



Heat-shock proteases promote survival of *Pseudomonas aeruginosa* during growth arrest

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Edited by Caroline S. Harwood, University of Washington, Seattle, WA, and approved January 8, 2020 (received for review July 14, 2019)

When nutrients in their environment are exhausted, bacterial cells become arrested for growth. During these periods, a primary challenge is maintaining cellular integrity with a reduced capacity for renewal or repair. Here, we show that the heat-shock protease FtsH is generally required for growth arrest survival of *Pseudomonas aeruginosa*, and that this requirement is independent of a role in regulating lipopolysaccharide synthesis, as has been suggested for *Escherichia coli*. We find that *ftsH* interacts with diverse genes during growth and overlaps functionally with the other heat-shock protease-encoding genes *hslVU*, *lon*, and *clpXP* to promote survival during growth arrest. Systematic deletion of the heat-shock protease-encoding genes reveals that the proteases function hierarchically during growth arrest, with FtsH and ClpXP having primary, nonredundant roles, and HslVU and Lon deploying a secondary response to aging stress. This hierarchy is partially conserved during growth at high temperature and alkaline pH, suggesting that heat, pH, and growth arrest effectively impose a similar type of proteostatic stress at the cellular level. In support of this inference, heat and growth arrest act synergistically to kill cells, and protein aggregation appears to occur more rapidly in protease mutants during growth arrest and correlates with the onset of cell death. Our findings suggest that protein aggregation is a major driver of aging and cell death during growth arrest, and that coordinated activity of the heat-shock response is required to ensure ongoing protein quality control in the absence of growth.

Pseudomonas aeruginosa | growth arrest | survival | proteostasis | FtsH

Most of our knowledge of bacterial physiology is derived from studying exponentially growing cells in nutrient-replete environments. While this study has provided a rich understanding of the complex and diverse processes at play during cellular growth and division, in many ways its scope is limited to what is likely just a fleeting period in the life of a bacterium. Awareness is mounting that most bacteria spend a majority of their lives not growing and dividing, as they quickly exhaust the nutrients in their environment and enter into a state of growth arrest (1–3). Yet comparatively few studies have investigated the molecular processes required for survival in this state, even though they are as important for evolutionary success as the ability to grow rapidly when nutrients once again become available (4). Moreover, successful strategies to survive extended periods of growth arrest can be important drivers in the emergence of antibiotic tolerance and resistance (5, 6), as well as in the establishment and persistence of biofilm communities (7–10). Thus, a better understanding of the underlying genetics and molecular mechanisms that promote survival during growth arrest is crucial for understanding how bacteria thrive in diverse ecological and clinical contexts.

Studies in *Escherichia coli* and other model species have provided valuable insight into the diverse strategies bacteria have evolved to survive growth arrest (3, 11, 12). These strategies are often dependent on the metabolic and biochemical capabilities of individual species, as well as on the nutrients available in the environment, highlighting the need to study growth arrest in evolutionarily and metabolically diverse bacteria under distinct

growth-arrest regimes. For example, bacteria use varied alternative substrates as sources of electron donors or acceptors to generate ATP during growth arrest, and reroute their metabolic pathways in diverse ways to improve the efficiency of ATP generation (3). This energy is needed to protect the integrity of cellular components, such as nucleic acids and proteins, that cannot be easily replaced by nutrient-limited cells and are essential for survival (2, 3). However, how cells efficiently coordinate maintenance needs in the face of slim energy resources is poorly understood.

To understand the mechanisms underpinning the growth-arrested state, we have used the opportunistic pathogen *Pseudomonas aeruginosa* as a model organism (13–16). *P. aeruginosa* is well-adapted for survival in a variety of nutrient-limited or otherwise hostile environments, ranging from open ocean and freshwater sources, to the interior of surface-attached biofilms, to chronic wounds and the lungs of patients with cystic fibrosis (17, 18). Despite superficial differences, these environments often share a common physiological constraint: They can be limited for electron donors or acceptors, compelling organisms that survive within them to hone sophisticated strategies to cope with periods of energy-limited growth arrest (19–25).

Previously, we identified the ATP-dependent membrane protease FtsH as one of only a few genes that confer a general fitness advantage to *P. aeruginosa* during energy-limited growth arrest, regardless of the limitation that prompted entry into this state (15). FtsH has been well-studied during growth, and is involved in quality control of membrane proteins (26), regulation of the heat-shock response (27), and fine-tuning of lipopolysaccharide (LPS) levels by regulated degradation of the LPS

Significance

This study reveals that protein degradation plays a major role in the survival of the opportunistic bacterial pathogen *Pseudomonas aeruginosa*. Loss of multiple proteases, better known for their roles in proteostasis in response to stresses such as heat shock, accelerates cell death during growth arrest. This finding, coupled to the fact that the accumulation of misfolded and aggregated proteins in aging in eukaryotic cells is well appreciated to contribute to cellular damage and senescence, suggests that a general role for proteases in preserving bacterial proteostasis during aging has been overlooked. Our findings have implications for the study and treatment of infectious disease and highlight potentially conserved functions for proteases in combatting aging from bacteria to humans.

Author contributions: D.W.B., D.A.-A., M.A.S., J.A.C., and D.K.N. designed research; D.W.B., D.A.-A., M.A.S., and J.A.C. performed research; D.W.B., D.A.-A., M.A.S., J.A.C., and D.K.N. analyzed data; and D.W.B. and D.K.N. wrote the paper.

The authors declare no competing interest.

This article is a PNAS Direct Submission.

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This article contains supporting information online at <https://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1912082117/-DCSupplemental>.

First published February 6, 2020.

biosynthetic enzymes LpxC and KdtA (28–34). This last activity is essential in *E. coli*, as loss of *ftsH* results in unbalanced lipid synthesis and lethal overproduction of LPS (28). Although many of the growth-related roles of FtsH are well characterized, its role during growth arrest in any organism has received less attention (35–37).

Recently, two studies (38, 39) attributed a role for FtsH in growth arrest survival through a novel cell death pathway in *E. coli*, in which dysregulation of an outer membrane (OM) signaling system putatively inhibits LpxC degradation by FtsH, resulting in overproduction of LPS and cell death upon nutrient depletion and growth arrest. Based on these results, we reasoned that one possibility FtsH is so important for survival of *P. aeruginosa* might be that FtsH plays a regulatory role during growth arrest by degrading LpxC, similar to its role in *E. coli*. However, given that LpxC levels are not regulated by FtsH in *P. aeruginosa* (37, 40), the plausibility of such a function was unclear.

Alternatively, several studies have pointed to a role for the heat-shock response during growth arrest in *E. coli*, suggesting that aggregation of misfolded or damaged proteins may become a critical problem for nongrowing cells (41–46). As a highly conserved component of the heat-shock response in many bacteria (32), we reasoned that FtsH might degrade these misfolded or damaged proteins during growth arrest in *P. aeruginosa*. Indeed, protein aggregation is a hallmark of aging across a wide range of cellular systems, from humans to bacteria (47–53), although whether protein aggregation represents a strictly pathological process has recently come into question (54–60). In this study, we sought to distinguish between these possibilities for the role of FtsH during growth arrest in *P. aeruginosa* using a genetic and cell biological approach.

Results

FtsH Maintains Cell Integrity during Growth Arrest. We use the general term “growth arrest” when referring to two states: When cells enter stationary phase after growth in rich medium (lysogeny broth, LB), and when cells growing exponentially in LB are shifted to a buffered minimal medium devoid of organic carbon (carbon starvation medium, CSM). We refer to the former growth-arrested state as “stationary phase” and the latter as “carbon starvation.”

We constructed an isogenic markerless deletion of the *ftsH* ORF in *P. aeruginosa* strain UCBPP-PA14 and observed the morphology of the mutant strain during growth and growth arrest. Loss of *ftsH* did not affect cell length compared to WT during exponential growth in LB at 37 °C (SI Appendix, Fig. S1A, 3 h). Cell length decreased gradually as cells entered stationary phase after ~8 h of growth, reaching a minimum of 2.8 μm for WT and 3.2 μm for Δ *ftsH*, respectively (SI Appendix, Fig. S1A, 8 h). WT cell length did not change over the next 20 h in stationary phase while Δ *ftsH* slowly increased in length to an average of 3.8 μm (SI Appendix, Fig. S1A, 28 h).

Death occurred after extended incubation in stationary phase for both WT and Δ *ftsH*, although Δ *ftsH* began to die earlier and to a greater extent than WT, becoming effectively unculturable after 60 h of total incubation time (Fig. 1A). Death of Δ *ftsH* also occurred after >24 h of growth arrest in CSM, with ~5- to 10-fold reduced viability after 96 h (Fig. 1B) and ~100-fold reduced viability after 10 d (SI Appendix, Fig. S2A). In contrast, WT maintained full viability after 96 h of carbon starvation (Fig. 1B) and had ~twofold reduced viability after 10 d (SI Appendix, Fig. S2A). Loss of *ftsH* also caused death when cells were starved for oxygen as a terminal electron acceptor or ammonium as a nitrogen source (SI Appendix, Fig. S2B and C). Integration of the *ftsH* ORF with its native promoter at the *glmS* locus on the chromosome fully rescued the viability defects in growth arrest, confirming the role of FtsH in survival (the complemented strain Δ *ftsH*/+*ftsH*) (Fig. 1A and B).

Most Δ *ftsH* cells lysed following death, as exhibited by a decrease in optical density and the appearance of only scattered “ghost” cells by microscopy (Fig. 1C and SI Appendix, Fig. S3). Before significant death and lysis of Δ *ftsH* occurred, however, we observed many intact cells with what appeared to be a “detached” inner membrane (IM) (Fig. 1C). This cell morphology was largely absent in the WT and the complemented strain at the same time points (Fig. 1C and D), and none of the strains displayed this morphology during growth. We stained carbon-starved Δ *ftsH* cells expressing cytoplasmic GFP with the OM dye FM 4-64 to confirm separation between the OM and IM (Fig. 1E). The morphology of growth-arrested Δ *ftsH* cells was strikingly similar to an LPS-overproducing strain of *E. coli* that also experienced rapid cell death during growth arrest (38). However, we ruled out the possibility of LPS overproduction as the cause of death of Δ *ftsH* in our experiments, as LPS levels were similar between WT and Δ *ftsH* during growth and growth arrest and cation supplementation did not prevent death (SI Appendix, Supplementary Text and Fig. S3). We conclude that FtsH is required to maintain cellular integrity during growth arrest, and that impaired cellular integrity in the absence of FtsH is not due to overproduction of LPS.

Identification of Genetic Interactions with *ftsH*. We performed transposon insertion sequencing (Tn-seq) in the Δ *ftsH* background to identify genes that interact with *ftsH* on a genome-wide scale (61). In Tn-seq, a pooled transposon mutant library is grown under a condition of interest and then sequenced at the transposon–genome junction to identify where transposons are inserted and to quantify their abundance in the total population. Mutant abundance is then compared to a control population to determine the relative fitness of insertion at a particular locus under the condition of interest.

Initially, we attempted Tn-seq under growth-arrest conditions in the Δ *ftsH* background using an experimental design similar to our previous study in the WT background (15). However, the reduced viability of the Δ *ftsH* background during growth arrest resulted in contaminating growth of the trace amounts of *E. coli* donor carried over from the original conjugation (Materials and Methods). To avoid this problem, we instead compared insertions between the WT and Δ *ftsH* transposon libraries after growth in liquid LB at 37 °C (Materials and Methods and Fig. 2A). We identified 36 genes that had a growth-aggravating interaction with *ftsH*: That is, mutations in these genes negatively affected fitness specifically in the Δ *ftsH* background, which was defined as >10-fold reduced insertions in the Δ *ftsH* background relative to the WT background ($P < 0.05$) (SI Appendix, Table S4 and Dataset S1). Aggravating interactions were detected among genes involved in cell envelope biogenesis and OM functions, as well as at the interface of carbohydrate and glycerophospholipid metabolism (SI Appendix, Fig. S4 and Table S4). Mutations in genes putatively involved in LPS biosynthesis (*ssg*, *wapH*), glycolysis (*tpiA*), fatty acid biosynthesis (*fabF1*), and phospholipid metabolism (*lptA*) were growth-aggravating in the Δ *ftsH* background, suggesting that although FtsH does not regulate LpxC in *P. aeruginosa*, it could still play an integral role in glycerophospholipid metabolism and cell envelope biogenesis.

Insertions in genes involved in posttranslational modification and protein turnover were also aggravating for growth, including the heat-shock proteases encoded by *hslVU* and *htpX* (62, 63) and the SsrA-binding protein encoded by *smgB* (64). We detected an aggravating interaction with the membrane-stress responsive two-component system encoded by *amgRS*, as observed previously in the *P. aeruginosa* strain PAO1 (65). AmgRS is a homolog of the CpxRA two-component system that regulates the envelope stress response in *E. coli* (66, 67), and loss of *amgRS* or *ftsH* in *P. aeruginosa* results in hypersensitivity to aminoglycoside-induced protein misfolding (65). Notably, *htpX* is regulated by

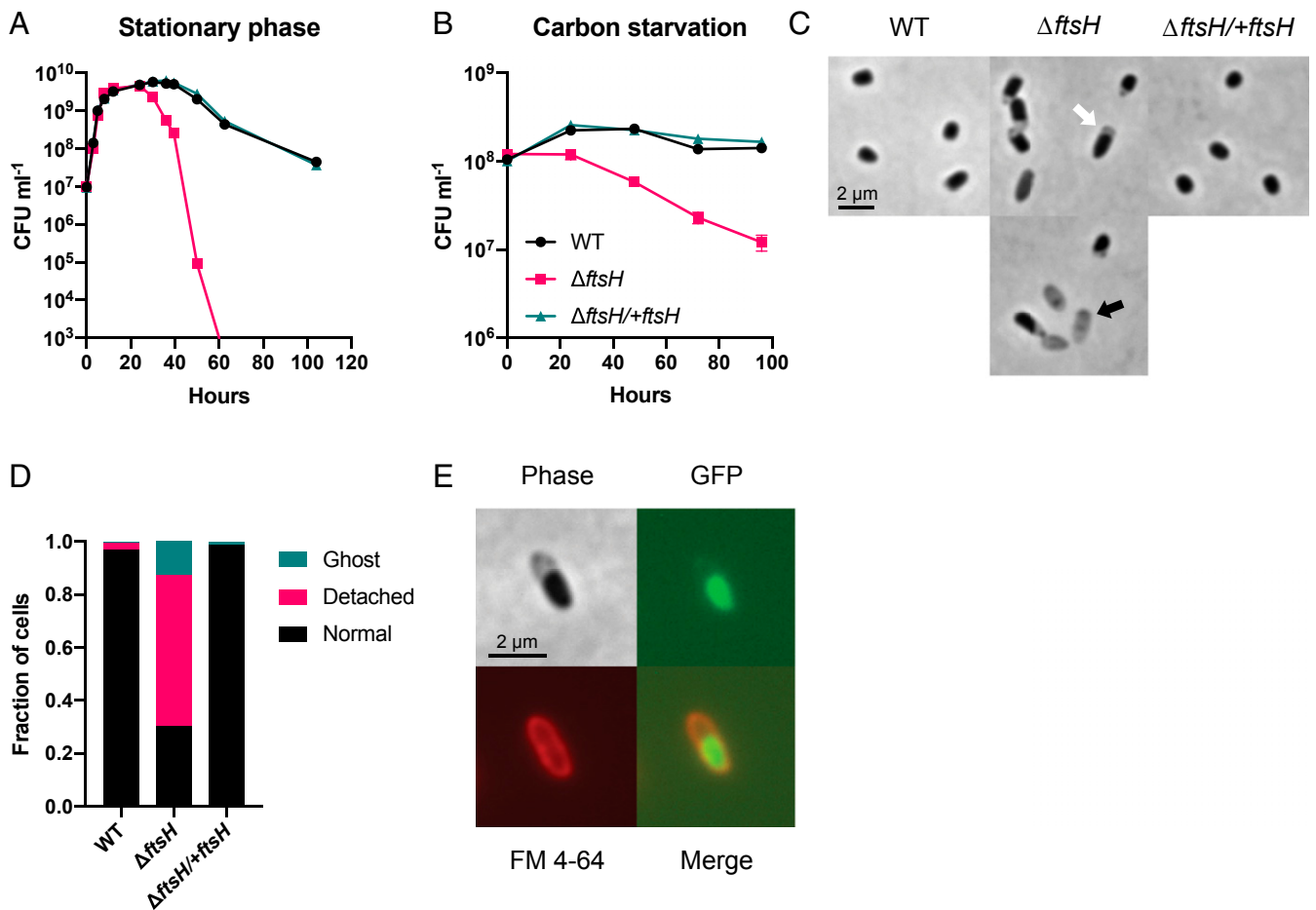


Fig. 1. FtsH maintains cell integrity during growth arrest. Loss of *ftsH* exacerbates cell death during stationary phase (A) and carbon starvation (B). Viability was below the limit of detection ($\sim 3 \times 10^2$ CFU mL⁻¹) for Δ ftsH at 60 h in A. Representative data from at least three independent experiments are shown in A and the averages and SD of biological replicates are shown in B ($n = 3$). (C) Characteristic morphology of Δ ftsH cells after 24 h of carbon starvation. The white arrow indicates a “detached” inner membrane and the black arrow indicates a “ghost cell.” (D) Quantification of the cellular morphologies described in C. A minimum of 300 cells were counted for each strain. (E) OM staining with FM 4-64 and cytoplasmic expression of GFP confirms that detachment occurs between the IM and OM.

AmgRS in *P. aeruginosa* and contributes to aminoglycoside resistance in conjunction with PA14_72930, another AmgRS-regulated gene of unknown function that was also aggravating for growth in our screen (SI Appendix, Table S4 and Dataset S1) (65).

We made isogenic markerless deletions of growth-aggravating genes from different functional categories to validate their interactions with *ftsH*. To this end, we made double mutants of *ftsH* with *tpiA*, *lptA*, and *fabF1* (involved in “carbohydrate and phospholipid metabolism”) (SI Appendix, Fig. S4) and with *hslVU*, *hlpX*, and *amgRS* (involved in “protein turnover”). The double mutants had significantly reduced growth rates in LB at 37 °C compared to the Δ ftsH single mutant, with the exception of Δ ftsH Δ hlpX (Fig. 2 B and C). The growth defects were greatest for Δ ftsH Δ fabF1, with a 63% reduction in growth rate compared to Δ ftsH, and for Δ ftsH Δ amgRS, with a 60% reduction in growth rate (Fig. 2 B and C). Together, these data validate the results of our Tn-seq experiment and confirm a role for FtsH in promoting diverse cellular processes during growth.

Importantly, all double mutants, with the exception of Δ ftsH Δ hslVU, showed similar survival profiles to the Δ ftsH single mutant, indicating that while most of these genes are vital for growth in the Δ ftsH background they do not affect survival during growth arrest (Fig. 2 D and E). In contrast, Δ ftsH Δ hslVU had a survival defect ~ 10 -fold more severe than Δ ftsH after 3 d of

carbon starvation (Fig. 2E), indicating that HslVU plays a backup role to FtsH during growth and growth arrest. This result prompted us to systematically investigate the role of the ATP-dependent heat-shock proteases during growth arrest.

Heat-Shock Proteases Promote Survival during Growth Arrest. The heat-shock proteases classically act in conjunction with protein-folding chaperones to maintain proteostasis after exposure to high temperature, ethanol, or heavy metal stress (68, 69). To more broadly characterize the role of the heat-shock response during growth arrest, we measured expression of the major ATP-dependent proteases and chaperones in WT PA14 during carbon starvation by qRT-PCR (Fig. 3A). Following a substantial decrease in expression within the first hour of carbon starvation, most heat-shock regulated genes are induced over the next 18 h, whereas expression of the growth-related gene *dnaA* does not increase over time.

We predicted that further genetic perturbation of the protease network in *P. aeruginosa* would exacerbate survival defects during growth arrest. To test this prediction, we made all combinatorial deletions of the ATP-dependent protease-encoding genes *ftsH*, *clpXP*, *lon*, and *hslVU*, and assessed survival of the mutant strains during carbon starvation (Fig. 3B). Single deletion of *lon*, like *hslVU*, did not cause a survival defect after 3 d of carbon starvation, while deletion of *clpXP* caused a similar survival defect as Δ ftsH (Fig. 3 B and C). We did not detect a fitness defect during

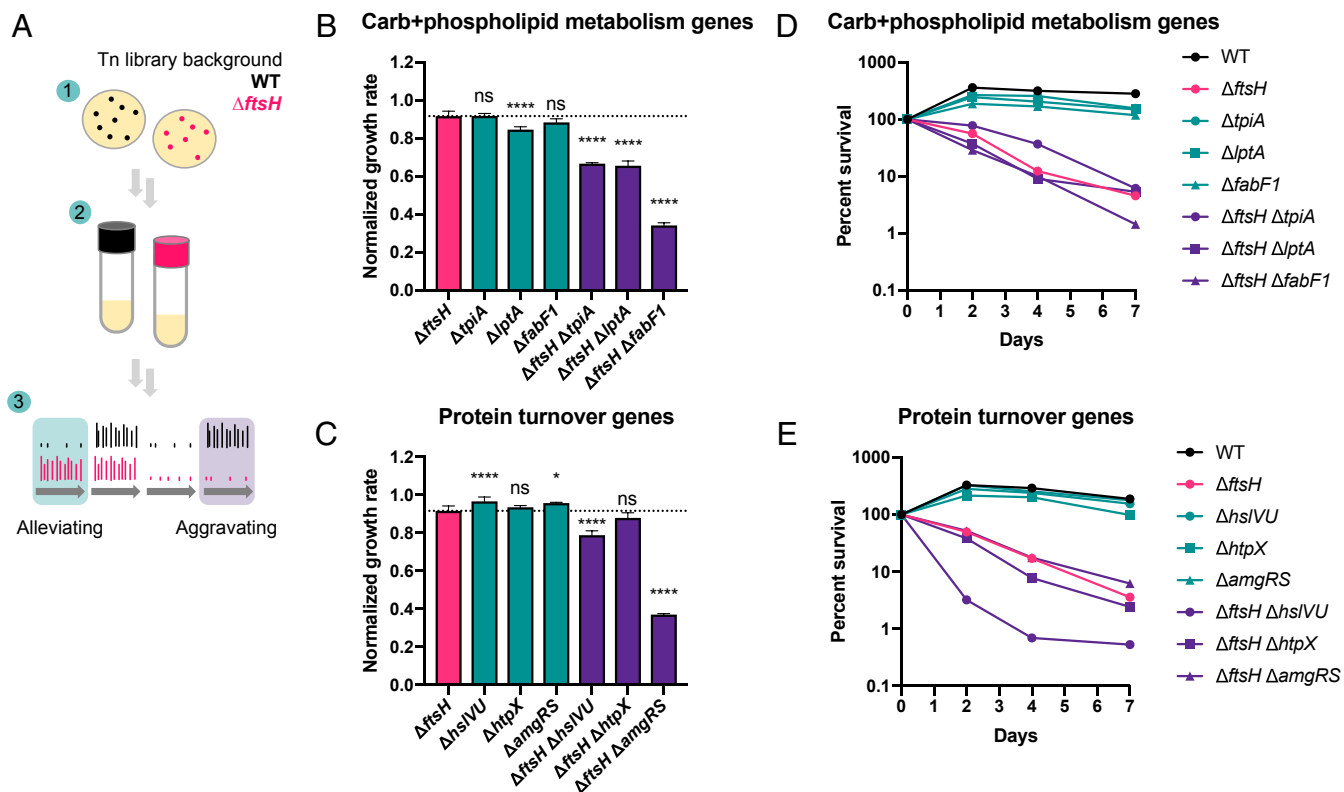


Fig. 2. Mutations in metabolic and protein turnover genes aggravate growth and survival defects of \DeltaftsH . (A) Experimental design of Tn-seq screen. 1) Transposon libraries were generated in the WT (15) and \DeltaftsH backgrounds by selection on LB agar plates plus antibiotics at 37 °C. 2) An aliquot of each library was grown approximately seven to eight generations in liquid LB. 3) The read counts of the WT and \DeltaftsH libraries were compared to identify genes that are alleviating or aggravating for growth in the \DeltaftsH background. (B and C) Growth rate of carbohydrate and phospholipid metabolism and protein turnover mutants relative to \DeltaftsH in LB at 37 °C. All growth rates were normalized to the WT growth rate in LB at 37 °C and the dotted line indicates the normalized \DeltaftsH growth rate. Results of one-way ANOVA with Dunnett's multiple comparisons test are denoted as follows: ns, not significant; * $P < 0.05$; **** $P < 0.0001$. Error bars show SD of biological replicates ($n \geq 3$). (D and E) Survival of mutants during carbon starvation. Representative data from at least two independent experiments are shown.

carbon starvation for *clpX* or *clpP* transposon mutants in our previous study because there were few insertions in these genes in the WT transposon library (15) (Dataset S2). Although $\Delta clpXP$ had a similar growth rate as \DeltaftsH at 37 °C (see below), we noticed that $\Delta clpXP$ colonies took longer to appear upon plating after growth arrest. We reasoned that the reduced insertion rate at the *clpXP* locus is due to an extended lag time of mutant cells before beginning to grow (70), which leads to these mutants being out-competed in the early stages of growth of the transposon libraries. These results emphasize that while Tn-seq is a powerful technique for studying single mutant fitness on a genome-wide scale, it frequently lacks resolving power for studying stress-related phenotypes of mutants with even a minor growth disadvantage under nonstressful conditions (71).

Unlike in the \DeltaftsH background, deletion of *hslVU* did not affect survival in the Δlon or $\Delta clpXP$ backgrounds, indicating that *hslVU* specifically backs up FtsH activity (Fig. 3B). Deletion of *lon* did not affect survival in the \DeltaftsH background, but exacerbated death in the $\DeltaftsH \Delta hslVU$ background, indicating that Lon plays a tertiary role in growth arrest survival (Fig. 3B). Deletion of *clpXP* also exacerbated death in the $\DeltaftsH \Delta hslVU$ background, as well as in the \DeltaftsH background (Fig. 3B and C). Thus, *ftsH* and *clpXP* appear to contribute independently to survival during growth arrest. Finally, the $\DeltaftsH \Delta hslVU \Delta lon \Delta clpXP$ quadruple mutant had a survival defect similar to the more sensitive triple mutants. However, suppressor mutations arise rapidly in this strain, making it difficult to determine an

accurate growth rate and survival phenotype for the parent strain.

Heat and Alkaline pH Exacerbate Death during Growth Arrest. The functional overlap of the heat-shock proteases during growth arrest suggests that they might act collectively to mitigate a common stress, which we hypothesized was caused by the accumulation of misfolded proteins. Thus, we tested the functional overlap between these proteases in two other conditions known to cause protein misfolding stress: Growth at high temperature or alkaline pH (72).

Functional overlap of the proteases during growth at high temperature was similar to that of growth arrest (Fig. 4A). While \DeltaftsH had only a minor growth defect compared to WT in LB at 37 °C (~8% reduced growth rate), this defect became more pronounced at 42 °C and 44 °C (~29% and ~41% reduced growth rate, respectively). Once again, we observed functional redundancy between *ftsH* and *hslVU*, with loss of both genes resulting in no growth at 42 °C and 44 °C. Unlike in growth arrest, however, the temperature sensitivity of \DeltaftsH was greater than that of $\Delta clpXP$, with the latter displaying temperature-sensitive growth only at 44 °C. The $\Delta hslVU$ and Δlon single mutants were also temperature-sensitive only at 44 °C. The pattern of overlap during growth in LB at pH 9 was strikingly similar to that of growth arrest (compare Fig. 4B and Fig. 3B), supporting the hypothesis that these two conditions invoke a common cellular crisis.

To determine whether the reduced growth rate of protease mutants at high temperature was caused by uniformly slower

to 37 °C increased the death rate of $\Delta ftsH$ during carbon starvation (Fig. 4D). Death of $\Delta ftsH$ and $\Delta clpXP$ in stationary phase was delayed in buffered LB (Fig. 4E and *SI Appendix, Fig. S6B*), indicating that a combination of growth arrest and alkaline pH likely causes the dramatically greater loss of viability in stationary phase LB cultures compared to carbon starvation in buffered minimal medium (compare Fig. 1 A and B) (73). Higher temperature also exacerbated growth defects at alkaline pH, as WT grew slower at 37 °C than at 30 °C in LB pH 9, and $\Delta ftsH$ failed to grow at 37 °C (Fig. 4F). In contrast, both strains grew faster at 37 °C than at 30 °C in neutral LB. Based on these data, we conclude that growth arrest, high temperature, and alkaline pH are synergistic stresses that require a coordinated response of the heat-shock proteases in order for cells to grow and survive.

Heat-Shock Proteases Delay Protein Aggregation during Growth Arrest. To indirectly assess the extent of protein misfolding during growth arrest, we visualized protein aggregates in single cells using fluorescently tagged IbpA, a small heat-shock protein that has been shown to bind to endogenous protein aggregates (74). IbpA fusion proteins constructed with monomeric fluorescent protein derivatives have been validated as a reliable reporter for monitoring protein aggregates in *E. coli* (58). A gene encoding IbpA-mVenus was cloned into the chromosome under the control of its native promoter in *P. aeruginosa*; when expressed, it rapidly formed fluorescent foci in both the WT and protease mutants upon heat shock (*SI Appendix, Fig. S7*). During exponential growth, and as cells entered stationary phase, IbpA-mVenus signal was predominantly low and diffusely distributed throughout the

cytoplasm, indicating that protein aggregation was minimal under nutrient-replete growth conditions (Fig. 5A). Importantly, diffuse labeling during growth demonstrates that our reporter is not inherently aggregation-prone, a common artifact in these types of experiments (58, 75). Both $\Delta ftsH$ and $\Delta ftsH \Delta hslVU$ developed one to two foci per cell during stationary phase that became brighter and more numerous over time, with an accelerated rate of foci formation in the double mutant. This pattern was also observed in WT, albeit with a delayed onset relative to the protease mutants. For all strains, the appearance of numerous foci correlated with cell death (Fig. 5B), and following death many cells lost fluorescence. Thus, aggregate formation is delayed, rather than completely prevented, by the activity of heat-shock proteases during growth arrest, and aggregate formation correlates with cell death in both the WT and protease-deficient strains.

Discussion

Bacteria must adequately maintain their cellular integrity during extended periods of starvation in order to survive and to resume growth when nutrients become available. In this study, we describe a role for FtsH in promoting diverse metabolic processes during growth of *P. aeruginosa* and in maintaining cellular integrity during growth arrest. Unlike its posited role in *E. coli* (38, 39), our findings suggest that FtsH does not play a regulatory role in growth arrest survival of *P. aeruginosa* per se. Rather, FtsH appears to act in conjunction with the other major ATP-dependent heat-shock proteases to ensure ongoing protein quality control in the absence of growth. Protein aggregation appears to become a major problem during growth arrest, and could be caused by

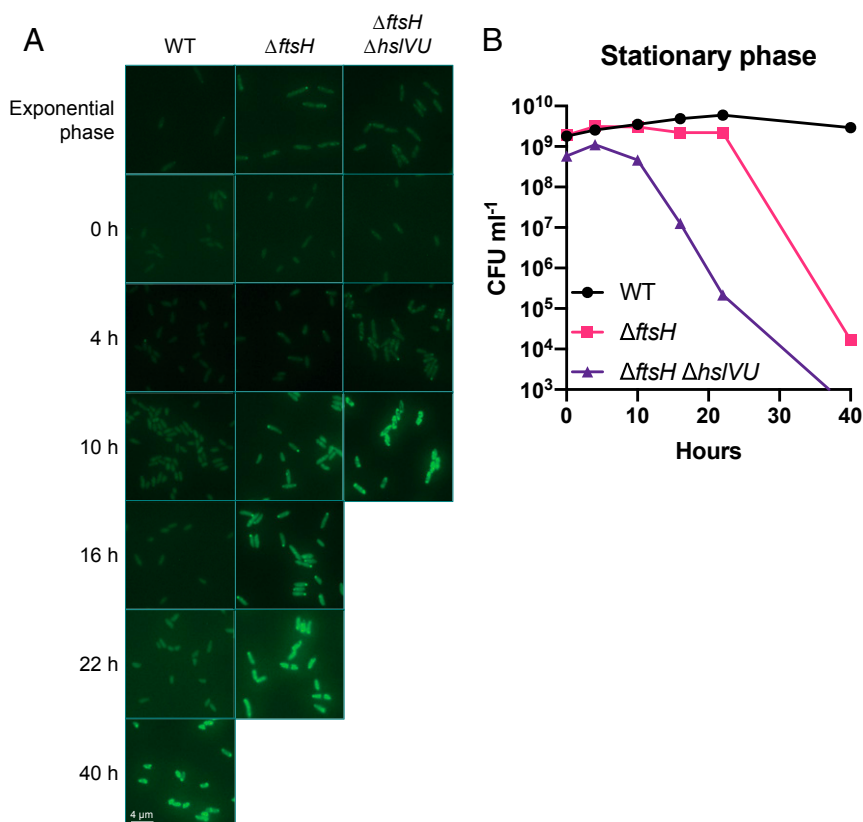


Fig. 5. Heat-shock proteases delay protein aggregation during growth arrest. (A) Fluorescence of cells expressing the IbpA-mVenus reporter during exponential growth and stationary phase at 37 °C; 0 h corresponds to the start of stationary phase (after 8 h of growth starting from an OD₅₀₀ of 0.01). All images were taken using the same exposure time (400 ms) and are displayed with the same brightness and contrast. (B) Viability measurements corresponding to the time points in A. Viability was below the limit of detection ($\sim 3 \times 10^2$ CFU mL⁻¹) for $\Delta ftsH \Delta hslVU$ at 40 h. Representative data from two independent experiments are shown.

numerous factors, including misfolding of nascent peptides due to reduced abundance or activity of cotranslational folding chaperones, or misfolding of bulk proteins over time due to sustained and cumulative damage in the absence of growth and turnover.

We observed differing phenotypic interactions between *ftsH* and diverse functional classes of genes during growth and growth arrest of *P. aeruginosa*, raising the question of whether FtsH has divergent roles in these two states. Mutations in carbohydrate and lipid metabolic genes exacerbated the growth defect of Δ *ftsH* but had no effect on Δ *ftsH* survival during growth arrest (Fig. 2 B and D), while mutations in heat-shock proteases exacerbated both the growth and survival defects of Δ *ftsH* (Fig. 2 C and E). What is the connection between FtsH and diverse metabolic pathways during growth? One possibility is that cells are more sensitive to protein misfolding and aggregation in the absence of FtsH-mediated protein quality control, which could interfere with the activity of growth-promoting metabolic enzymes. Evidence for this possibility is found in *Mycobacteria*, where disruption of proteostasis results in misfolding and reduced activity of multimodular lipid synthases (53). Another possibility is that, during growth, FtsH might still control flux through phospholipid and LPS metabolism in *P. aeruginosa* independent of a direct effect on the stability of LpxC. Recent studies have revealed complex cross-talk between phospholipid and LPS metabolism in *E. coli*, and point to a more nuanced role for FtsH in sensing and controlling flux through these pathways (76, 77). Perhaps an analogous role for FtsH exists in *P. aeruginosa*, with the regulatory architecture having diverged from that of *E. coli*.

Altered regulation of the heat-shock sigma factor σ^{32} in the absence of FtsH could also play a role during growth. During growth at moderate temperatures, σ^{32} levels are kept low by FtsH-mediated proteolysis in diverse species (31). Upon heat shock, σ^{32} becomes stabilized and induces expression of the heat-shock regulon. Overexpression of σ^{32} under noninducing conditions was shown to inhibit growth of *Caulobacter crescentus* by reprogramming gene expression away from a growth-promoting regime toward one favoring repair and maintenance (78). A similar problem could occur in *P. aeruginosa*, where loss of FtsH causes a global change in gene expression indirectly via elevated σ^{32} . In support of this possibility, overexpression of σ^{32} down-regulates expression of multiple fatty acid biosynthetic genes in PA14, including *fabF1* (79), whose deletion caused the most severe growth defect in the Δ *ftsH* background in our experiments (Fig. 2B). If growth-promoting pathways are generally down-regulated in Δ *ftsH*, then further perturbation to these pathways could severely compromise growth. This could explain why lipid metabolism genes are important for growth of Δ *ftsH* yet dispensable for survival, as fatty acid and phospholipid synthesis would no longer be required once cells have stopped growing.

Like FtsH, ClpXP has also been implicated in adaptation to and survival of growth arrest in diverse organisms (80–85). Genetic perturbation of protein-folding chaperones and σ^{32} in *E. coli* also causes a survival defect during growth arrest (41, 42). Furthermore, oxidatively modified proteins that accumulate during aerobic growth arrest cause specific induction of heat-shock genes in *E. coli* (43–46). Together, these findings point to a generalized role for the heat-shock response in ensuring survival of nongrowing bacteria. Outside of these studies, however, the importance of heat-shock genes is overlooked in the literature on growth arrest. Most reviews on bacterial growth arrest do not discuss a role for the heat-shock response, and most references to heat shock refer mainly to its namesake role in temperature stress. Strikingly, of all of the deletion strains of *P. aeruginosa* we have tested, loss of FtsH or ClpXP caused by far the most severe survival defect during carbon starvation (ref. 15 and this study). Survival was considerably worse in protease mutant strains than for a mutant lacking the stress sigma factor RpoS, which is generally considered one of the principle

molecular components required for adaptation to nutrient depletion (15, 86).

In contrast to the literature on growth arrest in bacteria, the importance of the heat-shock response and proteostasis in cellular aging is well described and appreciated in eukaryotic organisms (47, 48, 87–91). Defects in proteostasis are a hallmark of aging in eukaryotes, and our finding that growth-arrested bacteria sustain similar types of cellular damage as aging eukaryotic cells points to an equally important role for proteostasis in bacterial fitness. The role of proteostasis during bacterial growth arrest is in addition to its role during “replicative aging” of nutrient-replete, growing cells (50, 92–95). The latter phenomenon occurs when one daughter cell inherits the old cell pole while the other daughter is rejuvenated. Based on the literature and our data, we propose that cellular aging during growth arrest is at least equal in importance to replicative aging. Both replicative aging and aging in growth arrest are likely driven by the accumulation of protein aggregates (50, 52), and this process may be magnified in growth arrest due to the reduced capability of cells for growth and protein turnover.

The communal role of the heat-shock proteases during growth arrest is in contrast to the many varied and specific regulatory roles they play in the physiology of diverse bacteria. Perhaps growth arrest, along with high temperature and alkaline pH, was an important driver in the evolution of this ancestral proteolytic network to maintain protein quality control. Following its development, this redundant network could have been co-opted by different bacteria to regulate diverse cellular processes in addition to proteostasis, such as regulation of LPS and phospholipid metabolism by FtsH (32, 77), the DNA damage response by Lon and HslVU (96, 97), and RpoS stability, DNA replication, and cell division by ClpXP (98–100).

Our findings point to proteostasis as an important, yet underexplored, aspect of growth arrest physiology. If surviving growth arrest is as important for bacterial success in diverse ecological niches as we presume, this should compel us to investigate the role of proteostasis more thoroughly in these niches as a counterpart to the more commonly studied stresses that induce proteotoxicity, such as heat, peroxide, or antibiotic exposure during rapid growth. Indeed, opportunistic human pathogens like *P. aeruginosa* are unlikely to experience temperature stress much higher than 37 °C during chronic infection, a temperature at which *P. aeruginosa* grows optimally. On the other hand, cells can quickly become energy-limited for growth during chronic infection, and what was once an optimal growth temperature could now impede the survival of aging cells. Thus, proteases of the heat-shock response may hold potential as a therapeutic target for the treatment of *P. aeruginosa* and related bacterial infections.

Materials and Methods

Strains and Growth Conditions. The strains, plasmids, and primers used in this study are listed in *SI Appendix, Tables S1–S3*. *E. coli* and *P. aeruginosa* were grown in LB (Difco) or on LB agar plates with appropriate antibiotics at 30 °C or 37 °C for all cloning and strain construction purposes. All gene deletions in *P. aeruginosa* strain UCBBP-PA14 were made as described previously (15). The LpxC inhibitor CHIR-090 was purchased from ApexBio and the OM dye FM 4-64 was purchased from Life Technologies.

Growth Arrest Survival Assays. For studying growth arrest caused by carbon starvation, cells were grown in LB at 37 °C to an optical density at 500 nm (OD_{500}) between 0.5 and 1, pelleted, and resuspended in CSM, which is derived from a minimal medium without an added carbon source (13). In CSM, magnesium and sulfate are supplied in the form of $MgCl_2$ and $NaSO_4$, respectively, instead of the standard $MgSO_4$. This allows for magnesium concentrations to be titrated independently of sulfate when necessary. Starved cells were incubated aerobically at 37 °C with shaking at 250 rpm. For oxygen starvation, cells were resuspended in the same minimal medium plus 40 mM pyruvate and transferred into an anoxic glove chamber (Coy) containing an atmosphere of 15% CO_2 , 80% N_2 , and 5% H_2 and incubated

anaerobically at 33 °C without shaking. For nitrogen starvation, cells were resuspended in the same minimal medium plus 10 mM glucose and without NH₄Cl and incubated aerobically at 37 °C with shaking at 250 rpm. CFU mL⁻¹ were determined by viability plating over time in all conditions.

For studying growth arrest in stationary phase, overnight cultures grown in LB were back-diluted to an OD₅₀₀ of 0.01 in LB or LB plus 50 mM Mops (buffered LB) and grown at the indicated temperature with shaking at 250 rpm. CFU mL⁻¹ were determined by viability plating over time. Most strains reached stationary phase after ~8 h of growth.

LPS Measurements. Overnight cultures grown in 5 mL LB were back-diluted in triplicate to an OD₅₀₀ of 0.01 in 5 mL LB and grown at 37 °C with shaking at 250 rpm. An OD 0.5 mL⁻¹ equivalence of each culture was pelleted after 3 h of growth (corresponding to an OD₅₀₀ between 0.2 and 0.3; exponential phase) and after 25 h of growth (corresponding to an OD₅₀₀ between 4 and 5; stationary phase), and stored at -80 °C. LPS was extracted from frozen pellets as described previously (38). Gels were stained with Pro-Q Emerald 300 LPS Gel Stain Kit (Molecular Probes) according to the manufacturer's protocol and the LPS bands were visualized by UV transillumination.

Generation of the Transposon Library in *ΔftsH*. A transposon library was created in the *ΔftsH* background in the same manner as described for the WT strain (15). Briefly, the *ΔftsH* strain of *P. aeruginosa* and the *E. coli* strain SM10λ_{pir} carrying the transposon-bearing plasmid pIT2 were resuspended in 1 mL of LB from overnight streak plates on LB agar or LB agar plus carbenicillin (100 μg mL⁻¹), respectively. The resuspended cells were adjusted to an OD₅₀₀ of ~50 for *P. aeruginosa* and ~100 for *E. coli*. Next, 100 μL of each OD-adjusted strain was mixed together in an Eppendorf tube and two 50-μL aliquots of this mix were plated on sterile 0.2-μm filter discs placed on an LB agar plate. The conjugation spots were allowed to dry in a safety cabinet with laminar flow for 15 min and then the plate was incubated at 37 °C for 2.5 h. Following incubation, the two conjugation spots were resuspended in 3.5 mL of LB and 100-μL aliquots of this resuspension were plated on 33 LB agar plus tetracycline (60 μg mL⁻¹) and chloramphenicol (10 μg mL⁻¹) plates to select for transposon insertion mutants. The plates were incubated 27 h at 37 °C. Each plate yielded ~4,500 mutants for a total of ~150,000 mutants. Following incubation, colonies from all plates were pooled and resuspended in LB plus 15% glycerol. The density of the pooled library was adjusted to an OD₅₀₀ of 10 and stored as 1-mL aliquots at -80 °C.

Tn-Seq Sample Preparation and Data Analysis. Frozen aliquots of the WT and *ΔftsH* transposon libraries were thawed on ice for 15 min and diluted in 50 mL of LB to a concentration of ~6 × 10⁶ CFU mL⁻¹. The cultures were grown at 37 °C with shaking at 250 rpm for 3.5 h, corresponding to approximately three to four cell doublings. An aliquot of each culture was then diluted in 50 mL of LB plus chloramphenicol (10 μg mL⁻¹) to a concentration of ~6 × 10⁶ CFU mL⁻¹ and grown at 37 °C with shaking at 250 rpm for 3 h, corresponding to approximately four doublings. Following growth, an OD 4 mL⁻¹ equivalence of each LB plus chloramphenicol culture was pelleted and stored at -80 °C. Genomic DNA was extracted from the frozen samples for high-throughput sequencing, as described previously (15). Sequencing was performed at the Millard and Muriel Jacobs Genetics and Genomics Laboratory at the California Institute of Technology. Sequences were mapped to the UCBBP-PA14 genome sequence using Bowtie (101) and analyzed using the ARTIST Tn-seq analysis pipeline in MATLAB, as described previously (15, 102).

Growth-Rate Measurements. For batch growth-rate experiments, precultures of each strain were grown at 37 °C in LB then pelleted and resuspended at an OD₅₀₀ of 0.05 in LB. Next, 150-μL aliquots of the resuspensions were dispensed into a 96-well microtiter plate and 50 μL of mineral oil was added to each well to prevent evaporation. Plates were incubated in a BioTek plate reader at the indicated temperature (37 °C, 42 °C, 41 °C, or 44 °C) with medium shaking and reading at OD₅₀₀ every 15 min.

For microfluidic single-cell growth time-lapse experiments, precultures grown at 37 °C in LB were subcultured 1:200 in fresh LB and grown under the

same conditions to OD₅₀₀ ~1.0. Cells were then diluted in fresh LB at room temperature to OD₅₀₀ 0.01 and loaded into a Millipore CellASIC microfluidic chamber bacteria plate (Cat: B04A-03-5PK). The plate was transferred to a prewarmed OKOLabs incubation cage (37 °C or 41 °C) around a Nikon Ti2 Eclipse microscope stage. Single cells of each strain were trapped in separate chambers on the plate using the CellASIC ONIX2 microfluidic system according to the manufacturer's protocol. Fresh LB was flowed over the dispersed cells at a rate of ~2.5 μL h⁻¹. Phase-contrast images were taken with a 40×/0.95 NA objective every 5 min starting 30 min after the initial flow of fresh LB. Image analysis was performed in imageJ and data processing in Jupyter Lab. The imageJ plugin "StackReg" (<http://bigwww.epfl.ch/thevenaz/stackreg/>) was used to correct for drift in the time-lapse movies. Individual tracks of one or a few cells growing into isolated microcolonies were manually cropped out and inspected to ensure clonal growth. Binary masks of the cell area were generated for each frame and the total area per frame was quantified after excluding edge particles. Frames with zero area (due to the cell mass making transient contact with an edge) were removed and each track was divided by its first value to correct against initial cell area variability.

To calculate the growth rate, OD₅₀₀ and cell area data were processed using Python 3.6 with help from the following Python libraries: pandas, numpy, scipy, matplotlib, and seaborn. OD₅₀₀ data were smoothed using a savgol filter (scipy.signal.savgol_filter, window length: 9 time measurements, polyorder: 3). Outlier detection was performed by smoothing the averaged data for each genotype and identifying the data points >2 SDs away from the smoothed curve. Outliers were identified and replaced with the average smoothed value at their respective time points. Once outliers were mitigated, each individual sample was smoothed using the savgol filter. To estimate the growth rate for each sample, we searched for the line of best fit (fit via scipy.stats.linregress) with the largest slope using a sliding window of 250 min on the smoothed data. The sliding window was allowed to search intervals between 0 and 900 min.

qRT-PCR Measurements. Triplicate overnight cultures grown in 5 mL LB were back diluted to an OD₅₀₀ of 0.01 in 50 mL LB and grown at 37 °C with shaking at 250 rpm to an OD₅₀₀ of 0.7, pelleted, and resuspended to an OD₅₀₀ of 0.5 in CSM. Starved cells were incubated aerobically at 37 °C with shaking at 250 rpm for 1 h. Following this incubation, an OD 0.9 mL⁻¹ equivalence of each culture (1.8 mL) was pelleted after 0, 4, 8, and 18 h of carbon starvation and stored at -80 °C until RNA extraction. RNA extraction and qRT-PCR were performed as described previously (103). Briefly, RNA was extracted using the RNeasy Mini Kit (Qiagen) and genomic DNA was depleted using the TURBO DNA-free Kit (Invitrogen). cDNA was synthesized from TURBO DNase-treated RNA using the iScript cDNA synthesis kit (Bio-Rad) and the reactions were run using a Fast 7500 Real-Time PCR System machine (Applied Biosystems). Expression of each gene was normalized to the expression of the housekeeping gene *oprI*.

Routine Microscopy. Routine microscopy was performed using a Zeiss Axio Imager. An exposure time of 400 ms and 2,000 ms was used for cells expressing the lbpA-mVenus reporter in Fig. 5 and *SI Appendix, Fig. S7*, respectively. Within each experiment all images were adjusted for equal brightness and contrast using the image-processing software FIJI (104). Cell length in *SI Appendix, Fig. S1* was measured using SuperSegger (105).

Data Availability Statement. All strains used in the paper will be made available to readers.

ACKNOWLEDGMENTS. We thank members of the D.K.N. laboratory for thoughtful discussions and critical feedback on the manuscript, and Lisa Racki (The Scripps Research Institute) for the gift of pLREX97. This manuscript derives from a chapter in D.W.B.'s doctoral thesis from the California Institute of Technology. This work was supported by the Millard and Muriel Jacobs Genetics and Genomics Laboratory at the California Institute of Technology and by the NIH (1R01AI127850-01A1 and 1R21AI146987-01).

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