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Published in: Current Opinion in Microbiology

DOI: 10.1016/j.mib.2020.07.010

Publication date: 2020

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Document Version Peer reviewed version

Link to publication in Discovery Research Portal

Citation for published version (APA): Bergkessel, M. (2020). Regulation of Protein Biosynthetic Activity During Growth Arrest. *Current Opinion in Microbiology*, *57*, 62-69. https://doi.org/10.1016/j.mib.2020.07.010

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- 1 **Title:** Regulation of Protein Biosynthetic Activity During Growth Arrest
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7 Abstract: Heterotrophic bacteria grow and divide rapidly when resources are abundant. Yet

8 resources are finite, and environments fluctuate, so bacteria need strategies to survive when

- 9 nutrients become scarce. In fact, many bacteria spend most of their time in such conditions
- 10 of nutrient limitation, and hence they need to optimise gene regulation and protein
- biosynthesis during growth arrest. An optimal strategy in these conditions must mitigate the 11
- 12 challenges and risks of making new proteins, while the cell is severely limited for energy and
- 13 substrates. Recently, ribosome abundance and activity were measured in these conditions,
- 14 revealing very low amounts of new protein synthesis, which is nevertheless vital for survival.
- 15 The underlying mechanisms are only now starting to be explored. Improving our
- 16 understanding of the regulation of protein production during bacterial growth arrest could
- 17 have important implications for a wide range of challenges, including the identification of new
- 18 targets for antibiotic development.
- 19

20 What are the causes of growth arrest?

21 Many heterotrophic bacteria can grow and divide rapidly if their nutritional needs are met,

22 but this condition is the exception, not the rule. Even in nutrient-rich environments, bacterial

- 23 growth itself quickly depletes local resources and causes accumulation of waste products
- 24 that inhibit further rapid growth. For example, many species form dense biofilm communities,
- which are held together by self-produced matrices, and which quickly become self-limiting. 25
- 26 Experimental measurements and modelling in Pseudomonas aeruginosa biofilms suggest
- 27 that at thicknesses greater than about 40-70 microns, metabolism of the biofilm cells

28 completely depletes oxygen at the base of the biofilm, leading to near-zero growth rates[1,2].

- 29 Depending on the organism and environment, other macronutrients (e.g. carbon[3],
- 30 nitrogen[4], or phosphorus[5]), or micronutrients (such as iron[6]) may become limiting first,
- 31 but in all cases the lack of an essential substrate for new biosynthesis causes growth to
- 32 stop. Outside of biofilms, heterotrophic bacteria often exist in low-nutrient environments that
- 33 cause frequent growth arrest[7]. Other environmental stresses, such as reactive oxygen
- 34 species, high osmolarity, and unfavourable temperature, can suppress growth by directly
- 35 inhibiting ATP and/or protein synthesis[8-10]. Finally, bacteria (as well as fungi, plants, and
- 36 animals) have evolved a wide array of weapons that specifically target the energy
- 37 conservation or biosynthetic machinery of competing bacteria in order to inhibit their growth
- 38 (for example, see [11]). All of these challenges would hinder growth whether there were an
- 39 adaptive response from the challenged bacteria or not, but decades of work strongly suggest
- 40 that complex regulation is in place to coordinate stopping replication, repressing expression 41 of new biosynthetic machinery, and redirecting resources toward functions needed for
- 42
- survival (reviewed in [12,13]). Still, many questions remain about the mechanisms that allow
- 43 ongoing adjustments to protein biosynthetic rates and priorities during protracted growth

44 arrest – adjustments that, though small in magnitude, may make the difference between

- 45 survival and death.
- 46

47 Risks and benefits of protein biosynthesis during growth arrest

48 Regulation operating during growth arrest must weigh the risks of consuming extremely 49 limited resources against the benefits of making potentially useful new proteins. Many 50 studies of multiple organisms in different growth-arrest contexts have found that growth-51 arrested bacteria continue to make new proteins at low rates[14-20]. However, protein 52 biosynthesis requires networks of interconnected metabolic activities to supply the energy 53 and substrates required, and, especially if flux through these networks is inconsistent, activity can increase the vulnerability of the bacteria to damage (Fig. 1A). For example, the 54 55 redox reactions of the electron transport chain can release reactive oxygen species, 56 resulting in damage that is difficult to repair with limited resources[3,21]. Stalled, transcribing 57 RNA polymerases can increase the likelihood of some types of DNA damage by leaving 58 DNA-RNA hybrids that can be recognised improperly by DNA synthesis machinery[22-24], 59 and ongoing translation under starvation conditions can lead to protein aggregation[25], 60 requiring the activity of heat-shock proteases to maintain viability[26]. The presence of 61 antibiotic compounds exacerbates the risks associated with activity, as these compounds 62 largely subvert biosynthetic processes. Mutants that tolerate antibiotics better often show slower growth or increased lag times [27]. Indeed, the bacterial form best suited to long-term 63 64 survival of harsh, resource-depleted conditions is the spore, with metabolic activity very 65 close to zero (reviewed in [28]). 66 Despite this, even in organisms that can form spores as a response to extreme resource 67 limitation (such as Bacillus subtilis), a substantial subpopulation will instead maintain a

- 68 vegetative state that is metabolically and biosynthetically active at low rates[29].
- 69 Furthermore, researchers have observed that despite the risks of protein synthesis during
- growth arrest, inhibiting it decreases survival[14,30,31]. What benefits outweigh the risks of
 biosynthesis during resource limitation? Essential proteins sustain damage over time and
- 72 must be replaced[32]. Ongoing activity also allows defensive responses against specific
- 73 environmental threats, e.g. by upregulation of efflux pumps or DNA repair enzymes[31,33].
- And even in conditions of extreme resource limitation, cells encounter small amounts of
- vseful nutrients (from lysis of other bacteria, for example); taking advantage of them requires
- vptake and incorporation via biosynthesis[17,34]. Finally, when favourable conditions return,
- bacteria must be prepared to rapidly resume growth or risk being overrun by competitors.
- 78 Because growth-arrested bacteria are often exposed to fluctuating environments with
- opportunities and threats varying unpredictably over time, an ability to alter protein synthesis
- 80 activity in response to the environment while still remaining in a growth-arrested state would
- 81 likely be advantageous (Fig. 1B).
- 82

83 Ribosome dynamics during growth and growth arrest

- 84 Ribosomes are the engines of protein synthesis, so their abundance and activity are
- 85 important points of control of biosynthetic capacity, and accordingly, their regulation changes
- 86 dramatically as nutrient availability drops (Fig. 2). Elegant work over several decades has
- 87 shown how *Escherichia coli* tunes resource allocation to maximise ribosome biogenesis
- 88 when nutrients are plentiful, and diverts resources toward acquiring nutrients when those

- 89 nutrients become limiting ([35-37] and references therein). Earlier work compared ribosome
- 90 production and protein synthesis at fast (doubling time (τ) = 20 minutes) and slow
- 91 ($\tau = 100$ minutes) growth rates, but recently these quantitative analyses have been extended
- 92 to conditions that impose extremely slow ($\tau = 1440$ minutes, in chemostats) or no growth
- 93 (e.g. stationary phase or carbon starvation)[18,19,38].

94 Several interesting observations emerge from these studies. First, quantitative comparisons 95 across this expanded range of growth conditions highlight the striking magnitude of the 96 changes as cells enter growth arrest. The rate of new protein synthesis in growth arrest is 97 estimated to be approximately 1500-fold lower than during optimal growth (Fig. 2A), and the 98 average number of active ribosomes per cell approximately 100, compared to 62,000 at the 99 fastest growth rates (Fig. 2B). Changes of this magnitude have enormous impacts on the 100 biochemical and biophysical context in which the gene expression machinery operates, but 101 the mechanisms by which the machinery adjusts to these changes are not well understood.

102 Second, careful investigation of the translational responses to extreme limitation with 103 different limiting nutrients has revealed that ribosome dynamics are distinct, suggesting that 104 finely tuned regulation is important even though new protein synthesis is so strikingly 105 reduced. In carbon starvation, a large fraction (80%) of the ribosomes are sequestered in an 106 inactive state, while nitrogen limitation more dramatically represses the translation 107 elongation rate, and phosphorus limitation strongly reduces cells' ribosome content, at least 108 in part by degradation of ribosomes[18,38,39]. Growth-limiting osmotic and oxidative stresses also strongly repress translation elongation rates, requiring adjustments to 109 110 ribosome activities or amounts[8,9]. Maximising survival of growth arrest in a dynamic 111 environment likely requires complex regulation to identify and recycle damaged ribosomes, 112 safely sequester excess functional ribosomes, and occasionally produce new ribosomes as 113 conditions require. Consistent with this notion, multiple studies have shown that genes

- 114 involved in ribosome biogenesis are expressed during carbon- or energy-limited growth
- arrest, and that they are important for survival[14,15,40].

116 Ribosome dynamics have mostly been observed at the population level, so the distribution of active and inactive ribosomes across individual cells is not well studied. Multiple 117 118 mechanisms exist for sequestering ribosomes upon growth arrest, which could be used by 119 each cell to inactivate a consistent fraction of its ribosomes (reviewed in [41]). However, it is 120 also possible that translational activity is unequally distributed across the bacterial 121 population. Toxin-antitoxin systems have in fact been proposed to drive heterogeneity, when 122 stochastically arising imbalances allow the toxin to escape antitoxin control and target 123 biosynthetic machinery, including ribosomes, tRNAs, EF-Tu, and mRNAs (reviewed in [42]). 124 Two studies have investigated protein synthesis rates (in E. coli) and ribosome levels (in P. 125 aeruginosa) in single cells during starvation-induced growth arrest, to directly assess their 126 distributions[16,43]. In both cases, after initial large drops in ribosome abundance[43] or 127 protein synthesis rates[16] at the entry to growth arrest, the new levels were maintained by 128 most cells for several days. However, both studies also found outliers of high and low

- ribosome abundance and activity, suggesting that heterogeneity does arise and may play an
- important role. Within dense communities of sibling bacteria, selective pressure could
- 131 conceivably favour the uneven distribution of protein biosynthetic activity across individual132 cells to maximise survival of the population. Examples of metabolic coordination in biofilms
- 133 mediated by gated ion channels[4] and redox-active secondary metabolites[2] have been

- 134 described, but further explorations of mechanisms contributing to such coordination and their
- 135 impacts on protein biosynthesis are still needed.
- 136

137 Mechanisms for regulating protein biosynthesis during growth arrest

138 Mechanisms to dynamically adjust protein biosynthesis during growth arrest have been 139 studied relatively little, in part because of the difficulty of measuring low activity levels. Most 140 research has focused on the transition from rapid growth to growth arrest as nutrients are 141 depleted. The dominant pattern in this context is strong suppression of ribosome biogenesis and upregulation of genes involved in nutrient acquisition and general stress responses. Key 142 143 regulators of this transition (which have been recently comprehensively reviewed) include general stress sigma factors, which bias RNA polymerase (RNAP) activity toward stress-144 adaptive functions[44], and the modified nucleotides cAMP and (p)ppGpp, which bias RNAP 145 activity away from ribosomal RNA and protein genes[45,46]. The regulators playing 146 147 important roles after cells have entered into a protracted growth arrest are much less clear, 148 but (p)ppGpp likely plays an ongoing role even though its levels peak at the entry to growth 149 arrest[47]. Homologs of ReIA, which is activated by binding uncharged tRNAs and the 150 ribosome during translation[48], are major sources of (p)ppGpp in many organisms. Thus, 151 low levels of ongoing translation during growth arrest may contribute to ongoing adjustment of (p)ppGpp levels. Indeed, a Tn-Seq screen in Rhodopseudomonas palustris found that 152 both the ability to synthesise (p)ppGpp and genes for ribosome biosynthesis were important 153 154 for fitness throughout 20 days of carbon-limited growth arrest[14].

155 Even though (p)ppGpp has been studied extensively, the fact that its functions are carried 156 out differently in different organisms[49], and that it impacts many core cellular processes 157 simultaneously (reviewed in [50], Fig. 3), has prevented a full understanding of its direct 158 effects. Recent work has shown that these effects are even more widespread than 159 previously appreciated: a careful transcriptomic study showed that (p)ppGpp binding to RNA 160 polymerase in E. coli directly impacts expression levels for 757 genes[51], and a crosslinking approach identified 56 direct binding targets of (p)ppGpp, many of which are involved in 161 162 translation and nucleotide biosynthesis and were previously unknown[52]. The new insights 163 into the direct effects of (p)ppGpp on regulation of nucleotide biosynthesis are particularly 164 interesting and highlight the centrality of the nucleotide pools in cellular metabolic networks (Fig. 3). Levels are tightly controlled, but a notable shift in purine nucleotides toward 165 166 (p)ppGpp occurs at the entry to growth arrest[47,53], and complex crosstalk and feedback 167 extends these effects to numerous biosynthetic pathways. How cells might adjust gene expression activities in response to the nucleotide pools present in growth arrest, and how 168 they might restore the pools to resume active growth are not known. Within growth-arrested 169 170 states, perturbations to nucleotide pools appear to impact survival. For example, among 171 antibiotic-exposed persisters in *P. aeruginosa*, mutations inhibiting *de novo* pyrimidine biosynthesis decreased survival, but depleting ATP in those mutants restored survival[54]. 172 New tools to sensitively measure levels of specific nucleotides in single cells and in real time 173 174 will be extremely useful in dissecting these complex networks.

While mechanisms to downregulate protein biosynthesis during growth arrest are still not fully understood, mechanisms for transiently upregulating biosynthesis in this state are even less explored. In this context, "upregulation" is relative – mechanisms operating during growth arrest (potentially acting with only 100 active ribosomes available per cell) cannot drive high levels of expression of expetition, in checkute torme, but instead hole cells.

drive high levels of expression of anything, in absolute terms, but instead help cells

180 overcome challenges to gene expression imposed by nutrient limitation or by the activities of

- regulators like (p)ppGpp. One such challenge is that the substrates needed may not all be
- present in the environment at once. To take advantage of transient availability, many
- organisms have storage mechanisms for phosphorus (polyphosphate granules) and carbon
 (polyhydroxyalkanoate granules and glycogen). Production of these storage compounds is
- 185 induced at the onset of growth arrest[55,56], and genes for phosphorus and carbon storage
- 186 are under positive selection in an extremely low-nutrient environment[7]. Nitrogen is
- 187 abundant in proteins, and proteolysis and reuse of amino acids is important during growth
- 188 arrest, including both recycling material from dying cells in a starving population[17,20] and
- 189 degrading aggregated proteins within the cell[26].
- 190 Assuming that the substrates for biosynthesis can be scavenged from the environment or
- intracellular storage, the gene expression machinery must adapt to the repressive regulatory
- 192 context that occurs with growth arrest. A recent study of the stationary-phase nucleoid-
- associated protein Dps, which appears to bind the chromosome in a crystalline-like
- lattice[57], revealed that this growth-arrest adaptation may be less of a barrier to gene
 expression than previously assumed it has essentially no impact on RNAP's ability to
- 196 access and transcribe DNA[58]. Other work showed that a growth-arrest-specific
- 197 transcriptional regulator in *P. aeruginosa* (SutA) can directly enhance transcription initiation
- 198 of the rRNA genes; the effects were greatest in conjunction with the general stress sigma
- 199 factor[59]. This observation suggests that the stress-specific holoenzyme might be used for
- some transcription of housekeeping genes during growth arrest, which could be important if
- the housekeeping holoenzyme is largely sequestered by binding to the 6S small RNA[60].
- 202

203 Implications for antibiotic tolerance and persistence

204 One motivation for studying growth-arrested states has been the long-recognised correlation 205 between susceptibility to most commonly-used antibiotics and growth[3,61,62]. It is now 206 appreciated that non-growing cells that persist during antibiotic treatment maintain some 207 metabolic activity but strongly repress protein biosynthesis[63], and that multiple 208 mechanisms affecting biosynthetic activity give rise to complex population dynamics of 209 antibiotic killing[64]. A non-growing persister state also appears to protect Salmonella from 210 the host immune system, and requires some ongoing protein synthesis[65]. Recent efforts to standardise methods for investigating and describing non-growing states and their 211 212 susceptibilities[66] are helpful, and highlight the importance of developing a deeper 213 understanding of biosynthesis regulation under growth-arrested conditions. Some 214 distribution of biosynthetic activity levels will always exist for any population of bacteria, and 215 below a threshold, they are more likely to tolerate antibiotic exposure. The mean and shape 216 of the distribution, as well as the timescales over which an individual bacterium might 217 sample different regions of the distribution, are characteristics that likely have direct impacts on antibiotic susceptibility patterns. These characteristics are defined by the regulatory 218 219 networks that govern biosynthetic activity during growth arrest.

- A billion years of evolution have shaped these networks to balance the risks of ongoing protein synthesis against the benefits of renewing and adapting the proteome under severe resource limitation. Much still remains to be learned about the mechanisms responsible for this regulation in diverse organisms. By focusing research efforts on mechanisms that favour biosynthesis in addition to those that repress it during growth arrest, and seeking to
- 225 understand how these mechanisms are coordinated over time and across bacterial

- populations, we stand to improve our basic knowledge of bacterial life in natural
- environments, and also gain useful insights toward addressing the pressing challenges ofrising antibiotic resistance.
- 229

230 Acknowledgments

- 231 I would like to thank Dianne Newman, Phil Esra, and colleagues in the University of Dundee
- 232 Molecular Microbiology Division for helpful feedback on this manuscript. Financial support
- has been provided by the University of Dundee Institutional Strategic Support Funds.
- 234

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antibiotic exposure but are associated with disadvantages in utilising transiently available
nutrients, repairing damage, and resuming growth. In this example, a bacterium with a
moderate level of activity represents a "sweet spot" that achieves the highest fitness across
the series of events occurring, but in theory fitness could be further optimised by changing
activity levels in response to changes in conditions throughout growth arrest.

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510 Figure 2. Ribosome dynamics during growth and growth arrest. A) Box areas represent 511 total protein synthesis outputs per cell per unit time in E. coli at the maximum growth rate 512 (doubling time $(\tau)=20$ minutes), a slower growth rate ($\tau=100$ minutes) and stationary phase induced by carbon starvation. Rates for fast and slow growth were estimated by multiplying 513 514 the number of actively translating ribosomes per cell by the average elongation speed, using 515 numbers summarised in [67], and also generally agree with measurements made in [19]. 516 The rate estimate for stationary phase is based on measurements made in [19] and [17]. B) Proteome composition for *E. coli* cells at τ =20 minutes. τ =100 minutes. and stationary 517 518 phase. The area of the large box for each growth rate represents all the proteins (by mass) 519 in a cell. Each blue square represents 100 actively translating ribosomes; orange square=100 inactive ribosomes; red circle= 100 RNAPs actively transcribing rRNA; black 520 521 circle= 100 RNAPs actively transcribing something other than rRNA; grey circle= 100 522 inactive RNAPs. In this context, "inactive" includes ribosomes that are not yet mature, or are 523 in the process of terminating translation and being recycled to initiate a new protein, or are 524 sequestered, and RNAPs that are non-specifically associated with DNA, or bound at 525 promoters but not yet transcribing, or are sequestered. The area of each symbol is to scale. Numbers for fast and slow growth are from [67]. The ribosome fractions for stationary phase 526 527 are from [19]. The RNAP number in stationary phase is a rough estimate, by extrapolation 528 from measurements in [68] to stationary phase. The number of active RNAP in stationary 529 phase is not known but is likely small. The reallocation of proteome resources during the 530 transition from fast to slow growth is driven in large part by cAMP [35], while (p)ppGpp is a 531 major regulator of the transition to growth arrest [18,46]. 532

533 Figure 3: Nucleotide pools are tightly regulated, impact many biosynthetic processes, 534 and change in growth arrest. Sizes of boxes represent estimated sizes of nucleotide 535 pools, based on [53,69]. Direct synthesis reactions are represented by black arrows, 536 stimulatory effects are represented by green arrows, and inhibitory effects are represented by red symbols (nucleotides play roles both as regulators and as direct substrates for DNA 537 and RNA synthesis). The amounts of cAMP and the nucleotide sugars were not directly 538 539 measured at the entry to growth arrest. The synthesis of high amounts of (p)ppGpp can 540 divert a measurable fraction of purine nucleotides away from other processes, and then 541 actively represses new purine synthesis. Pyrimidine biosynthetic enzymes bind (p)ppGpp 542 and may be directly repressed but are also indirectly downregulated by a decrease in purine 543 levels, as purines are positive allosteric effectors. At the same time, consumption of 544 nucleotides by growth-related processes is also reduced, such that pyrimidine pools actually 545 slightly increase. How the cell adapts to the growth-arrest status of nucleotide pools and 546 fluxes, and how it escapes this state to return to active growth, are largely unknown.

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