Fundamental limits on the rate of bacterial cell division

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Abstract Recent years have seen a deluge of experiments dissecting the relationship between bacterial growth rate, cell size, and protein content, quantifying the abundances of single proteins across growth conditions with unprecedented resolution. However, we still lack a rigorous understanding of what sets the scale of these measurements why single protein abundances do (or do not) depend on growth rate. Here, we seek to quantitatively understand the scales of the observations in a collection of *Escherichia coli* proteomic data sets covering ≈ 4000 proteins and 31 growth conditions. We estimate the abundances of complexes needed for nutrient transport, energy generation, cell envelope biogenesis, and the processes of the central dogma, from which ribosome biogenesis emerges as a primary determinant of growth rate. We conclude by exploring a model of ribosomal regulation as a function of the nutrient supply, revealing a mechanism tying cell size and growth rate to ribosomal content.

Introduction

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The observed range of bacterial growth rates is enormously diverse. In natural environments, some microbial organisms might double only once per year (*Mikucki et al., 2009*) while in comfortable laboratory conditions, growth can be rapid with several divisions per hour (*Schaechter et al., 1958*). This six order-of-magnitude difference in time scales encompasses different microbial species and lifestyles, yet even for a single species such as *E. coli*, the growth rate can be modulated over a similar scale by tuning the type and amount of nutrients in the growth medium. This remarkable flexibility in growth rate illustrates the intimate relationship between environmental conditions and the rates at which cells convert nutrients into new cellular material – a relationship that has remained a major topic of inquiry in bacterial physiology for over a century (*Jun et al., 2018*).

As was noted by Jacques Monod, "the study of the growth of bacterial cultures does not constitute a specialized subject or branch of research, it is the basic method of Microbiology." Those words ring as true today as they did when they were written 70 years ago (*Monod*, 1949). Indeed,

the study of bacterial growth has undergone a renaissance. Many of the key questions addressed by the pioneering efforts in the middle of the last century can be revisited by examining them through the lens of the increasingly refined molecular census that is available for bacteria such as the microbial workhorse *Escherichia coli*. In this work, we explore an amalgamation of recent proteomic data sets to explore fundamental limits of bacterial growth.

Several of the evergreen questions about bacterial growth that were originally raised by microbiologists in the middle of the 20th century can now be reframed in light of this newly available data. For example, what biological processes set the absolute speed limit for how fast bacterial cells can grow and reproduce? How do cells alter the absolute numbers and relative ratios of their molecular constituents as a function of changes in growth rate or nutrient availability? In this paper, we address these two questions from two distinct angles. First, as a result of an array of high-quality proteome-wide measurements of the *E. coli* proteome under myriad growth conditions, we have a census that allows us to explore how the number of key molecular players change as a function of growth rate. This census provides a window into the question of whether the rates of central processes such as energy generation or DNA synthesis are regulated systematically as a function of cell growth rate by altering protein copy number in individual cells. Second, by compiling molecular turnover rate measurements for many of the fundamental processes associated with bacterial growth, we can make quantitative estimates to determine whether the observed protein copy numbers under varying conditions appear to be in excess of what would be minimally required to support cell growth at the observed rates.

In this paper, we make a series of order-of-magnitude estimates for the copy numbers and growth rate dependent expression of a variety of different processes, schematized in *Figure 1*, informed by the collection of proteomic data sets. We use these estimates to explore which, if any, of the hypothesis illustrated n *Figure 1* may act as molecular bottlenecks that limit bacterial growth. Specifically, we leverage a combination of *E. coli* proteomic data sets collected over the past decade using either mass spectrometry (*Schmidt et al., 2016*; *Peebo et al., 2015*; *Valgepea et al., 2013*) or ribosomal profiling (*Li et al., 2014*) across 31 unique growth conditions. Throughout, our estimates we consider a modest growth rate of $\approx 0.5 \, \text{hr}^{-1}$ corresponding to a doubling time of $\approx 5000 \, \text{seconds}$, as the the data sets heavily sample this regime. While we formulate point estimates for the complex abundances at this division time, we consider how these values will vary at other growth rates due to changes in cell size, surface area, and chromosome copy number (*Taheri-Araghi et al., 2015*).

Broadly, we find that for the majority of these estimates the protein copy numbers appear well-tuned for the task of cell doubling at a given growth rate. From our analysis, it emerges that translation, particularly of ribosomal proteins, is the most plausible candidate for a molecular bottleneck. We reach this conclusion by considering that translation is 1) a rate limiting step for the *fastest* bacterial division, and 2) a major determinant of bacterial growth across the nutrient conditions we have considered under steady state, exponential growth. This enables us to suggest that the long-observed correlation between growth rate and cell size (*Schaechter et al., 1958; Si et al., 2017*) can be simply attributed to the increased absolute number of ribosomes per cell under conditions supporting extremely rapid growth, a hypothesis which we formally mathematize and explore.

Uptake of Nutrients

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We begin our series of estimates by considering the critical transport processes diagrammed in Figure 1(A). In order to build new cellular mass, the molecular and elemental building blocks must be scavenged from the environment in different forms. Carbon, for example, is acquired via the transport of carbohydrates and sugar alcohols with some carbon sources receiving preferential treatment in their consumption (Monod, 1947). Phosphorus, sulfur, and nitrogen, on the other hand, are harvested primarily in the forms of inorganic salts, namely phosphate, sulfate, and ammonia (Jun et al., 2018; Assentoft et al., 2016; Stasi et al., 2019; Antonenko et al., 1997; Rosenberg et al., 1977; Willsky et al., 1973). All of these compounds have different permeabilities across the 85

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Box 1. The Rules of Engagement for Order-Of-Magnitude Estimates

This work relies heavily on so-called "back-of-the-envelope" estimates to understand the abundances and growth-rate dependences of a variety of molecular complexes. This moniker arises from the limitation that any estimate should be able to fit on the back of a postage envelope. Therefore, we must draw a set of rules governing our precision and sources of key values.

The rule of "one, few, and ten". The philosophy behind order-of-magnitude estimates is to provide a estimate of the appropriate scale, not a prediction with infinite accuracy. As such, we define three different scales of precision in making estimates. The scale of "one" is reserved for values that range between 1 and 2. For example, If a particular process has been experimentally measured to transport 1.87 protons for a process to occur, we approximate this process to require 2 protons per event. The scale of "few" is reserved for values ranging between 3 and 5. For example, we will often use Avogadro's number to compute the number of molecules in a cell given a concentration and a volume. Rather than using Avogadro's number as 6.02214×10^{23} , we will approximate it as 5×10^{23} . Finally, the scale of "ten" is reserved for values which we know within an order of magnitude. If a particular protein complex is present at 883 copies per cell, we say that it is present in approximately 103 copies per cell. These different scales will be used to arrive at simple estimates that report the expected scale of the observed data. Therefore, the estimates presented here should not be viewed as hardand-fast predictions of precise copy numbers, but as approximate lower (or upper) bounds for the number of complexes that may be needed to satisfy some cellular requirement. Furthermore, we use equality symbols (=) sparingly and frequently defer to approximation (≈) or scaling (\sim) symbols when reporting an estimate. When \approx is used, we are implicitly stating that we are confident in this estimate within a factor of a few. When a scaling symbol \sim is used, we are stating that we are confident in our estimate to within an order of magnitude. The BioNumbers Database as a source for values. In making our estimates, we often require approximate values for key cellular properties, such as the elemental composition of the cell, the average dry mass, or approximate rates of synthesis. We rely heavily on the BioNumbers Database (Milo et al., 2010) as a repository for such information. Every value we draw from

this reference in grey-boxes in each figure.

*Uncertainty in the data sets and the accuracy of an estimate.** The data sets presented in this work are the products of careful experimentation with the aim to report, to the best of their ability, the absolute copy numbers of proteins in the cell. These data, collected over the span of a few years, come from different labs and use different internal standards, controls, and even techniques (discussed further in Supplemental Section ??). As a result, there is notable disagreement in the measured copy numbers for some complexes across data sets. In assessing whether our estimates could explain the observed scales and growth-rate dependencies, we also considered the degree of variation between the different data sets. For example, say a particular estimate undercuts the observed data by an order of magnitude. If all data sets agree within a factor of a few of each other, we revisit our estimate and consider what me may have missed. However, if the data sets themselves disagree by an order of magnitude, we determine that our estimate is appropriate given the variation in the data.

this database has an associated BioNumbers ID number, abbreviated as BNID, and we provide

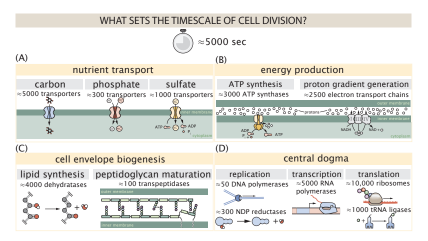


Figure 1. Transport and synthesis processes necessary for cell division. We consider an array of processes necessary for a cell to double its molecular components, broadly grouped into four classes. These categories are (A) nutrient transport across the cell membrane, (B) energy production (namely, ATP synthesis), (C) cell envelope biogenesis, and (D) processes associated with the central dogma. Numbers shown are the approximate number of complexes of each type observed at a growth rate of 0.5 hr⁻¹, or a cell doubling time of ≈ 5000 s.

cell membrane *Phillips* (2018) and most require some energetic investment either via ATP hydrolysis or through the proton electrochemical gradient to bring the material across the hydrophobic cell membrane. Given the diversity of biological transport mechanisms and the vast number of inputs needed to build a cell, we begin by considering transport of some of the most important cellular ingredients: carbon, nitrogen, oxygen, hydrogen, phosphorus, and sulfur.

The elemental composition of *E. coli* has received much quantitative attention over the past half century (*Neidhardt et al., 1991; Taymaz-Nikerel et al., 2010; Heldal et al., 1985; Bauer and Ziv, 1976*), providing us with a starting point for estimating the copy numbers of various transporters. While there is some variability in the exact elemental percentages (with different uncertainties), we can estimate that the dry mass of a typical *E. coli* cell is $\approx 45\%$ carbon (BioNumber ID: 100649, see *Box 1*), $\approx 15\%$ nitrogen (BNID: 106666), $\approx 3\%$ phosphorus (BNID: 100653), and 1% sulfur (BNID: 100655). In the coming paragraphs, we will engage in a dialogue between back-of-the-envelope estimates for the numbers of transporters needed to facilitate these chemical stoichiometries and the experimental proteomic measurements of the biological reality. Such an approach provides the opportunity to test if our biological knowledge is sufficient to understand the scale at which these complexes are produced. At the end of this section, we discuss physical limits as to the number of transporters that can be present, and comment on the plausibility of this process acting as a molecular bottleneck.

Nitrogen Transport

We must first address which elemental sources must require proteinaceous transport, meaning that the cell cannot acquire appreciable amounts simply via diffusion across the membrane. The permeability of the lipid membrane to a large number of solutes has been extensively characterized over the past century. Large, polar molecular species (such as various sugar molecules, sulfate, and phosphate) have low permeabilities while small, non-polar compounds (such as oxygen, carbon dioxide, and ammonia) can readily diffuse across the membrane. Ammonia, a primary source of nitrogen in typical laboratory conditions, has a permeability on par with water ($\sim 10^5$ nm/s, BNID:110824). In particularly nitrogen-poor conditions, *E. coli* expresses a transporter (AmtB) which appears to aid in nitrogen assimilation, though the mechanism and kinetic details of transport are still a matter of debate (*van Heeswijk et al., 2013; Khademi et al., 2004*). Beyond ammonia, another plentiful source of nitrogen come in the form of glutamate, which has its own

complex metabolism and scavenging pathways. However, nitrogen is plentiful in the growth conditions examined in this work, permitting us to neglect nitrogen transport as a potential rate limiting process in cell division in typical experimental conditions.

Carbon Transport

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We begin with the most abundant element in E. coli by mass, carbon. Using ≈ 0.3 pg as the typical F coli dry mass (BNID: 103904) we estimate that $\sim 10^{10}$ carbon atoms must be brought into the cell in order to double all of the carbon-containing molecules (Figure 2(A. top)). Typical laboratory growth conditions, such as those explored in the aforementioned proteomic data sets, provide carbon as a single class of sugar such as glucose, galactose, or xylose to name a few. F. coli has evolved myriad mechanisms by which these sugars can be transported across the cell membrane. One such mechanism of transport is via the PTS system which is a highly modular system capable of transporting a diverse range of sugars (Escalante et al., 2012). The glucose-specific component of this system transports ≈ 200 glucose molecules per second per transporter (BNID: 114686). Making the assumption that this is a typical sugar transport rate, coupled with the need to transport $\sim 10^{10}$ carbon atoms, we arrive at the conclusion that on the order of 1.000 transporters must be expressed in order to bring in enough carbon atoms to divide in 5000 s. diagrammed in the top panel of Figure 2(A). This estimate, along with the observed average number of the PTS system carbohydrate transporters present in the proteomic data sets (Schmidt et al., 2016; Peebo et al., 2015) Valgepea et al., 2013: Li et al., 2014), is shown in Figure 2(A). While we estimate 1500 transporters are needed with a 5000 s division time, we can abstract this calculation to consider any particular growth rate given knowledge of the cell density and volume as a function of growth rate and direct the reader to the Supplemental Information for more information. As revealed in Figure 2(A). experimental measurements exceed the estimate by several fold, illustrating that transport of carbon into the cell is not rate limiting for cell division. Abstracting this point estimate at 5000 s to a continuum of growth rates (grey line in *Figure 2*(A)) reveals an excess of transporters at other growth rates, though in rapid growth regimes, the abundance is below our simple estimate.

The estimate presented in *Figure 2*(A) neglects any specifics of the regulation of the carbon transport system and presents a view of how many carbohydrate transporters are present on average. Using the diverse array of growth conditions explored in the proteomic data sets, we can explore how individual carbon transport systems depend on the population growth rate. In *Figure 2*(B), we show the total number of carbohydrate transporters specific to different carbon sources. A striking observation, shown in the top-left plot of *Figure 2*(B), is the constancy in the expression of the glucose-specific transport systems. Additionally, we note that the total number of glucose-specific transporters is tightly distributed at $\approx 10^4$ per cell, the approximate number of transporters needed to sustain rapid growth of several divisions per hour. This illustrates that *E. coli* maintains a substantial number of complexes present for transporting glucose regardless of growth rate, which is known to be the preferential carbon source (*Monod, 1947*; *Liu et al., 2005*; *Aidelberg et al., 2014*).

It is now understood that a large number of metabolic operons are regulated with dual-input logic gates that are only expressed when glucose concentrations are low (mediated by cyclic-AMP receptor protein CRP) and the concentration of other carbon sources are elevated (*Gama-Castro et al., 2016; Zhang et al., 2014b*). A famed example of such dual-input regulatory logic is in the regulation of the *lac* operon which is only natively activated in the absence of glucose and the presence of allolactose, an intermediate in lactose metabolism (*Jacob and Monod, 1961*), though we now know of many other such examples (*Ireland et al., 2020; Gama-Castro et al., 2016; Belliveau et al., 2018*). This illustrates that once glucose is depleted from the environment, cells have a means to dramatically increase the abundance of the specific transporter needed to digest the next sugar that is present. Several examples of induced expression of specific carbon-source transporters are shown in *Figure 2*(B). Points colored in red (labeled by red text-boxes) correspond to growth conditions in which the specific carbon source (glycerol, xylose, or fructose) is present. These plots

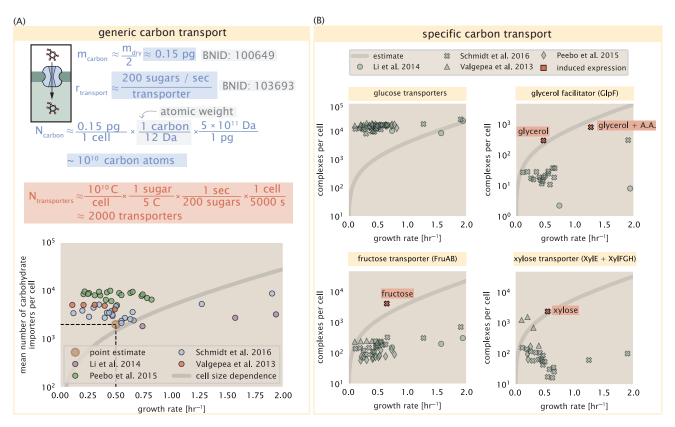


Figure 2. The abundance of carbon transport systems across growth rates. (A) A simple estimate for the minimum number of generic carbohydrate transport systems (top) assumes $\sim 10^{10}$ C are needed to complete division, each transported sugar contains ≈ 5 C, and each transporter conducts sugar molecules at a rate of ≈ 200 per second. Bottom plot shows the estimated number of transporters needed at a growth rate of ≈ 0.5 per hr (light-brown point and dashed lines). Colored points correspond to the mean number of complexes involved in carbohydrate import (complexes annotated with the Gene Ontology terms GO:0009401 and GO:0098704) for different growth conditions across different published datasets. (B) The abundance of various specific carbon transport systems plotted as a function of the population growth rate. The rates of substrate transport to compute the continuum growth rate estimate (grey line) were 200 glucose· s⁻¹ (BNID: 103693), 2000 glycerol·s⁻¹ (*Lu et al., 2003*), 200 fructose· s⁻¹ (assumed to be similar to PtsI, BNID: 103693), and 50 xylose·s⁻¹ (assumed to be comparable to LacY, BNID:103159). Red points and highlighted text indicate conditions in which the only source of carbon in the growth medium induces expression of the transport system. Grey line in (A) and (B) represents the estimated number of transporters per cell at a continuum of growth rates.

show that, in the absence of the particular carbon source, expression of the transporters is maintained on the order of $\sim 10^2$ per cell. However, when the transport substrate is present, expression is induced and the transporters become highly-expressed. The grey lines in *Figure 2*(B) show the estimated number of transporters needed at each growth rate to satisfy the cellular carbon requirement. It is notable that in all cases, the magnitude of induced expression (shown in red) falls close to the estimate, illustrating the ability of the cell to tune expression in response to changing environments. Together, this generic estimation and the specific examples of induced expression suggest that transport of carbon across the cell membrane, while critical for growth, is not the rate-limiting step of cell division.

We now turn our attention towards other essential elements, namely phosphorus and sulfur. Phos-

Phosphorus and Sulfur Transport

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phorus is critical to the cellular energy economy in the form of high-energy phosphodiester bonds making up DNA, RNA, and the NTP energy pool as well as playing a critical role in the post-translational modification of proteins and defining the polar-heads of lipids. In total, phosphorus makes up ≈3% of the cellular dry mass which in typical experimental conditions is in the form of inorganic phosphate. The cell membrane has remarkably low permeability to this highly-charged and critical molecule, therefore requiring the expression of active transport systems. In E. coli, the proton electrochemical gradient across the inner membrane is leveraged to transport inorganic phosphate into the cell (Rosenberg et al., 1977). Proton-solute symporters are widespread in E. coli (Ramos and Kaback, 1977; Booth et al., 1979) and can have rapid transport rates of 50 to 100 molecules per second for sugars and other solutes (BNID: 103159; 111777). As a more extreme example, the proton transporters in the F_1 - F_0 ATP synthase, which use the proton electrochemical gradient for rotational motion, can shuttle protons across the membrane at a rate of ≈ 1000 per second (BNID: 104890: 103390). In E. coli the PitA phosphate transport system has been shown to be very tightly coupled with the proton electrochemical gradient with a 1:1 proton; phosphate stoichiometric ratio (Harris et al., 2001: Feist et al., 2007). Taking the geometric mean of the aforementioned estimates gives a plausible rate of phosphate transport on the order of 300 per second. Illustrated in Figure 3(A), we can estimate that ≈ 200 phosphate transporters are necessary to maintain an \approx 3% dry mass with a 5000 s division time. This estimate is consistent with observation when we examine the observed copy numbers of PitA in proteomic data sets (plot in Figure 3(A)). While our estimate is very much in line with the observed numbers, we emphasize that this is likely a slight overestimate of the number of transporters needed as there are other phosphorous scavenging systems, such as the ATP-dependent phosphate transporter Pst system which we have neglected. Satisfied that there are a sufficient number of phosphate transporters present in the cell, we now turn to sulfur transport as another potentially rate limiting process. Similar to phosphate, sulfate is highly-charged and not particularly membrane permeable, requiring active transport. While there exists a H+/sulfate symporter in E. coli, it is in relatively low abundance and is not well characterized (Zhang et al., 2014a). Sulfate is predominantly acquired via the ATP-dependent ABC transporter CysUWA system which also plays an important role in selenium transport (Sekowska et al., 2000: Sirko et al., 1995). While specific kinetic details of this transport system are not readily available, generic ATP transport systems in prokaryotes transport on the order of 1 to 10 molecules per second (BNID: 109035). Combining this generic transport rate, measurement of sulfur comprising 1% of dry mass, and a 5000 second division time yields an estimate of \approx 1000 CysUWA complexes per cell (Figure 3(B)). Once again, this estimate is in notable agreement with proteomic data sets. suggesting that there are sufficient transporters present to acquire the necessary sulfur. In a similar spirit of our estimate of phosphorus transport, we emphasize that this is likely an overestimate of the number of necessary transporters as we have neglected other sulfur scavenging systems that are in lower abundance

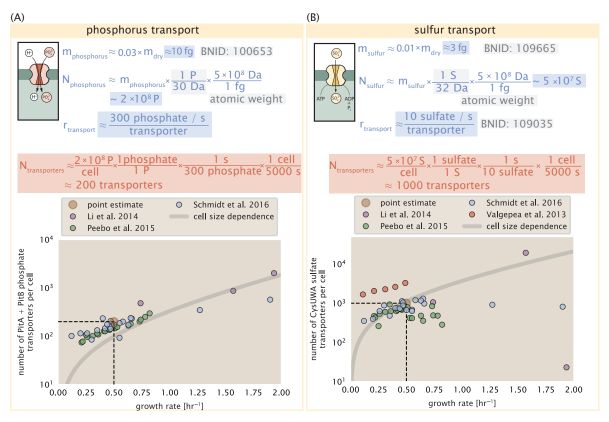


Figure 3. Estimates and measurements of phosphate and sulfate transport systems as a function of growth rate. (A) Estimate for the number of PitA phosphate transport systems needed to maintain a 3% phosphorus *E. coli* dry mass. Points in plot correspond to the the total number of PitA transporters per cell. (B) Estimate of the number of CysUWA complexes necessary to maintain a 1% sulfur *E. coli* dry mass. Points in plot correspond to average number of CysUWA transporter complexes that can be formed given the transporter stoichiometry [CysA]₂[CysU][CysW][Sbp/CysP]. Grey line in (A) and (B) represents the estimated number of transporters per cell at a continuum of growth rates.

64 Limits on Transporter Expression

So which, if any, of these processes may be rate limiting for growth? As suggested by Figure 2 265 (B), induced expression can lead to an order-of-magnitude (or more) increase in the amount of 266 transporters needed to facilitate transport. Thus, if acquisition of nutrients was the limiting state 267 in cell division, could expression simply be increased to accommodate faster growth? A way to 268 approach this question is to compute the amount of space in the bacterial membrane that could 269 be occupied by nutrient transporters. Considering a rule-of-thumb for the surface area of E. coli of 270 about 5 um² (BNID: 101792), we expect an areal density for 1000 transporters to be approximately 271 200 transporters/ um². For a typical transporter occupying about 50 nm²/dimer, this amounts to about only 1 percent of the total inner membrane (Szenk et al., 2017). In addition, bacterial cell 273 membranes typically have densities of 105 proteins/um² (Phillips, 2018), implying that the cell could accommodate more transporters of a variety of species if it were rate limiting. As we will see in the next section, however, occupancy of the membrane can impose other limits on the rate of energy production.

Translation and Ribosomal Synthesis

Lastly, we turn our attention to the process of synthesizing new proteins, translation. This process stands as a good candidate for potentially limiting growth since the synthesis of new proteins relies on the generation of ribosomes, themselves proteinaceous molecules. As we will see in the coming sections of this work, this poses a "chicken-or-the-egg" problem where the synthesis of ribosomes requires ribosomes in the first place.

We will begin our exploration of protein translation in the same spirit as we have in previous sections – we will draw order-of-magnitude estimates based on our intuition and available literature, and then compare these estimates to the observed data. In doing so, we will estimate both the absolute number of ribosomes necessary for replication of the proteome as well as the synthesis of amino-acyl tRNAs. From there we consider the limitations on ribosomal synthesis in light of our estimates on both the synthesis of ribosomal proteins and our earlier results on rRNA synthesis.

tRNA Synthetases

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We begin by first estimating the number of tRNA synthetases in *E. coli* needed to convert free amino-acids to polypeptide chains. Again using an estimate of $\approx 3 \times 10^6$ proteins per cell at a 5000 s division time (BNID: 115702) and a typical protein length of ≈ 300 amino acids (BNID: 100017), we can estimate that a total of $\approx 10^9$ amino acids are stitched together by peptide bonds.

How many tRNAs are needed to facilitate this remarkable number of amino acid delivery events to the translating ribosomes? It is important to note that tRNAs are recycled after they've passed through the ribosome and can be recharged with a new amino acid, ready for another round of peptide bond formation. While some *in vitro* data exists on the turnover of tRNA in *E. coli* for different amino acids, we can make a reasonable estimate by comparing the number of amino acids to be polymerized to cell division time. Using our stopwatch of 5000 s and 10^9 amino acids, we arrive at a requirement of $\approx 2 \times 10^5$ tRNA molecules to be consumed by the ribosome per second.

There are many processes which go into synthesizing a tRNA and ligating it with the appropriate amino acids. As we discussed previously, there appear to be more than enough RNA polymerases per cell to synthesize the needed pool of tRNAs. Without considering the many ways in which amino acids can be scavenged or synthesized *de novo*, we can explore ligation the as a potential rate limiting step. The enzymes which link the correct amino acid to the tRNA, known as tRNA synthetases or tRNA ligases, are incredible in their proofreading of substrates with the incorrect amino acid being ligated once out of every 10⁴ to 10⁵ events (BNID: 103469). This is due in part to the consumption of energy as well as a multi-step pathway to ligation. While the rate at which tRNA is ligated is highly dependent on the identity of the amino acid, it is reasonable to state that

the typical tRNA synthetase has charging rate of \approx 20 AA per tRNA synthetase per second (BNID: 105279).

We can make an assumption that amino-acyl tRNAs are in steady-state where they are produced at the same rate they are consumed, meaning that 2×10^5 tRNAs must be charged per second. Combining these estimates together, as shown schematically in *Figure 4*(A), yields an estimate of $\sim 10^4$ tRNA synthetases per cell with a division time of 5000 s. This point estimate is in very close agreement with the observed number of synthetases (the sum of all 20 tRNA synthetases in *E. coli*). This estimation strategy seems to adequately describe the observed growth rate dependence of the tRNA synthetase copy number (shown as the grey line in *Figure 4*(B)), suggesting that the copy number scales with the cell volume.

In total, the estimated and observed $\sim 10^4$ tRNA synthetases occupy only a meager fraction of the total cell proteome, around 0.5% by abundance. It is reasonable to assume that if tRNA charging was a rate limiting process, cells would be able to increase their growth rate by devoting more cellular resources to making more tRNA synthases. As the synthesis of tRNAs and the corresponding charging can be highly parallelized, we can argue that tRNA charging is not a rate limiting step in cell division, at least for the growth conditions explored in this work.

Protein Synthesis

With the number of tRNA synthetases accounted for, we now consider the abundance of the protein synthesis machines themselves, ribosomes. Ribosomes are enormous protein/rRNA complexes that facilitate the peptide bond formation between amino acids in the correct sequence as defined by the coding mRNA. Before we examine the synthesis of the ribosome proteins and the limits that may place on the observed bacterial growth rates, let's consider replication of the cellular proteome.

While the rate at which ribosomes translates is well known to have a growth rate dependence *Dai et al.* (2018) and is a topic which we discuss in detail in the coming sections. However, for the purposes of our order-of-magnitude estimate, we can make the approximation that translation occurs at a rate of ≈ 15 amino acids per second per ribosome (BNID: 100233). Under this approximation and assuming a division time of 5000 s, we can arrive at an estimate of $\approx 10^4$ ribosomes are needed to replicate the cellular proteome, shown in *Figure 4*(B). This point estimate, while glossing over important details such as chromosome copy number and growth-rate dependent translation rates, proves to be notably accurate when compared to the experimental observations (*Figure 4*(B)).

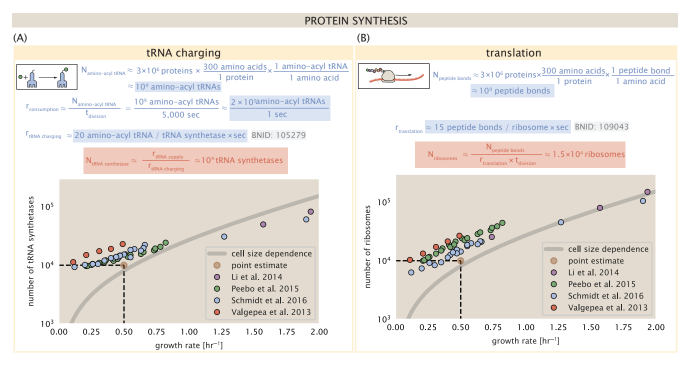


Figure 4. Estimation of the required tRNA synthetases and ribosomes. (A) Estimation for the number of tRNA synthetases that will supply the required amino acid demand. The sum of all tRNA synthetases copy numbers are plotted as a function of growth rate ([ArgS], [CysS], [GlnS], [GltX], [IleS], [LeuS], [ValS], [AlaS]_2, [AsnS]_2, [TyrS]_2, [TrpS]_2, [ThrS]_2, [SerS]_2, [ProS]_2, [PheS]_2[PheT]_2, [MetG]_2, [lysS]_2, [HisS]_2, [GlyS]_2[GlyQ]_2). (B) Estimation of the number of ribosomes required to synthesize 10^9 peptide bonds with an elongation rate of 15 peptide bonds per second. The average abundance of ribosomes is plotted as a function of growth rate. Our estimated values are shown for a growth rate of 0.5 hr⁻¹. Grey lines correspond to the estimated complex abundance calculated at different growth rates. See Supplemental Information XX for a more detail description of this calculation.

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