

# **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 measurements of glucose, organic acids, energy species, amino acids and the protein product, chloramphenicol acetyltransferase (CAT). The ensemble described the training data, with the exception of some of the amino acid dynamics. To gauge the performance of the cell free reaction, we compared the observed CAT carbon yield, with the maximum theoretical CAT carbon yield calculated using sequence specific flux balance analysis. The observed CAT yield was 45% of the maximum theoretical yield, suggesting CAT production could be further optimized. The metabolic flux distribution predicted by the dynamic model and flux balance analysis were significantly different. The ensemble of dynamic models predicted the majority of carbon flux was routed through glycolysis and the TCA cycle, while flux balance analysis predicted significant flux through the Entner-Doudoroff pathway. Local and global sensitivity analysis suggested CAT production was most sensitive to parameters and initial conditions directly associated with CAT synthesis, as well as GTP/GMP synthesis, amino acid synthesis, and to a lesser extent amino acid initial conditions. On the other hand, CAT production was robust to allosteric control parameters and the initial conditions of glucose and oxygen. Taken together, we presented the first dynamic model of *E. coli* cell free protein synthesis. This study provides a foundation for genome-scale, dynamic modeling of cell-free *E. coli* protein synthesis.

**Keywords:** Biochemical engineering, systems biology, cell free protein synthesis

## 1 Introduction

2 Cell-free systems offer many advantages for the study, manipulation and modeling of  
3 metabolism compared to *in vivo* processes. Central amongst these advantages is direct  
4 access to metabolites and the microbial biosynthetic machinery without the interference of  
5 a cell wall. This allows us to control as well as interrogate the chemical environment while  
6 the biosynthetic machinery is operating, potentially at a fine time resolution. Second,  
7 cell-free systems also allow us to study biological processes without the complications  
8 associated with cell growth. Cell-free protein synthesis (CFPS) systems are arguably the  
9 most prominent examples of cell-free systems used today [1]. However, CFPS is not new;  
10 CFPS in crude *E. coli* extracts has been used since the 1960s to explore fundamentally  
11 important biological mechanisms [2, 3]. Today, cell-free systems are used in a variety of  
12 applications ranging from therapeutic protein production [4] to synthetic biology [5]. Inter-  
13 estingly, many of the challenges confronting genome-scale kinetic modeling can poten-  
14 tially be overcome in a cell-free system. For example, there is no complex transcriptional  
15 regulation to consider, transient metabolic measurements are easier to obtain, and we  
16 no longer have to consider cell growth. Thus, cell-free operation holds several significant  
17 advantages for model development, identification and validation. Theoretically, genome-  
18 scale cell-free kinetic models may be possible for industrially important organisms, such  
19 as *E. coli* or *B. subtilis*, if a simple, tractable framework for integrating allosteric regulation  
20 with enzyme kinetics can be formulated.

21 Mathematical modeling has long contributed to our understanding of metabolism. Decades  
22 before the genomics revolution, mechanistically, structured metabolic models arose from  
23 the desire to predict microbial phenotypes resulting from changes in intracellular or extra-  
24 cellular states [6]. The single cell *E. coli* models of Shuler and coworkers pioneered the  
25 construction of large-scale, dynamic metabolic models that incorporated multiple, regu-  
26 lated catabolic and anabolic pathways constrained by experimentally determined kinetic

parameters [7]. Shuler and coworkers generated many single cell kinetic models, including single cell models of eukaryotes [8, 9], minimal cell architectures [10], as well as DNA sequence based whole-cell models of *E. coli* [11]. Conversely, highly abstracted kinetic frameworks, such as the cybernetic framework, represented a paradigm shift, viewing cells as growth-optimizing strategists [12]. Cybernetic models have been highly successful at predicting metabolic choice behavior, e.g., diauxie behavior [13], steady-state multiplicity [14], as well as the cellular response to metabolic engineering modifications [15]. Unfortunately, traditional, fully structured cybernetic models also suffer from an identifiability challenge, as both the kinetic parameters and an abstracted model of cellular objectives must be estimated simultaneously. However, recent cybernetic formulations from Ramkrishna and colleagues have successfully treated this identifiability challenge through elementary mode reduction [16, 17].

In the post genomics world, large-scale stoichiometric reconstructions of microbial metabolism popularized by static, constraint-based modeling techniques such as flux balance analysis (FBA) have become standard tools [18]. Since the first genome-scale stoichiometric model of *E. coli*, developed by Edwards and Palsson [19], well over 100 organisms, including industrially important prokaryotes such as *E. coli* [20] or *B. subtilis* [21], are now available [22]. Stoichiometric models rely on a pseudo-steady-state assumption to reduce unidentifiable genome-scale kinetic models to an underdetermined linear algebraic system, which can be solved efficiently even for large systems. Traditionally, stoichiometric models have also neglected explicit descriptions of metabolic regulation and control mechanisms, instead opting to describe the choice of pathways by prescribing an objective function on metabolism. Interestingly, similar to early cybernetic models, the most common metabolic objective function has been the optimization of biomass formation [23], although other metabolic objectives have also been estimated [24]. Recent advances in constraint-based modeling have overcome the early shortcomings of

the platform, including capturing metabolic regulation and control [25]. Thus, modern constraint-based approaches have proven extremely useful in the discovery of metabolic engineering strategies and represent the state of the art in metabolic modeling [26, 27]. However, genome-scale kinetic models of industrial important organisms such as *E. coli* have yet to be constructed.

In this study, we developed an ensemble of *E. coli* cell free protein synthesis (CFPS) models using the hybrid cell free modeling approach of Wayman et al [REFHERE]. Model parameters were estimated from measurements of glucose, organic acids, energy species, amino acids and the protein product, chloramphenicol acetyltransferase (CAT). The ensemble described the training data, with the exception of some of the amino acid dynamics. To gauge the performance of the cell free reaction, we compared the observed CAT carbon yield, with the maximum theoretical CAT carbon yield calculated using sequence specific flux balance analysis. The observed CAT yield was 45% of the maximum theoretical yield, suggesting CAT production could be further optimized. The metabolic flux distribution predicted by the dynamic model and flux balance analysis were significantly different. The ensemble of dynamic models predicted the majority of carbon flux was routed through glycolysis and the TCA cycle, while flux balance analysis predicted significant flux through the Entner-Doudoroff pathway. Local and global sensitivity analysis suggested CAT production was most sensitive to parameters and initial conditions directly associated with CAT synthesis, as well as GTP/GMP synthesis, amino acid synthesis, and to a lesser extent amino acid initial conditions. On the other hand, CAT production was robust to allosteric control parameters and the initial conditions of glucose and oxygen. Taken together, we presented the first dynamic model of *E. coli* cell free protein synthesis. We 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.

## Results

**Estimation of an ensemble of cell free protein synthesis models.** We used the hybrid cell free modeling framework of Wayman et al. to simulate the production of a model protein [REFHERE]. The cell-free *E. coli* metabolic model was constructed by removing the growth-associated processes from the model of Palsson and coworkers [19], and by adding reactions for the synthesis of chloramphenicol acetyltransferase (CAT), a model protein for which we have a comprehensive training dataset [28]. Thus, the model described core central carbon metabolism (glycolysis, pentose phosphate, Enter-Doudoroff, TCA cycle), as well as the synthesis of energy species, amino acids biosynthesis and degradation, and biosynthesis of the CAT protein. An ensemble of model parameters was estimated from dynamic 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. We generated an ensemble of  $N = 18,000$  parameter sets by minimizing the error between the training dataset and the metabolite concentrations predicted by the model. We defined the set with the lowest error value as the best-fit parameter set. [STATISTICS ON PARAMETERS].

The ensemble of models captured the time evolution of cell free CAT biosynthesis (Fig. 1 - 3). Glucose was exhausted with 3 hr [FILL ME IN]. The ensemble also captured the energy species dynamics, particularly the overall energy total (Fig. 1, top) and the totals by base . The ensemble and the best-fit set also predict some of the amino acid measurements, while failing to predict others (Fig. 3). the central carbon metabolism, including glucose uptake, CAT production, and the dynamics of the organic acid intermediates . Allosteric control is important to the dynamics of the organic acid intermediates, as without it several of the measurements are not captured by the ensemble or the best-fit set (Fig. 1, bottom). This is likely due to a structural deficiency in the model; in some cases, the consumption of an amino acid through CAT synthesis is not enough to ex-

plain the decrease shown in the data, and there are no other reactions that consume it. Thus, a more comprehensive biological description is needed to fully explain amino acid dynamics.

**Sensitivity analysis** We performed a local sensitivity analysis to determine the network reactions with the greatest effect on protein production and overall system state. CAT production was not sensitive to the CAT synthesis reaction, oxidative phosphorylation activity, and alanine synthesis, as well as various reactions in glycolysis, the TCA cycle, and amino acid synthesis and degradation.

**Maximum theoretical CAT yield showed CFPS can be optimized.** We calculated the carbon yield of CAT production for our experimental data and our best-fit parameter set as a function of the initial and final concentrations and the carbon numbers of CAT, glucose, and amino acids. The experimental data displayed a CAT yield of 0.0865, while the best-fit parameter set displayed a CAT yield of 0.0871. While the model ensemble described the experimental data, it was unclear whether the performance of the CFPS system was optimal. To address this question, we used ssFBA in combination with the cell-free metabolic network and a detailed promoter model under a T7 polymerase to compute the maximum theoretical carbon yield. However, we first validated the ssFBA approach by comparing an ensemble of simulated versus measured concentrations of CAT over a one hour period (Fig. 7A). The ensemble of 100 sets captured the CAT concentration profile which was randomly generated by sampling RNA polymerase levels, ribosome levels and elongation rates in a physiological range. We then used sequence-specific FBA to calculate a theoretical maximum CAT yield under four different cases: unconstrained, limited oxidative phosphorylation, bounded by transcription/translation rates, and bounded by experimental data (Fig. 7B). The theoretical maximum carbon yield of CAT was 0.349 for an unconstrained case and 0.194 for the transcription and translation constrained case. Thus, we showed that our experimental dataset and best-fit parameter set were each produc-

ing CAT at 25% of the theoretical maximum and 45% of a theoretical physiological case. Whereas, the case constrained by experimental data had a carbon yield of  $0.062 \pm 0.02$ , similar to the experimental yield. This allowed us to quantify the amount of carbon being diverted to byproducts, and suggests that there is potential for a doubling of CAT production by reducing this diversion of carbon. In comparing the flux distributions between the unconstrained and constrained cases (Fig. 8), both constrained cases heavily utilize the Entner–Doudoroff pathway which may be a first viable knockout to increase CAT yield.

**Sensitivity analysis on FBA system** We conducted global sensitivity on certain fluxes' upper bounds within a constraint-based FBA system, with protein export rate as the objective function.



## Discussion

In this study, we developed an ensemble of *E. coli* cell free protein synthesis (CFPS) models using the hybrid cell free modeling approach of Wayman et al [REFHERE]. Model parameters were estimated from measurements of glucose, organic acids, energy species, amino acids and the protein product, chloramphenicol acetyltransferase (CAT). The ensemble described the training data, with the exception of some of the amino acid dynamics. To gauge the performance of the cell free reaction, we compared the observed CAT carbon yield, with the maximum theoretical CAT carbon yield calculated using sequence specific flux balance analysis. The observed CAT yield was 45% of the maximum theoretical yield, suggesting CAT production could be further optimized. The metabolic flux distribution predicted by the dynamic model and flux balance analysis were significantly different. The ensemble of dynamic models predicted the majority of carbon flux was routed through glycolysis and the TCA cycle, while flux balance analysis predicted significant flux through the Entner-Doudoroff pathway. Local and global sensitivity analysis suggested CAT production was most sensitive to parameters and initial conditions directly associated with CAT synthesis, as well as GTP/GMP synthesis, amino acid synthesis, and to a lesser extent amino acid initial conditions. On the other hand, CAT production was robust to allosteric control parameters and the initial conditions of glucose and oxygen.

The cell free model ensemble described the training data with the exception of some of the amino acids. Specifically, adding more reactions that consume amino acids would improve the model's ability to predict those that show a decrease in the experimental data. Also, including specific transcription and translation steps for CAT would allow us to more accurately model the complexity and the resource cost of protein synthesis. Another area for future work is to more thoroughly sample parameter space. For the metabolites in the dataset, initial conditions were fixed at the initial data values. All other parameters were varied in a manner so as to best fit the dataset. However, the resulting ensemble may not

represent every biological or practical possibility. In a different region of parameter space, the system could behave differently, including the flux distribution through the network, the accuracy and spread of ensemble fits, the relative sensitivities, and the yield as a percentage of the theoretical maximum. Testing the model under a variety of conditions could strengthen or challenge the findings of this study. Further experimentation could also be used to gain a deeper understanding of model performance under a variety of conditions. Specifically, CAT production performed in the absence of amino acids could inform the system's ability to manufacture them, while experimentation in the absence of glucose or oxygen could shed light on how important they are to protein synthesis, and under which conditions. Finally, the approach should be extended to other protein products. CAT is only a test protein used for model identification; the modeling framework, and to some extent the parameter values, should be protein agnostic. An important extension of this study would be to apply its insights to other protein 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:

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

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

where  $\mathcal{R}$  denotes the number of reactions,  $\mathcal{M}$  denotes the number of metabolites and  $\mathcal{E}$  denotes the number of enzymes in the model. The quantity  $r_j(\mathbf{x}, \epsilon, \mathbf{k})$  denotes the rate of reaction  $j$ . Typically, reaction  $j$  is a non-linear function of metabolite and enzyme abundance, as well as unknown kinetic parameters  $\mathbf{k}$  ( $\mathcal{K} \times 1$ ). The quantity  $\sigma_{ij}$  denotes the stoichiometric coefficient for species  $i$  in reaction  $j$ . If  $\sigma_{ij} > 0$ , metabolite  $i$  is produced by reaction  $j$ . Conversely, if  $\sigma_{ij} < 0$ , metabolite  $i$  is consumed by reaction  $j$ , while  $\sigma_{ij} = 0$  indicates metabolite  $i$  is not connected with reaction  $j$ . Lastly,  $\lambda_i$  denotes the scaled enzyme degradation constant. The system material balances were subject to the initial conditions  $\mathbf{x}(t_o) = \mathbf{x}_o$  and  $\epsilon(t_o) = 1$  (initially we have 100% cell-free enzyme abundance).

The reaction rate was written as the product of a kinetic term ( $\bar{r}_j$ ) and a control term ( $v_j$ ),  $r_j(\mathbf{x}, \mathbf{k}) = \bar{r}_j v_j$ . In this study, we used either saturation or mass action kinetics. The control term  $0 \leq v_j \leq 1$  depended upon the combination of factors which influenced rate process  $j$ . For each rate, we used a rule-based approach to select from competing control factors. If rate  $j$  was influenced by  $1, \dots, m$  factors, we modeled this relationship as  $v_j = \mathcal{I}_j(f_{1j}(\cdot), \dots, f_{mj}(\cdot))$  where  $0 \leq f_{ij}(\cdot) \leq 1$  denotes a regulatory transfer function quantifying the influence of factor  $i$  on rate  $j$ . The function  $\mathcal{I}_j(\cdot)$  is an integration rule which maps the output of regulatory transfer functions into a control variable. Each regulatory

transfer function took the form:

$$f_{ij}(\mathcal{Z}_i, k_{ij}, \eta_{ij}) = k_{ij}^{\eta_{ij}} \mathcal{Z}_i^{\eta_{ij}} / (1 + k_{ij}^{\eta_{ij}} \mathcal{Z}_i^{\eta_{ij}}) \quad (3)$$

where  $\mathcal{Z}_i$  denotes the abundance factor  $i$ ,  $k_{ij}$  denotes a gain parameter, and  $\eta_{ij}$  denotes a cooperativity parameter. In this study, we used  $\mathcal{I}_j \in \{mean\}$  [? ]. If a process has no modifying factors,  $v_j = 1$ . We used multiple saturation kinetics to model the reaction term  $\bar{r}_j$ :

$$\bar{r}_j = k_j^{max} \epsilon_i \left( \prod_{s \in m_j^-} \frac{x_s}{K_{js} + x_s} \right) \quad (4)$$

where  $k_j^{max}$  denotes the maximum rate for reaction  $j$ ,  $\epsilon_i$  denotes the scaled enzyme activity which catalyzes reaction  $j$ , and  $K_{js}$  denotes the saturation constant for species  $s$  in reaction  $j$ . The product in Equation (4) was carried out over the set of *reactants* for reaction  $j$  (denoted as  $m_j^-$ ).

We added regulation to the network as informed by literature, for a total of 17 interactions. PEP was modeled as an inhibitor for phosphofructokinase [29, 30], PEP carboxykinase [29], PEP synthetase [29, 31], isocitrate dehydrogenase [29, 32], and isocitrate lyase/malate synthase [29, 32, 33], and as an activator for fructose-biphosphatase [29, 34–36]. AKG was modeled as an inhibitor for citrate synthase [29, 37, 38] and isocitrate lyase/malate synthase [29, 33]. 3PG was modeled as an inhibitor for isocitrate lyase/malate synthase [29, 33]. FDP was modeled as an activator for pyruvate kinase [29, 39] and PEP carboxylase [29, 40]. Pyruvate was modeled as an inhibitor for pyruvate dehydrogenase [29, 41, 42] and as an activator for lactate dehydrogenase [43]. Acetyl CoA was modeled as an inhibitor for malate dehydrogenase [29].

**Generation of model ensemble** We generated an ensemble of 100 diverse parameter sets via a Markov chain Monte Carlo random walk. Beginning with a single parameter

222 set as a starting point, we calculated its cost function, equal to the sum-squared-error  
 223 between experimental data and model predictions:

$$cost = \sum_{i=1}^{\mathcal{D}} \left( w_i \sum_{j=1}^{\mathcal{T}_i} abs \left( x_{ij}^{data} - x_i^{sim}|_{t(j)} \right) \right) \quad (5)$$

224 where  $\mathcal{D}$  denotes the number of datasets,  $w_i$  denotes the weight of the  $i$ th dataset,  $\mathcal{T}_i$   
 225 denotes the number of timepoints in the  $i$ th dataset,  $t(j)$  denotes the  $j$ th timepoint,  $x_{ij}^{data}$   
 226 denotes the value of the  $i$ th dataset at the  $j$ th timepoint, and  $x_i^{sim}|_{t(j)}$  denotes the sim-  
 227 ulated value of the metabolite corresponding to the  $i$ th dataset, interpolated to the  $j$ th  
 228 timepoint. We then perturbed model parameters:

$$k_i^{new} = k_i * exp(a r_i) \quad i = 1, 2, \dots, \mathcal{P} \quad (6)$$

229 where  $\mathcal{P}$  denotes the number of parameters, equal to 815, which includes 163 rate con-  
 230 stants, 163 enzyme degradation rate constants, 455 saturation constants, and 34 control  
 231 parameters,  $k_i^{new}$  denotes the new value of the  $i$ th parameter,  $k_i$  denotes the current value  
 232 of the  $i$ th parameter,  $a$  denotes a distribution variance, and  $r_i$  denotes a random sample  
 233 from the normal distribution. For each newly generated parameter set, we re-solved the  
 234 balance equations and calculated the cost function. All sets with a lower cost than the  
 235 previous set, and some with higher cost, were added to the ensemble. After generating  
 236 12,437 sets, we selected 100 sets with minimal correlation to each other so as to avoid  
 237 over-sampling any region of parameter space. The original 12,437-set ensemble had  
 238 a [mean,median,maximum] Pearson correlation coefficient [REFERENCE NEEDED?] of  
 239 [?] between pairs of sets; the 100-set ensemble had a [mean,median,maximum] Pearson  
 240 correlation coefficient of [?] between pairs of sets.

**Sensitivity analysis** We determined the reactions most important to protein production by computing the local sensitivity of CAT concentration to each rate constant in the network. Each sensitivity index was formulated as:

$$S_{ij} = \text{norm}(CAT(p_i, p_j, t) - CAT(\alpha * p_i, \alpha * p_j, t)) \quad i, j = 1, 2, \dots \mathcal{P} \quad (7)$$

where  $S_{ij}$  denotes the sensitivity of CAT production to the  $i$ th and  $j$ th parameters,  $CAT(p_i, p_j, t)$  denotes CAT concentration as a function of time and the  $i$ th and  $j$ th parameters,  $\alpha$  denotes the perturbation factor, equal to 1.01, and  $\mathcal{P}$  denotes the number of rate constants, equal to 163. In calculating the pairwise sensitivities, each parameter was perturbed by 1%; first-order sensitivities ( $i = j$ ) were subject to two 1% perturbations, equivalent to a perturbation of 2.01%.

**Sequence specific FBA and calculation of CAT yield** The yield on CAT production was calculated for each case as a ratio of carbon produced as CAT to carbon consumed as reactants (glucose and amino acids):

$$Yield = \frac{\Delta CAT \ C_{CAT}}{\sum_{i=1}^{\mathcal{R}} \max(\Delta m_i, 0) \ C_{m_i}} \quad (8)$$

where  $\Delta CAT$  denotes the amount of CAT produced,  $C_{CAT}$  denotes carbon number of CAT,  $\mathcal{R}$  denotes the number of reactants,  $\Delta m_i$  denotes the amount of the  $i$ th reactant consumed, never allowed to be negative, and  $C_{m_i}$  denotes the carbon number of the  $i$ th reactant. Because no data was available for arginine or glutamate, these reactants were left out of all three calculations. In the experimental case and the best-fit set case, yield was calculated by setting  $\Delta CAT$  equal to the final minus the initial CAT concentration and setting  $\Delta m_i$  equal to the initial minus the final reactant concentration. The theoretical yield was calculated using flux balance analysis (FBA) with a sequence-specific based

261 analysis on CAT. The sequence specific FBA [44] problem was formulated as:

$$\max_{\mathbf{w}} (w_{obj} = \boldsymbol{\theta}^T \mathbf{w})$$

$$\text{Subject to : } \mathbf{S}\mathbf{w} = \mathbf{0}$$

$$\alpha_i \leq w_i \leq \beta_i \quad i = 1, 2, \dots, \mathcal{R}$$

262 where  $\mathbf{S}$  denotes the stoichiometric matrix,  $\mathbf{w}$  denotes the unknown flux vector,  $\boldsymbol{\theta}$  denotes  
 263 the objective selection vector and  $\alpha_i$  and  $\beta_i$  denote the lower and upper bounds on flux  
 264  $w_i$ , respectively. The objective  $w_{obj}$  was to maximize the specific rate of CAT formation.  
 265 The specific glucose uptake rate was constrained to allow a maximum flux of 40 mM/hr  
 266 according to literature data; the specific amino acid uptake rates were also bound to allow  
 267 a maximum flux of 30 mM/hr, but did not reach this maximum flux. The transcription  
 268 and translation template reactions were added to the metabolic network and are based  
 269 off sequence specific analysis [44] involving transcription initiaion, transcription, mRNA  
 270 degradation, translation initiation, translation, and tRNA charging. The mRNA and protein  
 271 sequence of each protein was determined from literature. The transcription rate was  
 272 constrained as:

$$w_{tx} = RNAP \frac{v_{RNAP}}{l_{mRNA}} \left( \frac{Gene}{km + Gene} \right) P$$

273 where  $RNAP$  is the concentration of RNA polymerase,  $v_{RNAP}$  is the elongation rate (nu-  
 274 cleotides/hr) by the RNA polymerase,  $l_{mRNA}$  is the number of nucleotides in the mRNA,  
 275  $Gene$  is the gene concentration,  $km$  is the plasmid saturation coefficient, and  $P$  is the  
 276 promoter activity.

277 The promoter activity was formulated following Moon et al. for synthetic circuits as:

$$P = \frac{K_1}{1 + K_1}$$

where  $K_1$  represents the state of T7 RNA polymerase binding.

The translation rate was constrained as:

$$w_{tl} = K_P Ribo \frac{v_{Ribo}}{l_{protein}} [mRNA_{ss}]$$

where  $K_P$  is the polysome amplification constant,  $Ribo$  is the ribosome concentration,  $v_{Ribo}$  is the elongation rate (amino acids/hr) of the ribosome,  $l_{protein}$  is the number of amino acids in the protein of interest, and  $mRNA_{ss}$  is the mRNA concentration at steady state determined by the transcription rate divided by the degradation rate of mRNA. An ensemble of flux distributions was calculated for 100 sets by randomly sampling. Glucose, oxygen, and amino acids were modeled as being imported into the system, whereas CAT synthesis and metabolite byproduct formation was modeled as an export from the system. The rest of the network followed a pseudo steady-state assumption where all other metabolites were not allowed to accumulate; thus, the network could be solved by linear programming. The flux balance analysis problem was solved using the GNU Linear Programming Kit (v4.52) [45]. The solution flux vector was used to calculate the theoretical carbon yield of CAT for four different cases. For the unconstrained case, all rates were left unbounded. An ensemble of flux distributions was calculated by randomly sampling the maximum specific glucose uptake rate from 30 to 40 mM/hr determined from experimental data. For the limited oxidative phosphorylation case, an ensemble of flux distributions was generated by randomly sampling the maximum specific oxygen uptake rate to from 0 to 10 mM/hr. For the TXTL case (constrained by transcription and translation rates), an ensemble was generated by randomly sampling RNAP polymerase levels, ribosome levels, and elongation rates in a physiological range determined from literature as well as the specific glucose uptake rate from 30 to 40 mM/hr. For the case where the flux was constrained by experimental data, the lower and upper bounds where data was available were



301 randomly sampled in the physiological range of the data in addition to randomly sampling  
302 RNAP polymerase levels, ribosome levels, and elongation rates in a physiological range  
303 determined from literature.

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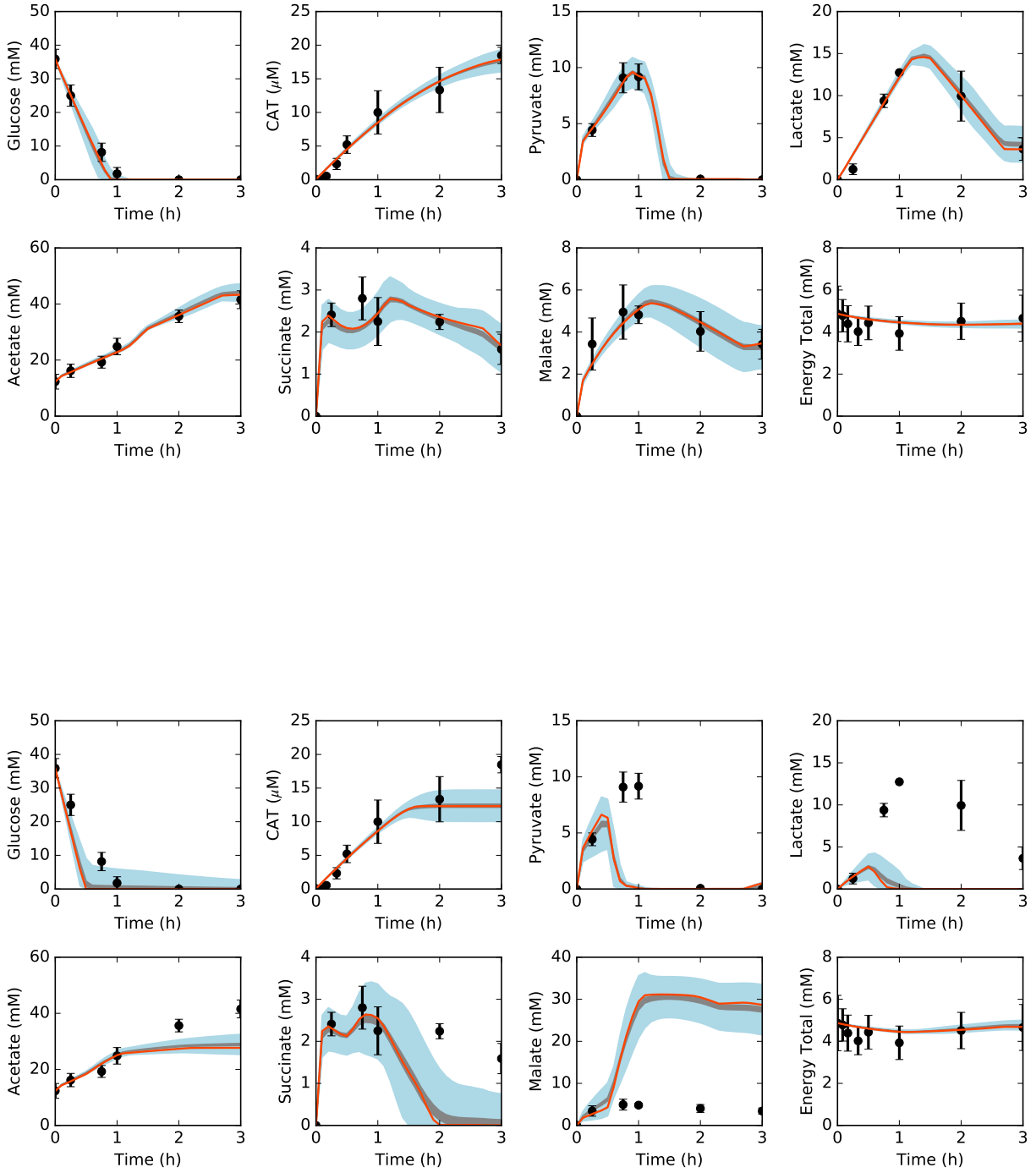
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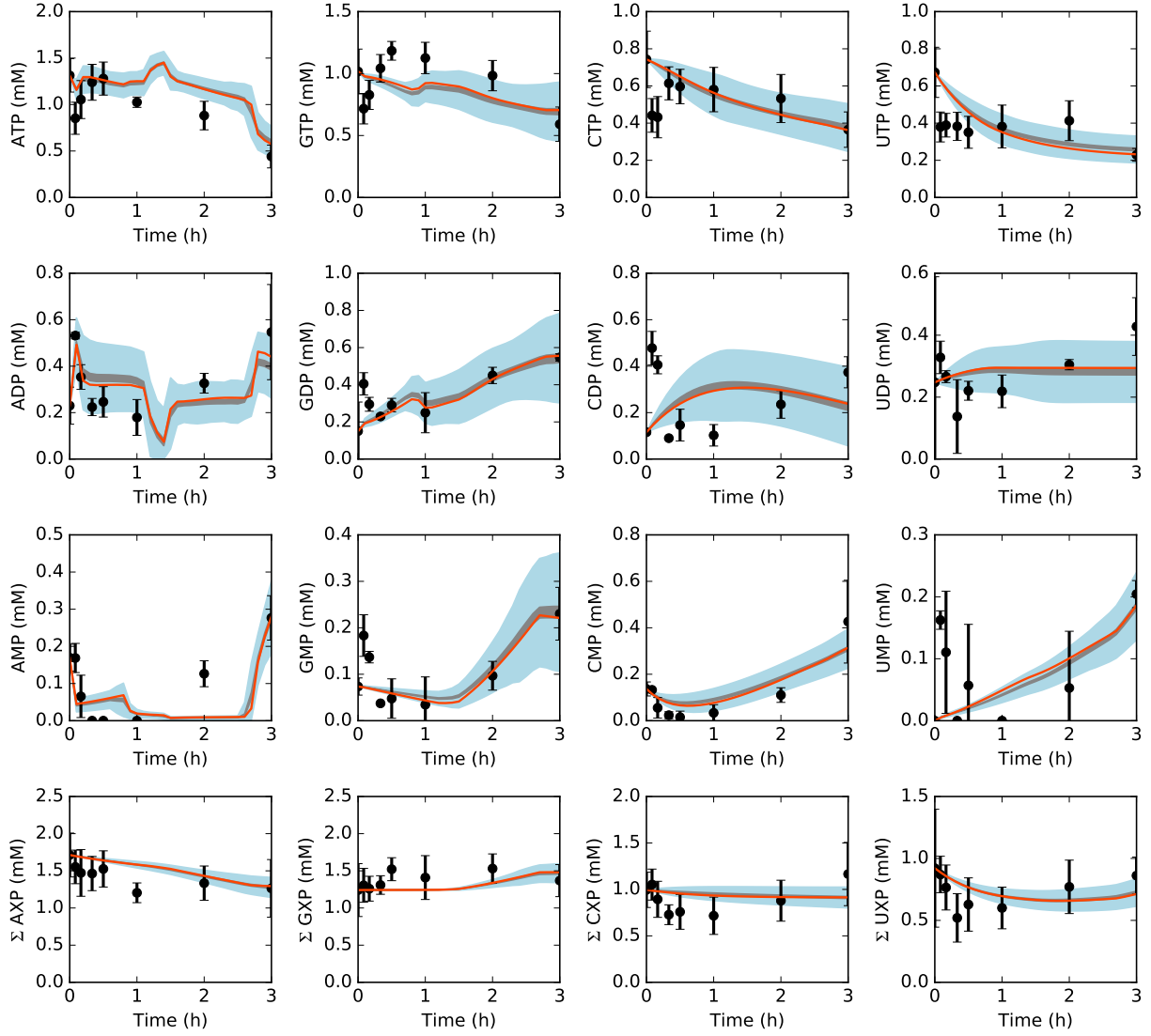
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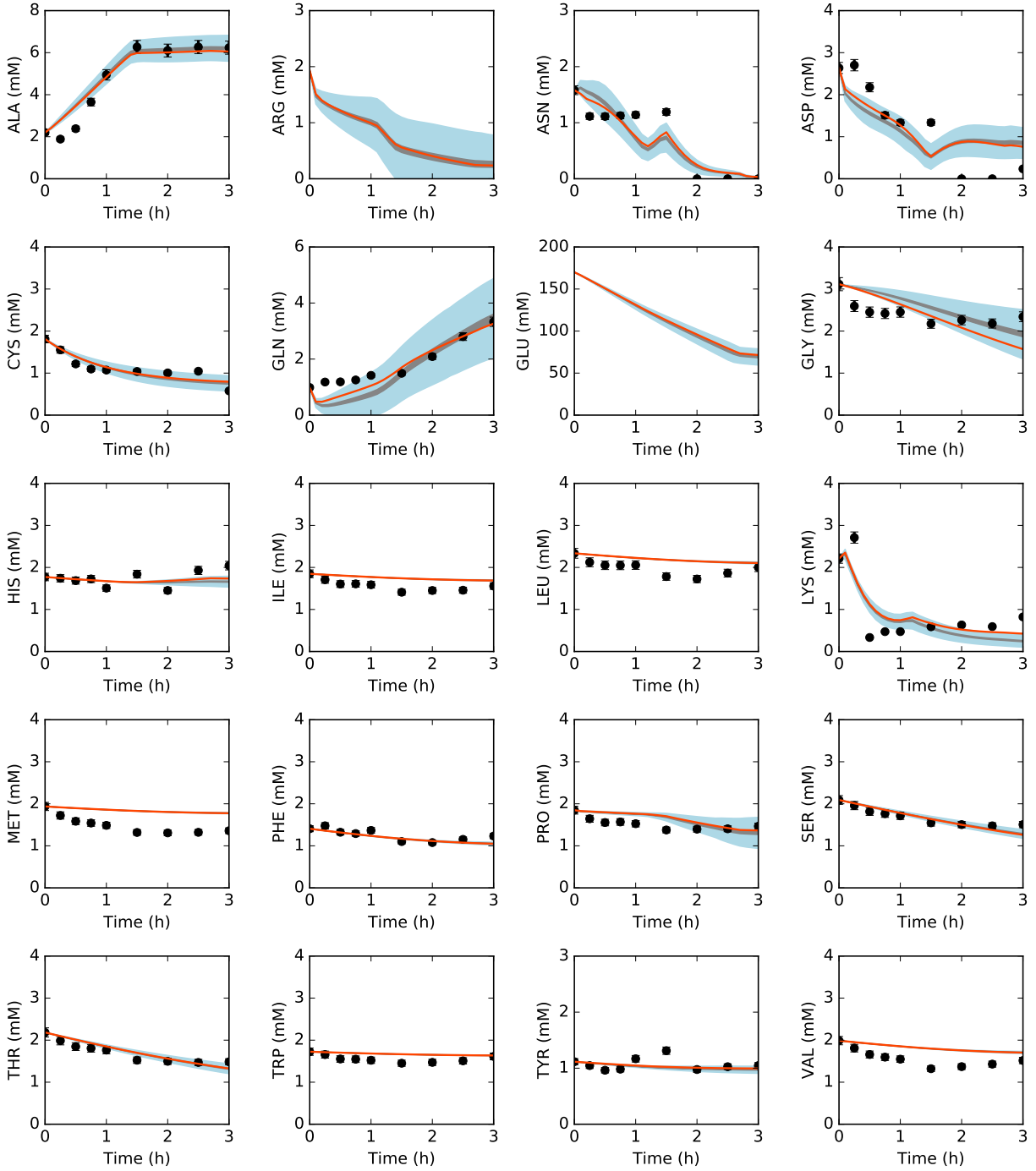


**Fig. 1:** Central carbon metabolism in the presence (top) and absence (bottom) of allosteric control, including glucose (substrate), CAT (product), and intermediates, as well as total concentration of energy species. Best-fit parameter set (orange line) versus experimental data (points). 95% confidence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

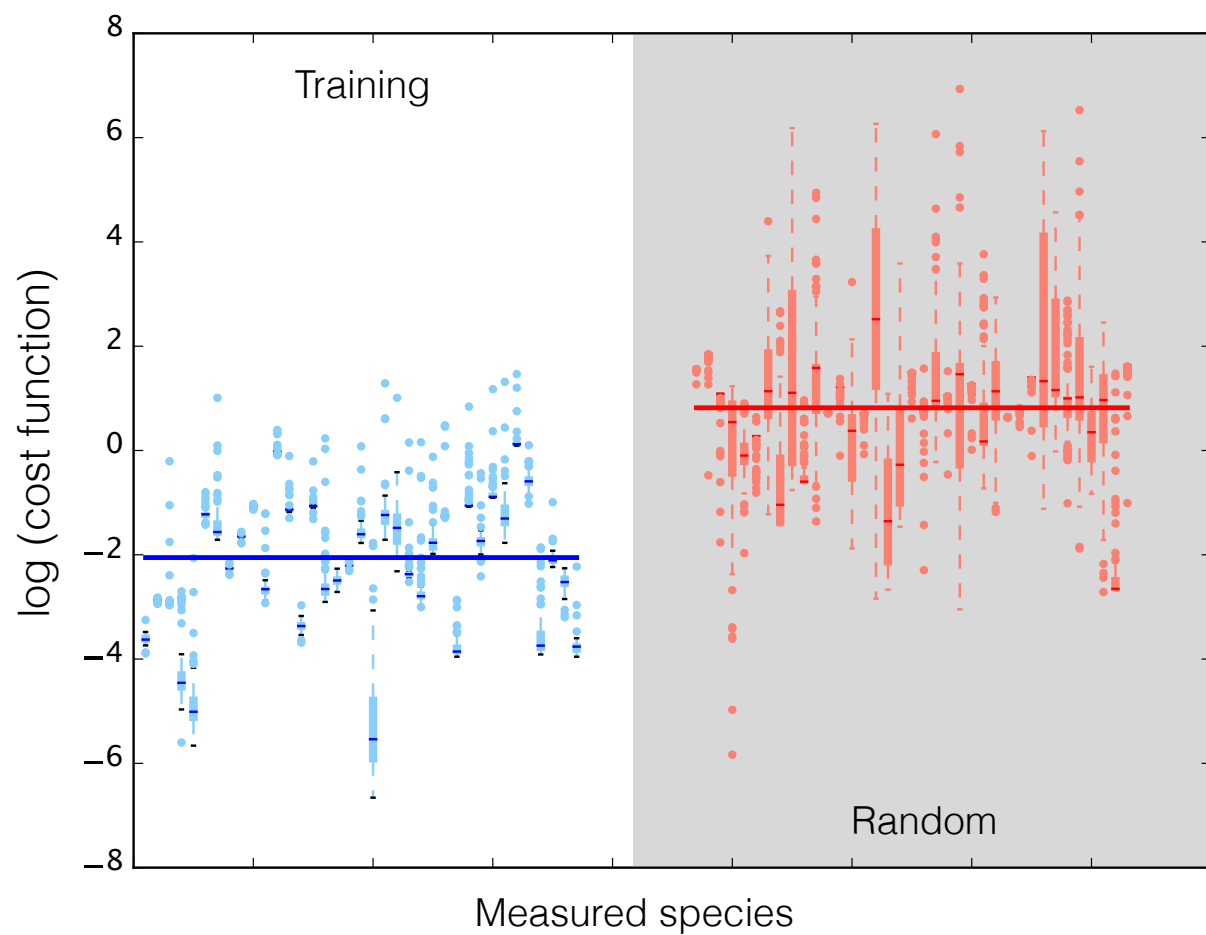




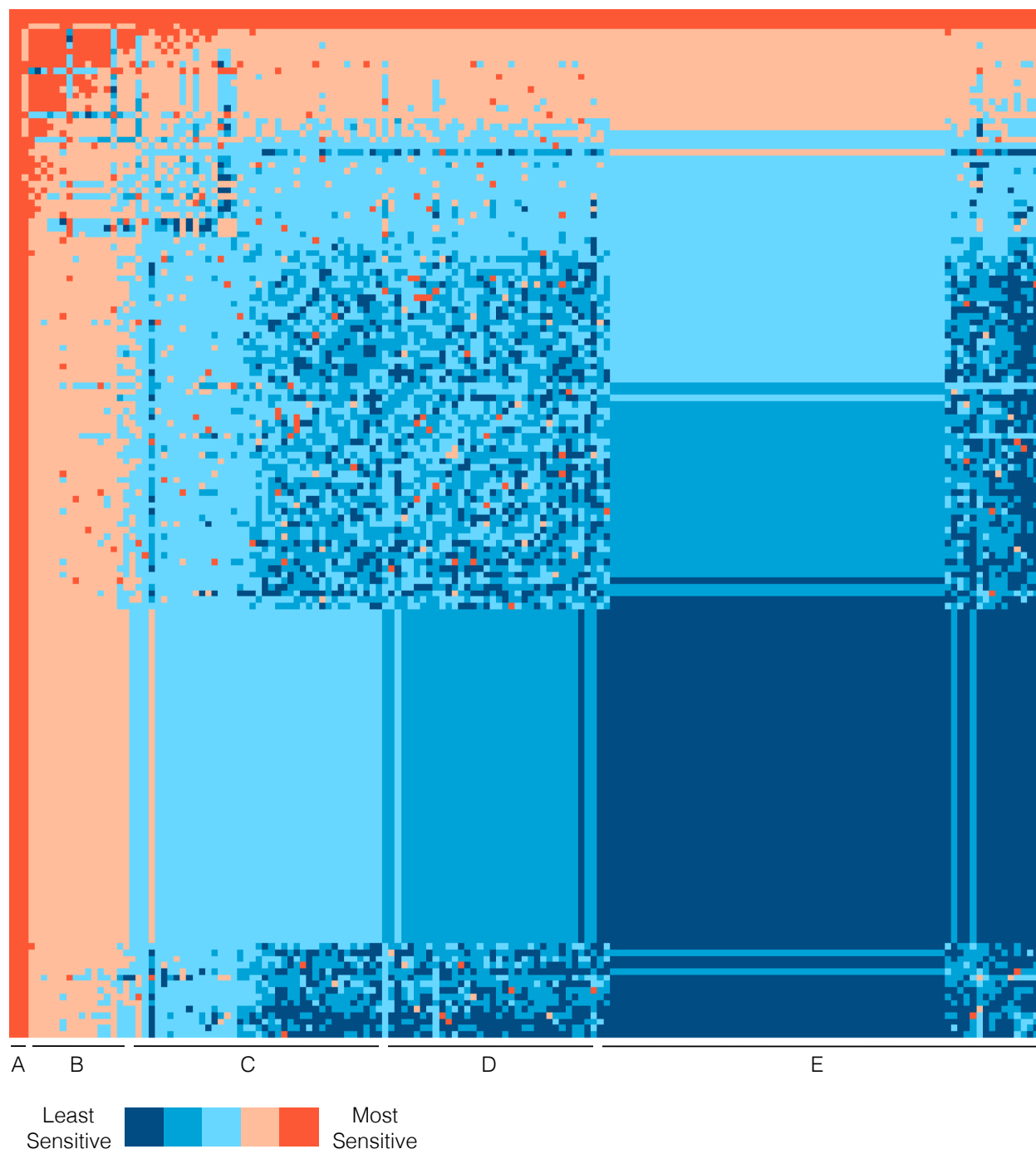
**Fig. 2:** Energy species and energy totals by base in the presence of allosteric control. Best-fit parameter set (orange line) versus experimental data (points). 95% confidence interval (blue shaded region) and 95% confidence interval of the mean (gray shaded region) over the ensemble of 100 sets.



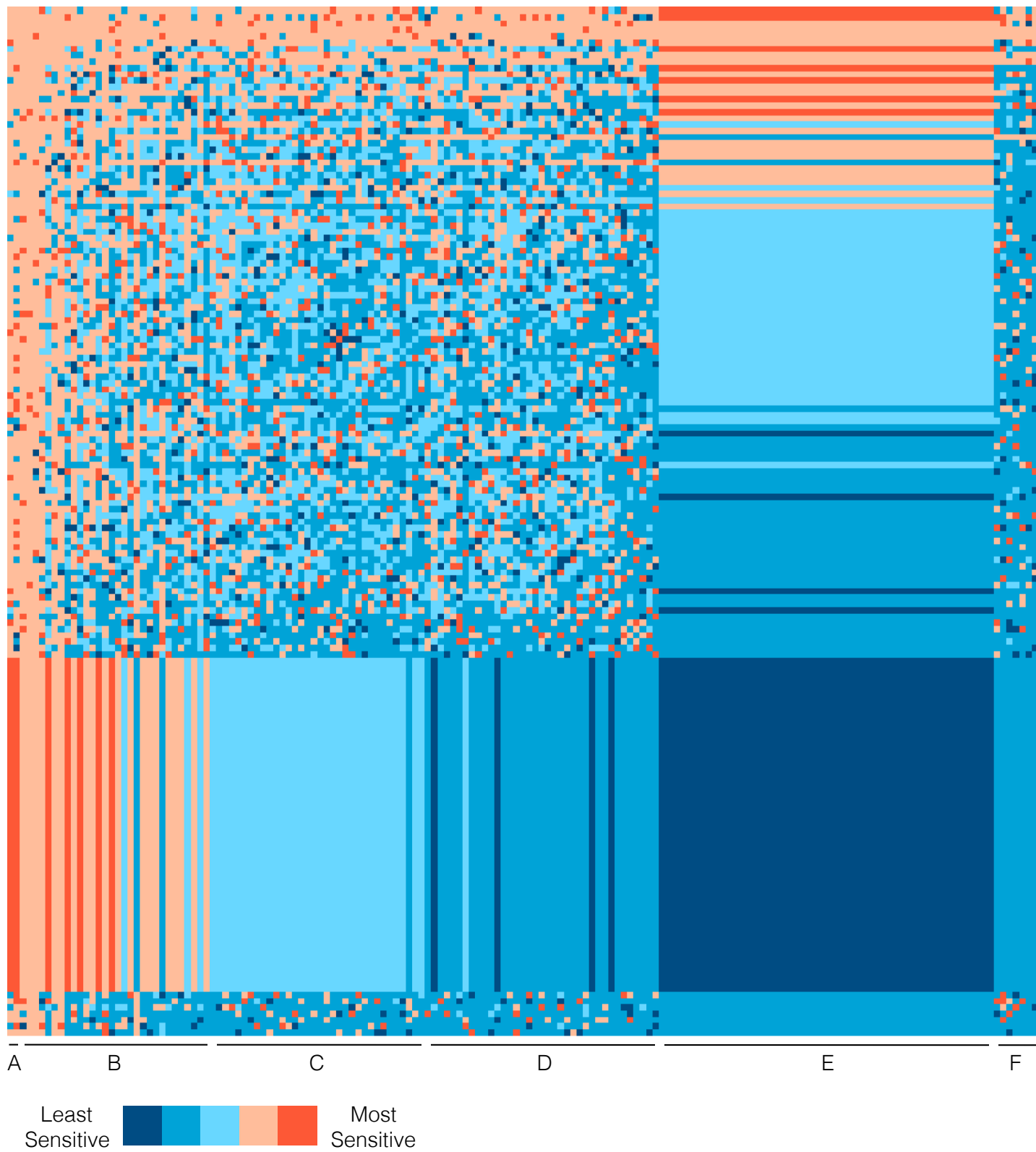
**Fig. 3:** Amino acids in the presence of allosteric control. Best-fit parameter set (orange line) versus experimental data (points). 95% confidence interval (blue shaded region) and 95% confidence interval of the mean (gray shaded region) over the ensemble of 100 sets.



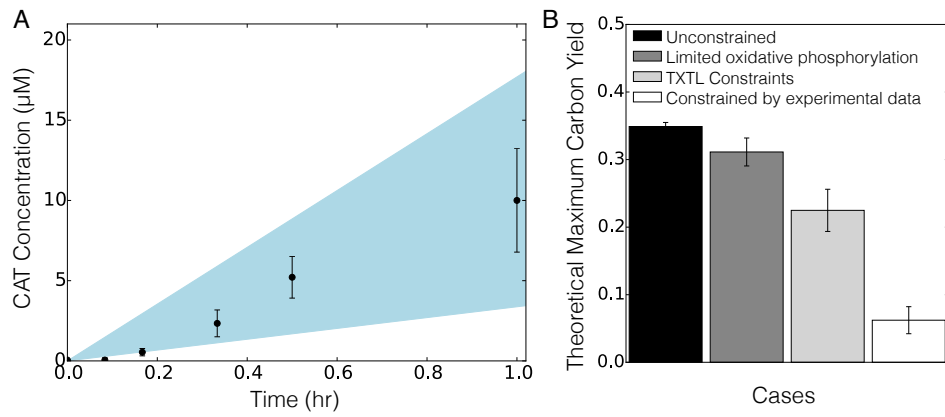
**Fig. 4:** Log of cost function across 37 datasets for data-trained ensemble (blue) and randomly generated ensemble (red, gray background). Median (bars), interquartile range (boxes), range excluding outliers (dashed lines), and outliers (circles) for each dataset. Median across all datasets (large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



**Fig. 5:** Normalized first-order and pairwise sensitivities of CAT production to rate constants.



**Fig. 6:** Normalized first-order and pairwise sensitivities of system state to rate constants. A (most sensitive parameters): hexokinase run by GTP, and lactate dehydrogenase forward reaction. B:



**Fig. 7:** Sequence specific flux balance analysis of CAT production and yield. **A.** 95% confidence interval of the ensemble (light blue region) for CAT concentration versus time. **B.** Theoretical maximum carbon yield of CAT calculated by ssFBA for four different cases: unconstrained (black), limited oxidative phosphorylation activity (gray), constrained by transcription and translation rates (light grey), and constrained by experimental data (white). Error bars represent standard deviation of the ensemble.

