- Molecular transport and packing underlie increasing ribosome productivity in faster
- 2 growing cells

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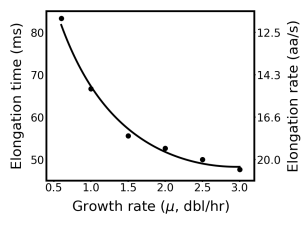
## **Summary**

Faster growing cells must make proteins more quickly. This occurs in part through increasing total ribosome abundance. However, the productivity of individual ribosomes also increases, almost doubling via an unknown mechanism. To investigate, we model both physical transport and chemical reactions among ensembles of individual molecules involved in translation elongation in *Escherichia coli*. We predict that the Damköhler number, the ratio of transport latency to reaction latency, for translation elongation is ~4; physical transport of individual ternary complexes accounts for ~80% of elongation latency. We also model how molecules pack closer together as growth quickens. Although denser cytoplasm both decreases transport distances and hinders motion, we predict that decreasing distance wins out, offering a simple mechanism for how individual elongating ribosomes become more productive as growth quickens. We also quantify how crowding imposes a physical limit on the performance of self-mixing molecular systems and likely undergirds cellular behavior more broadly.

**Keywords:** Systems biology, colloidal physics, physics of life, molecular crowding, protein synthesis, translation elongation, cytoplasm structure

#### Introduction

Protein synthesis is essential for cell maintenance and reproduction. For example, *Escherichia coli* (*E. coli*) cells synthesize the majority of their dry mass as protein every cell doubling. Accordingly, cells that grow more quickly must produce proteins more quickly. In quantitative detail, as *E. coli* growth speeds up five-fold protein synthesis across the entire cell increases 15-fold (Dennis & Bremer, 2008). Meanwhile, for the same growth rate increase, the quantity of ribosomes increases only nine-fold (**Figure S1**), suggesting that the absolute productivity of individual ribosomes must also somehow increase – almost doubling as growth quickens (**Figure 1**) (Bremer and Dennis, 1996; Dalbow and Young, 1975; Dennis and Bremer, 2008; Forchhammer and Lindahl, 1971; Klumpp et al., 2013; Pedersen, 1984; Young and Bremer, 1976). While it is easy to understand why having more translation machinery increases total protein synthesis capacity, it is not obvious how faster growing cells achieve the translation elongation rates needed to sustain growth.



**Figure 1. Individual ribosomes make proteins more quickly as growth quickens.** Total latency per peptide bond (left y-axis) or elongation rate (right y-axis) versus growth rate (x-axis). Experimental data from Bremer and Dennis, 1996; Dalbow and Young, 1975; Dennis and Bremer, 2008; Forchhammer and Lindahl, 1971; Klumpp et al., 2013; Pedersen, 1984; and Young and Bremer, 1976. Solid line is a second-order polynomial fit of experimental elongation rates.

Bremer & Dennis hypothesized that individual ribosome activity speeds up at faster growth rates owing to increased tRNA charging and also due to shifts in codon distribution among mRNA (Bremer and Dennis, 1996). However, subsequent work has shown that overall tRNA charging remains relatively constant across growth rate indicating that other mechanisms are likely at play (Avcilar-Kucukgoze et al., 2016). Another possibility is that the intrinsic chemical kinetics of peptide bond formation by the ribosome accelerate with increasing growth rates. For example, in exploring how to adapt chemical kinetic rates obtained from in vitro experiments for use with in vivo models, Rodnina and co-workers fit parameter values to data and showed that faster chemical kinetic rate constants could account for increased rates of peptide bond formation (Rudorf et al., 2014). However, the specific molecular mechanisms that might account for such parameter changes are unknown. As a third possibility, Hwa and co-workers hypothesized that physical processes could play a limiting role in determining the elongation rate of individual ribosomes (Klumpp et al., 2013). More specifically, by accounting for Brownian diffusion of ternary complexes via a growth-rate independent diffusion constant within a Michaelis-Menten kineticsbased model of translation elongation, Hwa and co-workers inferred that physical changes in cytoplasm could lead to changes in growth rate. Taken together, such studies suggest that both chemistry and physics likely play a role in the speedup of translation elongation.

However, understanding any potential speedup mechanism is challenging exactly because the chemistry and physics of translation elongation are complex and coupled. For example, the biochemical processes required are combinatorial: matching must take place between 42 unique ternary complexes and 64 possible triplet codons. Accordingly, any particular elongating ribosome may encounter numerous mismatching ternary complexes prior to a successful matching reaction. As a second example, the length- and time-scales of underlying processes span three and nine orders of magnitude, respectively; specifically, ternary complexes and ribosomes interact with surrounding biomolecules and each other over nanometers and nanoseconds but execute processes over microns and seconds. As a third example, while higher concentrations of ternary complexes might be expected to increase the frequency of encounters with ribosomes, the resulting increase in crowding might slow the physical search process. Such complexities are compounded by the fact that everything is happening in parallel among hundreds of thousands of self-mixing molecules in a growth-rate dependent and crowded cytoplasm.

We address these challenges by modeling both the physics and chemistry of translation elongation in a combined framework. To do so, we adapted an open-source simulation tool (Andrews et al., 2010) to more accurately represent transport and interactions among molecules comprising self-mixing systems. In our framework we explicitly represent the transport dynamics of individual biomolecules as they physically interact and chemically react, with nanometer and nanosecond resolution, to simulate processes spanning minutes in time. A key aspect of our approach is the robust modeling of Brownian motion and colloidal-scale particle interactions such that these molecules undergo the inertialess physical encounters appropriate to the colloidal regime (Ermak and McCammon, 1977; Heyes and Melrose, 1993; Zia, 2018). When combined with a well-known multi-step kinetic model for the reactions leading to peptide bond formation (Kothe et al., 2004) our framework enables analysis of the combined physical and chemical dynamics underlying translation elongation.

We employed our framework to explore how protein synthesis rates in *E. coli* should be expected to change with growth rate, directly representing the growth-rate dependent cytoplasm via first-principles modeling of physical and chemical dynamics without parameter fitting. Starting from

well-established measurements of macromolecular composition and physical properties of *E. coli* cytoplasm at varying growth rates, we demonstrate how well-known changes in the composition of cytoplasm are entirely sufficient to account for the speedup of translation elongation by individual ribosomes. We also identify the detailed contributions of transport and reaction to total elongation latency by monitoring the trajectories of and reactions between ternary complexes and ribosomes in simulation, finding that transport is the dominant component defining elongation latency. We find that physiological cytoplasmic crowding speeds up the transport mechanism and thus elongation rates overall. We confirm that the expected speedup due to crowding is insensitive to changes in chemical kinetics needed to exactly match observed elongation rates. Finally, we explore how still-greater crowding, beyond naturally observed limits, should lead to a collapse of the colloidal-scale transport speedup mechanism that ultimately limits the performance of self-mixing living systems.

## **Results**

## I. Embedding chemical kinetics within physical transport

We constructed a spatially resolved chemical and physical framework to model the combined roles of reaction chemistry and transport physics in translation elongation (**Figure 2A, 2B**), tracking the time spent by ternary complexes unbound and in motion (transport latency,  $\tau_{\text{transport}}$ ) as well as reacting with mismatching or matching ribosomes (reaction latency,  $\tau_{\text{rxn}}$ ) until a matching reaction successfully completes. Together, transport latency and reaction latency make up elongation latency ( $\tau_{\text{elong}}$ ).

- To estimate reaction latencies we represented the molecular reactions between ribosomes and ternary complexes following the individual chemical steps of protein synthesis, accounting for differences due to reactions involving cognate, near-cognate, and non-cognate ternary complexes (**Figure 2C**). We used well-established *in vitro* kinetic measurements to parameterize our model (**Table S5**) and developed physiologically accurate distributions of expected reaction latencies via a Markov-process (**Methods**). We analyzed the resulting reaction latency distributions, finding that matching reactions between cognate ternary complexes and ribosomes take 42 ms on average when successful (68% probability) (**Figure S2**). We also found that cognate ternary complexes can be rejected (32% probability), in which case reactions take 1.4 ms on average. Mismatching reactions involving near-cognate or non-cognate ternary complexes take on average 4.6 ms and 1.4 ms, respectively. We did not consider mis-incorporation events due to their low overall likelihood (<1% probability).
- Next, recognizing that translation elongation takes place within a crowded cytoplasmic milieu, we developed a molecular-mechanistic model for how protein synthesis occurs as a physical process. To start, we estimated the smallest volume of cytoplasm sufficient to enable protein synthesis. We assumed that the molecules required for protein synthesis are homogeneously distributed within the nucleoid-excluded cytoplasm. This volume is ultimately determined by the concentration of ternary complexes as the most-limiting species. So defined, each 'translation voxel' contains exactly one of each of 42 unique ternary complexes, one or more ribosomes, and many crowder
- proteins (average-sized molecules that represent all other surrounding proteins) (Figure 2D).
- Brownian diffusion allows each molecule to sample the voxel volume and encounter one another.
- We modeled diffusion explicitly as a random walk of each of the molecules throughout the voxel
- where, due to their finite size, they exclude one another entropically or, in the case of a ternary
- 134 complex and unbound ribosome pair, initiate a reaction (Methods; Table S4).

Due to non-uniform codon usage and non-uniform relative abundance of each type of ternary complex (**Table S6**) as well as stochastic variation in the physical distribution of translation molecules in cytoplasm, for any given cell-wide condition, individual translation voxels should be expected to vary in the exact combinations of unique ternary complexes and elongating ribosomes. For example, a translation voxel might contain more than one of a highly abundant tRNA. Accordingly, starting from our basic translation voxel platform we constructed ensembles of thousands of translation voxels to capture the natural distribution of chemical identities and spatial configurations that, together, better represent the natural variation expected within cytoplasm. We used these more-accurate voxel ensembles to examine the physical and chemical mechanistic relationship between growth rate and elongation rate by simulation (below).

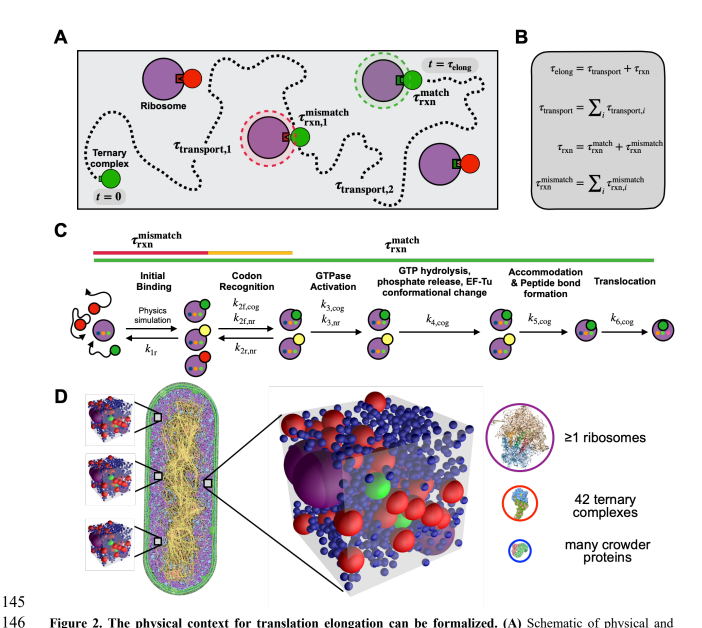


Figure 2. The physical context for translation elongation can be formalized. (A) Schematic of physical and chemical processes that contribute to translation elongation latency. Multiple transport and reaction steps (dashed line) may occur before a ternary complex (green/red) encounters and reacts with an unoccupied, matching ribosome (purple). The time ternary complexes spend unbound while searching for ribosomes is defined as transport latency  $(\tau_{\text{transport}})$  and the time ternary complexes spend bound in either mismatching (red shaded circle) or matching reactions (green shaded circle) is defined as reaction latency ( $\tau_{rxn}$ ). The time the entire process takes is defined as elongation latency ( $\tau_{\text{elong}}$ ). (B) Mathematical definitions of translation elongation latencies. Elongation latency ( $\tau_{\text{elong}}$ ) is the sum of transport latency ( $\tau_{transport}$ ) and reaction latency ( $\tau_{rxn}$ ), the latter of which is the sum of both mismatching ( $\tau_{rxn}^{mismatch}$ ) and matching  $(\tau_{rxn}^{match})$  reaction latencies. (C) Schematic of the kinetic mechanism of translation elongation within ribosomes (purple). Ternary complexes are either cognate (green), near-cognate (yellow), or non-cognate (red) to any particular ribosome, which determines kinetic rates. Mismatching reaction latency results from reversible reactions with non-cognate and near-cognate ternary complexes (red and yellow lines), while matching reaction latency results from cognate ternary complexes proceeding through the full kinetic process (green line). (D) Translation elongation is evaluated by constructing ensembles of statistically representative "translation voxels" that, in their minimal form, contain exactly 42 ternary complexes (cognate: green, non-cognate: red), at least one ribosome (purple), and numerous average-sized crowder proteins representing all other surrounding proteins (blue). Depiction of E. coli adapted with permission from Goodsell, 2009; molecular abundances adapted from literature (Main text and Methods).

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## II. Stoichiometric crowding accompanies faster protein synthesis

We gathered and analyzed well-established experimental data for cell mass, cell volume, and the sizes and abundances of ternary complexes, ribosomes, and crowder proteins in cells across growth rates (Dennis and Bremer, 2008; Volkmer and Heinemann, 2011; Dong et al., 1996; Pedersen et al., 1978; Schmidt et al., 2015; Woldringh and Nanninga, 1985) (Methods, Figure S1, Tables S1-S4). We noted that as growth rate increases (from  $\mu = 0.6$  to 3.0 dbl/hr) and translation elongation speeds up (from 12 to 21 aa/s), ternary complexes and ribosomes monotonically increase in number by nearly an order of magnitude, while crowder proteins monotonically increase by three-fold (Figure 3A). We described the coupled abundances and volume fraction of each constituent biomolecule as the 'colloidal stoichiometry' of the translation voxel; mathematically, the abundances and volume fraction of each constituent biomolecule *i* is described as  $N_i$  and  $\phi_i = V_i/V_{vox}$ , respectively, where  $V_i$  is the total volume occupied by a particular biomolecule species and  $V_{vox}$  is the total volume of the voxel. The colloidal stoichiometry of translation voxels – which captures both chemical and physical features of cytoplasm – changes with growth rate. Thus, we hypothesized that growth-rate dependent changes in colloidal stoichiometry might contribute to the speedup of translation elongation.

More specifically, our modeling revealed changes in the colloidal stoichiometry of translation voxels as growth quickens (0.6 to 3.0 dbl/hr): voxels shrink three-fold ( $V_{\text{vox}} = 10\text{E}5 \text{ nm}^3$ ) and become three-fold more crowded ( $\phi_{\text{vox}} = 0.13 \text{ to } 0.42$ ). However, the growth in packing fraction is not uniform across species: as voxels shrink in size the number of ribosomes doubles while the number of crowder proteins halves (**Figure 3B, 3D**). That is, increased total crowding is dominated by ribosomes: the volume fraction of ribosomes increases by seven-fold, more than double that of ternary complexes ( $\phi_{\text{rib}} = 0.03 \text{ to } 0.22$ ;  $\phi_{\text{tern}} = 0.04 \text{ to } 0.12$ ). Crowder proteins dominate the packing fraction at low growth rates but then plateau ( $\phi_{\text{crowder}} = 0.06 \text{ to } 0.10 \text{ for } \mu = 0.6 \text{ to } 2.0 \text{ dbl/hr}$ ;  $\phi_{\text{crowder}} = 0.10 \text{ to } 0.09 \text{ for } \mu = 2.0 \text{ to } 3.0 \text{ dbl/hr}$ ), thus contributing minimally to the overall increase in crowding (**Figure 3C**).

We referred to this growth-rate dependent change in colloidal stoichiometry as 'stoichiometric crowding,' which we expected should impact both the interactions and motion of translation molecules at different growth rates. For example, in the growth-rate trends noted above, as growth quickens ternary complexes and ribosomes should encounter each other more frequently relative to encountering crowders. As a second example, the distribution of molecule sizes matters: for a fixed total volume fraction the diffusion of individual particles is faster in a suspension of large versus small particles and, as the most dominant particle size shifts from smaller to larger, diffusion of all particles speeds up (Farris, 1968; Gonzalez et al., 2021; Lionberger, 2002); since ribosomes increase in their relative volume fraction with growth rate, voxels should mix more quickly for any given total volume fraction as growth quickens.

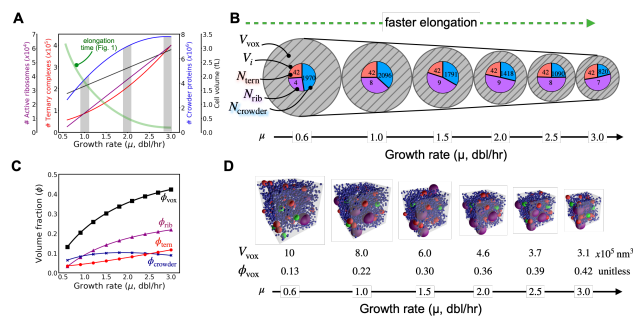


Figure 3. The relative abundances, concentrations, and volume fractions of translation molecules change as growth quickens. (A) Colloidal stoichiometry. Experimental observations of active ribosomes, ternary complexes, and crowder protein abundances in *E. coli*, as well as *E. coli* cell volume, reveal varying levels of increase with increasing growth rate (gray bars highlight values at particular growth rates). (B) Stoichiometric crowding. An abstracted representation of translation voxels as a function of growth rate reveals that differential changes in molecular abundances are accompanied by an overall increase in crowding (i.e., stoichiometric crowding). The volume of translation voxels ( $V_{\text{vox}}$ , hatched gray circles) decreases while the total volume of constituent biomolecules (concentric pie charts) remains relatively constant. The total number of each particular type of biomolecule species ( $N_{\text{tern}}$ ,  $N_{\text{rib}}$ , and  $N_{\text{crowder}}$ , shown within corresponding colors of the pie chart – red: ternary complexes, purple: ribosomes, blue: crowder proteins) in a given translation voxel as well as the total volume each biomolecule species occupies ( $V_i$ , the area of corresponding colors within the pie chart) change at different growth rates. (C) The volume fractions ( $\phi_i = V_i/V_{\text{vox}}$ ) of ribosomes ( $\phi_{\text{rib}}$ ), ternary complexes ( $\phi_{\text{tern}}$ ), and crowder proteins ( $\phi_{\text{crowder}}$ ) change differently with increasing growth rate, leading to an overall increase in the total volume fraction of translation voxels ( $\phi_{\text{vox}}$ ). (D) Representative snapshots of translation voxel simulations at increasing growth rates, along with their respective volumes ( $V_{\text{vox}}$ ) and volume fractions ( $\phi_{\text{vox}}$ ).

## III. Physical transport of ternary complexes accounts for most of elongation latency

We next sought to better understand and quantify any such impacts of crowding and composition on transport rates, reaction rates, and elongation rates overall. To do so we established a baseline quantification of the relative importance of physical transport to chemical reactions in setting elongation latency. We constructed translation voxels, from very simple to biologically faithful forms, and analyzed expected transport, reaction, and elongation latencies by simulation. From this we inferred the mechanisms by which colloidal stoichiometry regulates overall elongation latency as a function of growth rate. We used a slow growth rate (0.6 dbl/hr) as a benchmark, where bulk elongation takes about 87 ms on average (Figure 1).

More specifically, we first studied transport and reaction dynamics in detail using an idealized case in which only a single cognate ternary complex interacts with one matching ribosome (**Figure 4A**). Here, there is no competition with mismatching ternary complexes, no other molecules blocking the way, and the sought-after ribosome is unbound; rather, a lone ternary complex searches pure cytosolic fluid for a waiting matching ribosome, an idealized scenario often depicted in 'textbook' representations of translation elongation (e.g., Sannuga and Ramakrishnan, 2004).

- 230 As expected, we found that transport latency – the time a ternary complex spends not bound to a
- ribosome, diffusively searching for a match is nearly instantaneous ( $\overline{\tau}_{transport} = 0.08 \text{ ms}$ ) and 231
- that nearly all of the elongation process is taken up by reaction latency the time a ternary complex 232
- spends bound to ribosomes ( $\overline{\tau}_{rxn} = 42 \text{ ms}$ ) (Figure 4A). While such a result seems to support the 233
- conclusion that chemistry alone determines translation elongation rate (Figure 2C), the notion of 234
- 235 a two-molecule translation voxel operating at 3% volume fraction – far below physiological
- 236 conditions – is unrealistic (**Figure 3C**).
- 237 We next added a physiologically correct number of ternary complexes to the voxel (i.e., one
- 238 cognate, 41 non-cognate) such that each ternary complex competes to reach and bind to the
- ribosome (Figure 4B). We found that the total reaction time remains the same ( $\overline{\tau}_{rxn} = 42 \text{ ms}$ ). 239
- 240 However, the transport latency of the cognate ternary complex increases markedly ( $\overline{\tau}_{transport}$  =
- 251 ms) and is greater than reaction latency, supporting arguments that translation elongation 241
- 242 requires substantial reactant transport time. The increased transport time is also coupled to
- 243 reactions: a cognate ternary complex must "wait" to bind with the ribosome while that ribosome
- 244 is already bound to non-cognate ternary complexes; thus, there is an interplay between reactions
- 245 and transport.
- 246 We then added a physiologically correct number of ribosomes to the translation voxel, holding the
- 247 ternary complex population fixed at one cognate and 41 non-cognates (Figure 4C). Only one
- 248 ribosome was available for a matching reaction with the cognate ternary complex; the other
- 249 ribosomes were mismatching for all ternary complexes. Having just one matching ribosome means
- 250 we need track only the elongation events of a single ribosome, while simultaneously tracking
- 251 mismatching events at other ribosomes; this approximation provides a lower-bound estimate for
- 252 bulk elongation rates while allowing a more accurate accounting of transport and reaction effects
- 253 (below). We found by simulation that transport latency remains similar to voxels containing a
- single ribosome ( $\bar{\tau}_{transport} = 248 \text{ ms}$ ) because the single matching ribosome remains bound for 254
- almost the same amount of time, indicating that the cognate ternary complex still needs to 'wait' 255
- 256 nearly as long (i.e., the indirect impact of mismatching reactions). A slight decrease in transport
- 257 latency can be attributed to some non-cognate ternary complexes being bound by mismatching
- 258 ribosomes, meaning fewer non-cognate ternary complexes are available to occupy the matching
- ribosome. However, the reaction latency of the cognate ternary complex increases ( $\overline{\tau}_{rxn} = 63 \text{ ms}$ ) 259
- 260 due to the direct impact of mismatching reactions: cognate ternary complexes spend more time in
- 261 futile interactions with the more abundant mismatching ribosomes.
- 262 Next, we added a physiologically correct abundance of crowder proteins to the translation voxel
- resulting in further increases in both transport and reaction latencies ( $\overline{\tau}_{transport} = 396 \text{ ms}$ ,  $\overline{\tau}_{rxn} =$ 263
- 264 73 ms) (Figure 4D). These predicted increases arose because crowders increase the number of
- mismatching reactions by trapping non-cognate ternary complexes near ribosomes, which in turn 265
- 266 both promotes repeated mismatch reactions and reduces cognate ribosome availability.
- 267 Surprisingly, the direct impact of crowding on transport – slower diffusion – was minor compared
- 268 to the effects of increased mismatching.
- 269 Finally, we represented the expected statistical variation in cytoplasm by constructing thousands
- 270 of different voxels that, together, capture the physiological distribution of relative abundances of
- 271 translation molecules as reported in the literature (Methods). Specifically, in E. coli, there are 42
- 272 unique ternary complexes, each with its own abundance, and 64 codons, each with its own usage
- 273 rates, and these are present in many permissible combinatoric configurations in translation voxels

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throughout cytoplasm (Figure 2D). We randomly sampled all permissible configurations using reported E. coli codon usage and whole-cell tRNA abundances (Methods, Table S6, Figure S4). Recognizing that bulk elongation measurements correspond to the time needed to complete as many successful reactions as ribosomes are in a voxel, we computed the transport, reaction, and elongation latencies of each translation voxel from the time taken for a single matching reaction within the voxel. A weighted average of the per-ribosome latency for all permissible translation voxels corresponds to the typical time for just one matching reaction to occur in a voxel and thus provides a lower-bound estimate of bulk experimental elongation time as obtained from cellular measurements (Methods). While our prior simulations (Figures 4A - 4D) included only noncognate and cognate ternary complexes, our calculations of the ensemble latencies (Figure 4E) also included the more detailed classification of some ternary complexes as near-cognate, which affects system dynamics further because near-cognates are well-known to have a slower rejectiontime than non-cognates (Table S5, Figure S2). We monitored transport, reaction, and elongation latency during simulation in each of these thousands of voxels, and computed a weighted-average value for transport, reaction, and elongation latency. We found that the weighted-average transport, reaction, and elongation latencies decrease in the ensemble representation ( $\overline{\tau}_{transport}$  = 242 ms,  $\bar{\tau}_{rxn} = 56$  ms,  $\bar{\tau}_{elong} = 298$ ) (Figure 4E), compared to the single-voxel simulation (Figure 4D). This across-the-board decrease in latencies emerges naturally from the majority of voxels in which there is more than one cognate ternary complex, partly a result of the biological phenomenon of more frequently used codons being associated with more abundant cognate tRNAs (Figure S8).

Quantitatively, the Damköhler number – the ratio between the latency of transport and reaction (Da =  $\tau_{transport}/\tau_{rxn}$ ) – highlights the dependency of translation elongation latency on physical transport relative to chemical reaction. In our simplest model, Da ~ 0, suggesting that reaction latency dominates elongation latency. However, our increasingly accurate models estimate Da ~ 6, Da ~ 4, Da ~ 5, and Da ~ 4 respectively (**Figures 4B – 4E**). We thus concluded that processes that modulate transport latency play a dominant mechanistic role in regulating the overall speed of translation elongation.

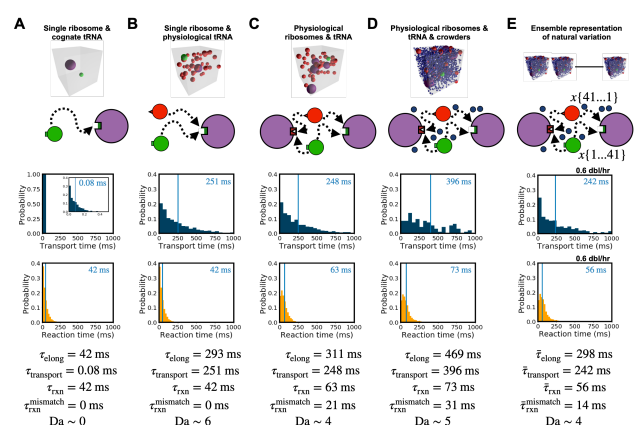


Figure 4. Most of the latency in translation elongation arises from physical transport of ternary complexes. (A-E) Simulation snapshots, model schematics, and simulation results (top to bottom) for increasingly realistic (left to right) translation voxels at growth rate  $\mu=0.6$  dbl/hr. In each plot the average latency is marked by a blue vertical line and displayed on the top right in milliseconds. (A) A highly simplified translation voxel containing only a single ribosome and cognate ternary complex. (B) A translation voxel containing a single ribosome and 42 ternary complexes. (C) A translation voxel with 42 ternary complexes and four ribosomes. (D) A translation voxel with 42 ternary complexes, four ribosomes, and 1970 crowder proteins. (E) An ensemble of translation voxels that capture the expected natural variation in cognate, near-cognate, and non-cognate ternary complexes due to non-uniform ternary complex and codon abundances coupled with spatial stochasticity.

#### IV. Stoichiometric crowding speeds up translation elongation

We returned to the puzzle of what mechanism(s) might cause the productivity of individual ribosomes to increase with increasing growth rates. We first evaluated the impact of stoichiometric crowding on transport latency by considering both molecule proximity (i.e., how close molecules are to one another) and molecular mobility (i.e., how fast molecules move). We also evaluated the impact of stoichiometric crowding on reaction latency by considering both local availability (i.e., to what extent ternary complexes are free from repeated mismatching reactions) and global availability (i.e., to what extent ternary complexes are free from mismatching reactions generally). Taken together we determined if and how each of these coupled physico-chemical mechanisms might regulate elongation latency as a function of growth rate.

We hypothesized that crowding should tend to reduce transport latency because ternary complexes need to search smaller volumes to find a matching ribosome. To explore this idea, we computed the average surface-to-surface distance between ternary complexes and their closest ribosomes (i.e., the shortest distance a ternary complex needs to travel to find a ribosome) across hundreds of

translation voxels at multiple growth rates (**Methods**). We found that stochiometric crowding brings ternary complexes and ribosomes five-fold closer on average (16 nm to 3 nm), which supports our hypothesis (**Figure 5A**, **left axis**). But stoichiometric crowding could also increase transport latency as ternary complexes become hindered in their motion and thus take longer to search. To explore this second idea, we examined the influence of stoichiometric crowding on molecule mobility by estimating the viscosity of cytoplasm as well as the hindered diffusivity of ternary complexes, ribosomes, and crowder proteins via simulation of hundreds of translation voxels at multiple growth rates (**Figure 5A**, **right axis**; **Figure S3**; **Methods**). We found that stochiometric crowding increases viscosity monotonically (1.0 to 2.4, normalized to viscosity at  $\mu = 0.6$  dbl/hr) while reducing diffusivity monotonically for all biomolecules (e.g., the diffusivity of ternary complexes,  $D_{\text{tern}}$ , slows from 35  $\mu$ m<sup>2</sup>/s to 16  $\mu$ m<sup>2</sup>/s), which would support the opposite conclusion: that crowding should hinder transport. Recognizing this competition between an increased proximity reducing transport latency and an increased viscosity increasing transport latency, we more systematically considered the contribution of each mechanism to transport as stoichiometric crowding increased due to increased cell growth rate.

For example, we simulated ensembles of translation voxels representing the full statistical distribution of ternary complexes and codon abundances from low- to high-growth rates (Methods). We found that transport latency monotonically decreases with stoichiometric crowding ( $\bar{\tau}_{transport} = 242$  ms to 83 ms) (Figure 5B). We deduced that, mechanistically, crowding drives faster transport because reducing the search distance between ternary complexes and ribosomes is more important than increased viscosity. However, while this net decrease in transport latency will decrease elongation latency overall, it could be that coupled changes in reaction latency either reverse or reinforce this trend.

Thus, we examined two mechanisms that could modulate reaction latency. First, we recalled that crowder proteins can induce repeat reactions by trapping ternary complexes and ribosomes together (Section III). These repeat reactions should reduce the local availability of ternary complexes, making it more difficult for ternary complexes to find matching ribosomes, driving up reaction latency. To examine whether increased stoichiometric crowding amplifies this effect, we tracked the number of times ternary complexes consecutively re-react with the same ribosome following a mismatching reaction across hundreds of translation voxels at varying growth rates. We were surprised to find that repeated reactions decrease five-fold (from  $\sim 10$  to  $\sim 2$  repeat reactions per ribosome on average) as growth rate quickens and total crowding increases (Figure **5C**, left axis). We resolved this apparent paradox by recognizing that the increased crowding arises primarily due to tighter packing of ribosomes, while the volume fraction of crowder proteins hardly changes. We deduced that this provides more local ribosome alternatives (i.e., higher local availability) for ternary complexes but with no increase in trapping by crowders (Figure S6). However, having more ribosomes, regardless of how well packed they are, provides more opportunities to preoccupy ternary complexes in mismatch reactions, reducing ternary complex global availability, which should drive up reaction latency. We found that the number of mismatching ribosomes in a voxel first increases and then decreases with growth rate, which should contribute an initial increase and then decrease in reaction latency as growth rate increases (Figure 5C, right axis). Taken together, the total impact on reaction latency depends on the relative strengths of each of these effects.

We next computed reaction latencies across our ensembles of translation voxels, capturing how physiological variation influences the competition between local and global availability. We found

- that total reaction latency increases ( $\overline{\tau}_{rxn} = 56 \text{ ms}$  to 70 ms) at low but increasing growth rates and then decreases monotonically thereafter ( $\overline{\tau}_{rxn} = 70 \text{ ms}$  to 52 ms) (**Figure 5D**). Low global
- availability of ternary complexes dominates at low growth rates, slowing reaction latency as
- growth rate increases. However, at higher growth rates, the increase in both global and local
- availability combine to drive down reaction latency. Overall, the growth-rate trend in reaction
- 376 latency (Figure 5D) follows ternary complex global availability (Figure 5C, right axis).
- Practically, even with a 25% increase that subsequently reverses, reaction latency changes only
- 378 7% over low-to-high growth rates, suggesting that transport plays the more substantial role in
- 379 speeding elongation.
- Indeed, the quantitative speedup of reaction latency with growth rate (Figure 5B) is minor
- compared to the corresponding speedup of transport latency (**Figure 5D**), indicating that transport
- mechanisms should be expected to dominate over reaction mechanisms in regulating the growth-
- 383 rate dependent productivity of individual ribosomes. Our ensemble simulations show that the
- dominance of transport manifests in the total elongation latency as a monotonic speedup of
- elongation with growth rate ( $\bar{\tau}_{elong} = 298 \text{ ms to } 135 \text{ ms}$ ), recovering the experimental trend of
- faster elongation at higher growth rates ( $\tau_{\text{elong}}^{\text{bulk}} = 83 \text{ ms to } 48 \text{ ms}$ ) (**Figure 5E**).
- Finally, although our model correctly predicts and recovers the qualitative behavior and overall
- trend (i.e., an increase in ribosome productivity with increasing growth rate), we noted that our
- 389 unfitted bottom-up modeling and simulations result in absolute predictions of translation
- elongation latencies that are ~3-fold too slow compared to experimental observations (**Figure 5E**).
- 391 Thus, we conducted sensitivity analyses in which chemical kinetic rates were fitted to match
- observed overall translation elongation latency (Supplement). We found that the speedup in
- translation elongation is insensitive to changes in the chemical kinetics of translation elongation
- 394 (**Figure S13**).

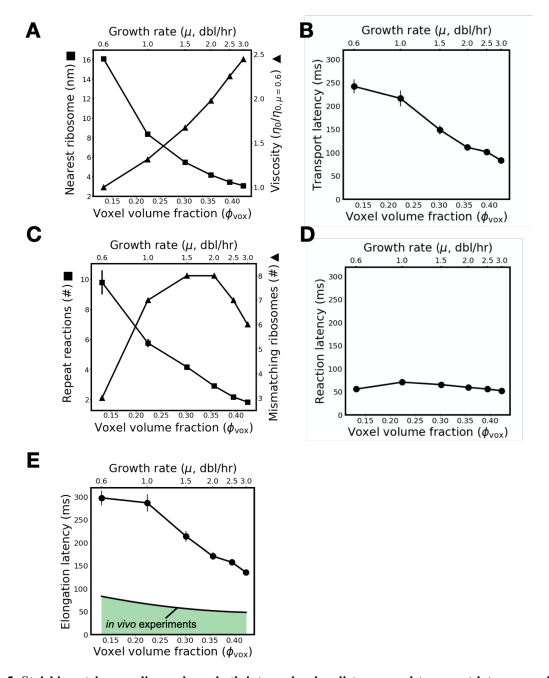


Figure 5. Stoichiometric crowding reduces both intermolecular distances and transport latency resulting in increasingly productive ribosomes as growth rate increases. (A) As crowding and growth rate increase (x-axes) ternary complexes become closer to their nearest ribosome (left y-axis) and translation voxel viscosity increases (right y-axis). Distance is reported as a surface-to-surface estimate. Viscosity is reported normalized to viscosity at  $\mu = 0.6$  dbl/hr. (B) Simulation results showing that transport latency (y-axis) decreases with increased crowding and growth rate (x-axes). (C) As crowding and growth rate increase (x-axes) the average number of repeat reactions between ternary complexes and ribosomes decreases (left y-axis) while the absolute number of mismatching ribosomes in a translation voxel first increases then decreases (right y-axis). (D) Simulation results showing that reaction latency (y-axis) first increases then decreases with increased crowding and growth rate (x-axes). (E) Simulation results showing that the predicted absolute elongation latency (filled circles, solid line) decreases with increased crowding and growth rate (x-axes). Experimentally measured per-ribosome elongation latency (solid line upon green area; replotting of Figure 1) also speeds up with growth rate but is faster than predicted across all growth rates. The standard errors in the estimate of the mean for all model results (A-E) are shown (error bars).

#### Discussion

The observed increase in individual ribosome activity as growth quickens has not been explained mechanistically (**Figure 1**). While many have focused attention on how translation initiation contributes to protein synthesis latency, we were intrigued by how the individually fast steps of translation elongation add up during translation elongation and should dominate overall process latency (**Supplement**). In the context of translation elongation, while prior studies of protein synthesis rates focused on chemical kinetic measurements or kinetics-based modeling, there have been persistent signals that Brownian diffusion of translation molecules plays a role in setting elongation rates.

With this in mind, we explored the idea that either chemistry or physics, or both, contribute to the speedup of translation elongation utilizing dynamic simulations. We proposed that reactions between ternary complexes and ribosomes are nontrivially coupled to their physical transport and that understanding this coupling is essential to explaining increased ribosome productivity at faster growth rates. To systematically interrogate the role of coupled physico-chemical processes in translation elongation, we adapted an open-source simulation tool to accurately represent transport of and interactions between translation molecules in cytoplasm. A key aspect of our approach is the robust modeling of Brownian motion and colloidal-scale particle interactions such that these molecules undergo the inertialess physical encounters appropriate to the colloidal regime. We defined translation voxels as naturally emergent from the constituent biomolecules required for translation and captured the natural distribution of chemical identities and spatial configurations of translation molecules in cytoplasm by constructing ensembles of thousands of voxels. We monitored in simulations the reactions and transport of molecules in these voxel ensembles to study the physical and chemical mechanistic relationships between growth rate and elongation rate (Figure 2).

We found that transport latency – the time ternary complexes spend searching for cognate ribosomes – is an essential component of elongation latency. Furthermore, we predict that transport latency dominates over reaction latency – the time ternary complexes spend reacting with ribosomes (Figure 4). By examining the elongation process as growth rate increases, we identified two competing mechanisms that underlie transport latency – proximity between ternary complexes and ribosomes that sets search distance, and cytoplasmic crowding that sets diffusive speed. Additionally, we observed that translation molecules become three-fold more crowded with increasing growth rate, suggesting that, beyond any absolute increase in the abundance of translation machinery, the machinery itself becomes packed closer together. The abundance and packing are physical as well as chemical (i.e., colloidal stoichiometry) and their changing due to changes in growth rate is a phenomenon we call stoichiometric crowding (Figure 3). We determined that the stoichiometric crowding associated with faster growth rates produces increased ribosome activity by reducing transport latency, revealing a mechanistic explanation for why individual ribosomes can produce proteins more quickly in faster-growing cells. In more detail, we predict that stoichiometric crowding reduces transport latency by increasing proximity: ternary complexes are closer to ribosomes, a change that outcompetes concomitant increases in viscosity and reaction latency to realize faster elongation (Figure 5).

We stress-tested our model and confirmed that the speedup of elongation requires physical transport, and that our prediction of speedup is robust to changes in the values of the input chemical parameters. We also found that increasing three-fold the values for all nine *in vitro* literature values for chemical kinetics parameters closes the quantitative gap between predicted and observed

elongation rates (**Figure S13**). This suggests the rather straightforward chemistry-only explanation for the gap: *in vitro* measurements being "off" by  $\sim 300\%$ , uniformly across all nine parameters. But, interestingly, we also found a 30-fold increase in only the ternary complex unbinding rate  $(k_{1r})$  – the only reaction that takes place exclusively outside the ribosome – could also close the quantitative gap (**Figure S13**, **Figure S9**), suggesting there may be a mechanism involved *in vivo* that quickens ternary complex exchange or one that obviates the need for fast rejection (i.e., a mechanism for favoring matching reactions near to the ribosome).

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Overall, our model reveals new opportunities for discovery. For example, better representation of electrostatic and hydrodynamic interactions or detailed molecular shape and orientation for site-specificity may be useful. More specifically, attractive interactions between the ribosomal L7/L12 domain and ternary complexes (Mustafi and Weisshaar, 2018) or between cognate ternary complexes and mRNA (Grosjean and Chantrenne, 1980) could have the effect of pre-loading or pre-sorting ternary complexes. As a second example, hydrodynamic models of small and large particles confined in a cavity show that both types of particles tend to concentrate near the cavity surface with minor impact on the mobility of small particles (Gonzalez et al., 2021), indicating that ternary complexes and ribosomes may concentrate by the cell membrane (not currently represented in our model) and effectively improve in proximity to each other.

More generally, our work supports exploration of the role of coupled, colloidal-scale physicochemical interactions in cytoplasm. For example, we predicted that ternary complexes and ribosomes should be up to five-fold closer together in faster growing cells (Figure 5A), a major shift in the colloidal-scale structure of cytoplasm that can be expected to modulate molecular interactions broadly across the cytoplasm. Such colloidal-scale structure is being increasingly measured experimentally (e.g., ribosome spatial positioning via cryo-electron tomography of entire cells) and merits increased attention for its role in cytoplasm behavior (Ortiz et al. 2006, Robinson et al. 2007). As a second example, the phenomenon of repeat reactions we found critical to the speedup of protein synthesis has also been identified as critical to efficient activation of Mitogen-Activated Protein (MAP) Kinases (Takahashi et al., 2010), suggesting a wider role for repeat reactions in cell functions. One can also infer the possibility that stoichiometric crowding with changing growth rate may impact cell signaling in general. As a third example, using our model we predicted that the viscosity of the nucleoid-excluded cytoplasm increases up to 2.5-fold with growth rate, which indicates a decrease in the mobility of all constituent molecules. Such a broad growth-rate dependent shift in colloidal-scale dynamics may suggest currently unappreciated forms of physical regulation in cells and motivates a renewed analysis of diffusive processes in cells with consideration to volume fraction and growth rate.

Advances in both computational modeling and experimental technique are needed to improve the accuracy of our predictions and promote broader exploration for how coupled colloidal-scale physics and chemistry in cytoplasm might regulate cellular behaviors. For example, dynamic simulation of the motion of solvent-suspended particles requires discretization of the time domain, where equations of motion are integrated forward in time. The selection of time step size impacts not only computational expense but also the fidelity of particle encounters where, for example, too-large timesteps can produce pathological displacements in response to steeply attractive or repulsive forces (e.g., the hard-sphere repulsion that represents entropic exclusion is singular at particle contact), a phenomenon that becomes more severe as crowding increases. Here, we performed a careful study to prioritize physical accuracy first and then optimize efficiency by, for example, developing a kinetic scaling method that leveraged the natural disparity between

diffusive and reactive timesteps (**Figures S3, S5, S10**). Even so, further capturing the complexity of cytoplasm (e.g., general protein-protein interactions, polysome dynamics, or cell-cycle dependency) will ultimately require modeling other microscopic forces at play in cytoplasm, including electrostatic or hydrodynamic interactions or membrane confinement, all of which lead to many-body interactions that increase computational expense. Modeling such forces, in addition to other molecular details like shape, softness, flexibility, and site-specificity, is becoming possible with other algorithms such as Stokesian dynamics for large or confined systems (Aponte-Rivera et al., 2018; Gonzalez et al., 2021; Maheshwari et al., 2019; Ouaknin et al., 2021; Zakhari et al., 2017), but will require substantial integration and iteration with experiments as well as improvements in computational efficiency to achieve accurate simulations over the timescales of cellular behavior (Endy and Brent, 2001). Capturing detailed molecular dynamics, such as those involved with ternary complex-ribosome binding and reactions, will also necessitate multi-scale modeling and experimentation from atomic to cellular scales.

In closing, we note that protein synthesis is inextricably tied to growth rate and fitness; cells cannot grow more quickly than they can reproduce their proteome, including the proteins that remake the proteome. Since stoichiometric crowding facilitates faster protein synthesis, could further crowding enable still-faster growth or, conversely, limit how fast cells can grow? To speculate, as a simple extension to our modeling, we projected what cytoplasm of faster-than-observed growing *E. coli* would look like (**Figure S11, Methods**). We found that *E. coli* would eventually reach maximum packing (i.e., with little to no space for molecular mixing) as growth rate continues to increase (**Figure 6A**). That such hypothetical growth rates are not observed suggests, among other possibilities, that as growth rate increases beyond the maximum observed, the beneficial effects of increased crowding (e.g., increased proximity) become outpaced by the deleterious effects of less available free volume (e.g., further increased viscosity) (**Figure 6B**). This hints that stoichiometric crowding may be linked to fitness and evolutionary fine tuning. If so, then we would expect that genes encoding currently unknown functions may serve to establish a physical, as well as chemical, basis for fitness (e.g., proteome polydispersity) undergirding cellular behavior broadly.

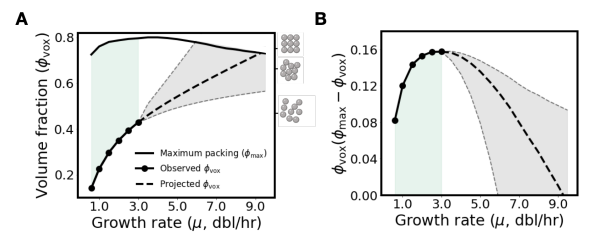


Figure 6. Stoichiometric crowding has diminishing returns that may impose a physical limit on growth rate. (A) Volume fraction of translation voxels at observed growth rates and projected growth rates compared to theoretical maximum random close packing (maximum packing changes with size polydispersity (Farr and Groot, 2009) and size polydispersity changes with growth rate). Bounds for volume fraction at projected growth rates are shown (gray dashed line and shading). Maximum packing increases across observed growth rates (green shading). The volume fraction for the most-crowded observed voxel ( $\phi_{vox} = 0.42$ ,  $\mu = 3.0$  dbl/hr) is shown (right axis, bottom schematic). For reference, the volume fractions at which long-term molecular motion in monodisperse suspensions is hindered or halted due to random close packing ( $\phi = 0.64$ ) or crystallization ( $\phi = 0.74$ ), respectively, are also shown (right axis, top and middle schematic). (B) The product of voxel volume fraction ( $\phi_{vox}$ ) and remaining available volume fraction ( $\phi_{max} - \phi_{vox}$ ) increases across observed growth rates (green shading) before decreasing across faster-than-observed growth rates.

#### Methods

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## I. Construction of a representative translation voxel

- We developed computational representations of translation voxels by analyzing the abundances
- and sizes of molecules comprising *E. coli* cytoplasm in relation to overall cell volume and mass.
- Where needed we inferred abundances across growth rates by fitting polynomials to reported
- measurements (described below).
- 550 I.1. Calculation of biomolecular abundances in cells
- We computed the average abundances of ribosomes  $(N_{\text{rib}})$  and ternary complexes  $(N_{\text{tern}})$  in single
- cells across physiological growth rates using existing data from literature (Table S1). We first
- calculated the total dry mass of crowder proteins in cytoplasm,  $M_{\text{cytoplasm,crowder}}$  (Equation 1).
- 554 Crowder protein mass encompasses the mass of all biomolecules in cytoplasm other than
- ribosomes, ternary complexes, mRNA, and DNA:

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$$M_{\text{cytoplasm,crowder}} = M_{\text{cytoplasm}} - M_{\text{tern}} * N_{\text{tern}} - M_{\text{rib}} * N_{\text{rib}} - M_{\text{cytoplasm, mRNA}} - M_{\text{cytoplasm,DNA}}$$
. (1)

- The masses of ternary complexes  $(M_{\text{tern}})$  and ribosomes  $(M_{\text{rib}})$  were specified using known
- 558 molecular structures and the total mass of mRNA, DNA, and cytoplasm
- $(M_{\text{cytoplasm,mRNA}}, M_{\text{cytoplasm,DNA}}, \text{ and } M_{\text{cytoplasm}}, \text{ respectively}) \text{ were taken from literature } (\textbf{Tables S4},$
- 560 S1). We then calculated the average effective spherical radius of crowder proteins ( $\overline{R}_{crowder}$ ), as
- well as the mass of resulting average-sized crowder proteins  $(\overline{M}_{crowder})$ , using single-cell E. coli
- mass spectrometry data and average protein density ( $\rho_{crowder}$ ) (CrowderProteinComputation.py):

$$V_{\text{crowder},i} = \frac{M_{\text{crowder},i}}{\rho_{\text{crowder}}}, \qquad (2)$$

$$\overline{R}_{\text{crowder}} = \sum_{i} \frac{N_{\text{crowder},i}}{N_{\text{crowder}}} \sqrt[3]{\frac{3V_{\text{crowder},i}}{4\pi}}, \qquad (3)$$

$$\overline{M}_{\text{crowder}} = \frac{4}{3}\pi(\overline{R}_{\text{crowder}}^3)(\rho_{\text{crowder}}). \tag{4}$$

- We obtained the mass and abundances of each protein in the E. coli cytoplasm  $(M_{crowder,i})$  and
- $N_{\text{crowder},i}$  respectively), as well as the total number of proteins ( $N_{\text{crowder}}$ ), from the mass
- 570 spectrometry measurements of Heinemann and colleagues (Schmidt et al., 2016). We also
- computed the volume of each protein in the E. coli cytoplasm ( $V_{\text{crowder},i}$ ) using these measurements.
- We then determined the abundance of crowder proteins in a cell as the ratio of total mass occupied
- by crowder proteins in cytoplasm (Equation 1) and the mass of average-sized crowder proteins
- 574 (Figure S1B),

$$N_{\text{crowder}} = \frac{M_{\text{cytoplasm,crowder}}}{\overline{M}_{\text{crowder}}} \,. \tag{5}$$

- 576 I.2. Calculation of translation voxel size and biomolecular abundances in translation voxels
- On a relative basis amino-acid specific ternary complexes are the least-concentrated molecules
- 578 involved in translation elongation and thus their concentration determines the minimum volume
- of cytoplasm capable of supporting protein synthesis. Therefore, we determined the size of
- translation voxels from our estimates of ternary complex abundances ( $N_{\text{tern}}$ ) and E. coli volume

- $(V_{\text{cell}})$  across growth rates (**Figure S1, Tables S1, S2**). More specifically, we defined a translation 581
- voxel to be the volume of cytoplasm that contains 42 ternary complexes ( $N_{\text{tern,vox}} = 42$ ), assuming 582
- 583 a spatially homogeneous distribution of ternary complexes within cytoplasm:

$$V_{\text{vox}} = \frac{N_{\text{tern,vox}} * V_{\text{cell}}}{N_{\text{tern}}} \,. \tag{6}$$

- 585 We then calculated the average number of crowder proteins ( $N_{\text{crowder, vox}}$ ) and ribosomes ( $N_{\text{rib, vox}}$ )
- within the translation voxel volume  $(V_{vox})$  assuming a homogeneous distribution of both species, 586
- but with ribosomes excluded by the nucleoid (Figure S1C): 587

$$N_{\text{crowder, vox}} = \frac{N_{\text{crowder}} * V_{\text{vox}}}{V_{\text{cell}}},$$

$$N_{\text{ribosome, vox}} = \frac{N_{\text{ribosome}} * V_{\text{vox}}}{V_{\text{cell}}(1 - \phi_{\text{nucleoid}})}.$$
(8)

$$N_{\text{ribosome, vox}} = \frac{N_{\text{ribosome}} * V_{\text{vox}}}{V_{\text{cell}} (1 - \phi_{\text{nucleoid}})}.$$
 (8)

- We estimated the volume fraction of the nucleoid ( $\phi_{\text{nucleoid}}$ ) based on published values (Table 590 591
- 593 I.3. Polynomial regression fitting

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We computed polynomial fits for ribosome abundances, ternary complex abundances, cell mass, cell volume, and nucleoid volume using a bootstrapping method that minimized the mean absolute error (Figure S1A, TranslationVoxelParameterization.py); we used mean absolute error instead of mean squared error to penalize all variation equally.

#### II. Calculation of translation voxel volume fractions and polydispersity

We calculated the volume fraction of ribosomes, ternary complexes, and crowder proteins (Figure 600 **3C**) using ribosome, ternary complex, and crowder protein abundances in translation voxels: 601 602 ribosome, ternary complex, and crowder protein single-molecule volumes; and translation voxel 603 volume:

$$\phi_{\rm rib} = \frac{N_{\rm rib, \, vox} * V_{\rm rib}}{V_{\rm vox}} \,, \tag{9}$$

$$\phi_{\text{tern}} = \frac{N_{\text{tern, vox}} * V_{\text{tern}}}{V_{\text{vox}}}, \qquad (10)$$

$$\phi_{\text{crowder}} = \frac{N_{\text{crowder, vox}} * \overline{V}_{\text{crowder}}}{V_{\text{vox}}}.$$
 (11)

- 607 ribosomes, ternary complexes, and  $(N_{\text{rib,vox}}, N_{\text{tern,vox}}, N_{\text{crowder,vox}})$  in translation voxels as well as the translation voxel volume  $(V_{\text{vox}})$ 608
- are as described in Equations 6, 7, and 8. The volume of a single average-sized crowder protein 609
- $(\overline{V}_{crowder})$  is defined as its mass divided by average protein density (**Equation 4**). We computed 610
- the volume of single ribosomes  $(V_{rib})$  and ternary complexes  $(V_{tern})$  based on their longest length 611
- (i.e., estimating the molecules as spheres with diameters equal to the longest length of the 612
- 613 molecules), which we measured from their detailed atomic resolution structures (PDB 4V4Q and
- 614 PDB 1B23, respectively) (**Table S4**).
- We calculated the size polydispersity, s, of translation voxels as in Fairhurst, 1999: 615

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$$s = \sqrt{\frac{\sum N_{i, \text{vox}} * \sum N_{i, \text{vox}} R_i^2}{\left(\sum N_{i, \text{vox}} R_i\right)^2} - 1}.$$
 (12)

Here,  $N_{i,vox}$  and  $R_i$  correspond to the translation voxel abundances and effective spherical radius of each biomolecule type, denoted by the subscript i, respectively (**Table S4**).

#### III. Simulation of translation voxels

We simulated transport and reaction of biomolecules within translation voxels using Brownian dynamics and single-molecule reaction kinetics, respectively. We implemented our simulations using "Colloidal Smoldyn", our adaptation of the open-source simulation software Smoldyn (Andrews et al., 2010). Colloidal Smoldyn accurately represents single-molecule resolution colloidal transport dynamics and reaction dynamics as described below.

- 626 III.1. Simulating colloidal transport dynamics
- Here, we briefly describe the theoretical framework that underlies Colloidal Smoldyn and then provide simulation implementation details. The transport dynamics of water-suspended particles of size ~ 1.5 nm to O(10) microns (colloids) have been thoroughly studied and characterized in the microhydrodynamics and colloid physics literature. Fluid motion generated by particle motion at these length scales is governed by the Stokes equations, while particle motion is described by the N-particle Langevin equation (Langevin, 1908),

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$$\boldsymbol{m} \cdot \frac{d\boldsymbol{U}}{dt} = \boldsymbol{F}^B + \boldsymbol{F}^P + \boldsymbol{F}^{ext} + \boldsymbol{F}^H, \tag{13}$$

a stochastic force balance, where particle momentum  $m \cdot dU/dt$  equates the sum of the forces F that act on the particles. Here m is a diagonal tensor containing the details of the mass of the particles and U is the particles' velocity vector. The forces F represent the stochastic forces  $F^B$  that give rise to Brownian motion, the deterministic interparticle forces  $F^P$  that here arise from a hard-sphere potential, the external forces  $F^{ext}$  that could represent either active motion or externally induced motion, and hydrodynamic drag forces  $F^H$  that arise due to the difference between the particle velocity U and imposed fluid motion  $u^\infty$ . Each of the vectors (U and F) has dimensions 3N describing the three-dimensional space in which all N particles move. In our system there are no external forces on the particles, thus  $F^{ext} = 0$ .

The Brownian force  $F_i^B$  on a given particle "i" arises from collisions between the colloidal particle and the solvent molecules as the fluid fluctuates thermally and follows Gaussian statistics,

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$$\overline{F_i^B} = \mathbf{0}$$

$$\overline{F_i^B(0)F_i^B(t)} = 2kT(6\pi\eta a_i)I\delta(t), \tag{14}$$

where k is the Boltzmann constant, T is the absolute temperature,  $\eta$  is the solvent viscosity,  $a_i$  is the particle radius, I is the identity tensor, and  $\delta(t)$  is the Dirac delta function. The overbar considers averages over times long compared to the solvent molecule timescale, such that a colloidal particle has undergone many random and decorrelated impacts from solvent molecules. Although the mean Brownian force is zero, its covariance,  $2kT(6\pi\eta a_i)I$ , has an amplitude set by the fluctuation-dissipation theorem, which dictates that colloidal motion due to thermal forces is dissipated viscously back into the solvent. In our system, we approximated the drag force on the colloid,  $(6\pi\eta a_i)I$ , to be constant, where each particle experiences solvent drag and excludes other particles entropically with finite size, but coupled hydrodynamic interactions are neglected. This

common simplification, the "freely draining model" is appropriate for gaining understanding of the basic colloidal physics of a suspension (Hoh and Zia, 2016a, 2016b; Russel, 1984; Russel et al., 1989; Zia and Brady, 2010) and is accurate in many physiological conditions. Incorporating hydrodynamic interactions in such models is routine (Banchio and Brady, 2003; Durlofsky et al., 1987; Ouaknin et al., 2021; Sierou and Brady, 2001) although computationally expensive and will be discussed in future work.

Interactions between colloidal particles can be attractive or repulsive and range from short- to long interparticle distances. In the present work, biomolecules are represented as hard spheres: they interact only at contact at which point we stipulate that they may not overlap. This can be represented mathematically via a hard-sphere potential that enforces an infinite penalty for overlap. The gradient of this entropic exclusion gives rise to a deterministic interparticle force between two particles of radius  $a_i$  and  $a_i$ 

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$$\boldsymbol{F}_{i}^{P} = \frac{a_{j}}{a_{j} + a_{i}} kT \delta \left[ r_{ij} - \left( a_{j} + a_{i} \right) \right] \hat{\boldsymbol{r}}_{ij}, \tag{15}$$

where  $r_{ij}$  is the interparticle distance between the particles' centers,  $\hat{r}_{ij}$  is the unit vector along the line that connects the particle centers, and the delta function  $\delta$  enforces the infinite penalty for overlap.

Finally, the hydrodynamic drag forces are given by the Stokes drag on each particle according to

$$\mathbf{F}_{i}^{H} = 6\pi\eta a_{i}(\mathbf{U}_{i} - \mathbf{u}^{\infty}). \tag{16}$$

Overall, the fluid motion is governed by Stokes' equations; hence, particle dynamics operate in the "overdamped limit" of the Langevin equation, where integrating twice over a time interval  $\Delta t$  large enough for a colloid to have received many uncorrelated solvent molecule impacts and to have relaxed their momenta ( $\Delta t \gg m/6\pi\eta a_i$ ) results in the displacement equation (see Ermak and McCammon, 1977 for further details)

$$\Delta \mathbf{x}_{i} = \mathbf{X}_{i}(\Delta t) + \frac{\mathbf{F}_{i}^{P}}{6\pi\eta a_{i}} \Delta t + \mathbf{u}^{\infty} \Delta t.$$
 (17)

Here,  $\Delta x_i$  is the change in position over a time step  $\Delta t$ , owing to the combined contributions of the Brownian force, the interparticle potential, and the imposed flow. In **Equation 17**, the Brownian displacement  $X_i(\Delta t)$  obeys Gaussian statistics given by

$$\frac{\overline{X}_{i} = \mathbf{0}}{\overline{X}_{i}(\Delta t)X_{i}(\Delta t)} = 2\mathbf{D}_{i}\Delta t,$$
(18)

where the amplitude of the co-variance in this freely draining model is set by the diffusion coefficient of a single particle defined by the familiar Stokes-Einstein relation,  $D_i = kT/6\pi\eta a_i I$  (Einstein, 1905; Stokes, 1850). In other approaches, the diffusion coefficient used to compute Brownian displacements is sometimes obtained from *in vivo* measurements in order to implicitly encode the effects of crowding but limits the applicability of the model to a specific temperature or crowding condition of the cell (among other issues). In our approach, the effects of crowding on particle motion emerge naturally from the excluded volume interactions enforced by the interparticle force (**Equation 15**): each particle undergoes a Brownian displacement commensurate with **Equation 18**, reflecting the dissipation of a thermal kick via its own solvent drag. However, over many displacements, a particle will encounter other particles and must "wait"

for them to move in order to sample that space, leading naturally to a slow-down of diffusion over longer distances through a crowded suspension. Changes in volume fraction thus seamlessly lead to changes in particle dynamics and interactions.

The hard-sphere interparticle force (**Equation 15**) imposes a singular force between particles at contact. The modeling of singular forces in dynamic simulation is intractable; at best, very steep repulsions can be captured. To overcome this limitation, rather than to impose a singularity, we utilized a "potential-free" algorithm (Foss and Brady, 2000; Heyes and Melrose, 1993), where once Brownian and imposed flow displacements are made, particles are permitted tiny overlaps which are then corrected. A pair of particles do not interact at all unless they come into direct contact or overlap and once they do interact, the hard-sphere displacement of each corrects any overlap:

$$\Delta \mathbf{x}_i^{HS} = \frac{a_j}{a_i + a_i} \left[ \Delta r_{ij} - \left( a_j + a_i \right) \right] H \left( a_j + a_i - \Delta r_{ij} \right) \hat{\mathbf{r}}_{ij}. \tag{19}$$

Here,  $\Delta r_{ij} - (a_j + a_i)$  is the overlap being corrected, the Heaviside step function H ensures that particles are displaced only when they overlap, and  $\hat{r}_{ij}$  is the unit vector along the line that connects the particle centers, giving the hard-sphere displacement along the line of centers. This approach is equivalent to applying to each particle an interparticle force  $F_i^P$  that is proportional to the overlap to ensure entropic exclusion. The physical consequence of the hard-sphere interaction and the overlap correction is a contribution to the osmotic pressure that arises from the finite volume particles occupy in the suspension (Brady, 1993).

Incorporating in **Equation 17** the potential-free algorithm and considering that in our system there are no imposed flows  $u^{\infty}$  gives a displacement equation of the form

$$\Delta \mathbf{x}_i = \Delta \mathbf{x}_i^{HS} + \mathbf{X}_i(\Delta t), \tag{20}$$

where  $\Delta x_i^{HS}$  is the displacement that enforces the hard-sphere potential. **Equations 18 – 20** constitute the framework underlying dynamic simulations in Colloidal Smoldyn, our adaptation of the original Smoldyn.

In our simulations, we chose a timestep  $\Delta t = 62$  picoseconds empirically by testing the temporal resolution necessary to recover expected mean-squared displacements over long timescales, which we computed using the open-source LAMMPS Molecular Dynamics Simulator, across the levels of crowding we modeled (**Figure S3**) (Plimpton, 1995). Practically, this timestep corresponds to an average Brownian displacement per-timestep of 0.1 nm for ribosomes, 0.14 nm for ternary complexes, and 0.25 nm for crowders (**Equation 18, Table S4**).

At each time step all particles have a Brownian displacement, after which the algorithm checks for overlaps between particles. In Colloidal Smoldyn, overlapping biomolecules are separated along their line of centers and placed at contact according to  $\Delta x^{HS}$  (**Equation 19**) and in accordance with well-established and experimentally validated models of colloidal physics (Heyes and Melrose, 1993). Our implementation is a direct modification to the physics represented by the original Smoldyn algorithm; the prior method over-corrects the overlap by separating the pair's surfaces by the full distance, meaning a gap is introduced almost as though the collision was being modeled as inertial (Andrews, 2017). In addition, the particles were moved not along their line of centers in the original algorithm but rather at a reflected angle again reminiscent of an inertial collision which is not appropriate for colloids in the overdamped dynamics limit. The original method of Smoldyn,

- 735 although intuitive, is physically incorrect for systems in which fluid motion is governed by Stokes 736 equations, and a strong drag force is exerted on the approaching molecules. In such systems inertia 737 is irrelevant, meaning that when two particles approach after receiving a Brownian kick they will 738 simply slow down until another Brownian displacement sets them in a different direction. Placing 739 particles at contact along the line of centers upon overlap approximates this entropic exclusion 740 interaction and the interparticle forces associated with this displacement contributes to the osmotic 741 pressure of the suspension. The Heyes and Melrose algorithm has been shown to recover 742 suspension osmotic pressure as predicted by statistical mechanics models (Foss and Brady, 2000; 743 Zia and Brady, 2012).
- 744 In simulations of crowded systems, correcting a set of overlaps typically produces additional new 745 overlaps, necessitating additional iterations of overlap correction. Two strategies are applied to 746 accurately process overlaps: first, utilizing short, discretized time steps to generate smaller and 747 fewer overlaps; second, executing several overlap-correction cycles. For computational efficiency, 748 the original Smoldyn algorithm executed only one round of overlaps, leaving many still-749 overlapping particles. Only in dilute systems is such an approximation physically accurate. In 750 colloidal Smoldyn, we conducted an optimization study to identify an appropriate balance of 751 efficiency and accuracy, from which we devised an algorithm with three iterations of overlap 752 resolution at the chosen timestep of  $\Delta t = 62$  picoseconds.
- In all our simulations, we avoided finite-size and boundary effects by implementing periodic boundary conditions.
- 755 III.2. Simulating reaction dynamics

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- In our simulations, after a ternary complex and unbound ribosome interact both biomolecules are converted to form a single bound-ribosome that either disassociates or forms a peptide bond following single-molecule rate kinetics. In the native Smoldyn algorithm, the distance at which chemical interactions are initiated between molecules is set a priori to be different than contact (e.g., molecules may be set to only react when halfway overlapping or conversely while still separated by several molecule diameters). In particular, the initiating distance is computed such that the overall reaction rate of all molecules in simulation is equal to experimentally-measured kinetic rates set for individual reactions (Andrews and Bray, 2004). This approximation is based on the Smoluchowski model for reactions and is only reasonable for dilute systems (or systems of point particles). To achieve a more physical representation for crowded colloidal systems (as in our modeling of transport) that allows ab initio estimates of latency without fitting, we further modified the Smoldyn algorithm in Colloidal Smoldyn to only allow for initiation of chemical reactions upon the physical (entropic) interaction of reactant biomolecules. We did not consider factors such as binding site orientation that would restrict reactions to only certain interactions or reaction activation energies that may need to be overcome prior to reaction. Instead, we made the approximation that all physical interactions between ternary complexes and ribosomes lead to reactions. Ribosomes and ternary complexes that disassociate following a reaction are placed at contact, congruent to the Brownian dynamics underlying our modeling of transport. This was a final modification to the Smoldyn algorithm, which places biomolecules at a distance following disassociation to artificially reduce repeat reaction probability based on fitting of experimental single molecule reaction kinetic parameters (an approximation that, as before, is reasonable only for dilute systems) (Andrews and Bray, 2004).
- We computed the time a ternary complex spends bound to a ribosome via a Markov process consisting of well-defined intra-ribosomal states and transition kinetic rates  $(k_i)$  (Kinz-Thompson

et al., 2016). Specifically, the algorithm we used to model biochemical reactions involving ternary complex-bound ribosomes is:

- 1. For each state transition  $i \rightarrow j$  out of the current state i, draw a transition dwell time,  $\tau_{i \to j} \leftarrow \exp\left(\frac{1}{k_{i \to i}}\right);$
- 2. Transition to state j corresponding to the fastest transition time,  $\operatorname{argmin}_{i}(\tau_{i \to i})$ ;
- 3. Draw an updated dwell time for the transition to state j,  $\tau_{i \to j} \leftarrow \exp\left(\operatorname{argmin}_{j}(\tau_{i \to j})\right)$ ;
- 4. Add the updated dwell time,  $\tau_{i \rightarrow i}$ , to total dwell time  $\tau_{tot}$ ;

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5. Repeat until either the ternary complex dissociates from the ribosome or a peptide bond is successfully formed.

Using our algorithm and reported kinetic rates, we computed the distributions of latencies for noncognate, near-cognate, and cognate ternary complexes reacting with ribosomes (Figure S2, Table S5, ReactionKineticsCalculations.ipynb). We did not consider the unlikely events of incorrect reactions leading to mis-incorporated amino acids ( $\leq 1\%$ ) or cognate ternary complexes being rejected following successful codon recognition (0.04%).

## IV. Simulation of translation voxels with varying composition

To measure the relative contributions of transport and reactions to protein synthesis rate, we simulated translation voxels with increasingly accurate composition ( $\mu = 0.6$  dbl/hr) (Results section III, ColloidalStoichiometryEffects.py). Specifically, we simulated five progressively accurate scenarios: (i) a single ribosome & cognate ternary complex matching pair; (ii) a matching pair surrounded by a physiological number of ternary complexes; (iii) a matching pair surrounded by physiological number of ternary complexes and ribosomes; (iv) a matching pair surrounded by physiological numbers of ternary complexes, ribosomes, and crowder proteins; and (v) a statistically representative ensemble of translation voxels each with a physiological number of ternary complexes, ribosomes, and crowder proteins. We performed 900 simulation replicates each for the first three cases and 100 simulation replicates for case (iv); we detail case (v) in the next section. Replicates were assigned random initial conditions, chosen using a Mersenne Twister random number generator with seeds set at multiples of five (i.e., 0, 5, 10, ...). For each replicate, we further randomly sampled 30,000 matching reaction latencies using a seed of zero.

#### V. Construction of statistically representative translation voxel ensembles

To construct statistically representative ensembles of translation voxels (used in Results sections III, IV, and V), we incorporated reported relative abundances of different types of ternary complexes and frequencies of codons among mRNA in E. coli at different growth rates (Table S6). We computationally constructed 100,000 translation voxels for cells in each of six different growth conditions (0.6, 1.0, 1.5, 2.0, 2.5, and 3.0 dbl/hr), sufficient to represent the statistical distribution of translation voxels across each condition. Individual translation voxels comprise different types of ternary complexes and codon-specific elongating ribosomes, randomly chosen using a Mersenne Twister random number generator with a seed of zero.

- 819 For each translation voxel, we randomly picked a single ribosome to track. We then classified 820 ternary complexes in each translation voxel as either non-cognate or cognate to the chosen 821 ribosome, leading to a growth-rate dependent distribution in the abundance of cognate ternary
- 822 (between zero translation voxels complexes to forty-two) across (Figure

CognatetRNADistributionCalculation.ipynb). For each of our six modeled growth conditions, we used the distribution of cognate ternary complexes calculated at the closest growth rate (measured at 0.4, 0.7, 1.6, 2.5 dbl/hr). The speed with which the single chosen ribosome in a translation voxel successfully finds and reacts with a cognate ternary complex provides a good lower bound for the bulk translation elongation rate; the bulk elongation rate corresponds to the speed with which as many peptide bonds are formed as number of ribosomes, and the speed with which a single ribosome finds and successfully reacts with a cognate ternary complex will typically be faster than the speed of as many successful reactions as ribosomes in the voxel.

## VI. Simulation of statistically representative translation voxels ensembles

To compute the transport, reaction, and elongation latencies of statistically representative ensembles of translation voxels (Results sections III, IV and V), we simulated translation voxels across the six different growth conditions (0.6, 1.0, 1.5, 2.0, 2.5, and 3.0 dbl/hr). Statistically representative ensembles of translation voxels correspond to the full set of possible translation voxels, meaning translation voxels can contain between zero to forty-two cognate ternary complexes for a single chosen ribosome (distributed as in **Figure S4**). For translation voxels containing between one to forty-two cognate ternary complexes belonging to cells growing at each of the six growth rates, we simulated 100 replicates starting from different random initial conditions ( $42 \times 6 \times 100 = 25,200$  total simulations). We set conditions for each replicate using the Mersenne Twister random number generator with seeds set as multiples of five (i.e., 0, 5, 10, ..., 495). Simulations were terminated when the ribosome being tracked successfully reacted with a cognate ternary complex.

VI.1. Post-simulation analysis of statistically representative translation voxel ensembles

For each translation voxel simulation (i), we computed the elongation latency  $(\tau_{elong,i})$ , transport latency  $(\tau_{\text{transport},i})$ , and reaction latency  $(\tau_{\text{rxn},i})$  of the cognate ternary complex that successfully reacted the ribosome being (StatisticallyRepresentativeTranslationVoxelAnalysis.ipynb). We incorporated the impact of near-cognate ternary complexes by scaling the time taken by a statistically accurate portion of noncognate reactions at the tracked ribosome. Specifically, we leveraged our calculation that translation voxels have eight near-cognate ternary complexes and 32 non-cognate ternary complexes, on average (Figure S4), and that near-cognates have an average latency of 4.6 ms while non-cognates have an average latency of 1.4 ms (Figure S2), to randomly scale non-cognate reaction times 3.3-fold with 20% probability. We note that this representation of near-cognates does not capture the impact of near-cognate ternary complexes on other ribosomes in the voxel; near-cognates could slightly reduce overall latencies by occupying mismatching ribosomes for longer than noncognate ternary complexes, allowing cognate ternary complexes to find their match more quickly.

We subsequently computed an overall transport latency ( $\tau_{transport}$ ), reaction latency ( $\tau_{rxn}$ ), and elongation latency ( $\tau_{elong}$ ) for each growth rate. We did so by calculating weighted averages of each of transport latency, reaction latency, and elongation latency acquired from translation voxel simulations for each particular growth rate ( $\mu$ ), averaging over replicates:

$$\tau_{\text{elong}}(\mu) = \frac{1}{100} \sum_{i}^{42} \sum_{i}^{100} p_i \tau_{\text{elong},i}, \qquad (21)$$

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$$\tau_{\text{transport}}(\mu) = \frac{1}{100} \sum_{i}^{42} \sum_{j}^{100} p_{i} \tau_{\text{transport}, i}, \qquad (22)$$

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$$\tau_{\text{rxn}}(\mu) = \frac{1}{100} \sum_{i}^{42} \sum_{j}^{100} p_i \tau_{\text{rxn},i}.$$
 (23)

The probability of each translation voxel configuration  $(p_i)$  is conditional on both the number of cognate ternary complexes in the particular translation voxel as well as growth rate (**Figure S4**). We did not consider the latency of translation voxels that contained zero cognate ternary complexes ( $\sim$ 22% of translation voxel instances), since such voxels would have infinite latency and are an artifact of constraining translation voxels to 42 total ternary complexes. In particular, if larger voxels with more than 42 ternary complexes are considered, the resulting proportion of cognate ternary complexes is similar but with fewer instances of zero cognates (e.g., we found that voxels with 42, 84 or 168 ternary complexes have the same number of average cognates when normalized by number of total ternary complexes, but have 22%, 10%, and 3% of instances with zero cognates respectively). Not considering the zero cognate ternary complex voxels thus provides a lower bound estimate of elongation, transport, and reaction latency.

# VII. Event-based stochastic simulations of statistically representative translation voxel ensembles

To measure the effect of removing transport physics from our simulations (Supplement), we developed an event-based stochastic simulation algorithm of statistically representative translation voxel ensembles (EventBasedStochasticSimulation.py). As in our other simulations, the ensemble of voxels captures the relative abundances of ternary complexes and frequencies of codons among mRNA, but unlike our other simulations, physical space is not represented.

In our stochastic simulation algorithm, all ribosomes in translation voxels are initialized as bound to randomly chosen ternary complexes. Each reacting ternary complex-ribosome pair is then assigned a time until either disassociation or successful amino acid incorporation, drawn from the distribution of non-cognate, near-cognate, and cognate reaction latencies we computed as in Methods Section III.2. (**Figure S2**). The simulation proceeds in an event-based fashion, iteratively transitioning to the next event that occurs in the translation voxel (i.e., the timestep of simulation is not fixed). Following a disassociation event, the disassociated ternary complex joins the available (unbound) ternary complex population, and the newly available ribosome instantly binds to a randomly chosen ternary complex. The simulation ends when a cognate ternary complex successfully reacts with a matching ribosome.

We computed the elongation latency at particular growth rates by simulating the statistical distribution of possible translation voxels (i.e., with the full permissible range of cognate ternary complexes, distributed as in **Figure S4**) and then averaging their resulting elongation latencies. For each growth rate and permissible number of cognate ternary complexes, we simulated 5000 replicate translation voxels with different random initial conditions. Conditions were set for each replicate using the Mersenne Twister random number generator with seeds set as multiples of five (i.e., 0, 5, 10, ..., 24995).

## VIII. Computational tools and costs

 All fixed time-step simulations of translation voxels were performed using Colloidal Smoldyn (based on Smoldyn v2.61) deployed on Amazon Web Services. Simulations required ~300,000 CPU-hours in total. Our longest simulations, for translation voxels at a growth rate of 0.6 dbl/hr, took up to ~3 weeks for some replicates, while our shortest simulations took seconds. The cost of all our simulations was approximately US \$10,000. Output file sizes for most simulation runs were small (<1 MB). All measurements and validation with LAMMPS were performed using the Sherlock High Performance Computing Cluster at Stanford University. Modeling, analysis, and event-based simulations were performed using Python 3.7.

#### IX. Acceleration of translation voxel simulations to reduce runtime and cost

Simulations of translation voxels were originally forecast to cost US \$6 million with the longest simulations taking ~36 years, making them intractable. To achieve feasible costs and run times, we implemented a procedure for accelerating our fixed-timestep simulations ~600-fold, reducing costs and run times as detailed above. In our acceleration procedure, kinetic rates of unbinding and codon recognition (i.e., the possible exits to the initially bound state) are increased 600-fold during simulations ( $k_1 = 717 \, \mathrm{s^{-1}}$  to  $430200 \, \mathrm{s^{-1}}$ , and  $k_{2f} = 1474 \, \mathrm{s^{-1}}$  to  $884400 \, \mathrm{s^{-1}}$ ). Simulations are run until completion following a successful match between a cognate ternary complex and matching ribosome. Subsequently, during post-simulation analysis, the time spent by ternary complexes in the initially bound state is re-scaled to be 600-fold slower, and re-scaled times are used to compute reaction, transport, and elongation latencies. Reaction latency is calculated as the time the cognate ternary complex spends bound in reactions; elongation latency is calculated as the total time the matching ribosome spends unbound or bound in reactions; and transport latency is calculated as the difference between elongation latency and reaction latency.

Our estimates of overall reaction, transport, and elongation latencies are not sensitive to this scaling procedure at the ~600-fold acceleration used (**Figure S5A-C**). This insensitivity is a result of unbinding kinetics remaining slow enough that, for a certain range of kinetic scaling, ternary complexes mix within the translation voxel in between unbinding events to a sufficiently similar extent (**Figure S5D**).

## X. Calculation of long-time self-diffusivity

We tracked the motion of individual biomolecules as they wandered far from their original positions, executing a random walk through cytoplasm. This sampling of many configurations in a voxel is termed the long-time self-diffusion  $(D_{\infty}^s)$  (referred to as diffusivity in the Results) and is a monotonically decreasing function of volume fraction at fixed molecule size polydispersity. We computed the long-time self-diffusion of particular biomolecule species (denoted by subscript i) at different growth rates by tracking the absolute position of biomolecules and computing their mean squared displacement over time (Results section IV):

$$D_{\infty,i}^{s} = \frac{1}{6} \frac{d}{dt} \langle \Delta \mathbf{x}_i \Delta \mathbf{x}_i \rangle.$$
 (24)

Here, the angle brackets signify an ensemble average over the motion of every biomolecule of a given species in a translation voxel.

## XI. Calculation of viscosity

 We calculated the viscosity of translation voxels at different growth rates (Results section IV) by performing shear rheology simulations in LAMMPS. For each growth rate, we initialized suspensions representative of multiple contiguous translation voxels. We imposed a simple shear flow on the suspensions at a constant shear rate in the x-direction ( $\dot{\gamma}_x$ ) and measured the resulting interparticle stress ( $\bar{\sigma}^p_{xy}$ ). The shear rate imposed was chosen to be small enough to remain in the linear-response regime (i.e., with insignificant deformation), allowing measurement of the intrinsic or so-called zero-shear viscosity ( $\eta_0$ ) (ViscosityCalculation.py) (Batchelor, 1977):

$$\frac{\eta_0}{\eta} = 1 + \frac{5}{2}\phi + \frac{\bar{\sigma}^p_{xy}}{\eta \dot{\gamma}_x} \ . \tag{25}$$

Here, the first two terms on the right-hand side of the equation are the Einstein viscosity that approximate the hydrodynamic contribution of particles to viscosity at equilibrium. The third term describes the interparticle contribution to viscosity and is equivalent to the Green-Kubo equilibrium interparticle contribution at the small shear rates used here.  $\bar{\sigma}^p_{xy}$  is computed as the xy-component of the interparticle stress  $\langle xF^p\rangle$ , where x corresponds to the position vectors of the particles and  $F^p$  is the (negative of) the gradient of a nearly hard-sphere, spherically symmetric repulsive potential.

## XII. Calculation of molecular proximity

We computed the proximity between ternary complexes and ribosomes at different growth rates (Results section IV). For each growth rate, we initialized 100 translation voxels with random initial spatial configurations, chosen using a Mersenne Twister random number generator with seeds set at multiples of five (i.e., 0, 5, 10, ..., 495). Following a brief equilibration period, we measured the distance from each ternary complex to its closest ribosome. Our reported values of proximity for any particular growth rate are an average of the minimal distance for all ternary complexes across all corresponding 100 replicate translation voxels.

#### XIII. Calculation of repeat reactions

We computed the average number of repeat reactions between ternary complexes and ribosomes at different growth rates (Results section IV). For each growth rate, we initialized 100 translation voxels with random initial spatial configurations, chosen using a Mersenne Twister random number generator with seeds set at multiples of five (i.e., 0, 5, 10, ..., 495). We subsequently tracked the number of times a ternary complex consecutively re-reacted with the same ribosome following a mismatching reaction within each translation voxel. Our reported values for repeat reactions for any particular growth rate are an average across all corresponding 100 replicates for the given growth rate.

## XIV. Chemical kinetics sensitivity analysis

To measure the sensitivity of our predicted elongation latencies to changes in chemical kinetics, we simulated the impact of slowing down or speeding up intra-ribosomal kinetic rates on elongation latency. To do so, we simulated ensembles of translation voxels as in Methods section VI while varying kinetic rates individually or all together and measuring the resulting elongation latency (**Figure S9**). Since the kinetic rate  $k_{1r}$  impacts the mix time of voxels (**Figure S5**), we varied the level of kinetic acceleration in our simulations for different  $k_{1r}$ , ensuring that our

translation voxels were simulated in regimes in which elongation latency is insensitive to changes in kinetic acceleration (our kinetic acceleration scheme is described in Methods section IX).

# XV. Calculation of maximum packing and projected growth rate voxel parameters

We computed the theoretical maximum packing for translation voxels between observed growth rates,  $\mu = 0.6$  dbl/hr to 3.0 dbl/hr, as well as faster hypothetical growth rates,  $\mu = 3.0$  dbl/hr to 8.0 dbl/hr, using the theoretical calculations of maximum packing for tridisperse systems from Farr and Groot, 2009. Our translation voxels are comprised of molecules having a 1:3:6.5 size ratio, which differs from the particle size ratio used by Far and Groot (1:3:9), giving an overprediction of our computed maximum packing of less than 10%.

To estimate ribosome abundances, ternary complex abundances, cell mass, cell volume, and nucleoid volume at hypothetical growth rates between 3.0 dbl/hr and 8.0 dbl/hr, we extrapolated from observed growth rates (Figure S1), guided by observed trends below 3.0 dbl/hr and allowing uncertainty while rejecting unphysical projections (e.g., negative cell mass and nucleoid volume fraction) (Figure S11, Figure 6). We calculated bounds by perturbing the extrapolated fits while still maintaining all expected trends (e.g., the lower bound of ribosome abundances never decreases with increasing growth rate). We computed volume fractions for translation voxels at hypothetical growth rates as in Methods Section II, setting upper and lower bounds by considering all permutations of fits for ribosome abundances, ternary complex abundances, cell mass, cell volume, and nucleoid volume fraction (ProjectedGrowthRateCalculations.ipynb).

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#### **Declaration of Interests**

The authors declare no competing interests.

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**Supplemental Information** 

Molecular transport and packing underlie increasing ribosome productivity in faster

growing cells

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#### Note S1. Sensitivity analysis and the impact of parameter values

We sought to understand if our prediction that the productivity of individual ribosomes increases due to stoichiometric crowding is sensitive to the fact that our unfitted model does not exactly match observations of absolute elongation latencies. One natural place to start is to make a change in the values of the chemical kinetic parameters used in our model, which are taken from *in vitro* measurements (**Table S5**). The rationale for modulating these parameters is that *in vitro* kinetic rates may differ from their *in vivo* values due to, for example, differences in salt concentrations *in vitro* compared to *in vivo* (**Figure S13A**). To test this idea, we implemented a uniform three-fold increase of all intra-ribosomal chemical kinetic rates in our model (**Figure S13C**). We then simulated ensembles of translation voxels at varying growth rates and measured the resulting elongation rates. This chemical-parameter change closed the quantitative gap: elongation latency ( $\tau_{\text{elong}} = 100 \text{ ms}$  at 0.6 dbl/hr to 45 ms at 3.0 dbl/hr) now matched experimental latency ( $\tau_{\text{elong}}^{\text{bulk}} = 83 \text{ ms}$  at 0.6 dbl/hr to 48 ms at 3.0 dbl/hr) (**Figure S13B**). Importantly, the speedup in elongation as growth quickens occurs even with the higher chemical kinetic rates used to fit the bulk experimental observations.

We also sought to test if our predicted increase in ribosome productivity requires physical transport. To do so we stress-tested the sensitivity of elongation to changes in transport by modeling instantaneous transport within our voxel ensembles (**Methods**). We found that the growth-rate dependence of elongation latency is lost when transport is modeled as an instantaneous process (**triangles in Figure S13B**, **Figures S6**, **S7**), demonstrating the essential role of transport physics in the growth-rate dependence of elongation rate, and also demonstrating the criticality of accurate chemical parameters in establishing absolute quantitative prediction of elongation latency.

### Note S2. Translation initiation latency relative to translation elongation latency

Translation initiation is slow compared to a single elongation step. Therefrom, one might naively expect initiation to be rate limiting to protein synthesis overall and, thus, expect that any increase in protein synthesis rates should mostly arise due to increases in the rate of initiation.

However, clues to the contrary are apparent. For example, there is overwhelming experimental evidence from work to optimize expression of heterologous proteins (Welch et al., 2009). Specifically, synonymous changes in coding sequences that require ribosomes to make the same protein via coding-sequence unique elongation processes produce dramatic changes in protein synthesis rates and abundances. That is, codon usage changes alone can result in changes in gene expression levels, ranging from undetectable to majority of cell protein. This strong impact on protein synthesis rate is solely due to impacts on translation elongation, not translation initiation. Studies of natural living systems further reveal a literature that advances how both translation initiation and elongation can limit protein synthesis rates, depending on conditions

1222 (Subramaniam et al., 2014; Vieira et al., 2016).

Nevertheless, it is important to consider initiation more formally and explain why elongation is most likely to underlie increases in overall protein synthesis rates. To this end, we return to the conventional view of translation elongation in which the correct ternary complex instantaneously presents itself to the ribosome exactly when needed (i.e., they appear instantaneously in exactly the right order). That is, a situation in which there is no additional latency due to physical transport, nor combinatorial sampling by and among competing ternary complexes (i.e., setting aside the fact that our work shows that most elongation latency arises from

- transport, not kinetics). In these unrealizable conditions, the rate of peptide bond formation is
- entirely determined by chemical kinetics and is about 42 ms per amino acid. By comparison,
- translation initiation should be much slower, taking about 1-12 seconds. But protein synthesis
- requires not just one elongation step but hundreds in series (i.e., the average protein in *E. coli* is
- 1234 ~333 amino acids). Considering elongation of the entire protein, we estimate ~14 seconds total
- just from chemical kinetic latency alone (42 ms per amino acid). Thus, it becomes apparent that,
- while the chemical kinetics of a single initiation event might be relatively slow, most of the time
- is spent in the elongation phase (e.g., 12 seconds for initiation vs. at least 14 seconds for
- 1238 elongation).

### Note S3. Estimating the evolutionary impact of increasing ribosome productivity

- To explore the potential evolutionary impact of increasing ribosome productivity, we estimated
- how much slower bacterial growth would be if ribosome productivity did not increase (i.e.,
- remained fixed) and only the abundances of translation machinery increased.
- For a cell to replicate, it must replicate all its protein machinery. Since a fraction of peptide bonds
- formed during cell replication belong to translation machinery, as individual cells grow, increased
- translation machinery should enable more total peptide bonds to be formed per unit time. We can
- estimate the lower and upper bounds of total peptide bond formation in the time cells typically
- take to double, and thus growth rate, by assuming that this fraction is 0 (i.e., a lower bound in
- which no new peptide bonds enable more translation) and by assuming this fraction is 1 (i.e., an
- 1249 upper bound in which all new peptide bonds are incorporated into ribosomes that enable more
- 1250 translation).
- 1251 As a lower bound estimate,

Lower bound total bond formation = 
$$N_{\text{rib}} * t_{\text{dbl}} * k$$
. (S1)

1253 As an upper bound estimate,

Upper bound total bond formation = 
$$aa_{rib} * N_{rib} * [exp(t_{dbl} * \frac{k}{aa_{rib}}) - 1].$$
 (S2)

- Here,  $t_{\rm dbl} = \mu/3600$  is the time in seconds per cell doubling and  $aa_{\rm rib} = 7336$  amino
- acids/ribosome is the number of amino acids in a ribosome.
- At  $\mu = 3.0$ , there are on average  $N_{\rm rib} = 62000$  actively elongating ribosomes per cell elongating
- 1258 at a rate of k = 21 amino acids/s, leading to:
- Lower bound total bond formation = 1.562,400,000 bonds, and
- 1260 Upper bound total bond formation = 13,660,863,648 bonds.
- As a sanity check, we can estimate the total bonds in a cell at  $\mu = 3.0$  dbl/hr from the total bonds
- composing crowder proteins and total (active and non-active) ribosomes (**Table S1, S4**): 7.78 x
- 1263  $10^6$  crowder proteins \* 264 aa/crowder protein + 73000 total ribosomes \*  $aa_{rib} = 2,589,448,000$
- bonds, which lies between the lower and upper bounds as expected.
- We can next consider the hypothetical case in which ribosome productivity at  $\mu = 3.0$  dbl/hr does
- not increase and instead remains fixed at that of  $\mu = 0.6$  dbl/hr, k' = 12 amino acids/s. In this
- case, we can estimate the lower bound and upper bound time needed for total bond formation by
- solving for  $t_{\rm dbl}$  in equations S1 and S2.
- 1269 In the lower bound case,

Lower bound 
$$t'_{dbl} = \frac{N_{rib} * k'}{\text{Lower bound total bond formation}}$$
. (S3)

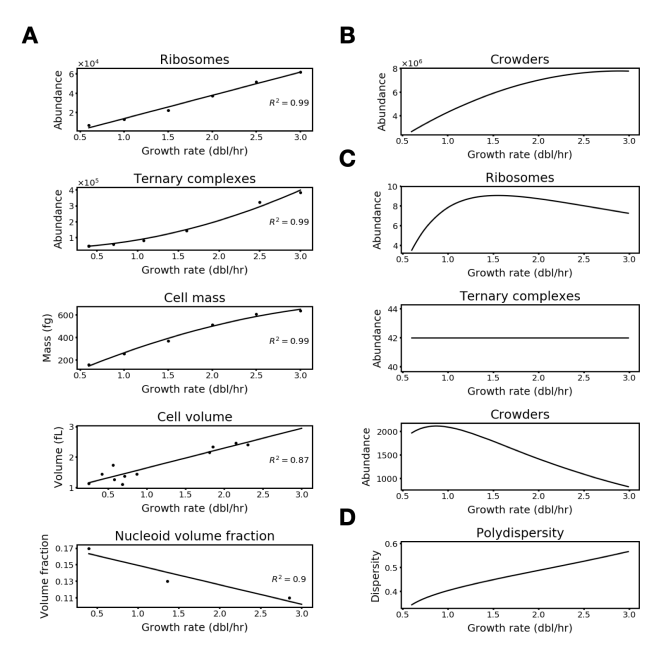
1271 In the upper bound case,

Upper bound 
$$t_{\rm dbl}' = \frac{aa_{rib}}{k'} * \ln\left(\frac{\text{Upper bound total bond formation}}{aa_{\rm rib} * N_{\rm rib}} + 1\right).$$
 (S4)

- 1273 This gives us,
- Lower bound  $t_{\rm dbl}' = 2100$  s, giving an effective  $\mu' = 1.7$  dbl/hr
- 1275 Upper bound  $t_{dbl}' = 2100$  s, giving an effective  $\mu' = 1.7$  dbl/hr
- 1276 Thus, we estimate that maximum observed cell doubling would proceed at ~1.7 dbl/hr instead of
- 3.0 dbl/hr in the hypothetical case in which ribosome productivity did not increase and remained
- 1278 fixed at that of  $\mu = 0.6$  dbl/hr.
- What would this difference mean practically from an evolutionary perspective? We can compare
- the expected growth of microbes with these different growth rates:

Fold-difference microbes = 
$$2^{(\mu-\mu')t}$$
. (S5)

- 1282 After t = 6 hours there would be ~200-fold more of the microbes with increased ribosome
- productivity, after 12 hours ~50,000-fold more, and after 24 hours 2.5 billion-fold more.



**Figure S1.** Cell and translation voxel parameters vary with growth rate. (A) Polynomial fits of reported cell parameter measurements (data points are taken from Tables S1, S2, and S3). (B) Estimate of crowder protein abundances in cells across growth rates. (C) Estimate of ribosome, ternary complex, and crowder protein abundances in translation voxels across growth rates. (D) Estimate of the polydispersity of translation voxels across growth rates.

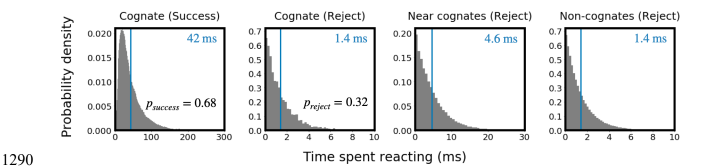


Figure S2. Ribosomal kinetics are dependent on whether a ternary complex is cognate, near-cognate, or non-cognate. (A-D): Distribution of time spent by tRNA within ribosomes, computed by sampling tRNA-ribosome reaction trajectories. (A) Cognate ternary complexes that successfully react (probability shown). (B) Cognate ternary complexes that are rejected (probability shown). (C) Near-cognate ternary complexes that are rejected (only rejections are considered). (D) Non-cognate ternary complexes that are rejected (only rejections are considered). In each plot, average latency is marked by a blue line and displayed on the top right.

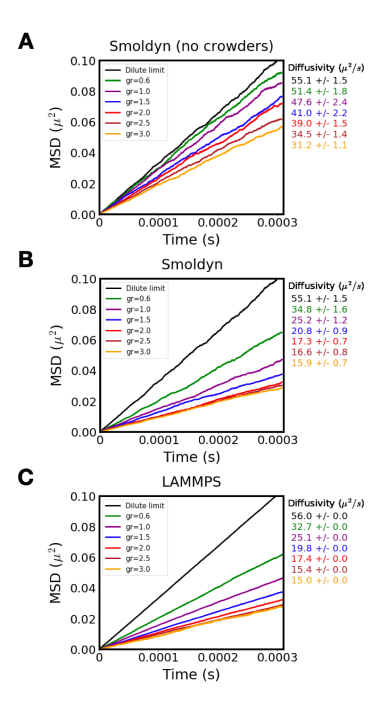


Figure S3. The long-time self-diffusivity of ternary complexes is dependent on crowding and growth rate. (A) Measurements of ternary complex mean squared displacement in translation voxels without crowder proteins using Smoldyn. (B) Measurements of ternary complex mean squared displacement in complete translation voxels (i.e., with crowder proteins) using Smoldyn. (C) Measurements of ternary complex mean squared displacement in complete translation voxels using LAMMPS. For all plots, in the dilute limit, ternary complex long-time diffusivity should equal its short-time diffusivity ( $D = 56 \mu m^2/s$ , Table S4).

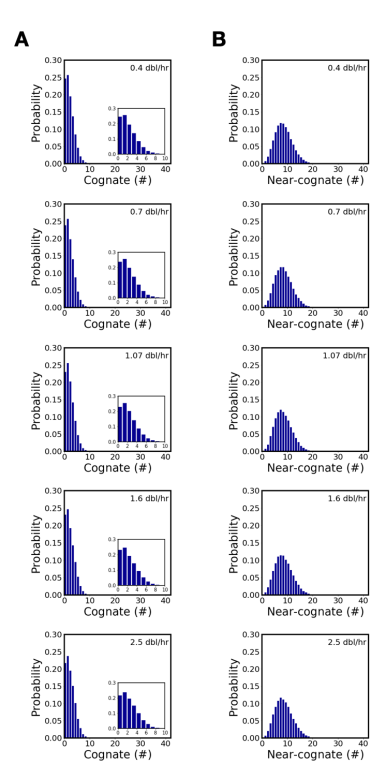


Figure S4. The number of cognate and near-cognate ternary complexes in a translation voxel follows a slightly growth-rate dependent probability distribution. (A) Distribution of cognate ternary complexes in translation voxels. (B) Distribution of near-cognate ternary complexes in translation voxels. Insets highlight the distribution between values of zero to ten cognate ternary complexes.

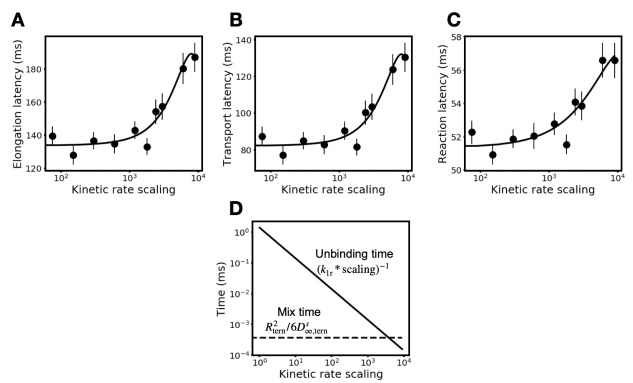


Figure S5. Predictions of translation latencies are insensitive to low and intermediate kinetic rate scaling due to differences in unbinding and mixing timescales. (A-C) Elongation latency, transport latency, and reaction latency are insensitive to kinetic rate scaling at a scaling of less than ~2000. Data shown for a translation voxel with growth rate  $\mu = 3.0$  dbl/hr. (D) The average time ternary complexes take to unbind from ribosomes (unbinding time) remains slower than the typical time ternary complexes take to diffuse their radius within a voxel (mix time, calculated using Equation 18 in the Methods). The mix time plotted is computed for the most crowded growth condition ( $\phi_{vox} = 0.42$ ,  $\mu = 3.0$  dbl/hr).

 $\begin{array}{c} 1313 \\ 1314 \end{array}$ 

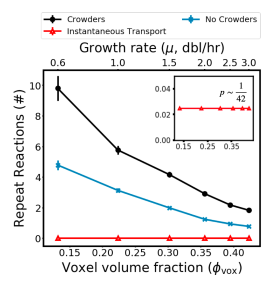


Figure S6. Repeat reactions between mismatching ternary complexes and ribosomes are sensitive to crowding and instantaneous transport. Average number of repeat reactions following initial reaction in complete translation voxels (black curve, filled circles), translation voxels with no crowder proteins (blue curve, x's), and translation voxels with instantaneous transport (red curve, triangles). Inset shows that simulations with instantaneous transport produce repeat reactions with uniform random likelihood across all growth rates.

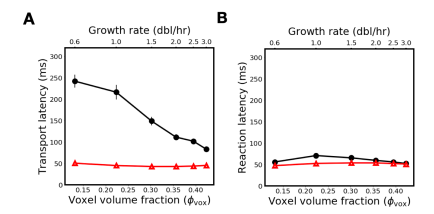


Figure S7. Growth rate dependence of transport and reaction latency is lost when transport is modeled as an instantaneous process. Simulations with instantaneous transport (red curve, triangles) produce (A) transport latencies and (B) reaction latencies that are uniform across growth rates. Original simulation results are shown for comparison (black curve).

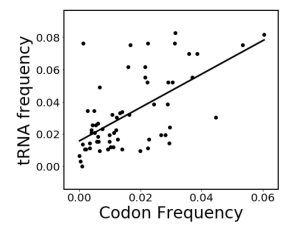


Figure S8. The frequency of a given codon and their corresponding tRNA are weakly correlated. Frequency data from Table S6.

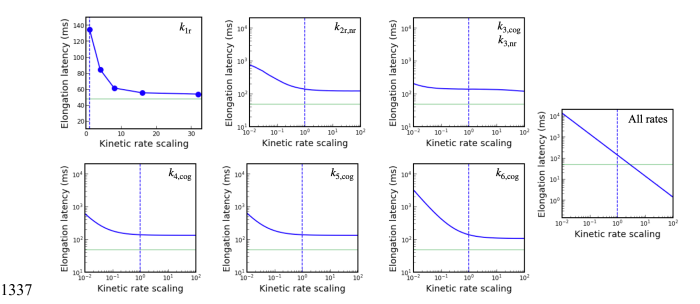


Figure S9. Our chemical kinetics sensitivity analysis shows that only changes in unbinding rate  $(k_{1r})$  or all kinetic rates together can speed up elongation to experimentally predicted latencies. Each plot shows the elongation latencies (solid blue line) that result from scaling one or more intra-ribosomal kinetic rates (specified in top right) to be faster or slower compared to baseline (dashed blue line). All simulations were for translation voxels at  $\mu = 3.0$  dbl/hr. Experimentally measured elongation latency at  $\mu = 3.0$  dbl/hr is shown for reference (green line). Scales are identical across plots except for the  $k_{1r}$  and All rates plots.

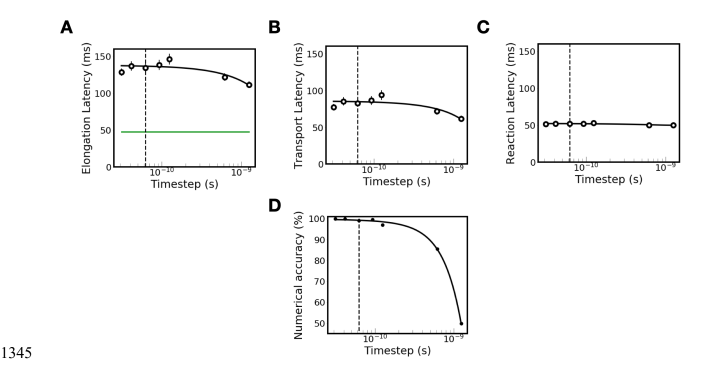


Figure S10. (A-C) Our simulation timestep sensitivity analysis shows that elongation latency, transport latency, and reaction latency each converge near our chosen timestep at the most crowded growth rate ( $\mu = 3.0$  dbl/hr). Experimentally measured elongation latency at  $\mu = 3.0$  dbl/hr is shown for reference in panel A (green line). (D) Numerical accuracy, computed as the percentage of entropic interactions between particles that are resolved, converges to 100% near our chosen timestep. All simulations were for translation voxels at  $\mu = 3.0$  dbl/hr. The dashed line corresponds to the timestep used for simulations throughout the paper ( $\Delta t = 62$  picoseconds).

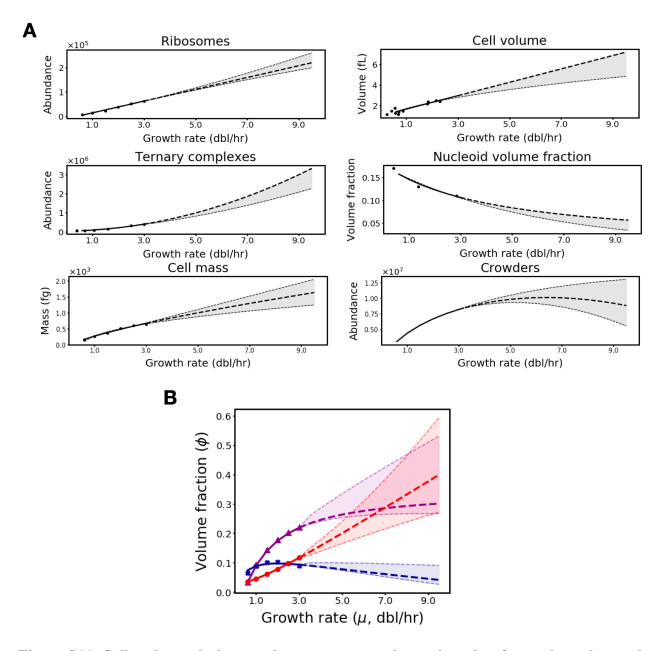


Figure S11. Cell and translation voxel parameters can be projected to faster-than-observed growth rates. (A) Extrapolated cell parameter fits (bolded dashed lines) projecting beyond observed growth rates (solid lines) for ribosomes (circle values from Table S1), ternary complexes (circle values from Table S1), cell mass (circle values from Table S1), cell volume (circle values from Table S2), nucleoid volume fraction (circle values from Table S3), and crowders. Perturbations to primary fits provide bounds (dashed lines and grey shaded region). (B) Extrapolated volume fractions (bolded dashed lines) projecting beyond observed growth rates (solid lines) for ribosomes (purple curve, triangles), ternary complexes (red curve, circles), and crowder proteins (blue curve, x's). Volume fractions are computed from cell parameter fits (Methods). Bounds for extrapolated volume fractions at hypothetical growth rates (light dashed lines and shading) are derived from permutations of the bounds for all cell parameters in panel A.

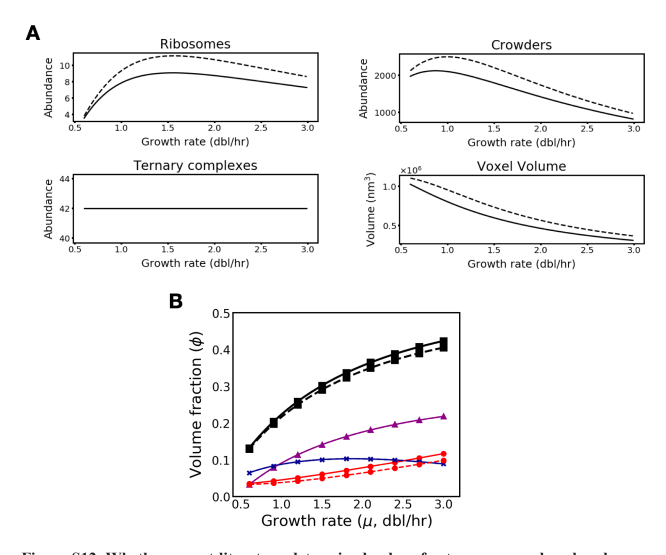


Figure S12. Whether or not literature-determined values for ternary complex abundances account for peptidyl-tRNA does not impact voxel composition or trends. (A) Key growth-rate dependent trends in translation voxel composition resulting from the modeling used throughout our manuscript (solid line), in which we interpret ternary complex abundance measurements as not including peptidyl-tRNA, are equivalent to those resulting from assuming that ternary complex abundance measurements do include peptidyl-tRNA (dashed line). (B) The volume fraction of ribosomes (purple curve, triangles), ternary complexes (red curve, circles), and crowder proteins (blue curve, x's), as well as total volume fraction (black curve, squares) are negligibly different for voxels constructed with the assumption that ternary complex measurements do not include peptidyl-tRNA (solid line) and the assumption that they do (dashed line).

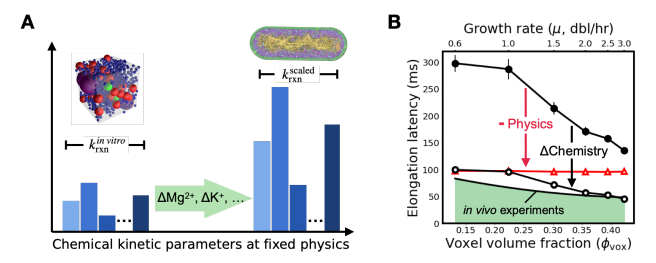


Figure S13. Faster elongation due to stoichiometric crowding is insensitive to changes in chemical kinetic parameters while sensitive to changes in transport physics. (A) Accuracy of chemical kinetics: cartoon schematic illustrating how differences in salt concentrations between *in vitro* conditions in which chemical kinetic rates were measured and *in vivo* conditions being modeled could lead to different and perhaps faster kinetic rates than used in our simulations. (B) Impact of faster chemistry, instantaneous physics: our simulations with three-fold faster kinetic rates produce elongation latency (open circles) that closes the quantitative gap between our original elongation latency prediction using published *in vitro* kinetic rates (solid circles) and *in vivo* measurements of per-ribosome elongation time (solid black line). The essentiality of physics in this agreement is demonstrated by the lack of faster elongation in simulations with instantaneous transport (red triangles). Depiction of *E. coli* in (A) adapted with permission from Goodsell, 2009.

Table S1. Estimation of elongating ribosome, ternary complex, and crowder protein abundances at varying growth rates

Estimation of elongating ribosome abundance								
Doublings/hr	$\mu = 0.6$	$\mu = 1.0$	$\mu = 1.5$	$\mu = 2.0$	$\mu = 2.5$	$\mu = 3.0$		
Ribosomes/cell <sup>a</sup>	8000	15000	26000	44000	61000	73000		
Elongating	7000	13000°	22000	37000	52000	62000		
Ribosomes/cell <sup>b</sup>								
Estimation of terna	ary complex	abundance						
Doublings/hr	$\mu = 0.4$	$\mu = 0.7$	$\mu = 1.07$	$\mu = 1.6$	$\mu = 2.5$	$\mu = 3.0$		
tRNA:ribosome	12.5	9.5	7.5	7.5	7.0	7.0		
ratio <sup>d</sup>								
Ribosomes/cell	5000e	$8000^{f}$	$15000^{\rm f}$	$26000^{\rm f}$	61000	73000		
tRNA/cell <sup>g</sup>	64000	76000	110000	190000	430000	510000		
Charged	48000	57000	83000	143000	323000	383000		
tRNA/cell <sup>h</sup>								
Ternary complex	48000	57000	83000	143000	323000	383000		
abundance <sup>i</sup>								
Estimation of crow								
Doublings/hr	$\mu = 0.6$	$\mu = 1.0$	$\mu = 1.5$	$\mu = 2.0$	$\mu = 2.5$	$\mu = 3.0$		
$M_{\rm cytoplasm}^{\rm j}$	167	267	383	530	628	659		
$M_{ m mRNA}^{ m j}$	0.5	0.9	1.5	2.6	3.6	4.3		
$M_{\rm DNA}^{\rm j}$	7.6	9.5	12.0	14.7	17.2	19.4		
$N_{\rm rib}^{\rm k}$	4000	14000	26000	38000	50000	62000		
$N_{\rm tern}^{}$	57000	86000	138000	208000	295000	399000		
N <sub>crowder</sub> k	$2.66 \times 10^6$	$4.30 \times 10^6$	$5.90 \times 10^6$	$7.02 \times 10^6$	$7.65 \times 10^6$	$7.78 \times 10^6$		

<sup>&</sup>lt;sup>a</sup>Values obtained from Dennis and Bremer, 2008.

<sup>b</sup>Calculated from ribosomes/cell using fraction of elongating ribosomes = 0.85 as reported in Dennis and Bremer, 2008.

°As a secondary check, we used single cell mass spectrometry data from Schmidt et al., 2016 to calculate the abundance of ribosomes at  $\mu = 0.83/hr$ . We estimated ~17,000 ribosomes (RibosomeAbundanceEstimation.ipynb) by log averaging the reported abundances of fifty-four ribosomal proteins. Using the value of ~77% of ribosomes elongating at steady state as reported by Forchhammer and Lindahl, 1971, our estimated ~17,000 ribosomes corresponds to ~13,000 elongating ribosomes, matching the value reported from Dennis and Bremer, 2008 at  $\mu = 1.0$  dbl/hr.

<sup>d</sup>Values obtained from Dong et al., 1996. The value at  $\mu = 3.0$  dbl/hr was inferred via extrapolation of their reported data. In our modeling, we interpreted these measurements as representing only ternary complexes that are free or within the A-site; this interpretation does not impact our modeling results (**Figure S12**).

- 1409 eWe estimated the number of ribosomes at  $\mu = 0.4$  dbl/hr using the reported measurements of tRNA
- and tRNA:ribosome ratios from Dong et al., 1996. Dong et al. performed six replicates of
- measurements at 0.4 dbl/hr, and so six data points were included in our fitting shown in Figure S1.
- 1412 <sup>f</sup>We estimated the number of ribosomes at each given growth rate using the number of ribosomes
- at the closest reported growth rate from the data of Dennis and Bremer, 2008.
- 1414 <sup>g</sup>Calculated as the product of the tRNA:ribosome ratio and the number of ribosomes/cell
- <sup>h</sup>Approximately 75%-80% of tRNA are typically charged with amino acids (Sørensen, 2001;
- 1416 Stenum et al., 2017). We assumed uniform charging of 75% across all growth rates and types of
- 1417 tRNA.
- <sup>i</sup>We assumed that all charged tRNA are found in ternary complexes (i.e., bound to GTP bound EF-
- Tu) based on the excess abundance of EF-Tu in cells reported by Schmidt et al., 2016 and the fast
- 1420 kinetic rates of EF-Tu binding to GTP and EF-Tu-GTP binding to charged tRNA reported by
- 1421 Burnett et al., 2013, 2014.
- <sup>j</sup>Values obtained from Dennis and Bremer, 2008. Units of fg/cell.
- 1423 \*Derived from the regression fit shown in Figure S1.

## Table S2. Estimation of cell volume at varying growth rates

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Doublings/hr	μ =	μ =	μ =	μ =	μ =	μ =
	0.25	0.42	0.56	0.58	0.68	0.71
Media <sup>a</sup>	Gal	Ac	GluN	Pyr	Fum	Suc
Volume (fL) <sup>b</sup>	1.9	2.4	2.9	2.1	2.4	2.3
Adjusted volume (fL) <sup>c</sup>	1.14	1.44	1.74	1.26	1.11	1.38

Doublings/hr	$\mu =$				
	0.87	1.81	1.85	2.15	2.3
Media <sup>a</sup>	Glc	Gly+AA	Man+AA	Glc+AA	LB
Volume (fL) <sup>b</sup>	2.4	3.2	3.9	4.1	1430
Adjusted volume (fL) <sup>c</sup>	1.44	2.15	2.34	2.46	2.4

<sup>a</sup>Gal = Galactose; Ac = Acetate; GluN = Glucosamine; Pyr = Pyruvate; Fum = Fumerate; Suc = Succinate; Glc = Glucose; Gly+AA = Glycerol + all amino acids; Man+AA = Mannose + all amino acids; Glc+AA = Glucose + all amino acids; LB = Lysogeny broth.

bValues obtained from Volkmer and Heinemann, 2011 using fluorescence microscopy.

<sup>c</sup>In Radzikowski et al., 2016, a subset of measurements from Volkmer and Heinemann, 2011

were redone at higher resolution using super-resolution imaging and found to be 30-50% smaller

than estimated by Volkmer et al., 2011. We thus adjusted volume measurements by Volkmer and

Heinemann, 2011, as discussed in Schmidt et al., 2016, to either the higher-fidelity

measurements if the conditions were explicitly tested or reduced measurements by 40% if not.

# Table S3. Estimation of nucleoid volume fraction at varying growth rates

Doublings/hr	$\mu = 0.4$	$\mu = 1.36$	$\mu = 2.85$
Media	Alanine	Glucose	Tryptone
Nucleoid volume (fL) <sup>a</sup>	0.08	0.14	0.30
Cell volume (fL) <sup>a</sup>	0.46	1.06	2.72
Nucleoid volume fraction <sup>b</sup>	0.17	0.13	0.11

<sup>&</sup>lt;sup>a</sup>Values obtained from Woldringh and Nanninga, 1985.

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<sup>&</sup>lt;sup>b</sup>Calculated by dividing the reported nucleoid volume by the reported cell volume.

### Table S4. Parameters not varying with growth rate

$M_{\rm rib}$	2300 kDa
$M_{ m tern}$	69 kDa
$M_{\mathrm{tRNA}}$	25 kDa
M <sub>amino acid</sub>	110 Da
$M_{\rm EF-Tu}$	43.2 kDa
$M_{ m GTP}$	523 Da
M <sub>crowder</sub> <sup>a</sup>	28.5 kDa
V <sub>crowder</sub> <sup>b</sup>	$3.35 \ \mu m^3$
$ ho_{ m crowder}^{ m c}$	1.410 g/cm <sup>3</sup>
$R_{ m crowder}^{ m d}$	2.0 nm
$R_{ m rib}^{ m de}$	13 nm
$R_{ m tern}^{ m df}$	5.9 nm
$k_B$	1.380649×10 <sup>-23</sup> J/K
T	37 C
η	0.6913 mPa*s
$D_{\text{tern}}$	56 μm <sup>2</sup> /s
$D_{ m rib}$	$25 \ \mu m^2/s$
$D_{ m crowder}$	$165 \ \mu m^2/s$

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<sup>a</sup>We computed the mass of individual crowder proteins as described in the methods. Given an average amino acid mass of 108 Da, as reported by Bremer and Dennis, 2008, the crowder mass corresponds to 264 amino acids on average. This protein length corresponds well to estimates in the literature of the average and median number of amino acids for a protein, 309 and 252 amino acids respectively (Ishihama et al., 2008), supporting our calculations.

bThe volume of a crowder protein was calculated as described in the methods.

<sup>c</sup>Value obtained from Fischer et al., 2004.

dBiomolecule radius was calculated as described in the methods. Note that we use R and a interchangeably.

eMeasured using PDB 4V4Q from Schuwirth et al., 2005 as described in the methods.

<sup>f</sup>Measured using PDB 1B23 from Nissen et al., 1999 as described in the methods.

Table S5. tRNA-ribosome reaction kinetic parameters for non-cognate, near-cognate, and cognate tRNA

Kinetic Rates (s <sup>-1</sup> )	$k_{1\mathrm{r}}^{\mathrm{a}^*}$	$k_{2\mathrm{f,cog,nr}}^{\mathrm{b*}}$	$k_{2r,\cos}^{b^*}$	$k_{2\mathrm{r,nr}}^{\mathrm{b*}}$	$k_{3,\cos}^{b^*}$	k <sub>3,nr</sub> <sup>b*</sup>	$k_{4,\cos}^{}^{}^{}$	$k_{5,\cos}^{\mathrm{d}}$	$k_{6,\cos}^{e}$
		1458	1.5	837	994	6.6			
		1535	2.7	837	2064	4.5			
		1535		837		2.4			
Data		2214		1305		0.8			
Data		837		1914		9.7			
		1535		994		13.8			
		1150							
		1535							
Average	718	1475	2.1	1120	1529	6.3	209	200	32
Standard deviation	226	392	0.9	429	757	4.9	11	40	-

<sup>&</sup>lt;sup>a</sup>Measurements from Gromadski and Rodnina, 2004

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<sup>e</sup>Borg and Ehrenberg, 2015 made several measurements in varying buffer conditions. We chose their measurement that most corresponds to cell-like conditions (Polymix buffer with 1 mM free  $Mg^{2+}$  and  $10\mu M$  EF-G).

\*Kinetic rates were adjusted from 20 C to 37 C using the methods from Rudorf et al., 2014 and measurements of  $k_{5,cog}$  at both 20 C and 37 C from Wohlgemuth et al., 2010.

<sup>&</sup>lt;sup>b</sup>Measurements from Gromadski et al., 2006

<sup>&</sup>lt;sup>1469</sup> <sup>d</sup>Measurements from Wohlgemuth et al., 2010

Table S6. Frequency of tRNA and codon usage at varying growth rates

Frequency of tRNA <sup>a</sup>							
Doublings/hr	$\mu = 0.4$	$\mu = 0.7$	$\mu = 1.07$	$\mu = 1.6$	$\mu = 2.5$		
Ala1B	0.051	0.054	0.060	0.056	0.059		
Ala2	0.009	0.010	0.010	0.010	0.010		
Arg2	0.074	0.068	0.065	0.076	0.072		
Arg3	0.010	0.012	0.007	0.008	0.006		
Arg4	0.013	0.011	0.012	0.010	0.010		
Arg5	0.006	0.007	0.007	0.008	0.006		
Asn	0.019	0.018	0.018	0.019	0.020		
Asp1	0.037	0.037	0.035	0.038	0.042		
Cys	0.025	0.022	0.022	0.023	0.020		
Gln1	0.012	0.012	0.016	0.010	0.012		
Gln2	0.014	0.014	0.014	0.017	0.018		
Glu2	0.073	0.072	0.070	0.078	0.082		
Gly2	0.033	0.033	0.033	0.036	0.031		
Gly3	0.068	0.071	0.072	0.064	0.070		
His	0.010	0.010	0.012	0.010	0.012		
Ile1	0.054	0.054	0.056	0.060	0.069		
Leu1	0.069	0.069	0.072	0.069	0.061		
Leu2	0.015	0.016	0.017	0.015	0.016		
Leu3	0.010	0.011	0.012	0.010	0.009		
Leu4	0.030	0.029	0.030	0.031	0.026		
Leu5	0.018	0.016	0.017	0.011	0.010		
Lys	0.030	0.031	0.031	0.028	0.029		
Met f1	0.019	0.022	0.025	0.020	0.028		
Phe	0.016	0.017	0.018	0.015	0.015		
Pro1	0.014	0.011	0.014	0.009	0.007		
Pro2	0.011	0.012	0.009	0.013	0.010		
Pro3	0.009	0.009	0.009	0.008	0.007		
Sec	0.003	0.004	0.004	0.004	0.003		
Ser1	0.020	0.025	0.023	0.022	0.020		
Ser2	0.005	0.005	0.005	0.004	0.004		
Ser3	0.022	0.020	0.020	0.018	0.016		
Ser5	0.012	0.012	0.012	0.011	0.012		
Thr1	0.002	0.002	0.003	0.001	0.001		
Thr2	0.009	0.009	0.009	0.009	0.009		
Thr3	0.017	0.017	0.017	0.015	0.016		
Thr4	0.014	0.014	0.014	0.015	0.019		
Trp	0.015	0.013	0.014	0.013	0.015		
Tyr1+Tyr2	0.031	0.029	0.027	0.032	0.026		
Val1	0.060	0.055	0.047	0.061	0.057		
Val2A+2B	0.020	0.020	0.022	0.020	0.019		

Frequency of	Frequency of codon usage (1x10 <sup>-3</sup> ) <sup>b</sup>							
Doublings/hr	$\mu = 0.4$	$\mu = 0.7$	$\mu = 1.07$	$\mu = 1.6$	$\mu = 2.5$			
GGG	4.81	4.26	3.57	2.79	2.36			
GGA	2.71	2.49	2.21	1.79	1.26			
GGU	38.29	39.18	40.49	42.27	45.55			
GGC	35.62	35.58	35.54	35.49	34.17			
GAG	16.57	16.78	17.04	17.31	16.97			
GAA	53.1	53.94	55.1	56.68	57.86			
GAU	24.25	23.43	22.4	21.08	19.27			
GAC	28.72	29.65	30.93	32.35	33.74			
GUG	21.4	20.34	18.93	17.74	14.98			
GUA	15.87	17.05	18.65	19.95	22.31			
GUU	31.31	33.1	35.63	38.14	43.18			
GUC	11.25	10.58	9.71	8.86	7.67			
GCG	30.33	29.55	28.45	27.29	24.11			
GCA	22.13	22.19	22.38	23.07	24.87			
GCU	28.85	30.31	32.41	34.79	39.49			
GCC	19.8	18.5	16.81	14.67	11.81			
AGG	0.09	0.07	0.05	0.03	0.03			
AGA	1.12	0.99	0.84	0.65	0.63			
AGU	3.99	3.55	3.01	2.38	2.19			
AGC	11.97	11.4	10.69	9.88	9.31			
AAG	12.08	12.76	13.74	14.89	17.22			
AAA	44.43	46.41	49.07	51.99	55.01			
AAU	9.79	8.88	7.79	6.43	5.61			
AAC	27.95	28.22	28.64	29.02	29.21			
AUG	22.37	22.36	22.34	22.3	21.67			
AUA	0.93	0.85	0.75	0.61	0.52			
AUU	21.38	20.45	19.26	17.72	15.79			
AUC	36.68	37.72	39.15	41.38	43.86			
ACG	7.53	6.95	6.21	5.2	4.17			
ACA	3.48	3.25	2.99	2.63	2.61			
ACU	13.88	15.1	16.76	18.31	20.64			
ACC	26.51	26.77	27.1	27.47	26.7			
UGG	9.76	9.28	8.69	8.01	7.03			
UGA	0.31	0.27	0.23	0.17	0.19			
UGU	4.23	3.97	3.64	3.24	2.76			
UGC	5.29	5.06	4.77	4.35	3.81			
UAU	10.68	9.9	8.9	7.85	6.72			
UAC	16.2	16.41	16.71	16.9	16.52			
UUG	6.63	6.22	5.72	4.93	4.27			
UUA	6.13	5.46	4.64	3.56	2.73			
UUU	12.55	11.54	10.3	8.72	7.92			
UUC	22.68	22.55	22.44	22.68	23.25			

UCG	6.05	5.41	4.58	3.75	2.51
UCA	3.89	3.54	3.09	2.55	1.98
UCU	13.12	13.54	14.14	14.84	16.33
UCC	11.15	11.57	12.09	12.34	11.68
CGG	1.75	1.52	1.23	0.9	0.62
CGA	1.32	1.17	0.99	0.75	0.67
CGU	31.12	33.46	36.61	39.6	43.82
CGC	22.25	22.31	22.39	21.76	20.59
CAG	29.24	28.8	28.33	27.69	27.28
CAA	10.19	9.65	8.98	7.99	7.01
CAU	9.23	8.72	8.11	7.23	6.78
CAC	13.9	13.9	13.91	14.08	14.21
CUG	60.13	60.62	61.29	61.59	60.75
CUA	2.15	1.87	1.53	1.09	0.82
CUU	5.7	5.22	4.64	4.01	3.86
CUC	6.19	5.91	5.52	5.03	4.09
CCG	29.51	29.22	28.88	28.91	28.82
CCA	6.52	6.47	6.4	6.08	5.18
CCU	4.99	4.9	4.79	4.62	4.38
CCC	3.32	2.77	2.1	1.4	1.09

<sup>&</sup>lt;sup>a</sup>Calculated by renormalizing the molar ratio measurements reported in Dong et al., 1996 for each growth rate

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<sup>&</sup>lt;sup>b</sup>Measurements from Dong et al., 1996