine, proline, and serine), organic acids (acetate only), and all other byproducts, as percentages of total carbon consumption (pyruvate, lactate, succinate, malate, and all other amino acids). Top right: Energy cost of CAT production, accumulation of amino acids (alanine, isoleucine, glutamine, proline, and tyrosine), accumulation of organic acids (pyruvate, lactate, acetate, succinate, and malate), and other byproducts, as percentages of total energy utilization from glucose. Bottom right: Energy cost of CAT production, accumulation of amino acids (alanine, glutamine, proline, and serine), accumulation of organic acids (acetate only), and other byproducts, as percentages of total energy utilization from pyruvate, lactate, succinate, and malate. Energy costs calculated in terms of equivalent ATP molecules.