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ORIGINAL ARTICLE

Evolutionary tradeoffs in cellular composition across diverse bacteria

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One of the most important classic and contemporary interests in biology is the connection between cellular composition and physiological function. Decades of research have allowed us to understand the detailed relationship between various cellular components and processes for individual species, and have uncovered common functionality across diverse species. However, there still remains the need for frameworks that can mechanistically predict the tradeoffs between cellular functions and elucidate and interpret average trends across species. Here we provide a comprehensive analysis of how cellular composition changes across the diversity of bacteria as connected with physiological function and metabolism, spanning five orders of magnitude in body size. We present an analysis of the trends with cell volume that covers shifts in genomic, protein, cellular envelope, RNA and ribosomal content. We show that trends in protein content are more complex than a simple proportionality with the overall genome size, and that the number of ribosomes is simply explained by cross-species shifts in biosynthesis requirements. Furthermore, we show that the largest and smallest bacteria are limited by physical space requirements. At the lower end of size, cell volume is dominated by DNA and protein content—the requirement for which predicts a lower limit on cell size that is in good agreement with the smallest observed bacteria. At the upper end of bacterial size, we have identified a point at which the number of ribosomes required for biosynthesis exceeds available cell volume. Between these limits we are able to discuss systematic and dramatic shifts in cellular composition. Much of our analysis is connected with the basic energetics of cells where we show that the scaling of metabolic rate is surprisingly superlinear with all cellular components.

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Introduction

An important question in evolution, ecology and astrobiology is: what fundamental limitations are present for the simplest organisms and what limitations might have been present when life emerged and started to evolve? It is important to understand how limitations differ for broad classes of organisms as this informs competitive dynamics and motivates evolutionary trajectories and transitions. Although speciation is the subtle process of unique evolutionary trajectories, it should be noted that these trajectories occur within boundaries defined by basic physical, energetic and chemical limitations. For example, scaling laws relating numerous organism features to body size often illustrate a shared physical limitation that, on the average, organizes

biological features (e.g., Niklas, 2004; Brown et al., 2004; West and Brown, 2005; DeLong et al., 2010). Recent work has shown that these scaling laws can be connected with metabolic partitioning to anticipate the smallest possible bacterium (Kempes et al., 2012). Indeed, in the fields of bioengineering, ecology and evolution, there are a large number of known scaling relationships, and mapping the entire space of physical, energetic and chemical limitations is a significant task of central consequence for evolution and ecology (Brown et al., 2004).

Here we present an analysis, which we verify using compiled data, of the changes in molecular, physiological and structural composition across the entire range of bacterial cell sizes. Specifically, we examine the total amount of the following components: DNA, ribosomes, proteins, cell membrane, tRNA and mRNA. For each of these features we often focus on power-law relationships for various organism features, and when two properties 'scale' with different exponents, this often implies the onset of a limitation at a given size which may have important evolutionary consequences. Using this framework

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we show that the interconnection between energetic, physical, informational (genomic), chemical and temporal processes leads to predictions of the upper and lower boundaries of bacterial size and defines the evolutionary flexibility for bacteria between these two bounds. This perspective, which spans a huge diversity of species and five orders of magnitude in cell volume, adds explicit cross-species and size-based dependencies to the large body of work on the interconnection between various cellular processes (e.g., Tempest and Hunter, 1965; Tempest et al., 1965; Simon and Azam, 1989; Bremer et al., 1996; Cayley and Record, 2003; Dethlefsen and Schmidt, 2007; Zaslaver et al., 2009; DeLong et al., 2010; Scott et al., 2010; Goehring and Hyman, 2012; Kempes et al., 2012; Turner et al., 2012; Lloyd, 2013; Burnap, 2015). Furthermore, our work connects these cross-species trends to previous considerations of space limitation for the smallest organisms (Knoll et al., 1999) and the recent observations of small environmental bacteria (e.g., Luef et al., 2015). Knowing these connections may help us to infer ecological trait tradeoffs, quantify the constraints in designing organisms or interpret long-standing environmental observations such as the Redfield ratio for nutrient proportionalities in the ocean (Geider and La Roche, 2002).

Finally, most of the cellular components analyzed here should have a deep connection with overall cellular metabolism. In some cases, such as the processes determined by growth rate (Kempes et al., 2012), metabolic rate explicitly constrains component requirements. However, we find that it is more difficult to determine what cellular features set overall metabolic rate, which scales superlinearly for bacteria in contrast to other classes of organisms (DeLong *et al.*, 2010). In fact, we find that there is no obvious connection between the scaling of any single cellular feature and the overall metabolic rate. highlighting the continuing mystery of metabolic scaling in bacteria. We also find increasing trends in the overall efficiency of bacteria defined as the metabolic power per genetic and protein content, which is surprising given previous considerations (Lane and Martin, 2010).

Materials and methods

Our overarching goal is to understand how the requirement for each cellular component depends on total cell size so that we can constrain the limits facing cells at various scales. We approach these constraints from both energetic perspectives and overall space constraints, the latter of which often derives from energetics via the requirements of certain rate processes (e.g., growth rates; Kempes et al., 2012). For the space constraints our analysis focuses on both the scaling of individual components and the sum total of all cellular components. Thus, it is useful to define the total volume of the

cell, $V_{\rm c}$, as

$$V_{\rm c} = V_{\rm w} + V_{\rm comp} \tag{1}$$

where $V_{\rm w}$ is the total volume of water in the cell and $V_{\rm comp}$ is the volume of cellular components defined as

$$V_{\rm comp} = V_{\rm DNA} + V_{\rm p} + V_{\rm r} + V_{\rm env} + V_{\rm tRNA} + V_{\rm mRNA}$$
 (2) where $V_{\rm DNA}$ is the total volume of DNA in the cell, $V_{\rm p}$ is the protein volume, $V_{\rm r}$ is the ribosomal volume, $V_{\rm env}$ is the volume of tRNA and $V_{\rm mRNA}$ is the total mRNA volume. One of our main efforts here is to determine the sizes at which single components or the total component volume limits cell size, and the combination of the both types of analysis allows us to infer tradeoffs across the range of bacteria. Additionally, it is useful within our considerations to define the volumetric fraction, $f_{\rm d}$, of the cell that is dry weight as

$$f_{\rm d} = \frac{V_{\rm comp}}{V_{\rm c}}.$$
 (3)

In general, we describe the scaling of a generic cell component (e.g., DNA) as $C_i = C_0 V_c^{\beta_i}$, where β_i is the exponent for volume (measured in m³ throughout this paper) dependence of the component. It is important to note that if $\beta_i < 1$ then component C_i will be a decreasing fraction of total cell volume with increasing cell size, while $\beta_i > 1$ leads to an increasing fraction of cell volume. In either case, if $\beta_i \neq 1$, then there will be a size at which the component volume equals total cell volume, implying a limitation because extrapolation beyond this point would produce $C_i > V_c$, which is impossible. It should also be noted that if some of the terms in V_{comp} scale with $\beta_i \neq 1$ then other terms cannot be simple power laws in order for the equalities in Equations (1) and (2) to hold. We will see that several aspects of the cell do not follow simple power-law relationships with overall cell volume. It should be noted, more generally, that if any component depends on V_c in a manner that is not a simple proportionality then the fractional composition of the cell will change across cell sizes, and many nonlinear functions will lead to critical sizes beyond which the volume of a component will exceed V_c , a physical impossibility that implies a constraint on size.

The basic perspective of physiological scaling is supported by many previous observations of organism features that follow power-law relationships; however, it should be noted that in many of the cases studied here more complicated relationships are warranted as illustrated, for example, by previous efforts on growth rates (Kempes *et al.*, 2012). Furthermore, many cellular features are expected to be connected mechanistically with others (e.g., growth rates and ribosome content), and Figure 1 illustrates these interconnections, along with the cascading dependence of many cellular features on total cell volume. We begin with the scaling of

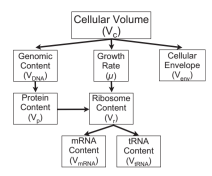


Figure 1 A schematic showing the proposed dependencies of various cellular features on cell volume where several of these relationships are connected to cell volume through several layers of dependencies. It should be noted that many of these features are intimately connected with cellular metabolism and energetics and that these connections have not been drawn in this figure. For example, previous work has shown that the growth rate, μ , is connected to the total cell size via the scaling of total metabolism and the energetics of basic maintenance requirements (Kempes et al., 2012).

cellular energetics given the central role that these will play in our analysis and the previous demonstrations that energetics can be used to derive other features such as growth rate (Kempes *et al.*, 2012).

Metabolic rate

Previous work has shown that metabolic rate, *B* (W), scales with body size according to

$$B = B_0 V_{\rm C}^{\beta_B} \tag{4}$$

where $\beta_B \approx 2$ in bacteria (DeLong *et al.*, 2010) ($\beta_B \approx 1.7$ for an OLS fit) and B_0 is a normalization constant with units W (m³Cell)^{- β_B}. Although most previous allometric studies focus on mass as the size unit for organisms, we use cell volume, V_c , here because this is typically what is directly measured in bacteria and, as can be inferred from our later findings, conversion to mass units can be complicated as cell density is not constant with cell size (we later find that density is greatest for the smallest and largest bacteria with a minimum value of 1.06 × 10⁶ g m⁻³ for an intermediate cell size of $4.92 \times 10^{-18} \,\mathrm{m}^3$; see Supplementary Figure S5). Furthermore, many previous studies report wet masses, which were the result of applying a constant density factor to measured volumes, so conversion to volume straightforward: $V_{\rm c} = m_{\rm w}/d_{\rm c}$ where units $d_c = 1.1 \times 10^6 \,\mathrm{g \, m^{-3}}$ is the previously used cell density assumed to be constant and $m_{\rm w}$ is the calculated wet mass from previous studies (e.g., West and Brown, 2005; Makarieva et al., 2008; DeLong et al., 2010). It should be noted that this conversion will not change the scaling exponents found in studies that use wet mass provided that these studies used a constant value for $d_{\rm c}$.

Growth rates

To derive growth rates, it has been demonstrated (Kempes *et al.*, 2012) that the metabolic scaling relationship from the preceding section can be combined with the concept of metabolic partitioning according to

$$B = E_{\rm m} \frac{dm}{dt} + B_{\rm m} m \tag{5}$$

where $B_{\rm m}$ (W g⁻¹) is the unit maintenance metabolism and $E_{\rm m}$ ($J g^{-1}$) is the unit cost of biosynthesis. Both of these parameters can be derived from bulk community energetic constants of the yield (e.g., g cells per mol resource) and maintenance (mols of resource consumed for survival alone) coefficients, measured, for example, in a chemostat and converted to energy units (e.g., J per mol of resource; see Kempes et al., 2012 for a more detailed presentation). The partitioned metabolism in Equation (5) has been used to derive a single-cell growth curve (Kempes et al., 2012), from which it is possible to determine the generation or division time, t_d (s), of a single cell, along with the cross-species population growth rate, $\mu \equiv \ln(2)/t_d$ (s⁻¹), of bacteria following the relationship

$$\mu = \frac{(B_{\rm m}/E_{\rm m})(1 - \beta_B)\ln[\varepsilon]}{\ln\left[\frac{1 - (B_{\rm m}/B_0)(V_{\rm c}d_{\rm c})^{1 - \beta_B}}{1 - \varepsilon^{1 - \beta_B}(B_{\rm m}/B_0)(V_{\rm c}d_{\rm c})^{1 - \beta_B}}\right]}$$
(6)

where $\varepsilon \approx 2$ is the ratio of the size of the cell at division compared with its initial size. Our efforts here largely concern the physiological requirements of a single cell, and thus most of our derivations (e.g., Supplementary Equations (S6)-(S47)) rely fundamentally on the division time t_d ; however, the final results appear as functions of $\ln(2)/t_d$ and so it is convenient to report equations in terms of μ . It should also be noted that we have presented this equation with the previous conversion (Kempes et al., 2012) between cell volume and wet mass explicitly stated, where d_c (g m⁻³) is the cell density. The relationship for μ in Equation (6) demonstrates an asymptotic behavior for the smallest bacteria that limits the smallest possible cells (Kempes et al., 2012) and approximates the power law

$$\mu \approx \mu_0 V_{\rm c}^{\beta_B - 1} \tag{7}$$

for larger cell volumes (see Supplementary Information) showing that growth rate increases rapidly as bacteria become larger. It should be noted that in this framework $V_{\rm c}$ is the only true variable while all of the other parameters are, on average, expected to be constants, which has been supported by data (Kempes $et\ al.$, 2012). However, it should be noted that the model could incorporate future discoveries of unknown dependencies of any of the constants; for example, $B_{\rm m}$ could have a complex response under extreme energy limitation, which is a regime that has not yet been well characterized. The overall growth rate, as represented by μ , defines the

2148

dominant rate processes of the cell, and we will later show that this connection is central to determining the number of ribosomes, tRNA and mRNA.

Given the strong dependence of growth rate on cell size, we should furthermore expect significant shifts in macromolecular content for bacteria of different volumes. This is consistent with previous work showing that the growth rate of bacteria is strongly connected with the relative abundance of various macromolecular components, most notably ribosomes, RNA and proteins (e.g., Bremer *et al.*, 1996; Zaslaver *et al.*, 2009; Scott *et al.*, 2010; Burnap, 2015).

Genome size

It is interesting then to consider how the genomic complexity of bacteria changes in comparison with the patterns in metabolic and growth rates. As bacteria are becoming larger how rapidly is their genome increasing? Previous efforts have shown that genome size has a strong scaling with cell size in bacteria (Shuter *et al.*, 1983; West and Brown, 2005; DeLong *et al.*, 2010) and here we consider

$$V_{\rm DNA} = D_0 V_{\rm c}^{\beta_{\rm D}} \tag{8}$$

where we later determine D_0 and β_D from a compilation of data and previous compilations.

Protein scaling

The genomic scaling law is expected to be a powerful tool for understanding basic constraints on bacteria because it should influence many other features of the cell. For example, the total volume of proteins can be related to genome length by knowing how many proteins are produced per gene:

$$V_{\rm p} = \frac{\overline{n}_{\rm p} \overline{v}_{\rm p} L}{\overline{I}_{\rm p}} \tag{9}$$

where \overline{v}_p (m³) is the volume of an average protein, \overline{l}_p (bp) is the average length, in nucleotides, of a protein encoding gene and \overline{n}_p (copies cell⁻¹) is the average copy number of a protein, and

$$L = \frac{V_{\rm DNA}}{\overline{v}_{\rm N}} \tag{10}$$

is the length of the genome given the average volume of a nucleotide $\overline{\nu}_N$ (m³) (see Supporting Information and Supplementary Figure S1) which allows us to write the volume-dependent form:

$$V_{\rm p} = \frac{\overline{n}_{\rm p} \overline{v}_{\rm p} D_0 V_{\rm c}^{\beta_{\rm D}}}{\overline{l}_{\rm p} \overline{v}_{\rm N}} \tag{11}$$

In Equation (11), \overline{v}_p , \overline{l}_p and \overline{n}_p could each scale with cell volume. This relationship for protein volume specifies the explicit relationship with protein copy number and the scaling of genome size with cell volume, however, for simplicity in presenting results

we consider and report the analogous scaling

$$V_{\rm p} = P_0 V_{\rm c}^{\beta_{\rm p}} \tag{12}$$

which can be easily connected with Equation (11).

Model of ribosome requirement

We derive the ribosome requirement by considering how many ribosomes are required to replicate all ribosomes and proteins within a division cycle while also replacing proteins and ribosomes that have been degraded. This perspective is complicated by the fact that as ribosomes and proteins are dynamically produced they contribute, respectively, to the biosynthetic capacity or biosynthetic requirements of a cell. This leads to a conceptually simple, but algebraically complicated derivation, which we present in the Supplementary Information, with the simple final ribosome requirement that

$$N_{\rm r} \ge \frac{\overline{l}_{\rm p} N_{\rm p} \left(\frac{\phi}{\mu} + 1\right)}{\frac{\overline{l}_{\rm r}}{\mu} - \overline{l}_{\rm r} \left(\frac{\eta}{\mu} + 1\right)} \tag{13}$$

where $l_{\rm r}$ is the average length of a ribosome in base pairs, $N_{\rm r}$ is the number or ribosomes in the cell, $r_{\rm r}$ (bp s⁻¹) is the maximum base pair processing rate of the ribosome which is assumed to be constant across both taxa and cell size, η (s⁻¹) and φ (s⁻¹) are specific degradation rates for ribosomes and proteins, respectively, and $N_{\rm p}$ is the total number of proteins given by

$$N_{\rm p} = \frac{V_{\rm p}}{\overline{V}_{\rm p}} = \frac{\overline{n}_{\rm p} D_0 V_{\rm c}^{\beta_{\rm D}}}{\overline{l}_{\rm p} \overline{v}_{\rm N}}$$
(14)

which depends on overall cell volume. The total number of proteins could also be found using Equation (12) as $N_{\rm p}=P_0V_{\rm c}^{\beta_{\rm p}}/\overline{v}_{\rm p}$ when $P_{\rm o}$ and $\beta_{\rm p}$ are known. In Equation (13) it is important to note that μ is a function determined solely by overall cell size given that the other parameters in Equation (6) are expected to be pure constants. In the Supplementary Information we present $N_{\rm r}$ with the dependence on $V_{\rm c}$, via μ and $N_{\rm p}$, explicitly stated (Supplementary Equations (S46) and (S47)). Our relationship for the number of ribosomes can be converted to the total volume of required ribosomes in the cell:

$$V_{\rm r} = \overline{V}_{\rm r} N_{\rm r} \tag{15}$$

where \overline{v}_r is the average volume of a ribosome (see Supplementary Information for the explicit dependence on V_c).

An alternative perspective, commensurate with our considerations of the other cellular components, is that the number of ribosomes follows a simple scaling with cell volume:

$$V_{\rm r} = R_0 V_{\rm c}^{\beta_{\rm R}} \tag{16}$$

and we also test this possibility when analyzing trends in the quantity of ribosomes.

Trends in other RNA

In relationship to the number of ribosomes we must also consider the requirements for tRNA and mRNA. The minimum number of tRNA and mRNA in the cell should be equal to the number of ribosomes so that every ribosome is engaged in protein synthesis and the cell is maximizing the biosynthetic rate. However, this does not consider the time scales required for mRNA and tRNA to find and 'react' with the ribosome. Another reasonable assumption is that there is some local concentration of tRNA and mRNA which allows the ribosome to operate at its maximum capacity. If this concentration is held constant, then the total number of tRNA and mRNA in the cell should be proportional to the number of ribosomes, as should the total volume of each of these components. Thus, we have

$$V_{\text{tRNA}} = \overline{V}_{\text{tRNA}} \overline{n}_{\text{tRNA}} N_{\text{r}} \tag{17}$$

where \overline{v}_{tRNA} is the average volume of a tRNA, and \overline{n}_{tRNA} is the average number of tRNA per ribosome, and

$$V_{\text{mRNA}} = \overline{V}_{\text{mRNA}} \overline{n}_{\text{mRNA}} N_{\text{r}} \tag{18}$$

where \overline{v}_{mRNA} is the average volume of an mRNA and \overline{n}_{mRNA} is the average number of mRNA per ribosome. It should be noted that Equations (17) and (18) both depend on cell volume because N_r is a function of V_c , and the explicit form is included in the Supplementary Information. Our estimates for the number of mRNA per ribosome are in fact close to one, implying that most ribosomes are engaged in protein synthesis ($\overline{n}_{mRNA} = 1.09$, see Supplementary Information).

Cellular envelope

Completing our analysis of cellular composition we consider the volume of the cellular envelope which we can approximate, considering a spherical cell, as

$$V_{\text{env}} = \left(V_{\text{c}} - \frac{4}{3}\pi \left[\left(\frac{3V_{\text{c}}}{4\pi}\right)^{1/3} - r_{\text{env}} \right]^{3} \right) \left(1 - \overline{p}_{\text{p}}\right)$$

$$\tag{19}$$

where $r_{\rm env}$ (m³) is the effective thickness of the cellular envelope and $\overline{p}_{\rm p}$ is the average percentage of the envelope that is occupied by proteins (we find that this is roughly equal to 0.15, see Supplementary Information). It should be noted that $r_{\rm env}$ will vary depending on whether the cell is Gram negative or positive.

Predicting limiting behavior

A key aspect of our scaling analysis is that scaling exponents that deviate from 1 imply that there is a distinct bounding size at which a given component will equal and then exceed the total cell volume. This represents an inferred lower bound for β <1 and an upper bound for β >1. For example, the

relationship for genome volume (Equation (8)) gives the size of a cell that is entirely filled by DNA, $V_{\rm DNA} = V_{\rm c}$, as

$$V_{\rm DNA}^* = D_0^{1/(1-\beta_{\rm D})} \tag{20}$$

and, from Equation (12), the volume at which proteins are predicted to entirely fill the cell would be given by

$$V_{\rm p}^* = P_0^{1/\left(1-\beta_{\rm p}\right)} \tag{21}$$

Similarly, a major prediction of the ribosome requirements (Equation (13)) is the appearance of two asymptotes at both the small and large end of bacterial sizes. The small-end asymptote is characterized by the previous observation that the division rate will become zero for a cell volume of roughly $1.45 \times 10^{-20} \, \mathrm{m}^3$ (Kempes *et al.*, 2012). The large-end asymptote is characterized by the point at which the time to divide is not sufficient to replicate just the ribosomes, and in the context of Equation (13) this is the point when the denominator is less than or equal to zero leading to

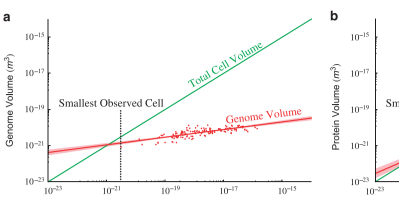
$$\mu \le \frac{\overline{r}_{r} - \eta \overline{l}_{r}}{\overline{l}_{r}} \tag{22}$$

which we term the 'ribosome catastrophe'.

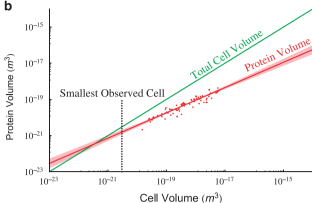
Beyond these single component limits we also consider the limitation faced by the combination of all overall cellular components given in Equation (2). Here it should be noted that some of the subcomponents follow power laws, others do not (e.g., RNA components), and the overall relationship, which represents a sum, has a more complicated nonlinear form than a simple power law. However, it is still possible to solve for the point at which $V_{\text{tot}} = V_{\text{c}}$, where it should be noted that the errors associated with each scaling may become amplified at the largest and smallest scales. In plotting the relationships, either predicted or fit, for single components and the total component volume we extrapolate the fits to regions where these volumes would exceed the total cell volume. This is physically impossible, but we make these extrapolations to show the scale of the challenge faced at certain sizes in terms of component requirements.

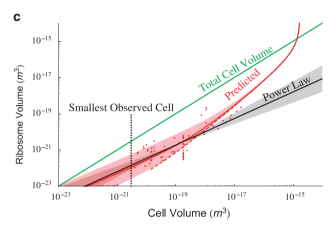
Results and discussion

Basic scaling relationships and component trends Figure 2a gives our own compilation (which includes data from Shuter et al., 1983; West and Brown, 2005; DeLong et al., 2010) of genome size, $V_{\rm DNA}$, showing that the scaling follows Equation (8) with $D_0=3.0\times 10^{-17}~({\rm m}^3\,{\rm DNA}~({\rm m}^3{\rm Cell})^{-\beta_{\rm D}})$ and $\beta_{\rm D}=0.21\pm0.03$ (see Supplementary Information for the 95% confidence intervals on all reported normalizations constants such as D_0). As previously noted (DeLong et al., 2010), this scaling shows that genome size is increasing much less steeply than metabolic



Cell Volume (m3)





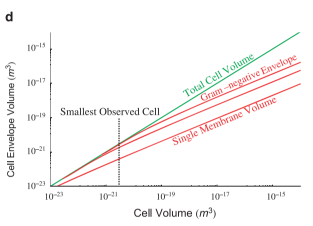


Figure 2 The cell-volume-dependent scaling of total (a) genome volume, (b) protein volume, (c) ribosome volume and (d) cellular envelope volume. In each plot compiled data are given as red points along with predictions or model fits in red and the 95% confidence intervals as shaded regions around each curve. The green curves represent the total cell volume (one-to-one line) for reference, and the volume of the smallest observed cell is noted by the black dashed line (Seybert et al., 2006; Luef et al., 2015). In (c) the black curve is a best-fit power law, the red curve is the prediction from Equations (13)–(15) and the dashed line represents a pure prediction for the lower bound on the number of ribosomes given measured values for degradation rates and our previous model of μ . For (d) the volumes of the cellular envelope are given for an average Gram-negative and -positive bacterium along with the volume of a single membrane. Please see the Supplementary Information for a summary of the data compilations.

rate, which is perhaps surprising, although it is not clear how the number of genes should explicitly determine overall metabolism. For example, perhaps metabolic rate should be related instead to the number of copies of certain pathway genes or to the concentration of specific proteins in the cell (Burnap, 2015).

Our analysis, which includes data compiled in Tempest and Hunter (1965), Tempest et~al. (1965), Simon and Azam (1989), Dethlefsen and Schmidt (2007), Milo (2013) and Valgepea et~al. (2013), shows that the scaling for the volume of protein compared with that of overall cell volume is also less than linear, with $P_0=3.42\times 10^{-7}~(\rm m^3 Protein~(\rm m^3 Cell)^{-\beta_p})$, and $\beta_p=0.70\pm 0.06~(\rm Figure~2b)$. Considering the observed scaling for protein volume and DNA volume in the context of Equation (11), we calculate that $\frac{\overline{n}_p \overline{\nu}_p}{\overline{l}_p}$ scales like $V_c^{0.49}$. It is likely that $\overline{\nu}_p$ and \overline{l}_p are roughly constant across cells (e.g., the average gene length has been found to be approximately invariant across all bacteria (Xu et~al., 2006)) and, therefore, this result implies that the average copy number of

proteins is likely increasing as cells become larger. Thus, not only are the number of unique proteins increasing for larger cells and genomes but the number of times that each protein is copied is also increasing.

The departure of protein scaling from genome scaling gives an example of a cellular feature that is not simply predicted by a proportionality with overall genome size (e.g., the surprising scaling of $V_{\rm p} \propto L^{3.33}$ given $\beta_{\rm p}/\beta_{\rm D} \approx 3.33$). Given that total protein content is expected to play a strong role in determining the number of ribosomes (Equation (13)) and the overall RNA content of the cell, we should also expect these features to have a complicated connection with the genomic complexity of a bacterium.

We predict the relationship between the total volume of ribosomes and overall cell volume from Equations (13)–(15) using the asymptotic form of μ (see Supplementary Information for explicit forms) and measured values for the degradation rates ϕ and η . This prediction should represent a true lower bound, and in Figure 2c this curve (dashed line)

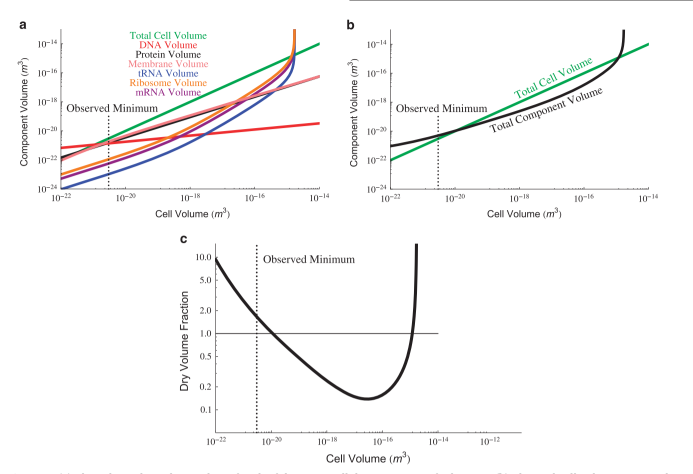


Figure 3 (a) The volume-dependent scaling of each of the major cellular components for bacteria. (b) The total cell volume compared with the volume of all cellular components as a function of cell size. (c) The fraction of total cell volume that is occupied by the essential components. It should be noted that in each of these plots we have extrapolated curves to regions that are not physically possible (such as the dry fraction exceeding 1) in order to illustrate crossings that represent limiting sizes, and to show the increasing challenges faced by bacteria beyond these critical values.

tracks, but is consistently smaller than the observed volume of ribosomes from our cross-species compilation of published data, which includes Bremer et al. (1996), Fegatella et al. (1998), Seybert et al. (2006) and Luef et al. (2015). We also find a relationship $_{
m for}$ the number of ribosomes using a best fit of the degradation rates with the power-law approximation of μ (see Supplementary Information), which accurately captures the crossspecies trends in ribosome volume in Figure 2c. We find that this fitted prediction, which relies on only a single free parameter, has a very similar goodness of fit to a simple power law, where we find that the best fit to Equation (16) is given by $R_0 = 1.54 \times 10^{-7}$ (m³Ribosomes (m³Cell)^{- β_R}) and $\beta_R = 0.73 \pm 0.15$. It should be noted that the best fit of our full model for ribosome composition makes predictions for ranges of cell size where we do not have data to test the model. Some of our later conclusions and inferences rely on these predictions, and thus provide a set of testable and open hypotheses, such as the number of ribosomes found at larger bacterial sizes or the exact size at which the 'ribosome catastrophe' occurs.

Shifts in cellular composition and the smallest bacteria Our analysis now provides the size dependence for the major cellular components across the domain of bacteria. This includes the prediction for the volume of the envelope considering three cases: only a single membrane (organisms without a cell wall) and the effective thickness for Gram-negative or -positive bacteria (Figure 2d), and predictions for *tRNA* and *mRNA* given the volume of ribosomes.

Of critical interest is how the relative cellular composition changes across species of diverse size and what tradeoffs and limitations we might be able to infer from these shifts. Figure 3a shows all of the cross-species trends for various cellular components compared with the overall cell volume. The compiled trends show that the composition of the cell greatly shifts across the range of bacterial body sizes. The smallest cell volumes are dominated by DNA and, to a lesser extent, cell membrane and protein, while the largest bacteria are composed of mostly ribosomes, tRNA and mRNA. These changes in composition are accompanied by shifts in the dry fraction of cells, as discussed later, and the overall

2152

cell density (see Supplementary Figure S5 and related discussion).

These trends also predict at what scale certain components would become limiting. For example, the relationship for genome volume gives the mass of a cell that is entirely filled by DNA as

$$V_{\rm DNA}^* = 1.14 \pm 0.30 \times 10^{-21} \rm m^3 \tag{23}$$

which is expectedly smaller than the smallest observed organism by a factor of approximately three (Table 1). This result shows that for the smallest cells there is not much room beyond the basic requirements for the genome. Similarly, the volume at which proteins are predicted to entirely fill the cell would be given by

$$V_{\rm p}^* = 3.38 \pm 4.72 \times 10^{-22} {\rm m}^3 \tag{24}$$

which is about an order of magnitude smaller than the smallest cell. The cellular envelope volume that completely fills the cell can be found using Equation (19) given values for the envelope thickness. For a Gram-negative bacterium, the envelope would fully occupy the cell at a size of 7.44×10^{-22} m³, for a Gram-positive, at a size of 2.91×10^{-23} m³, and for a single membrane (organisms lacking a cell wall), at a size of 1.01×10^{-25} m³. These results show that for the smallest cells the envelope represents a significant portion of the overall volume. In order to reduce the required envelope volume, the smallest cells are expected to minimize surface area by becoming increasingly spherical, which agrees well with the mostly spherical cell shapes reported by Luef et al. (2015), and are expected to reduce envelope thickness or layers, consistent with Mycoplasma that lack a cell wall.

The above analysis is useful for understanding which individual components become limiting at the smallest cell sizes. More generally, we can understand the overall space constraints of the cell at the small end of life by considering the total volume from Equation (2). This summed component volume will equal the total cellular volume for a size of 1.02×10^{-20} m³ for a Gram-positive bacterium, which is comparable to the previous energetic prediction (Kempes et al., 2012; Figure 3b) and larger than the smallest observed and hypothesized cells by roughly a factor of two (Knoll et al., 1999; Seybert et al., 2006; Luef et al., 2015). Considering only a single membrane would give a minimal size of 4.10×10^{-21} m³, which closely matches measurements for the smallest bacteria (Table 1). Considerations of the space for the smallest cells is a topic that has been deeply considered in the past and previous estimates agree with our prediction for a lower bound, where it was estimated that the minimum size required for a cell of modern biochemical complexity should fall in the range of 4.19×10^{-21} – 1.41×10^{-20} m³ (Knoll *et al.*, 1999). Furthermore, this result, and our space constraint estimate from $V_{\text{comp}} = V_{\text{tot}}$, also compare well with the limit anticipated from energetic considerations (Kempes et al., 2012), thus highlighting that multiple constraints are likely limiting the possibility of becoming smaller at the smallest scale of life.

Our result shows that the average cross-species trends converge to predict the smallest cell at the appropriate size range while still being an accurate predictor of cellular composition as cells move orders of magnitude away from this lower bound (Figures 2 and 3). This is remarkable because our analysis shows that average trends in cellular composition across the diversity of bacteria are consistent with the scale of the lower limits of life. However, these predictions also highlight necessary tradeoffs for the smallest cells, where the total requirements accurately predict volume observed minimum size only for a reduced cellular envelope of a single membrane. This may not be biologically feasible and thus other compositional

Table 1 Comparisons of estimates and measurements for the smallest bacterium

	Minimum size estimate	Type of estimate
Space limitations from this study		
For a Gram-positive bacterium	$1.02 \times 10^{-20} \text{ m}^3$	Cross-species prediction and theory
For bacterium with a minimal membrane	$4.10 \times 10^{-21} \text{ m}^3$	
Previous measurements and estimates		
Ground water microbes (Luef et al., 2015)	4.00×10^{-21} to 1.3×10^{-20} m ³	Measured
Mycoplasma pneumoniae (Seybert et al., 2006)	3.00×10^{-21} to 2.4×10^{-19} m ³	Measured
Energetic/growth limitations (Kempes et al., 2012)	$1.45 \times 10^{-20} \text{ m}^3$	Cross-species prediction and theory
NRC basic metabolism and components (Knoll <i>et al.</i> , 1999)	$4.19 \times 10^{-21} \text{ m}^3$	Calculation of average biochemical properties
Size at which individual components would fill the entire cell	volume	
Genome scaling	$1.14 \times 10^{-21} \text{ m}^3 (8.66 \times 10^{-22}, 1.46 \times 10^{-21})^a$	
Protein scaling	$3.38 \times 10^{-22} \text{ m}^3 (6.90 \times 10^{-23}, 6.75 \times 10^{-22})$	Cross-species prediction and theory
Single membrane scaling	$1.01 \times 10^{-25} \text{ m}^3$	
Ribosome scaling	$3.99 \times 10^{-26} \text{ m}^3 (2.03 \times 10^{-28}, 1.17 \times 10^{-23})$	

^aThe bracketed values, (,), denote the 95% confidence interval.

tradeoffs may be required for the smallest cells. For example, the genome size for Mycoplasma genitalium is 5.8×10^5 bp (Moya et al., 2009), which is about half the size of the prediction from the average trend for a bacterium of this volume. Similarly, at the smallest sizes, it is possible that cells could evolve to have lower copy number of proteins per gene, or have smaller average protein sizes—all to accommodate cell volume constraints.

Our own previous cross-species considerations (Kempes et al., 2012) predict a smallest bacterium that compares well with the mean cell size of Luef et al. (2015) and Seybert et al. (2006) but is about four to five times larger than the smallest cells from these studies (Table 1). This difference highlights that the smallest organisms may deviate from average energetic properties, such as unit maintenance costs, in addition to compositional shifts, though some of these deviations would require a cellular biochemistry that differs from modern examples. For example, previous estimates for possible primitive cells would lead to a size of 3.59×10^{-24} m³ (Knoll *et al.*, 1999), which is not much bigger than the size at which the cell volume is entirely filled by a single membrane $(1.01 \times 10^{-25} \text{ m}^3)$.

Although the data presented here are for the domain of bacteria, we may find similar constraints in archaea. For example, the smallest observed archaea are roughly comparable to the smallest bacteria in volume ($\approx 3.41 \times 10^{-20}$ m³; Huber *et al.*, 2002; Comolli *et al.*, 2009) and have similar genome sizes (0.5 megabases (Huber *et al.*, 2002)) and ribosome counts (≈ 92 ; Comolli *et al.*, 2009) at this size.

RNA limitations and the largest bacteria

For the largest bacteria, the dominant cell components are determined by the increasing need for processes related to transcription and translation to match increasing growth rates. As discussed earlier, this eventually leads to a 'ribosome catastrophe', where a finite size would require an infinite number of ribosomes, tRNA and mRNA. The limits anticipated in our model for the volume of ribosomes can be seen in the prediction curve of Figure 2c, where the entire cell volume would be filled by ribosomes, catastrophe', 'ribosome at a $1.39 \pm 0.03 \times 10^{-15}$ m³. This is in contrast to the smallest bacteria, where the need for ribosomes, tRNA and mRNA are predicted to diminish based on the decreasing growth rates. Previous predictions predict that growth rate goes to zero at a size of $1.45 \times 10^{-20} \text{ m}^3$ (Kempes *et al.*, 2012).

Expanding from the number of ribosomes to the total RNA content, the prediction from Equation (18) compares well with measurements for mRNA in *Escherichia coli* and *Mycoplasma pneumoniae* (see Supplementary Information for the value of the constants in Equations (17) and (18)). For *E. coli*, previous measurements give a range of 1380

(Neidhardt et al., 1996) to $\approx 10\,000$ mRNA per cell (see compilation in the Supplementary Information of Lu et al., 2007), compared with our estimate of 2540 to 12 257 using a cell volume between roughly 0.7 and 6 µm³ (Dethlefsen and Schmidt, 2007; Milo, 2013), and dividing Equation (18) by $\overline{\nu}_{tRNA}$. For M. pneumoniae, the mRNA quantities are two orders of magnitude lower than E. coli at around 10 per cell (Maier et al., 2011), which is similar to the lower bound of our estimate of 68-1159 given the considerable range in observed cell size of 0.005–0.24 µm³ (Seybert et al., 2006).

Similarly, the number of tRNA per cell in *E. coli* has been observed to range between 52 000 and 375 000 (Jakubowski and Goldman, 1984; Neidhardt *et al.*, 1996; Mackie, 2013), where we would predict a range of 23 561 to 113 707 based on the range of cell volumes used above. For *M. pneumoniae*, observations give 190 tRNA per cell (Maier *et al.*, 2011), which is similar to our prediction of 631 to 10 755. It should be noted that in both cases we overestimate the number of tRNA and mRNA in *Mycoplasma* which is likely due to the considerable space constraints faced by these minimal organisms, as discussed earlier.

Parallel to our analysis of the smallest cells, the total component volume also sets an upper bound on size, where $V_{\rm comp} = V_{\rm tot}$ at a size of $1.19 \times 10^{-15}~{\rm m}^3$. This limit is dominated by the 'ribosome catastrophe', where the cell will require more ribosomes than can fit in its volume in order for biosynthesis to keep up with increasing growth rate. It should be noted that this predicted upper bound is four orders of magnitude smaller than the largest observed bacterium of $\approx 4.19 \times 10^{-12}$ m³ (Schulz et al., 1999). However, this giant sulfur bacterium (Thiomargarita namibiensis) is dominated by vacuoles used for nutrient storage, which does not represent the metabolic active or relevant volume considered in our model of the required components (Schulz et al., 1999). If we remove the vacuole volume (approximately 98% of the total cell volume; Schulz et al., 1999) then we are left with a size of 1.05×10^{-14} m³, which is only an order of magnitude larger than our predicted upper bound. It is interesting to consider what tradeoffs these cells have made to avoid the 'ribosome catastrophe', and one possibility is that the environment regulates growth at a slower rate than cell size would dictate.

Similar to our biosynthetic limit, previous work has shown that there is a linear increase in the fraction of promoter activity devoted to ribosomes with increasing growth rate (Zaslaver *et al.*, 2009). If the observed trend is extrapolated, then at a growth rate of 2.9 divisions per hour all promoter activity would be devoted to ribosomes, which is not feasible, and this would correspond to a cell volume of 2.1×10^{-17} m³. This predicted upper bound is smaller than our space limitation and highlights the multiple constraints that could be faced for the largest bacteria. Both of these upper bounds on cell

volume correspond to the rough size at which there is an observed transition to eukaryotic life, along with corresponding and dramatic shifts in growth and metabolic rates (DeLong *et al.*, 2010; Kempes *et al.*, 2012). Thus, this major transition may in part be motivated by the challenges related to biosynthesis and ribosome requirements.

Free volume

In between this lower and upper limit, changes in composition can be most easily understood by considering the total physical space required for essential components compared with the remaining volume. Considering the average composition trends that we have discussed thus far, the fraction of the cell that is dry weight (Equation (3)) is 1 for the smallest bacteria and decreases quickly for larger cells until reaching a minimum around 0.13 at a size of 2.83×10^{-17} m³, before increasing again towards the largest cell (Figure 3c). From this relationship we can see that bacteria of intermediate size have the most free volume, and the maximum of this volume occurs around the size of many well-studied species such as $E.\ coli.$

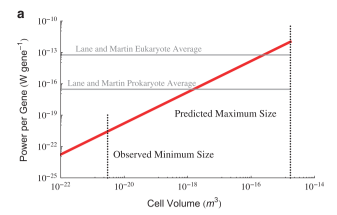
This free volume, which is an inferred and testable prediction from our model, has important evolutionary and ecological consequences because it implies that there is a range of cell sizes that may have the most flexibility due to the lack of physical space constraints. This flexibility could manifest as an increase in the copies of specific proteins in stressful conditions, greater ability to buffer waste products or plasticity in increasing the number of ribosomes for faster growth, among many other possible benefits. This is in sharp contrast to the smallest cells that barely have enough space for basic biochemical requirements. An important avenue of future research is to examine whether and how individual taxa allocate 'free volume' differently to various components or functions, and whether such allocation can be understood as an adaptation to specific niches or conditions.

However, it is also important to consider, as part of this 'free volume', the volume of water necessary for cellular reactions and internal transport. For example, if all of the free volume were dedicated to water, and given the size of E. coli, we would predict that 67-76% of the cell is filled with water, in good agreement with previous estimates of (Neidhardt et al., 1996). This comparison implies that all of the free volume could reasonably be occupied by water, and that evolutionary flexibility at these cell sizes could depend on the degree to which the water content is constrained. Our predictions suggest that the smallest and largest cells have lower water content, which is consistent with previous efforts that describe a cytoplasm with high molecular packing densities (e.g., von Hippel and Berg, 1989; Cayley and Record, 2003; Golding and Cox, 2006; Burnap, 2015). Similarly, these changes in water content may alter processes related to the solvent capacity or diffusive, sub-diffusive or active transport in the cell (e.g., von Hippel and Berg, 1989; Cayley and Record, 2003; Errington, 2003; Golding and Cox, 2006; Burnap, 2015), and it is of potential future interest to understand whether there are specific and calculable requirements for water content in cells of different size in connection with cellular composition. Furthermore, these results imply that the application of a constant dry weight ratio to diverse species may often be inaccurate.

Energetic limitations and complexities

Previous studies have highlighted that prokaryotes have a superlinear scaling between metabolic rate and body size. This observation has been successfully used to derive cross-species trends in growth rates and a limit where growth rate should go to zero in bacteria (Kempes et al., 2012). However, the superlinear scaling of metabolism is still not fundamentally understood and here we are surprised to find that every cellular feature scales sublinearly with overall cell volume. This implies that metabolism is not a simple proportionality of any one cellular feature. Figure 4b gives the scaling of the total metabolic rate as a function of the volume of each cellular component where it can be seen that all of the scalings are superlinear. Taken together these observations suggest that metabolic rate is a complicated emergent property of some measure of increasing cellular complexity. In particular, the scaling of metabolic rate with genome size stands out with an exponent close to 8, and it may be the case that metabolism is a complicated function of the growing complexity of the metabolic network which adds capacity quickly with the addition of novel proteins (DeLong et al., 2010). Metabolism also goes up steeply (exponents close to 2.4) with the total envelope volume and total number of proteins which also makes sense given that the membrane surface area controls ATP synthesis (Lane and Martin, 2010) and the number of proteins are the actual components of the metabolic network. The scaling of proteins scales closely to a power-law approximation of the growth rate equation (6) (an exponent of 0.65 compared with 0.70). Since growth rate is set by the overall scaling of metabolic rate (Kempes et al., 2012), it is possible that the protein concentration could be the dominant factor controlling both metabolic rate and growth rate, albeit not via any direct proportionality. Yet the explicit connection between any of these components to the scaling of metabolic rate remains unclear.

Furthermore, these scalings also highlight the unexpected result that cellular efficiency, defined as the metabolic power per unit of component (Lane and Martin, 2010), is increasing for every cellular component. It has previously been pointed out that the plasma membrane serves as the only region for ATP synthesis in bacteria, and since this surface area



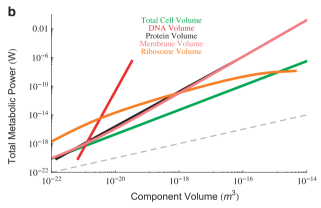


Figure 4 (a) The estimated scaling of metabolic rate per gene as a function of overall cell volume calculated from the scaling of genome size here and the data from DeLong et al. (2010). Previous average values from Lane and Martin (2010) for prokaryotes and eukaryotes are both shown. It can be seen that the previous prokaryote average value agrees with the scaling for the middle range of bacteria, and that bacterial values are close to the eukaryotic average value for the largest bacteria. Surprisingly, bacteria are increasing the metabolic rate per gene with a scaling exponent of 1.49. (b) Scaling of the total cellular metabolism as a function of the total volume of each cellular component. It can be seen that all of these relationships scale with an exponent greater than 1 (linear scaling is indicated by the gray dashed line), implying that metabolism is not a simple proportionality of any single cellular component. This suggests that the way in which cellular components are combining to produce superlinear scaling in cells is a complicated and emergent phenomena. Most notably, metabolic rate scales with total genome volume with an astonishing power of ≈ 8 .

scales sublinearly with volume it will be outpaced by anything that is proportional to volume (Lane and Martin, 2010). Previous analyses have thus suggested that bacteria are becoming less efficient on a perprotein or per-gene level (Lane and Martin, 2010). However, the surprising superlinear scaling of metabolic rate and the sublinear scaling of both genome size and protein content lead to an increasing efficiency for both components. Figure 4a gives the power per gene as a function of cell size showing that it is increasing superlinearly across bacteria. This scaling agrees with the average values from Lane and Martin (2010) for mid-range bacteria and as bacteria grow larger the values approach the reported eukaryotic average (Lane and Martin, 2010) at the upper range of bacteria. If bacteria were able to grow larger they would be expected to continue to increase the power per gene and overtake the eukaryotes. Thus, it would seem that bacteria are not limited by an energetic efficiency challenge but rather by an energetic surplus that demands ever faster rates of biosynthesis and eventually leads to a space limitation via the packing of ribosomes as discussed earlier. It may be that the evolutionary transition to eukaryotes was motivated by biosynthetic and space constraints resulting from a fast metabolism rather than to increased metabolic capacity. In fact, eukaryotes have a metabolism that scales with cell size following a smaller exponent than bacteria (DeLong et al., 2010). These results are counterintuitive and bring up deep questions about the explicit connection between membrane areas and metabolic rate: if mitochondria evolved to increase the ATP synthesis area and provide an advantage over bacteria (Lane and Martin, 2010), then how and why can metabolic rate scale superlinearly in bacteria when the surface area is growing sublinearly with body size?

Concluding remarks

In general, we find that cellular components follow strong trends, often power laws, with overall cell size. From the perspective of size limitations of cells, these cross-species trends, along with previous work (Knoll et al., 1999; Kempes et al., 2012), show that, on average, the smallest cells face challenges related to growth rate, energetic constraints and the physical space required to contain basic components. Our work here demonstrates how various cellular components are connected with one another and provides a foundation for understanding tradeoffs in physiology as related to cell size. Moving forward it is important to connect this work with detailed perspectives of how cell size and composition respond to environmental conditions and stress (e.g., Chien et al., 2012).

Furthermore, while we can predict many crossspecies trends, such as the number of ribosomes or the growth rate (Kempes et al., 2012), several basic scaling relationships remain unexplained in bacteria such as the scaling of genome size and the overall metabolic rate (West and Brown, 2005; DeLong et al., 2010). These relationships imply surprising consequences for the cell such as the increasing power per gene or protein as cells become larger based on the superlinear scaling of metabolism compared with the sublinear scaling of all other cellular components. To truly understand the full set of limitations that constrain the smallest cells and define species tradeoffs, we will need to continue to mechanistically underpin these observed cross-species trends. In doing so, we may be able to gain further insight into key evolutionary bifurcations, adaptation to various niches and better understand the requirements for originating and sustaining simple life.

Conflict of Interest

The authors declare no conflict of interest.

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