

Heterogeneity in surface sensing suggests a division of labor in *Pseudomonas aeruginosa* populations

Catherine R Armbruster^{1†}, Calvin K Lee^{2,3,4†}, Jessica Parker-Gilham¹, Jaime de Anda^{2,3,4}, Aiguo Xia⁵, Kun Zhao^{6,7}, Keiji Murakami⁸, Boo Shan Tseng⁹, Lucas R Hoffman^{1,10}, Fan Jin^{5,11}, Caroline S Harwood¹, Gerard CL Wong^{2,3,4}, Matthew R Parsek^{1,12*}

¹Department of Microbiology, University of Washington, Seattle, United States; ²Department of Bioengineering, University of California, Los Angeles, Los Angeles, United States; ³Department of Chemistry and Biochemistry, University of California, Los Angeles, Los Angeles, United States; ⁴California NanoSystems Institute, University of California, Los Angeles, Los Angeles, United States; ⁵Hefei National Laboratory for Physical Sciences at the Microscale, University of Science and Technology of China, Hefei, China; ⁶Key Laboratory of Systems Bioengineering (Ministry of Education), School of Chemical Engineering and Technology, Tianjin University, Tianjin, China; ⁷Collaborative Innovation Centre of Chemical Science and Engineering, Tianjin University, Tianjin, China; ⁸Department of Oral Microbiology, Institute of Biomedical Sciences, Tokushima University Graduate School, Tokushima, Japan; ⁹School of Life Sciences, University of Nevada, Las Vegas, United States; ¹⁰Department of Pediatrics, University of Washington, Seattle, United States; ¹¹Institute of Synthetic Biology, Shenzhen Institutes of Advanced Technology, Chinese Academy of Sciences, Shenzhen, China; ¹²Integrative Microbiology Research Centre, South China Agricultural University, Guangzhou, China

*For correspondence:
parsem@uw.edu

†These authors contributed equally to this work

Competing interests: The authors declare that no competing interests exist.

Funding: See page 25

Received: 11 January 2019

Accepted: 08 June 2019

Published: 10 June 2019

Reviewing editor: Alexandre Persat, EPFL, Switzerland

© Copyright Armbruster et al. This article is distributed under the terms of the [Creative Commons Attribution License](#), which permits unrestricted use and redistribution provided that the original author and source are credited.

Abstract The second messenger signaling molecule cyclic diguanylate monophosphate (c-di-GMP) drives the transition between planktonic and biofilm growth in many bacterial species. *Pseudomonas aeruginosa* has two surface sensing systems that produce c-di-GMP in response to surface adherence. Current thinking in the field is that once cells attach to a surface, they uniformly respond by producing c-di-GMP. Here, we describe how the Wsp system generates heterogeneity in surface sensing, resulting in two physiologically distinct subpopulations of cells. One subpopulation has elevated c-di-GMP and produces biofilm matrix, serving as the founders of initial microcolonies. The other subpopulation has low c-di-GMP and engages in surface motility, allowing for exploration of the surface. We also show that this heterogeneity strongly correlates to surface behavior for descendent cells. Together, our results suggest that after surface attachment, *P. aeruginosa* engages in a division of labor that persists across generations, accelerating early biofilm formation and surface exploration.

DOI: <https://doi.org/10.7554/eLife.45084.001>

Introduction

Pseudomonas aeruginosa is an opportunistic pathogen that engages in a range of surface-associated behaviors and is a model bacterium for studies of surface-associated communities called biofilms. Biofilms are dense aggregates of cells producing extracellular matrix components that hold the

eLife digest Bacteria can adopt different lifestyles, depending on the environment in which they grow. They can exist as single cells that are free to explore their environment or group together to form 'biofilms'. The bacteria in biofilms stick to a surface, and produce a slimy 'matrix' that covers and thereby protects them. Biofilms have been found in lung infections that affect people with the genetic disorder cystic fibrosis, and can also form on the surface of medical implants. Because the biofilm lifestyle protects bacteria from the immune system and antimicrobial drugs, learning about how biofilms form could help researchers to discover ways to prevent and treat such infections.

Many bacteria switch between the free-living and biofilm lifestyles by altering their levels of a signaling molecule called cyclic diguanylate monophosphate (called c-di-GMP for short). Bacteria living in biofilms have much higher levels of c-di-GMP than their free-living counterparts, and bacteria that have high levels of c-di-GMP produce more biofilm matrix.

Bacteria called *Pseudomonas aeruginosa* use a protein signaling complex called the Wsp system to sense that they are on a surface and increase c-di-GMP production. Questions remained about how quickly this change in production occurs, and whether bacteria pass on their c-di-GMP levels to the new descendant cells when they divide.

Armbruster et al. monitored individual cells of *P. aeruginosa* producing c-di-GMP as they began to form biofilms. Unexpectedly, not all cells increased their c-di-GMP levels when they first attached to a surface. Instead, Armbruster et al. found that there are two populations – high and low c-di-GMP cells – that each perform complementary and important tasks in the early stages of biofilm formation.

The high c-di-GMP cells represent 'biofilm founders' that start to produce the biofilm matrix, whereas the low c-di-GMP cells represent 'surface explorers' that spend more time traveling along the surface. Armbruster et al. found that the Wsp surface sensing system generates these two populations of cells. Moreover, the c-di-GMP levels in a bacterial cell even affect the behavior of the descendant cells that form when it divides. This effect can persist for several cell generations.

More work is needed to examine exactly how the biofilm founders and surface explorers interact and influence how biofilms form, and to discover if blocking c-di-GMP signaling prevents biofilm formation. This could ultimately lead to new strategies to prevent and treat infections in humans.

DOI: <https://doi.org/10.7554/eLife.45084.002>

community together. The biofilm mode of growth is beneficial for bacteria in that it allows cells to maintain close proximity to nutrients, promotes exchange of genetic material, and confers cells protection from a variety of chemical and environmental stresses (e.g. nutrient limitation, desiccation, and shear forces), as well as engulfment by protozoa in the environment or by phagocytes in a host (**Davey and O'toole, 2000**). Collectively, these advantages make biofilm formation integral to prokaryotic life.

The secondary messenger signaling molecule cyclic diguanylate monophosphate (c-di-GMP) controls the transition between the planktonic to the biofilm mode of growth. In many bacterial species, including *P. aeruginosa*, elevated c-di-GMP results in repression of flagellar motility genes, while promoting expression of genes involved in producing a biofilm matrix (**Römling et al., 2013**). The *P. aeruginosa* biofilm matrix is composed of a combination of polysaccharides (including Pel and Psl), proteins (including the adhesin CdrA), and extracellular DNA (**Ma et al., 2009**). Biofilm matrix production is an energetically costly process that is regulated at multiple levels (**Wei and Ma, 2013**). The *cdrA*, *pel* and *psl* genes are all transcriptionally induced under conditions of high c-di-GMP (**Starkey et al., 2009**).

For many species, the initial step in biofilm formation involves adherence of free swimming planktonic cells to a surface and the initiation of surface sensing. *P. aeruginosa* has at least two distinct surface sensing systems, the Wsp and the Pil-Chp systems, that when activated, lead to biofilm formation. The Wsp system senses an unknown surface-related signal (recently proposed to be membrane perturbation [**Chen et al., 2014**]) through WspA, a membrane-bound protein homologous to methyl-accepting chemotaxis proteins (MCPs). Activation of this system stimulates phosphorylation of the diguanylate cyclase WspR, which leads to the formation of aggregates of phosphorylated

WspR (WspR-P) in the form of visible subcellular clusters. This aggregation of WspR-P potentiates its activity, increasing c-di-GMP synthesis (*Huangyutitham et al., 2013*). In comparison, the Pil-Chp chemosensory-like system initiates a hierarchical cascade of second messenger signaling in response to a surface (*Luo et al., 2015*). First, an increase in cellular cAMP levels occurs through activation of the adenylate cyclase CyaB by the Pil-Chp complex. This increases expression of genes involved in type IV pilus biogenesis, including PilY1. PilY1 is associated with the type IV pilus and harbors a Von Willebrand motif, which is involved in mechanosensing in eukaryotic systems (*Kuchma et al., 2010*). Thus, it has been proposed that this protein may be involved in the mechanosensing of surfaces (*Persat et al., 2015*). The output of this second signal is through the diguanylate cyclase, SadC, resulting in an increase in cellular c-di-GMP levels. Unlike the Wsp system, which localizes laterally along the cell (*O'Connor et al., 2012*), PilY1 is required to be associated with polarly-localized type IV pili in order to stimulate c-di-GMP production (*Luo et al., 2015; Kuchma et al., 2010*), suggesting that *P. aeruginosa* deploys both polar and laterally localized systems to promote c-di-GMP synthesis in response to a surface.

Bacteria in biofilms have long been appreciated to exhibit phenotypic heterogeneity due to chemical variation within the biofilm itself, including gradients of oxygen (*Wessel et al., 2014*), nutrients (*Schreiber et al., 2016*), and pH (*Vroom et al., 1999*). These environmental conditions are sensed by individual bacterial cells, leading to differential gene expression and metabolic activities even within a genetically homogeneous population (*Stewart and Franklin, 2008*). Specifically, the term 'division of labor' refers to cases where genetic or phenotypic heterogeneity results in subpopulations of cells cooperating to perform distinct tasks that provide an overall fitness benefit to the population (*West and Cooper, 2016*). Through task allocation, subpopulations of cells can engage in behaviors that are impossible to perform simultaneously (e.g. bet-hedging strategies between biofilm and planktonic cells [*Lowery et al., 2017*]), energetically costly to switch between (e.g. task-switching [*Goldsby et al., 2012*]), or are metabolically incompatible (e.g. photosynthesis and nitrogen fixation in cyanobacteria [*Rossetti et al., 2010*]). In particular, there is a rich body of literature demonstrating that genetic (*Kim et al., 2016; Dragoš et al., 2018; Klausen et al., 2003*) and phenotypic variation (*Klauck et al., 2018; Serra et al., 2015; Haagenzen et al., 2007*) in surface motility and polysaccharide production among individual bacterial cells within a biofilm can represent a division of labor that is required to achieve the architecture and structural integrity of the biofilm matrix (*van Gestel et al., 2015*).

Beyond heterogeneity as a result of variation in environmental signals, recent single-cell analyses have revealed that c-di-GMP signaling can drive phenotypic heterogeneity among populations of single cells exposed to the same environmental inputs. Planktonic *P. aeruginosa* have been shown to achieve heterogeneity among very low levels of c-di-GMP through asymmetrical partitioning of a diguanylate cyclase during cell division, leading to diverse swimming motility behaviors (*Kulasekara et al., 2013*). More recently, this same asymmetrical cell division mechanism was shown to generate two populations of *P. aeruginosa*, one piliated and one flagellated, that are each required for efficient tissue colonization (*Laventie et al., 2019*). Together, these studies support a role for c-di-GMP heterogeneity in generating diverse bacterial behaviors during both biofilm and planktonic growth.

Here, we examined the dynamics of c-di-GMP production and bacterial surface motility at the single-cell level during early stages of biofilm formation. We used a plasmid-based, transcriptional reporter of intracellular c-di-GMP to follow the downstream fate of cells producing varying levels of c-di-GMP in response to surface attachment. Within a clonal population of *P. aeruginosa*, we found that levels of c-di-GMP vary among individual cells as they sense a surface, leading to a division of labor between two energetically costly behaviors associated with early biofilm formation: surface exploration and polysaccharide production.

Results

Cellular c-di-GMP levels rapidly increase upon surface attachment

We initially compared levels of c-di-GMP between *P. aeruginosa* PAO1 cells growing attached to a silicone surface and subjected to constant flow for 4 hr to those grown planktonically for 4 hr. As expected, we observed that PAO1 cellular c-di-GMP levels are 4.4-fold higher (± 0.78 SD, $N = 3$,

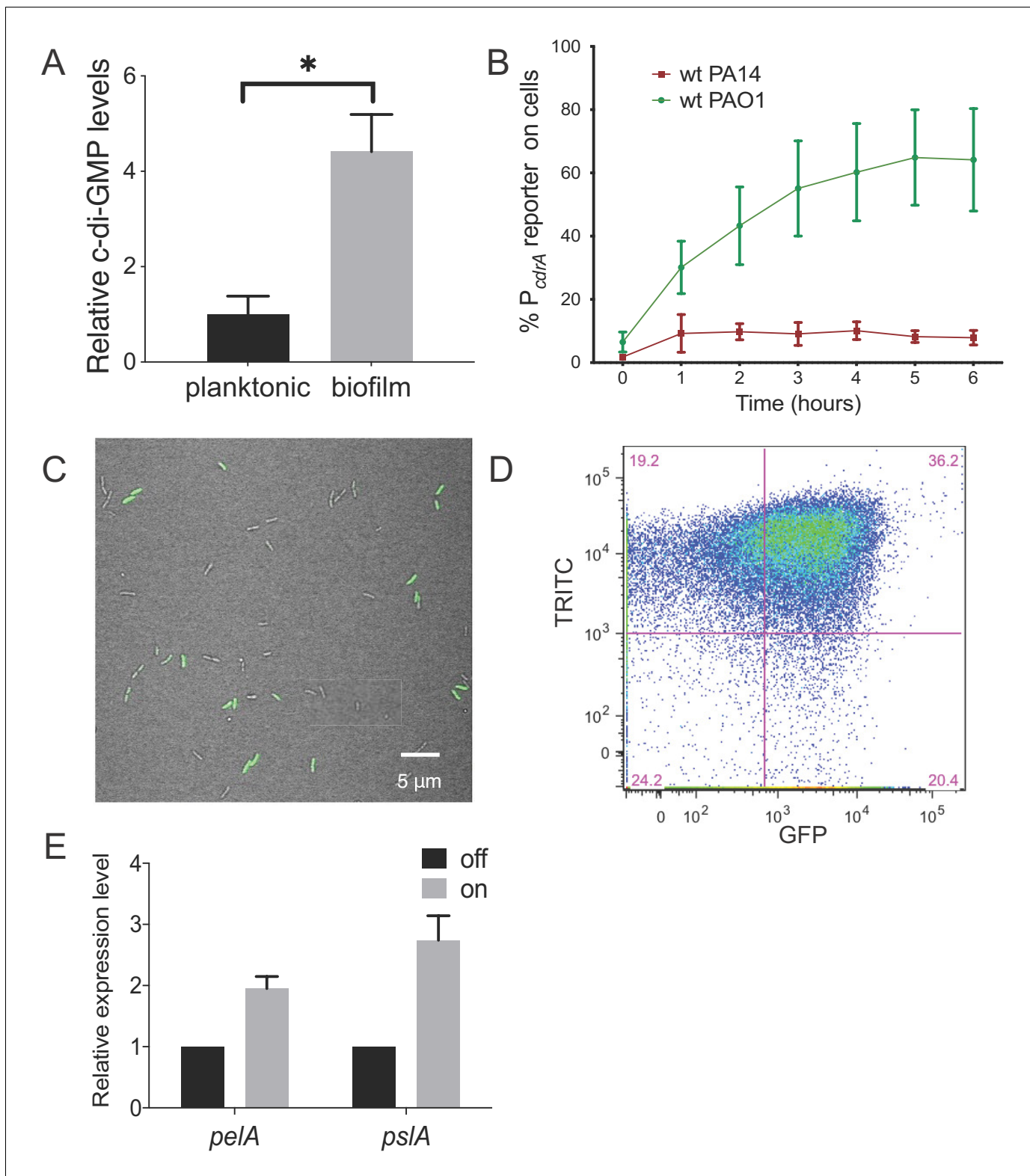


Figure 1. Heterogeneity in cellular levels of c-di-GMP during early *P. aeruginosa* biofilm formation. (A) c-di-GMP levels are elevated rapidly upon association of *P. aeruginosa* PAO1 cells with a surface. Relative levels of intracellular c-di-GMP in wild type PAO1 cells grown either planktonically or after 4 hr of attachment to a silicone tube. Values are normalized to the average concentration of c-di-GMP in planktonic cells, in pmol c-di-GMP/mg total protein as determined by LC-MS/MS, and presented as mean and SD. * $p < 0.05$ by T-test, $N = 3$. **Figure 1—figure supplement 1** shows the Pel polysaccharide operon is transcriptionally activated almost 10-fold compared to planktonic cells within 30 min of surface attachment. (B) Two commonly **Figure 1 continued on next page**

Figure 1 continued

studied *P. aeruginosa* lab strains, PAO1 and PA14, differentially activate the c-di-GMP reporter during surface sensing. Wild type PAO1 or PA14 cells harboring the c-di-GMP reporter (pP_{cdrA}::gfpASV) were grown to mid-log phase in planktonic culture, then inoculated into a flow cell and supplied with 1% LB medium. Surface attached cells were imaged immediately after inoculation (time 0 hr), and hourly for 12 hr. The c-di-GMP reporter is activated in a subset of wild type PAO1 cells within 1 hr of surface attachment and remains activated in approximately 60% of PAO1 cells during the first 6 hr of attachment. In PA14, the c-di-GMP reporter is activated in a smaller proportion of attached cells compared to PAO1. Data points are mean percentage of reporter activated cells from each time point across at least three biological replicates, with standard deviation. **Figure 1—figure supplement 4** shows an additional c-di-GMP responsive transcriptional reporter (using the *siaA* promoter) is also responsive to Wsp-dependent changes in cellular levels of c-di-GMP. **(C)** Wild type PAO1 cells display heterogeneity in c-di-GMP reporter activity after 6 hr of surface attachment. Confocal microscopy image of wild type PAO1 P_{cdrA}::gfpASV grown in 1% LB after 6 hr of surface attachment during a time course flow cell experiment. Bright field (gray) and GFP (green) channels are merged. Wild type PAO1 P_{cdrA}::gfpASV was grown in 1% LB and imaged by CSLM. **Figure 1—figure supplement 2** shows additional representative timecourse images of PAO1. **(D)** Psl exopolysaccharide production is enriched in the population of cells with high c-di-GMP. Representative scatterplot of reporter activity versus Psl lectin binding in wild type PAO1 harboring the pP_{cdrA}::gfpASV reporter grown for four hours in LB before surface attached cells were harvested, stained with the lectin, washed, and counted by flow cytometry. **(E)** Subpopulations of PAO1 cells with high and low c-di-GMP reporter activity are physiologically distinct. Cells with higher c-di-GMP reporter activity have increased expression of Pel and Psl biosynthetic machinery genes. After 4 hr of attachment to glass, wild type PAO1 cells were separated by flow-assisted cell sorting (FACS) into a population of cells with high (on) and low (off) c-di-GMP reporter activity, then qRT-PCR was performed to quantify expression of Pel and Psl exopolysaccharide biosynthesis genes. Levels of expression of Pel or Psl mRNA were normalized to the off population. *p<0.05 by T-test, N = 3 biological replicates. **Figure 1—figure supplement 3** shows controls for validating the protocol to monitor pP_{cdrA}::gfpASV by flow cytometry. **Figure 1—source data 2** shows by flow cytometry that Psl and Pel polysaccharide production is highest in cells with high pP_{cdrA}::gfpASV reporter activity.

DOI: <https://doi.org/10.7554/eLife.45084.003>

The following source data and figure supplements are available for figure 1:

Source data 1. Raw data for fluorescence intensities.

DOI: <https://doi.org/10.7554/eLife.45084.014>

Source data 2. Distributions of median fluorescence intensity values of PAO1 WT pP_{cdrA}::gfpASV cells obtained via bootstrap sampling.

DOI: <https://doi.org/10.7554/eLife.45084.005>

Figure supplement 1. The c-di-GMP-regulated promoter of the Pel polysaccharide operon is transcriptionally activated almost 10-fold compared to planktonic cells within 30 min of attachment of PAO1 to a silicone tube.

DOI: <https://doi.org/10.7554/eLife.45084.006>

Figure supplement 2. Representative time course images showing the c-di-GMP reporter (P_{cdrA}::gfpASV) transitioning from inactive upon initial attachment of wild type PAO1 (0 hr) to active in a subpopulation of cells between 1 and 12 hr during a flow cell experiment.

DOI: <https://doi.org/10.7554/eLife.45084.007>

Figure supplement 3. Fluorescence intensities of individual wild type PAO1 pP_{cdrA}::gfpASV cells imaged over 12 hr following surface attachment in a flow cell experiment (n ≥ 3 experiments per time point).

DOI: <https://doi.org/10.7554/eLife.45084.008>

Figure supplement 4. P-values for Kruskal-Wallis test for comparing medians observed in **Figure 1—figure supplement 3**, followed by Tukey's post-test for multiple comparisons.

DOI: <https://doi.org/10.7554/eLife.45084.004>

Figure supplement 5. Development of a protocol to monitor pP_{cdrA}::gfpASV using an LSRII flow cytometer.

DOI: <https://doi.org/10.7554/eLife.45084.009>

Figure supplement 6. The *siaA* promoter, regulated by c-di-GMP binding to FleQ, is responsive to Wsp-dependent changes in cellular levels of c-di-GMP.

DOI: <https://doi.org/10.7554/eLife.45084.010>

Figure supplement 7. Psl and Pel polysaccharide production is highest in cells with high c-di-GMP as measured by the pP_{cdrA}::gfpASV reporter.

DOI: <https://doi.org/10.7554/eLife.45084.011>

Figure supplement 8. Development of a protocol to sort biofilm cells by pP_{cdrA}::gfpASV using flow assisted cell sorting (FACS).

DOI: <https://doi.org/10.7554/eLife.45084.012>

Figure supplement 9. Flow cytometry control depicting the specificity of TRITC-HHA lectin for Psl.

DOI: <https://doi.org/10.7554/eLife.45084.013>

p≤0.05) after 4 hr of growth attached to a surface compared to planktonic growth (**Figure 1A**). Because direct measurement of c-di-GMP by LC-MS/MS is limited by our ability to generate enough biomass at earlier time points, we used qRT-PCR to monitor *pel* transcript levels as a readout of c-di-GMP. We found that after just 30 min of surface attachment, *pelA* transcript levels had increased almost 10-fold compared to planktonically grown cells (**Figure 1—figure supplement 1**). This is consistent with previously published literature showing that transcription of the *pel* operon is

directly and positively controlled by high cellular levels of c-di-GMP (*Hickman and Harwood, 2008; Baraquet et al., 2012*).

The $P_{cdrA}::gfp$ reporter suggests heterogeneity in c-di-GMP levels during surface sensing

Next, we sought to visualize early c-di-GMP signaling events at the single cell level. To this end we used a plasmid-based, c-di-GMP responsive transcriptional reporter, $pP_{cdrA}::gfpASV$ (*Rybtke et al., 2012*) in two commonly-studied *P. aeruginosa* strains, PAO1 and PA14. Planktonic cells (a condition where the reporter is inactive due to low c-di-GMP levels) were used to inoculate flow cell chambers. We imaged individual cells of each reporter strain hourly for up to 6 hr after surface attachment (*Figure 1B* and *Figure 1—figure supplement 2*). As expected, we saw minimal GFP fluorescence at the 0 hr time point (right after surface attachment). However, by 1 hr, the reporter was activated in a subset of surface attached cells, as defined by background subtracted GFP fluorescence ≥ 321 fluorescence units (referred to as reporter ‘on’ subpopulations). Interestingly, between 4 and 6 hr post inoculation, we consistently observed that the c-di-GMP reporter was only active in a subset of cells in both strains (*Figure 1C*). In PA14, the reporter was activated in 10% of the population over 6 hr, whereas PAO1 displayed greater reporter activity, with 40–60% of the cells displaying reporter activity through 12 hr (*Figure 1B*). An analysis of single cell fluorescence supported these observations. We plotted each cell’s individual fluorescence values over time and observed populations of cells with low and high c-di-GMP reporter activity at each timepoint after the 0 hr (*Figure 1—figure supplement 3*). We also observed long tails of high GFP fluorescence, particularly at 2 and 4 hr. This suggests that the reporter ‘on’ subpopulation represents cells with a range of high c-di-GMP levels. This wide range of high reporter activity cells between 2 and 4 hr could be indicative of an early ‘spike’ in c-di-GMP production that levels off over time (*Figure 1—figure supplement 3*).

We next examined the change in distribution of fluorescence intensity over time for single cells by bootstrap sampling of the single cell fluorescence intensity values (*Figure 1—figure supplement 4*). The purpose of this bootstrapping analysis is to examine whether the distributions of fluorescence intensity differ at each time point. We found that the median fluorescence intensity was significantly different at every time point except between 4 and 6 hr, and between 8, 10, and 12 hr (*Figure 1—source data 2*). Together, these single cell analyses support a model in which c-di-GMP signaling is initiated rapidly upon surface attachment in the first 4–6 hr for a subpopulation of attached cells, while the rate of c-di-GMP increase tends to level off at later timepoints.

We confirmed the microscopy results from comparing PAO1 and PA14 reporter fluorescence using flow cytometry to assess the proportion of attached cells that were fluorescent (*Figure 1—figure supplement 5D, E*). To be sure that the promoter of *cdrA* is representative of c-di-GMP-regulated gene expression, we replaced P_{cdrA} with the promoter of *siaA*, a gene that is also highly expressed under conditions of elevated c-di-GMP (*Starkey et al., 2009; Baraquet and Harwood, 2016*). We found that $pP_{siaA}::gfpASV$ reporter activity resembled that of $pP_{cdrA}::gfpASV$ in response to a surface (*Figure 1—figure supplement 6*). Thus, reporter activity is indeed linked to cellular levels of c-di-GMP.

Cyclic di-GMP heterogeneity leads to phenotypic diversification at early stages of biofilm formation

We then wanted to confirm that subpopulations of surface-attached *P. aeruginosa* cells with high and low c-di-GMP reporter activity are truly physiologically distinct from one another. We used TRITC-labeled lectins to stain for two c-di-GMP-induced exopolysaccharides, Psl and Pel (*Zhao et al., 2013; Jennings et al., 2015*), the presence of which is indicative of biofilm formation by PAO1 and PA14, respectively. After 4 hr of attachment to glass, we observed an enrichment of TRITC-conjugated lectin staining in the population of cells with high c-di-GMP reporter activity (*Figure 1D* and *Figure 1—figure supplement 7*), demonstrating that the subpopulation of cells with high c-di-GMP is producing more exopolysaccharide than their low c-di-GMP counterparts. This correlation was weaker for Psl than Pel, probably due to the fact planktonic populations can make low levels of Psl (though not the case for Pel) (*Wei and Ma, 2013*). As a complementary approach, we separated 4 hr surface-grown cells of the reporter strain into reporter ‘on’ and ‘off’ subpopulations using flow-assisted cell sorting (FACS; *Figure 1—figure supplements 8 and 9*). We then applied

qRT-PCR to compare *Pel* and *Psl* transcript levels in these two populations. Both the *pel* and *psl* operon transcripts were elevated in the reporter 'on' subpopulation, relative to the reporter 'off' subpopulation (**Figure 1E**). These data support that, with respect to c-di-GMP signaling, there are at least two distinct subpopulations that arise shortly after surface attachment.

The Wsp system is required for surface sensing

We next evaluated the relative contributions of the Wsp and Pil-Chp surface sensing systems to surface-induced c-di-GMP production. Strains with mutations in the Pil-Chp chemosensory system were not significantly defective in surface sensing activity. Deletion of the diguanylate cyclase activated through the Pil-Chp system (PAO1 $\Delta sadC$) and the gene encoding the putative sensor PilY1 (PAO1 $\Delta pilY1$) did not significantly influence reporter activity in response to a surface (**Figure 2—figure supplement 1A,B**). Whereas both the *SadC* and *PilY1* mutants displayed wild type levels of reporter activity, a mutant lacking the main Type IV pilus filament protein (PAO1 $\Delta pilA$) did show a statistically significant defect in reporter activity by 6 hr (**Figure 2—figure supplement 1B**; $p < 0.05$ by T-test). We then mutated the c-di-GMP cyclase gene, *wspR*, to inactivate the Wsp system. In addition, we deleted the gene encoding the methylesterase *wspF*, which locks the system into the active state, regardless of whether cells are surface-associated. We found that the PAO1 $\Delta wspR$ strain exhibited extremely low levels of reporter activity during the first 6 hr after surface attachment (**Figure 2A** and **Figure 2—figure supplement 2**). Complementation of PAO1 $\Delta wspR$ restored wild type levels of activity at all time points (**Figure 2—figure supplement 3**). As expected, PAO1 $\Delta wspF$ had a high proportion of reporter active cells (**Figure 2A**). We repeated these experiments in the lab strain PA14 and saw a similar trend for Wsp mutants (**Figure 2—figure supplement 4**). Kulesekara et al. (**Kulasekara et al., 2013**) used a FRET-based c-di-GMP reporter to show that planktonic *P. aeruginosa* has heterogeneous, albeit very low, concentrations of c-di-GMP which it achieves through asymmetrical partitioning of a phosphodiesterase (PA5017, also called *Pch* or *DipA*) to the flagellated cell pole during cell division. However, we found no evidence of heterogeneity in c-di-GMP within the planktonic cell inoculum contributing to our observations. We ruled out the phosphodiesterase (PA5017) as responsible for the heterogeneity we see during surface sensing by examining a deletion mutant of PA5017 and showing that this strain still exhibited heterogeneity during surface sensing (**Figure 2—figure supplement 5**).

Since the Pil-Chp surface sensing apparatus is polarly localized and the Wsp system is localized laterally along the length of the cell body, we examined whether reporter activity correlated with polar versus lateral attachment to the surface. We found that reporter activity was very low in polarly attached cells, while cells attached along the entire length of the cell body displayed a higher proportion of reporter-activated cells at that same time point (**Figure 2B**). (Our analysis does not account for the time period each cell spends in either the polarly or non-polarly attached states.) This finding is consistent with the localization of the Wsp system and its role for early c-di-GMP signaling during surface sensing.

Heterogeneity in c-di-GMP levels among cells correlates with Wsp system activity

The specific activity of purified WspR increases as a function of WspR concentration when the protein is treated with beryllium fluoride to mimic phosphorylation, supporting the idea that formation of subcellular clusters of WspR-P potentiates its diguanylate cyclase activity and leads to elevated c-di-GMP (**Huangyutitham et al., 2013**). Fewer than 1% of wild-type cells grown in broth have a visible WspR-YFP cluster. However, after a short period of growth on an agar surface, WspR-YFP clusters were visible in 30–40% of wild type PAO1 cells, and this is dependent on sensing by the membrane-bound protein WspA, which is laterally distributed in cells (**Kuchma et al., 2010**). To directly link WspR cluster formation with diguanylate cyclase activity at the cellular level and with surface sensing, we constructed a version of the c-di-GMP reporter that expresses mTFP1 instead of GFP ($pP_{cdrA}::mTFP1$) to avoid the issue of spectral overlap with WspR-YFP. We monitored reporter activity in two point mutants of WspR (L170D and E253A) that are driven by an inducible promoter, translationally fused to eYFP, and have been previously shown to form large subcellular WspR clusters in a higher percentage of cells than wild-type WspR. The WspR[L170D] protein is highly active for c-di-GMP production, and it forms subcellular clusters in about 75% of agar surface-grown cells.

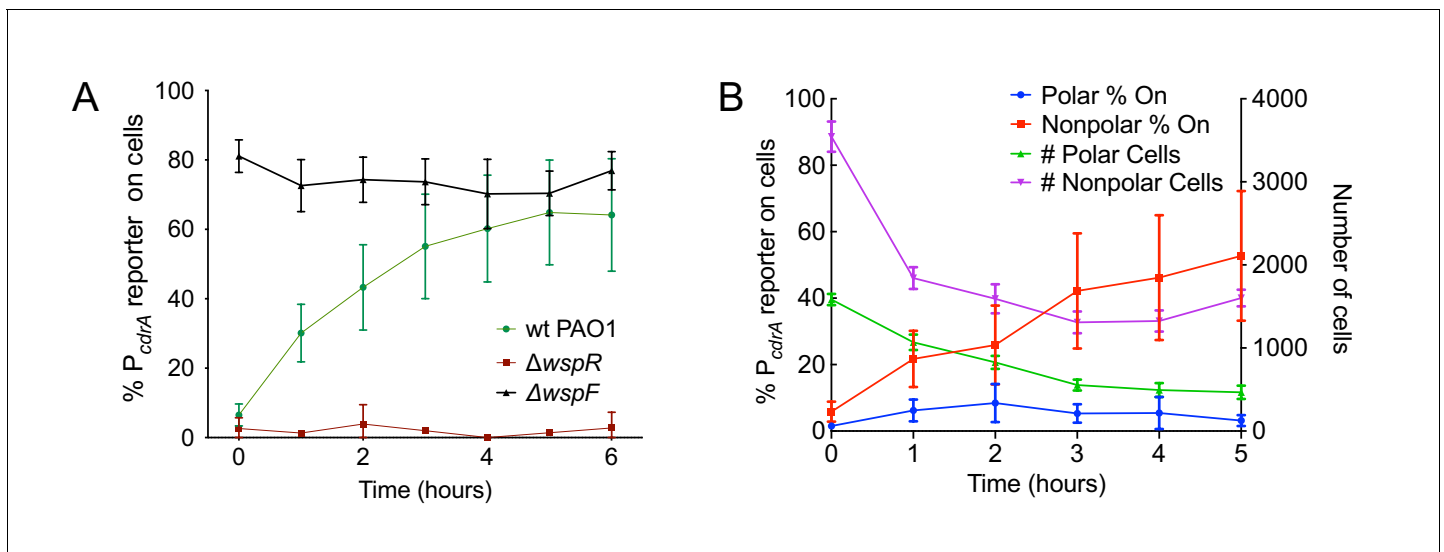


Figure 2. The Wsp system generates heterogeneity in cellular levels of c-di-GMP during early *P. aeruginosa* biofilm formation. (A) The Wsp system is required for activation of the $pP_{cdrA}::gfpASV$ reporter during surface sensing. Six hour time course plot of the average percentage of surface-attached cells from either wild type PAO1 (green), PAO1 $\Delta wspR$ (red), or PAO1 $\Delta wspF$ (black) in which the $pP_{cdrA}::gfpASV$ reporter had turned 'on' at each hour. Cells were identified as 'on' if their average GFP fluorescence was greater than twice the average background GFP fluorescence of the image. Error bars = standard deviation. $N \geq 3$ biological replicates. See **Figure 2—figure supplement 1** for the same timecourse using mutants in the Pil-Chp surface sensing system. **Figure 2—figure supplement 2** shows representative images from **Figure 2A**. **Figure 2—figure supplement 3** shows that complementing the *wspR* mutant restores wild type levels of reporter activity. **Figure 2—figure supplement 4** shows that the lab strain PA14 also displays Wsp-dependent c-di-GMP heterogeneity. **Figure 2—figure supplement 5** shows that PA5017 (*Pch*) is not required for heterogeneity during surface sensing. (B) Laterally attached cells have higher c-di-GMP levels than polarly attached cells. Five hour time course plot depicting, on the left axis, the percentage of $pP_{cdrA}::gfpASV$ reporter 'on' cells that were either polarly (blue) or laterally (red) attached to the surface of a glass coverslip in a flow cell at each hour. The right axis depicts the total number of polar (green) and laterally attached (purple) cells at each time point. Cells were identified as $pP_{cdrA}::gfpASV$ reporter 'on' if their average GFP fluorescence was greater than 321 fluorescence units. Error bars = standard deviation. $N = 4$ biological replicates.

DOI: <https://doi.org/10.7554/eLife.45084.015>

The following figure supplements are available for figure 2:

Figure supplement 1. Mutants predicted to inactivate the Pil-Chp surface sensing system largely retain $pP_{cdrA}::gfpASV$ reporter activity during the first six hours of surface sensing.

DOI: <https://doi.org/10.7554/eLife.45084.016>

Figure supplement 2. The $pP_{cdrA}::gfpASV$ reporter is sensitive to Wsp-dependent variation in c-di-GMP during surface sensing.

DOI: <https://doi.org/10.7554/eLife.45084.017>

Figure supplement 3. Complemented diguanylate cyclase mutants display wild type levels of $pP_{cdrA}::gfpASV$ reporter activity.

DOI: <https://doi.org/10.7554/eLife.45084.018>

Figure supplement 4. Activity of the $pP_{cdrA}::gfpASV$ reporter in strain PA14 is dependent on the Wsp system.

DOI: <https://doi.org/10.7554/eLife.45084.019>

Figure supplement 5. A mutation in PA5017 (also referred to as *pch* or *dipA*) does not impact the heterogeneity observed in surface sensing.

DOI: <https://doi.org/10.7554/eLife.45084.020>

A *WspR*[E253A] point mutation abolishes diguanylate cyclase activity, but this protein still forms clusters in about 70% of surface-grown cells (*Huangyutitham et al., 2013*). As expected, in the presence of inducer, we observed a large increase in c-di-GMP reporter activity in *WspR*[L170D], but not *WspR*[E253A] (**Figure 3A,B**). We then asked whether the heterogeneity in reporter activity in response to surface attachment correlates with *WspR* clustering in the *WspR*[L170D] strain. We found that $pP_{cdrA}::mTFP1$ activity was significantly higher in cells with at least one subcellular *WspR*-eYFP focus in the *WspR*[L170D] strain compared to cells without a *WspR*-eYFP focus (**Figure 3C** and **Figure 3—figure supplement 1**). These data indicate that the heterogeneity observed in c-di-GMP signaling after surface attachment is due to the heterogeneity in the activity of the Wsp system, as reflected by subcellular clustering of active *WspR*-P.

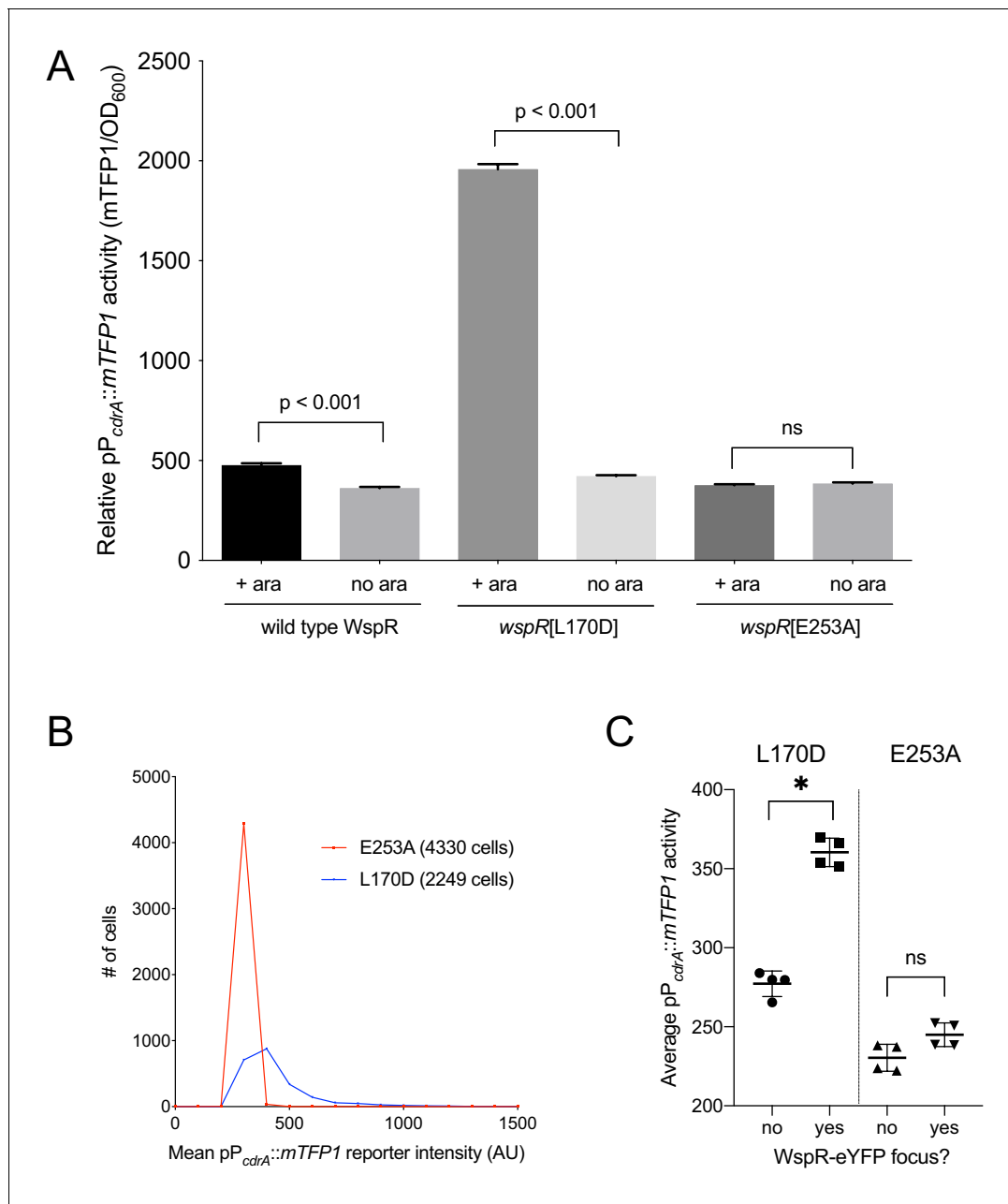


Figure 3. Activity of the pP_{cdrA}::mTFP1 reporter is dependent on the ability of WspR to produce c-di-GMP. (A) The pP_{cdrA}::mTFP1 reporter is active in surface grown cells with functional, arabinose-inducible alleles of WspR when arabinose is added to the media. Wild type WspR represents the strain PAO1 Δ wspR attCTX::wspR-eYFP. *wspR*[L170D] represents the strain PAO1 Δ wspR attCTX::wspR[L170D]-eYFP, which produces large subcellular clusters of WspR and grows as rugose small colonies on LB with 1% arabinose, a phenotype that is indicative of high intracellular c-di-GMP. *wspR*[E253A] represents the strain PAO1 Δ wspR attCTX::wspR[L170D]-eYFP cells, which forms large subcellular WspR clusters, but does not produce c-di-GMP via WspR due to the point mutation located in its active site. Cells were grown on LB agar plates with 100 μ g/mL gentamicin, and in the presence or absence of 1% arabinose. Cells were resuspended in PBS and mTFP1 fluorescence and OD₆₀₀ were measured. Relative pP_{cdrA}::mTFP1 reporter activity is the level of mTFP1 fluorescence normalized to OD₆₀₀. Asterisk indicates statistical significance by Student's t-test ($p < 0.001$) in six technical replicates. Error bars = standard deviation. (B) The pP_{cdrA}::mTFP1 reporter displays heterogeneity in a strain with a functional WspR (*wspR*[L170D]) and is consistently dark in a strain with inactive WspR. Histogram displaying the distribution of average cellular levels of mTFP1 fluorescence from expression of the pP_{cdrA}::mTFP1 reporter in either the PAO1 Δ wspR attCTX::wspR[L170D]-eYFP (blue) or PAO1 Δ wspR attCTX::wspR[E253A]-eYFP (red) backgrounds. (C) Cells with subcellular clusters of functional WspR have higher levels of c-di-GMP reporter activity than cells without a WspR focus or cells without a functional WspR. L170D refers to a point mutant in WspR that forms large subcellular clusters and retains diguanylate cyclase activity: WspR[L170D]-eYFP in the strain PAO1 Δ wspR attCTX::wspR[L170D]-eYFP. E253A refers to a point mutant in WspR that forms large subcellular clusters but does not retain diguanylate cyclase activity: WspR[E253A]-eYFP in the strain PAO1 Δ wspR attCTX::wspR[E253A]-eYFP. Asterisk indicates statistical significance. *Figure 3 continued on next page*

Figure 3 continued

significance by Student's t-test ($p < 0.001$) in four replicates, n.s. = not significant. Error bars = standard deviation. See **Figure 3—figure supplement 1** for a representative image of WspR[L170D]-eYFP foci.

DOI: <https://doi.org/10.7554/eLife.45084.021>

The following source data and figure supplement are available for figure 3:

Source data 1. Figure 3B source data.

DOI: <https://doi.org/10.7554/eLife.45084.023>

Figure supplement 1. A representative image depicting subcellular foci of WspR[L170D]-eYFP in cells with high $pP_{cdrA::mTFP1}$ reporter activity.

DOI: <https://doi.org/10.7554/eLife.45084.022>

For phenotypic heterogeneity to represent a division of labor, it must result in a fitness benefit to the population. Therefore, we next asked whether the observed heterogeneity in c-di-GMP signaling in response to a surface has a meaningful influence on biofilm formation. This was particularly important since previously published results indicated that a *wspR* mutation had only a small impact on biofilm production (Kulasakara et al., 2006). However, these studies assessed biofilm formation at later stages of biofilm growth that were well beyond initial surface attachment. Therefore, we chose to compare a *wspR* mutant to wild type at earlier biofilm stages. We performed *in vitro* biofilm assays and observed that a PAO1 Δ *wspR* mutant was defective for biofilm formation relative to wild type PAO1 at 2, 4, and 6 hr post-attachment (Figure 4A). However, at later stages of development (24 hr), the *wspR* mutant caught up and produced similar amounts of biofilm biomass relative to wild type. Complementation of the Δ *wspR* strain *in trans* restored wild type levels of biofilm formation at all time points (Figure 4A). These data suggest that the Wsp system rapidly responds to surface contact to generate elevated levels of c-di-GMP in a subpopulation of cells, which accelerates biofilm production. Given the importance of c-di-GMP signaling in biofilm production, the fact that the Δ *wspR* strain can ultimately attain wild-type levels of biofilm biomass suggests that one of the many other known c-di-GMP cyclases present in *P. aeruginosa* may ultimately compensate for c-di-GMP production in the absence of WspR.

Cyclic di-GMP heterogeneity leads to diversification in surface exploration at the lineage level

We hypothesized that heterogeneity in c-di-GMP signaling dictated by the Wsp complex could impact the surface behavior of the two observed subpopulations. We predicted that the subpopulation of cells with high c-di-GMP after surface attachment would produce biofilm matrix exopolysaccharides and contribute to initial microcolony formation, while the cells with low c-di-GMP would exhibit increased surface motility and detachment, which is known to be inhibited by exopolysaccharide production. To test this hypothesis, we tracked both reporter activity and surface behavior for cells within a single field of view for 40 hr. From our single-cell tracking data, we generated family trees across at least four generations of cells, using a previously described technique (Lee et al., 2018). We tracked the time-averaged $P_{cdrA::gfpASV}$ reporter activity ($I_{c-di-GMP}$), surface motility behavior (F_{motile} , defined as the fraction of time that cells are motile), and detachment behavior (tree asymmetry λ , where $\lambda = 0$ represents both daughter cells remaining attached to the surface and $\lambda = 1$ represents when one daughter cell detaches or travels outside the field of view).

In *P. aeruginosa*, surface exploration is mainly accomplished by type IV pili-mediated twitching motility, and does not appear to be influenced by levels of intracellular c-di-GMP when analyzing single cells (Ribbe et al., 2017). Interestingly, when analyzing correlations between c-di-GMP and motility for entire lineages in family trees, we found clear inverse correlations between $I_{c-di-GMP}$ and F_{motile} (Figure 4B, $\rho = -0.53$, $p = 0.0012$) and between $I_{c-di-GMP}$ and λ (Figure 4C, $\rho = -0.45$, $p = 0.0068$), suggesting that c-di-GMP levels are strongly inversely correlated with surface motility behavior and detachment behavior over multiple generation of cells. Analyzing these correlations across multiple cell divisions is important because it tells us when and how bacteria respond to c-di-GMP signaling events in terms of their surface motility and detachment. When looking at correlations between c-di-GMP and surface motility, the time between seeing a signaling event (i.e., rise or drop in reporter fluorescence intensity) and seeing a response (i.e., change in motility) can span a broad range (from a few minutes to well over a cell division time). Using a cell's entire lineage history

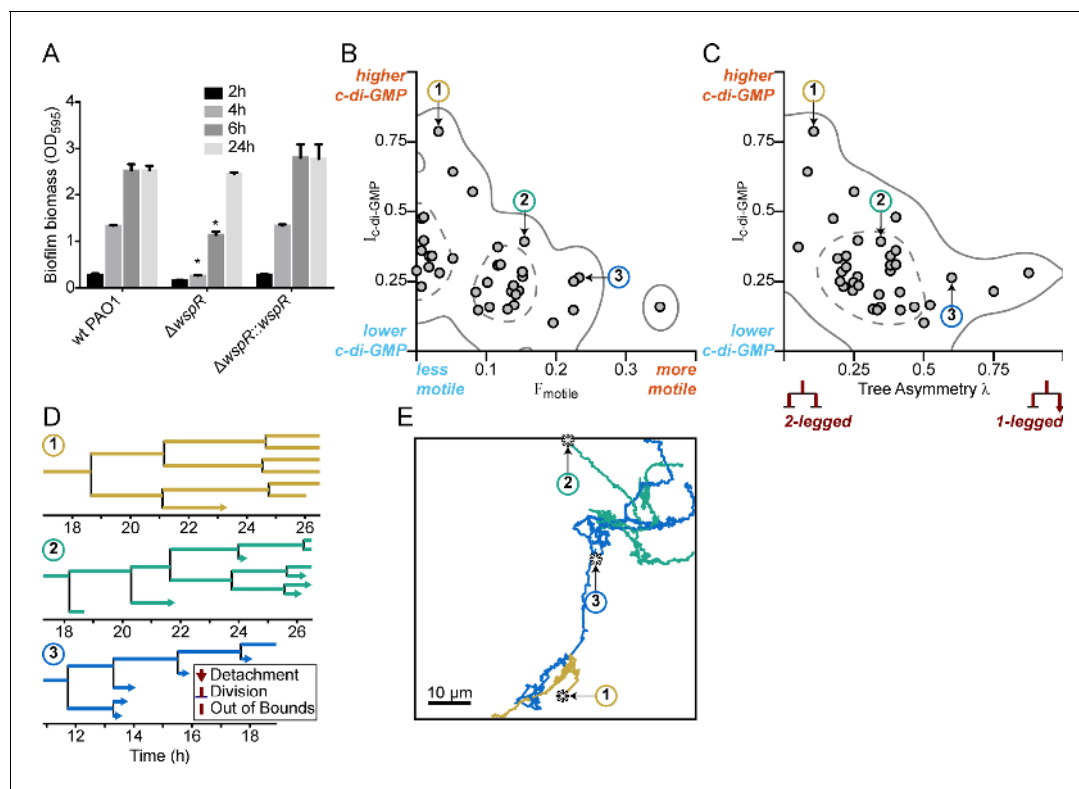


Figure 4. Multigenerational c-di-GMP levels within populations of surface-attached wild type PAO1 cells are inversely correlated with surface motility and detachment. (A) The Wsp surface sensing system is involved in the early stages of biofilm formation in PAO1. Static biofilm assay performed in wild type PAO1, a single deletion mutant of *wspR*, and the PAO1 Δ *wspR* mutant complemented with *wspR*. Between 4 and 6 hr, PAO1 Δ *wspR* shows a defect in surface attachment and biofilm formation relative to the wild type. However, after 24 hr, PAO1 Δ *wspR* formed equal biofilm biomass compared to wild type. Plotted values are the mean of 6 technical replicates and error is standard deviation. Asterisk indicates a statistically significant change in biomass relative to wild type PAO1 at each time point (Student's t test; $p < 0.05$). (B) Plot of $I_{c\text{-di-GMP}}$ vs F_{motile} for individual wild type PAO1 families. $I_{c\text{-di-GMP}}$ is the relative normalized c-di-GMP reporter intensity averaged across all members of a family. F_{motile} is the fraction of time that cells in a family are motile (specifically surface translational motility). Each circle represents an individual family ($N = 35$) with at least four tracked generations. Solid lines represent the 95% probability bounds and dashed lines represent the 50% probability bounds, calculated via kernel density estimation. Spearman correlation: $\rho = -0.53$, $p = 0.0012$. (C) Plot of $I_{c\text{-di-GMP}}$ vs tree asymmetry λ for individual wild type PAO1 families. Colored numbers indicate the same three families from (B) and (D). Tree asymmetry λ quantifies the detachment behavior of family trees as follows. $\lambda = 0$ corresponds to ideal trees with purely 'two-legged' division-branching, when both daughter cells remain attached to the surface. $\lambda = 1$ corresponds to ideal trees with purely 'one-legged' division-branching when one daughter cell detaches or travels outside the field of view. Points here are the same families as in (B). Solid lines represent the 95% probability bounds and dashed lines represent the 50% probability bounds, calculated via kernel density estimation. Spearman correlation: $\rho = -0.45$, $p = 0.0068$. (D) Family trees of the same three representative wild type PAO1 families indicated in (B) and (C). Time 0 hr is the start of the dataset recording. Lengths of horizontal lines on the plots are proportional to time spent in each generation. Horizontal lines that end with arrows are detachment events, lines that intersect with a vertical line are division events, and lines that end without a marker are out-of-bound events where we lose track of the bacterium (moving out of the field of view or reaching the end of the recording; represented as moving outside the XYT limits of the dataset boundaries). Vertical lines are arbitrarily spaced to show all the descendants. Colors represent the families in (B) and (C). (E) Spatial trajectories of the three representative families. Asterisks (*) represent the initial location of the founder cell. Scale bar 10 μm . The families are color coded as in the previous panels.

DOI: <https://doi.org/10.7554/eLife.45084.024>

The following source data and figure supplement are available for figure 4:

Source data 1. MATLAB data for WT and Δ *wspR* families.

DOI: <https://doi.org/10.7554/eLife.45084.026>

Figure supplement 1. The Δ *wspR* mutant does not have correlations between c-di-GMP, surface motility, and detachment.

DOI: <https://doi.org/10.7554/eLife.45084.025>

(i.e., tracking daughter cells across cell divisions) will capture all of these events, whereas using single cell history (i.e., within one generation) will only capture a portion of them. For example, this lineage-level tracking of the influence of signaling events on bacterial behavior was recently shown for correlations between cyclic AMP signaling and *P. aeruginosa* surface motility (Lee et al., 2018). In this study, Lee et al. found that correlations between cell motility and signaling activity were stronger when they took into account lineage history, rather than using single cell history. However, if there are enough instances where the time between a signaling event and a cell's corresponding behavioral response are within a single generation, then correlations can still be found when using single cell history. For wild type PAO1 (WT), we observe weaker correlations when looking at individual cells in these lineages. $I_{c\text{-di-GMP}}$ vs F_{motile} for single cells had a Spearman correlation value $\rho = -0.40$ ($p < 0.0001$), which is smaller than the lineage-level correlation value, suggesting that lineage-level correlations are stronger.

To illustrate these correlations, we chose three representative families, with either high, intermediate, or low $I_{c\text{-di-GMP}}$ and plotted their family trees (Figure 4D) and spatial trajectories (Figure 4E). Families with the highest $I_{c\text{-di-GMP}}$ had the lowest F_{motile} and λ (Family 1, Figure 4B–E). In these families, daughter cells remained attached following cell division, exhibited continuously elevated c-di-GMP, did not move appreciable distances on the surface, and ultimately produced small microcolonies. In contrast, families of cells with low $I_{c\text{-di-GMP}}$ had the highest F_{motile} and λ . For these families, daughter cells frequently detached or traveled outside the field of view, had lower c-di-GMP levels, traveled larger distances on the surface, and ultimately did not form microcolonies (Family 3, Figure 4B–E).

Tracking and lineage analyses were also performed on a PAO1 mutant with the Wsp system inactivated (PAO1 ΔwspR). We observed that the range of tree asymmetry values is like that of WT and that the ΔwspR mutant eventually reaches WT c-di-GMP levels despite initially being lower, which is consistent with the observation that the mutant eventually forms WT-like biofilms. What our analysis revealed about the ΔwspR strain was quite unexpected: The WspR mutant has overall lower surface motility and, importantly, lacks correlations between c-di-GMP, surface motility, and detachment behavior (both at the level of lineages and at the level of single cells). For single cell surface motility, 23% of ΔwspR mutant cells (48 of 210 cells) have non-zero F_{motile} (the metric for surface motility) compared to 44% of WT cells (251 of 565 cells; Figure 4—figure supplement 1). The overall lowered surface motility and the lack of correlations between c-di-GMP and surface motility in the ΔwspR mutant suggest that the Wsp system is involved in translating the heterogeneous c-di-GMP signaling events into the corresponding motility-related responses for cells and their progeny. Therefore, the Wsp system's multi-generational temporal propagation of surface sensing signaling and behavior is important for the generation of heterogeneous populations of surface motile and immotile cells during early biofilm formation.

Use of an optogenetic reporter to control bacterial surface behavior at the single-cell level

One important question is what happens to early biofilm development if we were to effectively remove heterogeneity in c-di-GMP output rooted in the WspR surface sensing system. To address this question, we used a strain in which c-di-GMP production could be easily controlled using an optogenetic system. The precise control of c-di-GMP expression in individual cells was made possible by the use of a chimeric protein that fused a diguanylate cyclase domain to a bacteriophytochrome domain. Flow chambers were seeded with the optogenetic strain encoding a heme oxygenase (*bphO*) and light-



Video 1. Single cells are precisely illuminated by ATIM via *in situ* analysis and tracking of bacteria. The left panel shows the merged images of *gfpASV* and mCherry fluorescence microscopy images over time. The right panel shows the merged images of red LED projected patterns and bright field images corresponding to the left panel. The fluorescence intensity of *gfpASV* in the illuminated cells and their offspring (colored red in right panel) is significantly increased after using ATI for 460 mins. In contrast, the *gfpASV* fluorescence intensity of the un-illuminated cells remains low and these cells remain motile.

DOI: <https://doi.org/10.7554/eLife.45084.032>

responsive diguanylate cyclase (*bphS*) (Ryu and Gomelsky, 2014). We initially characterized the optogenetic strain and verified the reporter activity increased with exposure of the optogenetic strain to red light (Figure 5—figure supplement 1) and that the laser light did not impact growth or motility (data not shown). Following validation of the strain, cells inoculated on a glass surface were tracked and continuously stimulated with red-light over ~8 hr using adaptive tracking illumination microscopy (ATIM), which allows for precise stimulation of the initial attached cells and their offspring and ensures sustained intracellular c-di-GMP production for a fixed number of surface cell generations (Figure 5—figure supplement 2). Cellular lineages (a cell and all of its offspring) and c-di-GMP reporter activity were continually monitored for at least 12 hr. Families that were not stimulated with light demonstrated a heterogeneous surface response (Video 1 and Figure 5B,D) similar to that of Families 1–3 in Figure 4B–E. Some lineages were dominated by surface explorers, whereas others were seen to commit to microcolony formation. In contrast, in families stimulated with light for more than one generation, the resulting c-di-GMP production artificially forced lineages to have low surface motility and commit to microcolony production (Video 1 and Figure 5A,C) similar to that of Family one in Figure 4B–E. Families stimulated with light in this manner had higher $I_{c\text{-di-GMP}}$ and lower λ values than those that were not stimulated (Figure 5—figure supplement 3). We also found that optogenetic control of c-di-GMP results in phenotypes that are consistent with the wild-type behavior presented in Figure 4, with illuminated cells (high c-di-GMP) displaying the least motility and control (non-illuminated) cells displaying comparatively greater surface motility (Figure 5—figure supplement 3). Interestingly, families stimulated with light for one generation or less are not significantly different from un-illuminated controls (data not shown). Our data show that the generation of c-di-GMP can deterministically lead to the creation of an entire lineage of sessile cells with post-division surface persistence, low motility, and initiation of microcolony formation. Altogether, these results show that c-di-GMP levels, surface motility, and detachment are inversely correlated at the lineage level, and that the time scale for this occurs over multiple generations.

Discussion

Collectively, our data show that heterogeneity in cellular levels of c-di-GMP generated by the Wsp system in response to surface sensing, leads to two physiologically distinct subpopulations that each contribute to surface colonization. Phenotypic heterogeneity of single cells is a common phenomenon in bacteria that can be beneficial at the population level by allowing a single genotype to survive sudden environmental changes. Sources of phenotypic heterogeneity among genetically homogeneous populations include bistability (Dubnau and Losick, 2006) and stochasticity (Elowitz et al., 2002) of gene expression, unequal partitioning of proteins during cell division due to low abundance (Elowitz et al., 2002), epigenetic modifications resulting in phase variation (Casadesús and Low, 2006), or through asymmetrical cell division (Kulasekara et al., 2013; Laventie et al., 2019). In this study, we show that the Wsp system generates heterogeneity in c-di-GMP signaling, and it is never fully activated in 100% of wild-type, surface-attached cells. One possible outcome of phenotypic heterogeneity is a division of labor between costly behaviors that support the growth and survival of the population (Ackermann, 2015). We found that abolishing c-di-GMP heterogeneity through inactivation of WspR leads to defects in early biofilm formation. This suggests that the subpopulations of high c-di-GMP, polysaccharide producers and low c-di-GMP, surface explorers are both required for efficient biofilm formation and that they represent a division of labor during early biofilm formation.

The data show that Wsp-generated c-di-GMP heterogeneity results in phenotypic changes for entire family lineages of cells. It is interesting that correlations between c-di-GMP, surface motility, and surface detachment probability are stronger when considered for an entire lineage in a bacterial family tree, but weaker when considered at the individual cell level. We think that this difference in the strength of correlations between c-di-GMP and bacterial behavior at the lineage versus single cell level is biologically meaningful and likely reflects the complex relationship between c-di-GMP signaling and type IV pili-mediated motility (Ribbe et al., 2017; Jain et al., 2012; Jain et al., 2017). The time between initiation of a signaling event (i.e., increased intracellular c-di-GMP) and the associated response (i.e., attenuation of motility or initiation of polysaccharide production) can span a large range, depending on the behavior. When examining polysaccharide production, signal propagation appears to be quick, with high c-di-GMP cells initiating exopolysaccharide production within minutes of *P. aeruginosa* encountering a surface. However, when examining a different bacterial

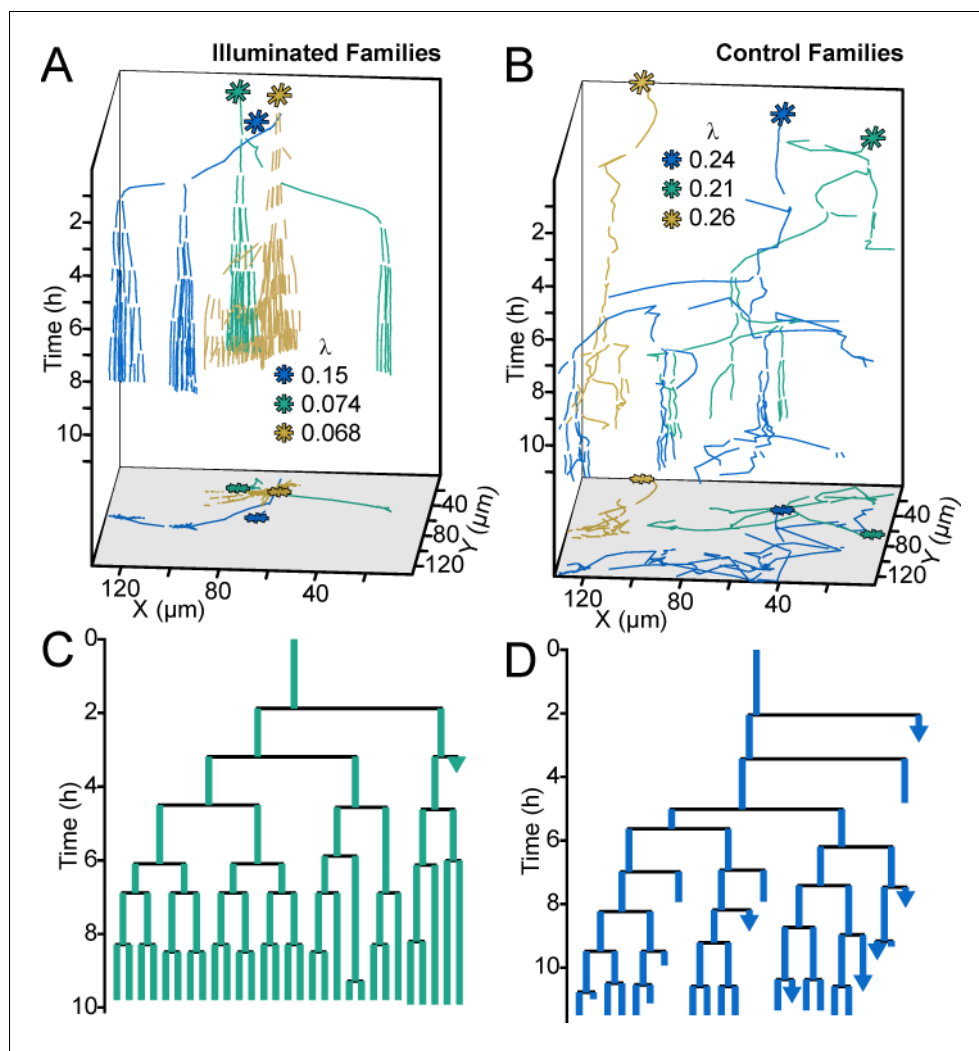


Figure 5. Optogenetic control of c-di-GMP production drastically affects family architecture and surface motility. (A,B) Spatiotemporal plot of 3 illuminated families (A) and three control families (B). The individual cell tracks in the 3D plot are projected onto the XY plane as spatial trajectories. As in **Figure 4**, λ is a measure of tree asymmetry, with higher values indicating more cells traveling outside the field of view or detaching. In A, the illuminated families tend to be sessile, as expected for cells with high c-di-GMP. In B, control cells are more motile than the illuminated cells in A. (C,D) Family trees of a single corresponding family in (A) and (B), where the color corresponds to the same family. Illuminated cells (C) tend to stay adhered across multiple generations, whereas control cells (D) display more surface motility and detachments. See **Video 1** for a representative video of the optogenetic reporter experiment. **Figure 5—figure supplement 1** is a control experiment showing that c-di-GMP levels increase in response to red light intensity in the optogenetic reporter strain. **Figure 5—figure supplement 2** shows a schematic of the ATIM apparatus. See **Figure 5—figure supplement 3** for the data from **Figure 5** overlaid onto **Figure 4C**, showing that optogenetic-controlled families follow the trend of family behavior observed in wt PAO1 cells.

DOI: <https://doi.org/10.7554/eLife.45084.027>

The following source data and figure supplements are available for figure 5:

Source data 1. MATLAB data for optogenetic and WT families.

DOI: <https://doi.org/10.7554/eLife.45084.031>

Figure supplement 1. Activity of the c-di-GMP reporter of PAO1-*bphS-P_{cdrA}*-GFP-mCherry was dependent on red-light intensity.

DOI: <https://doi.org/10.7554/eLife.45084.028>

Figure supplement 2. Using Adaptive Tracking Illumination Microscopy (ATIM) to exactly illuminate single *P. aeruginosa* cells on surface.

Figure 5 continued on next page

Figure 5 continued

DOI: <https://doi.org/10.7554/eLife.45084.029>

Figure supplement 3. Optogenetic-controlled families follow the trend of family behavior observed in wt PAO1 cells, with illuminated families resembling the high c-di-GMP matrix producers and control families resembling low c-di-GMP surface explorers.

DOI: <https://doi.org/10.7554/eLife.45084.030>

behavior—surface motility—we found the strongest correlation between c-di-GMP signaling and cellular behavior at the lineage level. For many cells, there was a lag between when c-di-GMP first increased and surface motility decreased. While a mother cell may still be surface motile upon initiating c-di-GMP signaling, we found that following cell division, daughter cells with high c-di-GMP eventually became immotile. Thus, whereas an increase in c-di-GMP relatively quickly results in increased biofilm matrix production, this same increase in c-di-GMP likely indirectly influences surface motility. Supporting this, *P. aeruginosa* is known to produce Psl polysaccharide while engaging in type IV pili mediated motility across a surface, leaving behind a trail of Psl (Zhao *et al.*, 2013). Finally, this apparent multigenerational influence of c-di-GMP signaling on bacterial behavior resembles the recently observed multigenerational memory of cAMP signaling in *P. aeruginosa* (Lee *et al.*, 2018). Additional work is needed to determine whether surface-naïve daughter cells have any ‘memory’ of surface attachment by a mother cell, through the maintenance of elevated c-di-GMP across one or more cell divisions. Another future direction of this work is to examine whether other bacterial signaling modalities, including other nucleotide or non-nucleotide signaling systems (e.g. ppGpp or quorum sensing), may exhibit similar multigenerational features.

We observed that the Pil-Chp surface sensing system did not play a role in early c-di-GMP signaling in our experimental system. There are numerous potential explanations for this. For example, it is possible that the Pil-Chp system is the dominant surface sensing mechanism under different environmental conditions than the ones we used for this study. Alternatively, the Pil-Chp system might not contribute to general cytoplasmic pools of c-di-GMP (for which the reporter is sensitive) and instead participates in specific localized c-di-GMP signaling events. In support of this notion, inactivation of individual diguanylate cyclases is well known to lead to distinct changes in c-di-GMP-regulated behaviors (Kulasakara *et al.*, 2006; Merritt *et al.*, 2010). It is interesting to note that whereas the other Pil-Chp inactivation mutants tested did not have a phenotype, the *pilA* deletion mutant strain displayed a slight defect in surface sensing in our study, although why is currently unknown.

If we overwhelm WspR-generated c-di-GMP heterogeneity by using optogenetically-induced sustained c-di-GMP production, we find that phenotypic heterogeneity is lost, and that illuminated cells deterministically become sessile and form microcolonies. Interestingly, our optogenetic experiments show that sustained c-di-GMP production for more than one generation is required before commitment to the sessile lifestyle. This observation is consistent with the fact that we see stronger correlations between c-di-GMP levels and motility behavior at the lineage level compared to the individual cell level. Moreover, since the Wsp surface sensing system generates heterogeneous c-di-GMP levels, this requirement of sustained c-di-GMP production for more than one generation is inherently difficult for wild-type cells to meet, and virtually guarantees the simultaneous existence of motile and sessile subpopulations.

The heterogeneity we observed in Wsp signaling shares many similarities with phenotypic heterogeneity generated from other c-di-GMP signaling (Petersen *et al.*, 2019) and quorum sensing systems (Grote *et al.*, 2015; Ramalho *et al.*, 2016). Nucleotide second messenger and quorum sensing (QS) signaling systems are traditionally thought to coordinate cellular behavior in response to information regarding the cell’s environment. However, rather than functioning to initiate a completely homogeneous response at the population level to environmental conditions, a growing body of literature suggests that a common theme of these signaling systems is that they introduce some level of behavioral heterogeneity (Grote *et al.*, 2015). For example, QS-induced phenotypic heterogeneity in *Vibrio harveyi* is attributable to variability in the phosphorylation state of LuxO and influences bioluminescence and biofilm formation (Anetzberger *et al.*, 2009). In *P. aeruginosa* and *Caulobacter crescentus*, heterogeneity in the very low levels of c-di-GMP present during planktonic growth is achieved through asymmetrical cell division and influences swimming motility (Kulasekara *et al.*, 2013). Thus, *P. aeruginosa* appears to have at least two distinct mechanisms of

generating c-di-GMP heterogeneity, which it employs during different modes of growth. In the case of the Wsp system, this phenotypic heterogeneity, which has been 'hardwired' into the structure of the Wsp surface sensing network, allows for a division of the labor during early biofilm formation, with one subpopulation committing to initiating the protective biofilm lifestyle, while the other subpopulation is free to explore the surface and potentially colonize distant, perhaps more favorable, locations.

Materials and methods

Key resources table

Reagent type (species) or resource	Designation	Source or reference	Identifiers	Additional information
Strain, strain background (<i>Pseudomonas aeruginosa</i>)	PAO1	PMID: 111024		The version of PAO1 used in this study can be obtained from Matthew Parsek's laboratory.
Recombinant DNA reagent	pPcdrA::gfpASV	PMID: 22582064		
Recombinant DNA reagent	PAO1 attCTX::bphS attMiniTn7::mCherry pPcdrA::gfpASV	This study	NA	This strain can be obtained from Fan Jin's laboratory.
Commercial assay or kit	QuikChange Lightning Site-Directed Mutagenesis Kit	Agilent Technologies	210519	
Commercial assay or kit	Gateway BP Clonase II Enzyme mix	ThermoFisher Scientific	11789020	
Commercial assay or kit	LR Clonase II Plus enzyme	ThermoFisher Scientific	12538120	
Commercial assay or kit	QIAquick gel extraction kit	Qiagen	28115	
Commercial assay or kit	QIAquick PCR purification kit	Qiagen	28104	
Commercial assay or kit	Antarctic phosphatase	New England BioLabs	M0289S	
Commercial assay or kit	TRIzol LS	ThermoFisher Scientific	10296010	
Commercial assay or kit	RQ1 RNase-Free DNase	Promega	M6101	
Commercial assay or kit	iTaq Universal SYBR Green One-Step kit	Bio-rad	172-5150	
Chemical compound, drug	TRITC Conjugated Wisteria floribunda lectin	EY laboratories	R-3101-1	
Chemical compound, drug	TRITC Conjugated Hippeastrum hybrid Lectin (<i>Amaryllis</i>)	EY laboratories	R-8008-1	
Software, algorithm	Volocity Image Analysis Software	Quorum Technologies Inc	Version 6.00	
Software, algorithm	NIS-Elements AR	Nikon	Version 4.00	
Software, algorithm	MATLAB code for tracking experiments	This study	Version R2015a	The code and datasets are available to download as source data files associated with Figures 1, 4 and 5.

Continued on next page

Continued

Reagent type (species) or resource	Designation	Source or reference	Identifiers	Additional information
Software, algorithm	Statistics and Machine Learning Toolbox for MATLAB	MathWorks	Version R2015a	
Other	Flow cell for time course experiments	University of Iowa Machine Shop	Standard dimension flow cell	Flow cell dimensions: 5 mm x 35 mm x 1 mm
Other	Flow cell for cell tracking experiments	PMID: 18770573	Standard dimension flow cell	This flow cell can be ordered from Department of Systems Biology, Technical University of Denmark.

Bacterial strains and growth conditions

The strains, plasmids, and primers used in this study are listed in **Table 1**. *Escherichia coli* and *P. aeruginosa* strains were routinely grown in Luria–Bertani (LB) medium and on LB agar at 37°C. For the flow cell experiments, *P. aeruginosa* was grown in either LB or FAB minimal medium supplemented with 10 mM or 0.6 mM glutamate at room temperature ([Zhao et al., 2013](#)). For flow cytometry experiments, *P. aeruginosa* was grown in either LB medium or in Jensen’s defined medium with glucose as the carbon source (a growth medium in which Pel is more abundantly produced than in LB) ([Jennings et al., 2015](#)). For the tube biofilm and c-di-GMP measurements, *P. aeruginosa* strains were grown in Vogel-Bonner Minimal Medium (VBMM; [Vogel and Bonner, 1956](#)). Antibiotics were supplied where necessary at the following concentrations: for *E. coli*, 100 µg/mL ampicillin, 10 µg/mL gentamicin, and 10 or 60 µg/mL tetracycline; for *P. aeruginosa*, 300 µg/mL carbenicillin, 100 µg/mL gentamicin, and 100 µg/mL tetracycline. *P_{cdrA}::gfp_{ASV}* reporter and vector control plasmids were selected with 100 µg/mL gentamicin for *P. aeruginosa* strains and 10 µg/mL gentamicin for *E. coli*.

PAO1 Δ *pilY1* was constructed using two-step allelic exchange following conjugation of wild type PAO1 with *E. coli* S17.1 harboring pENTRPEX18Gm:: Δ *pilY1* (a gift from Joe Harrison) as previously described ([Hmelo et al., 2015](#)). PAO1 Δ *pilY1* was identified by colony PCR using primers PAO1p*iY1*-SEQ-F and PAO1p*iY1*-SEQ-R. PAO1 Δ *dipA* was constructed similarly by conjugation of wild type PAO1 with *E. coli* S17.1 harboring pENTRPEX18Gm:: Δ *dipA* (a gift from Joe Harrison). PAO1 Δ *dipA* was identified by colony PCR using primers PAO1dipA-SEQ-F and PAO1dipA-SEQ-R. PA14 Δ *wspR* and Δ *wspF* deletion mutants were confirmed by PCR using primers PA14wspR-SEQ-F and PA14wspR-SEQ-R or PA14wspF-SEQ-F and PA14wspF-SEQ-R, respectively.

To create MPAO1 attTn7::P(A1/04/03)::GFPmut, the miniTn7 from pBT270 was integrated into the chromosome of *P. aeruginosa* PAO1 with the helper plasmid pTNS2, as previously described ([Choi and Schweizer, 2006](#)). pBT270 was created by introducing the constitutive A1/04/03 promoter ([Lanzer and Bujard, 1988](#)) and removing the trc promoter from pBT223 using the Quik-Change Lightning Kit (Agilent Technologies) and the oligonucleotides OBT314 and OBT315. pBT223 was constructed via recombineering of pBT200, pUC18-miniTn7T2-Gm-GW, and pBT212 using Multisite Gateway technology (Invitrogen). pBT212 was constructed by cloning the *gfpmut3* from AKN66 using OBT268 and OBT269, and recombining the PCR product with pDONR221 P1-P5r.

Construction of optogenetic, c-di-GMP reporter strain in *P. aeruginosa*

Chromosomal insertion of *bphS* was achieved using the mini-CTX system and these strains were marked with different fluorescent proteins by mini-Tn7 site-specific transposition essentially as previously described ([Choi and Schweizer, 2006](#); [Hoang et al., 2000](#)). First, a *bphS* fragment obtained from the plasmid pIND4 was cloned into the vector mini-CTX2 with the PA1/04/03 promoter upstream of the MCS via a two-piece ligation. The constructed plasmid was electroporated into PAO1 and the corresponding recombinant strain was identified by screening on LB agar plates containing 1 mM IPTG and 100 µg/mL tetracycline. Then, the strains were electroporated with a pFLP2 plasmid and distinguished on LB agar plates containing 5% (w/v) sucrose for the excision of the resistance marker. The c-di-GMP reporter plasmid and mCherry/EGFP marked *bphS* mutants were

Table 1. Strains, primers, and plasmids used in this study.

		Reference
<i>P. aeruginosa</i> strains		
PAO1	wild-type	Holloway et al., 1979
PA14	wild-type	Rahme et al., 1997
PAO1 Δ wspF	markerless, in frame deletion of WspF	Hickman et al., 2005
PAO1 Δ wspF Δ pelA Δ pslBCD	markerless, in frame deletions of WspF, PelA, and PslBCD genes	Rybtke et al., 2012
PAO1 Δ wspR	markerless, in frame deletion of WspR	Hickman, 2005
PAO1 Δ pilY1	markerless, in frame deletion of PilY1	this study
PAO1 Δ sadC	markerless, in frame deletion of SadC	Irie et al., 2012
PAO1 Δ pilA	markerless, in frame deletion of PilA	Shrout et al., 2006
PAO1 Δ dipA	markerless, in frame deletion of DipA	this study
PAO1 Δ wspR attCTX::PwspA::wspR	PAO1 Δ wspR complemented with WspR under control of the Wsp operon promoter and including intergenic region upstream of WspR	Gift from Yasuhiko Irie
PAO1 Δ sadC attCTX::sadC	PAO1 Δ sadC complemented with SadC under control of its native promoter	Gift from Yasuhiko Irie
MPAO1 attTn7:: P(A1/04/03)::GFPmut	wild type MPAO1 constitutively expressive stable GFP	this study
PA14 Δ wspF	markerless, in frame deletion of WspF	Gift from Caroline Harwood
PA14 Δ wspR	markerless, in frame deletion of WspR	Gift from Caroline Harwood
PAO1 Δ wspR attCTX:: PBAD-wspR-eYFP	markerless, in frame deletion of WspR with arabinose-inducible, C-terminally eYFP-tagged wild type WspR allele	Huangyutham et al., 2013
PAO1 Δ wspR attCTX:: PBAD-wspR[L170D]-eYFP	markerless, in frame deletion of WspR with arabinose-inducible, C-terminally eYFP-tagged WspR[L170D] allele	Huangyutham et al., 2013
PAO1 Δ wspR attCTX:: PBAD-wspR[E253A]-eYFP	markerless, in frame deletion of WspR with arabinose-inducible, C-terminally eYFP-tagged WspR[E253A] allele	Huangyutham et al., 2013
<i>P. aeruginosa</i> reporter strains		
PAO1 pMH489		Rybtke et al., 2012
PAO1 pPcdrA::gfpASV		Rybtke et al., 2012
PAO1 pPsiaA::gfpASV		this study
PA14 pMH489		this study
PA14 pPcdrA::gfpASV		this study
PAO1 Δ wspF pMH489		this study
PAO1 Δ wspF pPcdrA::gfpASV		this study
PAO1 Δ wspF pPsiaA::gfp		this study

Table 1 continued on next page

Table 1 continued

	Reference	
PAO1ΔwspFΔpelCΔpslD pMH489	this study	
PAO1ΔwspFΔpelCΔpslD pPcdrA::gfpASV	this study	
PAO1ΔwspR pMH489	this study	
PAO1ΔwspR pPcdrA::gfpASV	this study	
PAO1ΔwspR pPsiaA::gfp	this study	
PAO1ΔpilY1 pMH489	this study	
PAO1ΔpilY1 pPcdrA::gfpASV	this study	
PAO1ΔsadC pMH489	this study	
PAO1ΔsadC pPcdrA::gfpASV	this study	
PAO1ΔpilA pMH489	this study	
PAO1ΔpilA pPcdrA::gfpASV	this study	
PAO1ΔdipA pMH489	this study	
PAO1ΔdipA pPcdrA::gfpASV	this study	
PAO1ΔwspR attCTX::PwspA::wspR pMH489	this study	
PAO1ΔwspR attCTX::PwspA::wspR pPcdrA::gfpASV	this study	
PAO1ΔsadC att::sadC pMH489	this study	
PAO1ΔsadC att::sadC pPcdrA::gfpASV	this study	
PA14 ΔwspF pMH489	this study	
PA14 ΔwspF pPcdrA::gfpASV	this study	
PA14 ΔwspR pMH489	this study	
PA14 ΔwspR pPcdrA::gfpASV	this study	
PAO1ΔwspR attCTX::PBAD-wspR-eYFP pPcdrA::mTFP1	this study	
PAO1ΔwspR attCTX::PBAD-wspR[L170D]-eYFP pPcdrA::mTFP1	this study	
PAO1ΔwspR attCTX::PBAD-wspR[E253A]-eYFP pPcdrA::mTFP1	this study	
PAO1 attCTX::bphS attMiniTn7::mCherry pPcdrA::gfpASV	this study	
PAO1 attMiniTn7::mCherry pPcdrA::gfpASV	this study	
E. coli strains		
<i>E. coli</i> S17.1 pENTRPEX18Gm::ΔpilY1	conjugation proficient <i>E. coli</i> harboring <i>pilY1</i> deletion allele	Gift from Joe Harrison
<i>E. coli</i> S17.1 pENTRPEX18Gm::ΔdipA	conjugation proficient <i>E. coli</i> harboring <i>dipA</i> deletion allele	Gift from Joe Harrison
<i>E. coli</i> DH5α pUC18-miniTn7T2-PcdrA-RBSg10L-gfpAGA	source of PcdrA-RBSg10L	this study
<i>E. coli</i> DH5α pBBR1MCS5-PcdrA::RBSg10L::mTFP1	referred to as 'pPcdrA::mTFP1'	this study
<i>E. coli</i> DH5α pPsiaA::gfp	plasmid-based, fluorescent <i>siaA</i> transcriptional reporter	this study

Primers

Table 1 continued on next page

Table 1 continued

		Reference
PAO1pilY1-SEQ-F	CTACTACGAGACCAATAGCGTC	this study
PAO1pilY1-SEQ-R	GTCGATGTCCACCAGGTTCTTC	this study
PAO1dipA-SEQ-F	GATACGCTTAACCTGGGCCCTG	this study
PAO1dipA-SEQ-R	CTTTTCTTGGTGAGGATTCAGAAC	this study
PA14wspR-SEQ-F	GCTTCCTCACCATCGCCC	this study
PA14wspR-SEQ-R	CAGGTCGTCCAGGGTTTCC	this study
PA14wspF-SEQ-F	CTCACGGTGCGTGAGCTG	this study
PA14wspF-SEQ-R	GGTCCTGGAGGATCACCG	this study
SacI – PcdrA - F	GGGGAGCTC GTATGGAA GGTTCCTTGGCGG	this study
SOE-PcdrA-RBSg10L - R	ctcctgcacctgtcaccat GGATATATCTCCTTCTTAAAG	this study
mTFP1 - F	atggtagcaaggcgaggag	this study
KpnI - mTFP1 – R	GGGGTACC ttactgtacagctcgcc	this study
BamH1-Psia-F	GGG GGATCC GGCAGCGGCAACCGCCTCTG	this study
SiaA-BamH1-R	CCC GGATCC CAACCCCAAGTTCGCCCCAT	this study
M13F(–21)	TGTAAAACGACGGCCAGT	GeneWiz
M13R	CAGGAAACAGCTATGAC	GeneWiz
ampR-F-qPCR	GCG CCA TCC CTT CAT CG	Colvin et al., 2011
ampR-R-qPCR	GAT GTC GAC GCG GTT GTT G	Colvin et al., 2011
pslA-F-qPCR	AAG ATC AAG AAA CGC GTG GAA T	Colvin et al., 2011
pslA-R-qPCR	TGT AGA GGT CGA ACC ACA CCG	Colvin et al., 2011
pelA-F-qPCR	CCT TCA GCC ATC CGT TCT TCT	Colvin et al., 2011
pelA-R-qPCR	TCG CGT ACG AAG TCG ACC TT	Colvin et al., 2011
rplU-F-qPCR	CGC AGT GAT TGT TAC CGG TG	Colvin et al., 2011
rplU-R-qPCR	AGG CCT GAA TGC CGG TGA TC	Colvin et al., 2011
OBT268	GGGGACAACCTTTGTATACAAAG TTGTAATATAGAGGGACAAACTC AAGGTCATTGCAAGAGTGGC CTTTATGATTGACCTTCTCCGG TTAATACGACCGGGATAACTCCAC TTGAGACGTGAAAAAGAGGAGTA TTCATGCGTAAAGGAGAAGAA CTTTTCACTGGAG	This study
OBT269	GGGGACAAGTTTGTACAAAAAAGCA GGCTCGGCTTATTTGTATAGTTCAT CCATGCCATGTGTAATC	This study
OBT314	CAGGTCGACTCTAGAGGATCCCCATC AGAAAATTTATCAAAAAGAGTGTGACT TGTGAGCGGATAACAATGATACTTAGATT CAATTGTGAGCGGATAACAATTTACACA CATCTAGAATTAAGAGGAGAAATTA GCATGGTGAGCAAGGGCGAGGAG	Zhao et al., 2013
OBT315	CTCCTCGCCCTTGTCTCACCATGCTTAA TTTCTCCTCTTTAATTCTAGATGTGT GAAATTGTTATCCGCTCACAATTGAA TCTAAGTATCATTGTTATCCGC TCACAAGTCAACTCTTTTT GATAAATTTCTGATGGGGAT CCTCTAGAGTCGACCTG	Zhao et al., 2013

Table 1 continued on next page

Table 1 continued

		Reference
pPcdra::gfpASV	Pcdra reporter with short halflife GFP	Rybtke et al., 2012
pENTRPEX18Gm::ΔpilY1	suicide plasmid containing <i>pilY1</i> deletion construct for use in PAO1	Gift from Joe Harrison
pENTRPEX18Gm::ΔdipA	suicide plasmid containing <i>dipA</i> deletion construct for use in PAO1	Gift from Joe Harrison
pBBR1MCS5	broad host range vector that is stable in <i>P. aeruginosa</i> , GentR	Elzer et al., 1995
pUC18-miniTn7T2-Pcdra-RBSg10L-gfpAGA	source plasmid containing promoter of <i>cdrA</i> with enhanced ribosomal binding site	this study
pNCS-mTFP1	source plasmid containing mTFP1	Allele Biotech
pBBR1MCS5-Pcdra::RBSg10L::mTFP1	teal fluorescent protein version of Pcdra reporter	this study
pPsiaA::gfp	PsiaA reporter expressing stable GFP, constructed using pMH487 plasmid	this study
pBT270	miniTn7 transposon with <i>gfpmut3</i> driven by the A1/04/03 promoter; Apr, Gmr	This study
pTNS2	T7 transposase expression vector	Choi and Schweizer, 2006
pBT223	miniTn7 transposon with <i>gfpmut3</i> driven by the <i>trc</i> promoter; Apr, Gmr	This study
pBT212	A Gateway compatible plasmid containing <i>gfpmut3</i> flanked by attR5 and attL1 recombination sites; Kmr	This study
pBT200	A Gateway compatible plasmid containing the <i>trc</i> promoter flanked by attL2 and attL5 recombination sites; Knr	Zhao et al., 2013
pUC18-miniTn7T2-Gm-GW	A Gateway compatible mini-Tn7 based vector; Cmr, Apr and Gmr;	Zhao et al., 2013
AKN66	source for <i>gfpmut3</i>	Lambertsen et al., 2004
pDONR221 P1-P5r	A Gateway compatible vector with attP1 and attP5r recombination sites and <i>ccdB</i> ; Knr and Cmr	Invitrogen

DOI: <https://doi.org/10.7554/eLife.45084.033>

constructed as described above. The c-di-GMP reporter plasmid (P_{cdrA::gfpASV}) was electroporated into the mCherry-marked strain harboring *bphS* to monitor the intracellular c-di-GMP level.

To validate the optogenetic reporter strain in **Figure 5—figure supplement 1**, strains were grown on LB agar plates at 37°C for 24 hr from frozen stocks. Monoclonal colonies were inoculated and cultured with a minimal medium (FAB) at 37°C overnight, adding 1 μM FeCl₃ and 30 mM glutamate as the carbon source, until the culture reached an OD₆₀₀ of approximately 2.1. Then, the bacterial culture was further diluted (1:100) in fresh FAB medium to OD₆₀₀ 0.5. When required, gentamicin was added to medium at 30 μg/mL. Plates and tubes were wrapped with aluminum foil to achieve a dark condition. Finally, the culture was diluted (1:50) in fresh FAB medium and 6 μL

diluted culture was spotted onto an FAB agarose (2%) pad, with 30 mM glutamate and 1 μ M FeCl₃. The agarose pad was pressed on a coverglass before cells were illuminated.

Cyclic di-GMP measurement and qRT-PCR of tube biofilms

Measurement of c-di-GMP in tube biofilm cells was performed as previously described (Colvin *et al.*, 2011). Transcriptional analysis of PelA expression in tube biofilms was performed as described in the 'FACS and qRT-PCR of c-di-GMP reporter cells' section.

Crystal violet attachment assays

Crystal violet assays were performed essentially as previously described to measure biofilm biomass, except using gentle washing after 2–6 hr of static incubation (Armbruster *et al.*, 2016). To measure biofilm biomass at 24 hr, the crystal violet assay was performed as previously described without gentle washing (Colvin *et al.*, 2012).

Flow cell time course experiments and confocal microscopy

P. aeruginosa cells harboring the pP_{cdrA::gfp}_{ASV} reporter plasmid or a promoterless vector control (pMH489) were grown to mid-log in LB with 100 μ g/mL gentamicin (Gm100) from LB Gm100 plates or from overnight broth cultures in FAB +10 mM glutamate. Mid-log cells were back diluted into 1% LB or FAB +0.6 mM glutamate and flow chambers were inoculated at a final OD₆₀₀ 0.1 and inverted for 10 min to allow cells to attach before induction of flow. Clean media was used to wash non-attached cells by flow at 40 mL per hour for 20 min. Flow was then reduced to a final constant flow rate of 3 mL per hour and bacteria were imaged immediately on a Zeiss LSM 510 scanning confocal laser microscope (t = 0 hr). Flow cells were incubated at a constant flow rate at room temperature and imaged hourly for up to 24 hr. For every strain and time point, 5 fields of view and a minimum of 300 cells were captured using identical microscope settings to image GFP fluorescence across all experiments. Images were analyzed using Volocity software (Improvision, Coventry, UK). We binned cells by their mean GFP fluorescence intensity per pixel, in increments of 20 fluorescence units, and determined the cut-off bin that corresponded to cells clearing produced GFP when images were examined by eye. Therefore, cells were counted as pP_{cdrA::gfp}_{ASV} reporter 'on' if their mean GFP fluorescence intensity per pixel was ≥ 321 fluorescence units. For all summary figures depicting the percentage of cells with the reporter 'on', data are presented in terms of the percentage of cells with an average GFP fluorescence per pixel of ≥ 321 fluorescence units (pP_{cdrA::gfp}_{ASV} reporter 'on'). Microscopy images were artificially colored to display GFP fluorescence as green.

Construction of pP_{siaA::gfp}

A region 259 bp upstream through 21 bp into the coding sequence of *siaA* was amplified from PAO1 genomic DNA using primers BamH1-Psia-F and SiaA-BamH1-R, then gel purified using a QIAquick gel extraction kit (Qiagen, Hilden, Germany) digested with BamH1, then column purified with a QIAquick PCR purification kit (Qiagen, Hilden, Germany) to remove BamH1. The GFP expression vector pMH487, which contains the *gfp*_{mut3} gene with an RNase III splice site and lacking a promoter (Borlee *et al.*, 2010), was digested with BamH1, treated with Antarctic phosphatase (New England Biolabs, Ipswich, MA), then column purified with a QIAquick PCR purification kit (Qiagen, Hilden, Germany) to remove BamH1. The *PsiaA* allele was ligated into digested pMH487, then transformed into *E. coli* DH5 α , purified, and sequenced using primer M13F(-21) (Genewiz). The reporter pP_{siaA::gfp} was electroporated into *P. aeruginosa* as previously described and maintained under gentamycin selection at 100 μ g/mL.

Multi-generation single cell tracking of type IV motility and c-di-GMP reporter activity

Wild type PAO1 harboring the pP_{cdrA::gfp}_{ASV} reporter was grown shaking for 20 hr in FAB media with 6 mM glutamate. The flow cell inoculum was prepared by diluting the culture to a final OD₆₀₀ of 0.01 in FAB with 0.6 mM glutamate. The flow cell inoculum was injected into the flow cell (Department of Systems Biology, Technical University of Denmark) and allowed to incubate for 10 min at 30° C prior to flushing with media at 30 mL/h for 10 min. Experiments were performed under a flow rate of 3 mL/hour for a total of 40 hr.

Images were acquired with an Olympus IX81 microscope equipped with a Zero Drift Correction autofocus system, a 100 \times oil objective with a 2 \times multipler lens, and an Andor iXon EMCCD camera using Andor IQ software. Bright-field images were recorded every 3 s and GFP fluorescence every 15 min. Acquisition continued for a total recording time of 40 hr, which resulted in approximately 48000 bright-field images, and 160 fluorescence images.

Images were analyzed in MATLAB to track bacterial family trees, GFP fluorescence, and surface motility essentially as previously described (Lee *et al.*, 2018) with the following modifications. Image analysis, family tracking and manual validation, family tree plotting, and tree asymmetry λ calculations were performed as previously described (Lee *et al.*, 2018) without modification. GFP fluorescence intensities were normalized by calculating the distribution of intensities per cell per frame (extracted by using the binary image as a mask) and then setting the minimum and maximum intensities to the 1st and 99th percentiles of this distribution for each dataset. $I_{c-di-GMP}$ (relative normalized c-di-GMP reporter intensity) was calculated by averaging the normalized fluorescence intensities across all members of a family. F_{motile} (fraction of time that cells in a family are motile) was calculated as follows. For each family, every cell trajectory in the family was divided into time intervals. For each time interval, presence or absence of motility was determined using a combination of metrics, including Mean Squared Displacement (MSD) slope, radius of gyration, and visit map. MSD slope quantifies the directionality of movement relative to diffusion. Radius of gyration and visit map are different metrics for quantifying the average distance traveled on the surface. F_{motile} was then calculated by the fraction of these time intervals that have motility. This calculation was modified from the 'TFP activity metric' previously described (Lee *et al.*, 2018).

Setup of Adaptive Tracking Illumination Microscopy

Figure 5—figure supplement 2 shows a schematic of the Adaptive Tracking Illumination Microscopy (ATIM) setup. An inverted fluorescent microscope (Olympus, IX71) was modified to build the ATIM. The modification includes: 1) a commercial DMD-based LED projector (Gimi Z3) was used to replace the original bright-field light source, in which the original lenses in the projector were removed and three-colored (RGB) LEDs were rewired to connect to an external LED driver (ThorLabs) controlled by a single chip microcomputer (Arduino UNO r3); 2) the original bright-field condenser was replaced with an air objective (40 \times NA = 0.6, Leica); and 3) an additional 850 nm LED light (ThorLabs) was coupled to the illumination optical path using a dichroic mirror (Semrock) for the bright-field illumination. The 850 nm LED bright field light source does not affect optogenetic manipulation. An inverted fluorescent microscope (Olympus, IX71) equipped with a 100 \times oil objective and a sCMOS camera (Zyla 4.2 Andor) was used to collect bright-field images with 0.2 frame rate. The bright-field images were further analyzed to track multiple single cells in real time using a high-throughput bacterial tracking algorithm coded by Matlab. The projected contours of selected single cells were sent to the DMD (1280 \times 760 pixels) that was directly controlled by a commercial desktop through a VGA port. The manipulation lights were generated by the red-color LED (640 nm), and were projected on the single selected cells in real time through the DMD, a multi-band pass filter (446/532/646, Semrock) and the air objective. Our results indicated that feedback illuminations could generate projected patterns to exactly follow the cell movement (Figure 5 – Supplemental 2B) or single cells divisions (Figure 5 – Supplemental 2C) in real time.

Manipulation of c-di-GMP expression in single initial-attached cells

The bacterial strain PAO1-*bphS-PcdrA-GFP-mCherry* was inoculated into a flow cell (Denmark Technical University) and continuously cultured at 30.0 \pm 0.1 $^{\circ}$ C by flowing FAB medium (3.0 mL/h). The flow cell was modified by punching a hole with a 5 mm diameter into the channel, and the hole was sealed by a coverslip that allows the manipulation light to pass through. An inverted fluorescent microscope (Olympus, IX71) equipped with a 100 \times oil objective and a sCMOS camera (Zyla 4.2 Andor) was used to collect bright field or fluorescent images with 0.2 or 1/1800 frame rate respectively. The power density of the manipulation lights was determined by measuring the power at the outlet of the air objective using a power meter (Newport 842-PE). GFP or mCherry was excited using a 480 nm or 565 nm LED lights (ThorLabs) and imaged using single-band emission filters (Semrock): GFP (520/28 nm) or mCherry (631/36 nm). Initial-attached cells were selected to be manipulated using ATIM with the illumination at 0.05 mW/cm², which allowed us to compare the results arising

from illuminated or un-illuminated mobile cells in one experiment. The c-di-GMP levels in single cells were gauged using the ratio of GFP and mCherry intensities.

Lectin staining and flow cytometry

Glass culture tubes were inoculated with 1 mL of *P. aeruginosa* in LB or Jensen's minimal media at an OD₆₀₀ 0.8 and incubated statically at 37°C for 4 hr. Non-adhered cells were removed by washing three times with 2 mL sterile phosphate buffered saline (PBS). Biofilm cells were harvested by vortexing in 1 mL PBS with tetramethylrhodamine (TRITC) conjugated lectins (TRITC-labeled WFL lectin (100 µg/mL; Vector Laboratories) for Pel, TRITC-labeled HHA (100 µg/mL; EY Laboratories) for Psl) and incubated on ice for 5 min. Cells were washed 3 times to remove non-adhered lectin, resuspended in PBS, and immediately analyzed for GFP and TRITC fluorescence on a BD LSRII flow cytometer (BD Biosciences). Events were gated based on forward and side scatter to remove particles smaller than a single *P. aeruginosa* cell and large aggregates.

We used PAO1 cells that did not express GFP (wild type PAO1; **Figure 1—source data 2**) or constitutively expressed GFP (PAO1 Tn7::P(A1/04/03)::GFPmut; **Figure 1—source data 2**) to define a gate for high GFP fluorescence. We validated this gate using a strain in which we expect very high levels of reporter activity (surface grown PAO1 $\Delta wspF\Delta pelA\Delta pslBCD$ harboring pPcdrA::gfp_{ASV}) and saw that 91.6% of cells had high GFP levels (**Figure 1—source data 2**), in agreement with our flow cell characterization of this strain (**Figure 2A**). We determined gating for TRITC using cells that had not been stained with TRITC-conjugated lectin (**Figure 1—figure supplement 6A**), as well as two strains that overproduced either Psl (**Figure 1—figure supplement 6B**) or Pel (**Figure 1—figure supplement 6C**) that were stained with the appropriate TRITC-conjugated lectin. Our flow cytometry gating procedure accurately gated 99.7% of wild type PAO1 cells (without the PcdrA reporter or lectin-staining) as low GFP and low TRITC (**Figure 1—figure supplement 6D**).

FACS and qRT-PCR of c-di-GMP reporter cells

Static biofilm reporter cells were grown as described above and harvested without lectin staining. Cells were fixed with 6% paraformaldehyde for 20 min on ice, then rinsed once with sterile PBS prior to analysis with a FACSArialI (BD Biosciences, San Jose, CA). Events were gated first to remove debris and large cellular aggregates, and then gated into cells with low and high GFP fluorescence intensity. The low GFP gate was drawn using wild type PAO1 cells without the gfp gene (**Figure 1—figure supplement 7A**) and the high GFP gate was drawn using both PAO1 Tn7::P(A1/04/03)::GFPmut (**Figure 1—figure supplement 7B**) and PAO1 $\Delta wspF\Delta pelA\Delta pslBCD$ PcdrA::gfp_{ASV} reporter (**Figure 1—figure supplement 7C**). As expected, wild type PAO1 pPcdrA::gfp_{ASV} reporter cells that had been harvested after 4 hr of surface attachment to glass in static LB liquid culture displayed subpopulations of high GFP, reporter 'on' cells (30.8% of the population) and 'off' (57.2%) cells (**Figure 1—figure supplement 7D**), whereas this same strain grown to mid-log planktonically in LB displayed mostly reporter 'off' cells (**Figure 1—figure supplement 7E**). Cells were sorted at 4°C by flow assisted cell sorting (FACS) to collect 100,000 events into TRIzol LS (Thermo Fisher Scientific, Waltham, MA). RNA was extracted from sorted cells by boiling immediately for 10 min and following the manufacturer's instructions for RNA isolation. DNA was digested by treating with RQ1 Dnase I (Promega, Madison, WI) and samples were checked for genomic DNA contamination by PCR to detect *rplU*. Expression of *pelA*, *pslA*, and *ampR* was measured by quantitative Reverse Transcriptase PCR (qRT-PCR) using the iTaq Universal SYBR Green One-Step kit (Biorad, Hercules, CA) and a CFX96 Touch Real-Time PCR detection system (Bio-Rad, Hercules, CA). The $\Delta\Delta C_q$ was calculated for three independent samples of sorted wild type PAO1 PcdrA::gfp_{ASV} reporter biofilm cell populations by normalizing *PelA* and *PslA* to relative levels of *AmpR* expression. Data were presented as the average fold change in *PelA* or *PslA* expression in the PcdrA::gfp_{ASV} sorted 'on' population (high GFP) relative to the 'off' population (low GFP) for the three biological replicates.

WspR-YFP foci and pPcdrA::mTFP1 reporter

A version of the pPcdrA reporter was constructed in the pBBR1MCS5 plasmid to express mTFP1 instead of GFP, for use with YFP-tagged WspR proteins. The PcdrA promoter and an enhanced ribosomal binding site from the gene 10 leader sequence of the T7 phage (g10L) was amplified from pUC18-miniTn7T2-PcdrA-RBSg10L-gfp_{PAGA} using primers SacI-PcdrA-F and SOE-PcdrA-RBSg10L-R.

The primers mTFP1-F and KpnI-mTFP1-R were used to amplify the mTFP1 gene from plasmid pNCS-mTFP1 (Allele Biotech, San Diego, CA). The $P_{cdrA::RBSg10L::mTFP1}$ allele was constructed by SOE-PCR using primers SacI-PcdrA-F and KpnI-mTFP1-R, then pBBR1MCS5 and the SOE PCR product were doubly digested with SacI/KpnI. Digested pBBR1MCS5 was treated with Antarctic phosphatase, then both digests were gel purified and ligated. The ligation was transformed into *E. coli* DH5 α , and plasmid from clones growing on LB with 10 μ g/mL gentamycin were sequenced with primers M13F and M13F(-21) (GeneWiz). Fluorescence of the $pP_{cdrA::mTFP1}$ reporter was measured in Wsp mutants in a fluorimeter (BioTek Synergy H1 Hybrid Reader, BioTek Instruments, Inc, Winooski, VT, USA) and in flow cells to confirm its activity resembled that of $pP_{cdrA::gfp_{ASV}}$. The $pP_{cdrA::mTFP1}$ reporter was electroporated into *P. aeruginosa* strains with the native WspR deleted and harboring an arabinose-inducible copy of WspR-YFP on its chromosome (*Huangyutitham et al., 2013*). Cells were grown on LB agar plates with 100 μ g/mL gentamycin and 1% arabinose for 10 hr, then transferred to an agar pad for imaging of Differential Interference Contrast (DIC), YFP, and TFP. WspR-YFP foci and mTFP1 fluorescence was imaged using a Nikon Ti-E inverted wide-field fluorescence microscope with a large-format scientific complementary metal-oxide semiconductor camera (sCMOS; NEO, Andor Technology, Belfast, United Kingdom) and controlled by NIS-Elements. WspR-YFP foci were detected and $pP_{cdrA::mTFP1}$ reporter activity were analyzed using NIS-Elements AR software (Nikon Instruments, Melville, NY, USA). Regions of interest (ROI) corresponding to individual cells were determined using DIC images and the average mTFP1 fluorescence was measured within these ROIs. Next, we used essentially the same protocol as previously described for detecting WspR-eYFP foci (*Huangyutitham et al., 2013*), by examining the ratio of the maximum eYFP signal to the mean eYFP signal for each ROI. We verified by eye that the previously determined cut-off ratio of 1.7 accurately represented cells with at least one visible WspR-eYFP focus (*Huangyutitham et al., 2013*).

Acknowledgements

We thank Drs. Julie Cass and Paul Wiggins providing the wide-field microscope and CMOS camera to image WspR-eYFP clusters, and Drs. Joe J Harrison and Yasuhiko Irie for the gift of bacterial strains.

Additional information

Funding

Funder	Grant reference number	Author
National Institutes of Health	T32GM007270	Catherine R Armbruster
Charlie Moore Endowed Fellowship		Catherine R Armbruster
Army Research Office	W911NF1810254	Calvin K Lee Gerard CL Wong Matthew R Parsek
National Institutes of Health	K22AI121097	Boo Shan Tseng
National Institute of General Medical Sciences	GM56665	Caroline S Harwood
National Natural Science Foundation of China	21474098	Fan Jin
Fundamental Research Funds for the Central Universities	WK2340000066	Fan Jin
National Institutes of Health	K24HL141669	Lucas R Hoffman
National Institutes of Health	5R01AI077628	Matthew R Parsek
National Natural Science Foundation of China	21774117	Fan Jin
National Natural Science Foundation of China	21522406	Fan Jin

Fundamental Research Funds for the Central Universities	WK3450000003	Fan Jin
National Institutes of Health	1R01AI143730-01	Calvin K Lee Gerard CL Wong
National Institutes of Health	R01AI143916	Matthew R Parsek

The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

Author contributions

Catherine R Armbruster, Conceptualization, Data curation, Formal analysis, Supervision, Funding acquisition, Investigation, Visualization, Methodology, Writing—original draft, Project administration, Writing—review and editing; Calvin K Lee, Conceptualization, Data curation, Formal analysis, Investigation, Visualization, Methodology, Writing—review and editing; Jessica Parker-Gilham, Data curation, Formal analysis, Investigation, Visualization; Jaime de Anda, Data curation, Formal analysis, Investigation, Methodology; Aiguo Xia, Data curation, Formal analysis, Visualization; Kun Zhao, Data curation, Investigation, Methodology; Keiji Murakami, Data curation; Boo Shan Tseng, Conceptualization, Supervision, Funding acquisition, Investigation, Methodology, Writing—review and editing; Lucas R Hoffman, Supervision, Funding acquisition, Methodology, Writing—review and editing; Fan Jin, Conceptualization, Resources, Formal analysis, Supervision, Funding acquisition, Investigation, Methodology, Writing—review and editing; Caroline S Harwood, Conceptualization, Resources, Funding acquisition, Methodology, Writing—review and editing; Gerard CL Wong, Conceptualization, Resources, Supervision, Funding acquisition, Methodology, Project administration, Writing—review and editing; Matthew R Parsek, Conceptualization, Supervision, Funding acquisition, Methodology, Project administration, Writing—review and editing

Author ORCIDs

Catherine R Armbruster  <https://orcid.org/0000-0003-0795-802X>

Calvin K Lee  <https://orcid.org/0000-0001-6789-0317>

Boo Shan Tseng  <http://orcid.org/0000-0001-7563-0232>

Fan Jin  <http://orcid.org/0000-0003-2313-0388>

Matthew R Parsek  <https://orcid.org/0000-0003-2932-7966>

Decision letter and Author response

Decision letter <https://doi.org/10.7554/eLife.45084.039>

Author response <https://doi.org/10.7554/eLife.45084.040>

Additional files

Supplementary files

- Source code 1. Source code for fluorescence statistical tests.
DOI: <https://doi.org/10.7554/eLife.45084.034>
- Source code 2. Source code for family tree plots.
DOI: <https://doi.org/10.7554/eLife.45084.035>
- Source code 3. Source code for optogenetic family tree plots.
DOI: <https://doi.org/10.7554/eLife.45084.036>
- Transparent reporting form
DOI: <https://doi.org/10.7554/eLife.45084.037>

Data availability

Source data files and/or MATLAB code have been provided for Figures 1, 3, 4, and 5.

References

- Ackermann M.** 2015. A functional perspective on phenotypic heterogeneity in microorganisms. *Nature Reviews Microbiology* **13**:497–508. DOI: <https://doi.org/10.1038/nrmicro3491>, PMID: 26145732
- Anetzberger C, Pirch T, Jung K.** 2009. Heterogeneity in quorum sensing-regulated bioluminescence of *Vibrio harveyi*. *Molecular Microbiology* **73**:267–277. DOI: <https://doi.org/10.1111/j.1365-2958.2009.06768.x>, PMID: 19555459
- Armbruster CR, Wolter DJ, Mishra M, Hayden HS, Radey MC, Merrihew G, MacCoss MJ, Burns J, Wozniak DJ, Parsek MR, Hoffman LR.** 2016. *Staphylococcus aureus* Protein A Mediates Interspecies Interactions at the Cell Surface of *Pseudomonas aeruginosa*. *mBio* **7**. DOI: <https://doi.org/10.1128/mBio.00538-16>, PMID: 27222468
- Baraquet C, Murakami K, Parsek MR, Harwood CS.** 2012. The FleQ protein from *Pseudomonas aeruginosa* functions as both a repressor and an activator to control gene expression from the *pel* operon promoter in response to c-di-GMP. *Nucleic Acids Research* **40**:7207–7218. DOI: <https://doi.org/10.1093/nar/gks384>, PMID: 22581773
- Baraquet C, Harwood CS.** 2016. FleQ DNA binding consensus sequence revealed by studies of FleQ-Dependent regulation of biofilm gene expression in *Pseudomonas aeruginosa*. *Journal of Bacteriology* **198**:178–186. DOI: <https://doi.org/10.1128/JB.00539-15>, PMID: 26483521
- Borlee BR, Goldman AD, Murakami K, Samudrala R, Wozniak DJ, Parsek MR.** 2010. *Pseudomonas aeruginosa* uses a cyclic-di-GMP-regulated adhesin to reinforce the biofilm extracellular matrix. *Molecular Microbiology* **75**:827–842. DOI: <https://doi.org/10.1111/j.1365-2958.2009.06991.x>, PMID: 20088866
- Casadesús J, Low D.** 2006. Epigenetic gene regulation in the bacterial world. *Microbiology and Molecular Biology Reviews* **70**:830–856. DOI: <https://doi.org/10.1128/MMBR.00016-06>, PMID: 16959970
- Chen AI, Dolben EF, Okegbe C, Harty CE, Golub Y, Thao S, Ha DG, Willger SD, O'Toole GA, Harwood CS, Dietrich LE, Hogan DA.** 2014. *Candida albicans* ethanol stimulates *Pseudomonas aeruginosa* WspR-controlled biofilm formation as part of a cyclic relationship involving phenazines. *PLoS Pathogens* **10**:e1004480. DOI: <https://doi.org/10.1371/journal.ppat.1004480>, PMID: 25340349
- Choi KH, Schweizer HP.** 2006. mini-Tn7 insertion in bacteria with single attTn7 sites: example *Pseudomonas aeruginosa*. *Nature Protocols* **1**:153–161. DOI: <https://doi.org/10.1038/nprot.2006.24>, PMID: 17406227
- Colvin KM, Gordon VD, Murakami K, Borlee BR, Wozniak DJ, Wong GC, Parsek MR.** 2011. The *pel* polysaccharide can serve a structural and protective role in the biofilm matrix of *Pseudomonas aeruginosa*. *PLoS Pathogens* **7**:e1001264. DOI: <https://doi.org/10.1371/journal.ppat.1001264>, PMID: 21298031
- Colvin KM, Irie Y, Tart CS, Urbano R, Whitney JC, Ryder C, Howell PL, Wozniak DJ, Parsek MR.** 2012. The *pel* and *psl* polysaccharides provide *Pseudomonas aeruginosa* structural redundancy within the biofilm matrix. *Environmental Microbiology* **14**:1913–1928. DOI: <https://doi.org/10.1111/j.1462-2920.2011.02657.x>, PMID: 22176658
- Davey ME, O'toole GA.** 2000. Microbial biofilms: from ecology to molecular genetics. *Microbiology and Molecular Biology Reviews* **64**:847–867. DOI: <https://doi.org/10.1128/MMBR.64.4.847-867.2000>, PMID: 11104821
- Dragoš A, Kiewewalter H, Martin M, Hsu CY, Hartmann R, Wechsler T, Eriksen C, Brix S, Drescher K, Stanley-Wall N, Kümmerli R, Kovács ÁT.** 2018. Division of labor during biofilm matrix production. *Current Biology* **28**:1903–1913. DOI: <https://doi.org/10.1016/j.cub.2018.04.046>, PMID: 29887307
- Dubnau D, Losick R.** 2006. Bistability in bacteria. *Molecular Microbiology* **61**:564–572. DOI: <https://doi.org/10.1111/j.1365-2958.2006.05249.x>, PMID: 16879639
- Elowitz MB, Levine AJ, Siggia ED, Swain PS.** 2002. Stochastic gene expression in a single cell. *Science* **297**:1183–1186. DOI: <https://doi.org/10.1126/science.1070919>, PMID: 12183631
- Elzer PH, Kovach ME, Phillips RW, Robertson GT, Peterson KM, Roop RM.** 1995. In vivo and in vitro stability of the broad-host-range cloning vector pBBR1MCS in six *Brucella* species. *Plasmid* **33**:51–57. DOI: <https://doi.org/10.1006/plas.1995.1006>, PMID: 7753908
- Goldsby HJ, Dornhaus A, Kerr B, Ofria C.** 2012. Task-switching costs promote the evolution of division of labor and shifts in individuality. *PNAS* **109**:13686–13691. DOI: <https://doi.org/10.1073/pnas.1202233109>, PMID: 22872867
- Grote J, Krysciak D, Streit WR.** 2015. Phenotypic heterogeneity, a phenomenon that may explain why quorum sensing does not always result in truly homogenous cell behavior. *Applied and Environmental Microbiology* **81**:5280–5289. DOI: <https://doi.org/10.1128/AEM.00900-15>, PMID: 26025903
- Haagensen JA, Klausen M, Ernst RK, Miller SI, Folkesson A, Tolker-Nielsen T, Molin S.** 2007. Differentiation and distribution of colistin- and sodium dodecyl sulfate-tolerant cells in *Pseudomonas aeruginosa* biofilms. *Journal of Bacteriology* **189**:28–37. DOI: <https://doi.org/10.1128/JB.00720-06>, PMID: 17041046
- Hickman JW, Tifrea DF, Harwood CS.** 2005. A chemosensory system that regulates biofilm formation through modulation of cyclic diguanylate levels. *PNAS* **102**:14422–14427. DOI: <https://doi.org/10.1073/pnas.0507170102>
- Hickman JW, Harwood CS.** 2008. Identification of FleQ from *Pseudomonas aeruginosa* as a c-di-GMP-responsive transcription factor. *Molecular Microbiology* **69**:376–389. DOI: <https://doi.org/10.1111/j.1365-2958.2008.06281.x>, PMID: 18485075
- Hmelo LR, Borlee BR, Almlad H, Love ME, Randall TE, Tseng BS, Lin C, Irie Y, Storek KM, Yang JJ, Siehnel RJ, Howell PL, Singh PK, Tolker-Nielsen T, Parsek MR, Schweizer HP, Harrison JJ.** 2015. Precision-engineering the

- Pseudomonas aeruginosa* genome with two-step allelic exchange. *Nature Protocols* **10**:1820–1841. DOI: <https://doi.org/10.1038/nprot.2015.115>, PMID: 26492139
- Hoang TT**, Kutchma AJ, Becher A, Schweizer HP. 2000. Integration-proficient plasmids for *Pseudomonas aeruginosa*: site-specific integration and use for engineering of reporter and expression strains. *Plasmid* **43**:59–72. DOI: <https://doi.org/10.1006/plas.1999.1441>, PMID: 10610820
- Holloway BW**, Krishnapillai V, Morgan AF. 1979. Chromosomal genetics of *Pseudomonas*. *Microbiology Reviews* **43**:73–102.
- Huangyutitham V**, Güvener ZT, Harwood CS. 2013. Subcellular clustering of the phosphorylated WspR response regulator protein stimulates its diguanylate cyclase activity. *mBio* **4**:e00242. DOI: <https://doi.org/10.1128/mBio.00242-13>, PMID: 23653447
- Irie Y**, Borlee BR, O'Connor JR, Hill PJ, Harwood CS, Wozniak DJ, Parsek MR. 2012. Self-produced exopolysaccharide is a signal that stimulates biofilm formation in *Pseudomonas aeruginosa*. *PNAS* **109**:20632–20636. DOI: <https://doi.org/10.1073/pnas.1217993109>
- Jain R**, Behrens AJ, Kaefer V, Kazmierczak BI. 2012. Type IV pilus assembly in *Pseudomonas aeruginosa* over a broad range of cyclic di-GMP concentrations. *Journal of Bacteriology* **194**:4285–4294. DOI: <https://doi.org/10.1128/JB.00803-12>, PMID: 22685276
- Jain R**, Sliusarenko O, Kazmierczak BI. 2017. Interaction of the cyclic-di-GMP binding protein FimX and the type 4 pilus assembly ATPase promotes pilus assembly. *PLoS Pathogens* **13**:e1006594. DOI: <https://doi.org/10.1371/journal.ppat.1006594>, PMID: 28854278
- Jennings LK**, Storek KM, Ledvina HE, Coulon C, Marmont LS, Sadvovskaya I, Secor PR, Tseng BS, Scian M, Filloux A, Wozniak DJ, Howell PL, Parsek MR. 2015. Pel is a cationic exopolysaccharide that cross-links extracellular DNA in the *Pseudomonas aeruginosa* biofilm matrix. *PNAS* **112**:11353–11358. DOI: <https://doi.org/10.1073/pnas.1503058112>, PMID: 26311845
- Kim W**, Levy SB, Foster KR. 2016. Rapid radiation in bacteria leads to a division of labour. *Nature Communications* **7**:10508. DOI: <https://doi.org/10.1038/ncomms10508>, PMID: 26852925
- Klauck G**, Serra DO, Possling A, Hengge R. 2018. Spatial organization of different sigma factor activities and c-di-GMP signalling within the three-dimensional landscape of a bacterial biofilm. *Open Biology* **8**:180066. DOI: <https://doi.org/10.1098/rsob.180066>, PMID: 30135237
- Klausen M**, Aaes-Jørgensen A, Molin S, Tolker-Nielsen T. 2003. Involvement of bacterial migration in the development of complex multicellular structures in *Pseudomonas aeruginosa* biofilms. *Molecular Microbiology* **50**:61–68. DOI: <https://doi.org/10.1046/j.1365-2958.2003.03677.x>, PMID: 14507363
- Kuchma SL**, Ballok AE, Merritt JH, Hammond JH, Lu W, Rabinowitz JD, O'Toole GA. 2010. Cyclic-di-GMP-mediated repression of swarming motility by *Pseudomonas aeruginosa*: the pilY1 gene and its impact on surface-associated behaviors. *Journal of Bacteriology* **192**:2950–2964. DOI: <https://doi.org/10.1128/JB.01642-09>, PMID: 20233936
- Kulasakara H**, Lee V, Brencic A, Liberati N, Urbach J, Miyata S, Lee DG, Neely AN, Hyodo M, Hayakawa Y, Ausubel FM, Lory S. 2006. Analysis of *Pseudomonas aeruginosa* diguanylate cyclases and phosphodiesterases reveals a role for bis-(3'-5')-cyclic-GMP in virulence. *PNAS* **103**:2839–2844. DOI: <https://doi.org/10.1073/pnas.0511090103>, PMID: 16477007
- Kulasekara BR**, Kamischke C, Kulasekara HD, Christen M, Wiggins PA, Miller SI. 2013. c-di-GMP heterogeneity is generated by the chemotaxis machinery to regulate flagellar motility. *eLife* **2**:e01402. DOI: <https://doi.org/10.7554/eLife.01402>
- Lambertsen L**, Sternberg C, Molin S. 2004. Mini-Tn7 transposons for site-specific tagging of Bacteria with fluorescent proteins. *Environmental Microbiology* **6**:726–732. DOI: <https://doi.org/10.1111/j.1462-2920.2004.00605.x>, PMID: 15186351
- Lanzer M**, Bujard H. 1988. Promoters largely determine the efficiency of repressor action. *PNAS* **85**:8973–8977. DOI: <https://doi.org/10.1073/pnas.85.23.8973>, PMID: 3057497
- Laventie BJ**, Sangermani M, Estermann F, Manfredi P, Planes R, Hug I, Jaeger T, Meunier E, Broz P, Jenal U. 2019. A Surface-Induced asymmetric program promotes tissue colonization by *Pseudomonas aeruginosa*. *Cell Host & Microbe* **25**:140–152. DOI: <https://doi.org/10.1016/j.chom.2018.11.008>, PMID: 30581112
- Lee CK**, de Anda J, Baker AE, Bennett RR, Luo Y, Lee EY, Keefe JA, Helali JS, Ma J, Zhao K, Golestanian R, O'Toole GA, Wong GCL. 2018. Multigenerational memory and adaptive adhesion in early bacterial biofilm communities. *PNAS* **115**:4471–4476. DOI: <https://doi.org/10.1073/pnas.1720071115>, PMID: 29559526
- Lowery NV**, McNally L, Ratcliff WC, Brown SP. 2017. Division of labor, Bet Hedging, and the evolution of mixed biofilm investment strategies. *mBio* **8**:e00672. DOI: <https://doi.org/10.1128/mBio.00672-17>, PMID: 28790201
- Luo Y**, Zhao K, Baker AE, Kuchma SL, Coggan KA, Wolfgang MC, Wong GC, O'Toole GA. 2015. A hierarchical cascade of second messengers regulates *Pseudomonas aeruginosa* surface behaviors. *mBio* **6**. DOI: <https://doi.org/10.1128/mBio.02456-14>, PMID: 25626906
- Ma L**, Conover M, Lu H, Parsek MR, Bayles K, Wozniak DJ. 2009. Assembly and development of the *Pseudomonas aeruginosa* biofilm matrix. *PLoS Pathogens* **5**:e1000354. DOI: <https://doi.org/10.1371/journal.ppat.1000354>, PMID: 19325879
- Merritt JH**, Ha DG, Cowles KN, Lu W, Morales DK, Rabinowitz J, Gitai Z, O'Toole GA. 2010. Specific control of *Pseudomonas aeruginosa* surface-associated behaviors by two c-di-GMP diguanylate cyclases. *mBio* **1**. DOI: <https://doi.org/10.1128/mBio.00183-10>, PMID: 20978535
- O'Connor JR**, Kuwada NJ, Huangyutitham V, Wiggins PA, Harwood CS. 2012. Surface sensing and lateral subcellular localization of WspA, the receptor in a chemosensory-like system leading to c-di-GMP production. *Molecular Microbiology* **86**:720–729. DOI: <https://doi.org/10.1111/mmi.12013>, PMID: 22957788

- Persat A**, Inclan YF, Engel JN, Stone HA, Gitai Z. 2015. Type IV pili mechanochemically regulate virulence factors in *Pseudomonas aeruginosa*. *PNAS* **112**:7563–7568. DOI: <https://doi.org/10.1073/pnas.1502025112>, PMID: 26041805
- Petersen E**, Mills E, Miller SI. 2019. Cyclic-di-GMP regulation promotes survival of a slow-replicating subpopulation of intracellular *Salmonella* Typhimurium. *PNAS* **116**:6335–6340. DOI: <https://doi.org/10.1073/pnas.1901051116>
- Rahme LG**, Tan M-W, Le L, Wong SM, Tompkins RG, Calderwood SB, Ausubel FM. 1997. Use of model plant hosts to identify *Pseudomonas aeruginosa* virulence factors. *PNAS* **94**:13245–13250. DOI: <https://doi.org/10.1073/pnas.94.24.13245>
- Ramalho T**, Meyer A, Mückl A, Kapsner K, Gerland U, Simmel FC. 2016. Single cell analysis of a bacterial Sender-Receiver system. *PLOS ONE* **11**:e0145829. DOI: <https://doi.org/10.1371/journal.pone.0145829>, PMID: 26808777
- Ribbe J**, Baker AE, Euler S, O'Toole GA, Maier B. 2017. Role of cyclic Di-GMP and exopolysaccharide in type IV pilus dynamics. *Journal of Bacteriology* **199**:e00859. DOI: <https://doi.org/10.1128/JB.00859-16>, PMID: 28167523
- Römling U**, Galperin MY, Gomelsky M. 2013. Cyclic di-GMP: the first 25 years of a universal bacterial second messenger. *Microbiology and Molecular Biology Reviews* **77**:1–52. DOI: <https://doi.org/10.1128/MMBR.00043-12>, PMID: 23471616
- Rossetti V**, Schirmer BE, Bernasconi MV, Bagheri HC. 2010. The evolutionary path to terminal differentiation and division of labor in cyanobacteria. *Journal of Theoretical Biology* **262**:23–34. DOI: <https://doi.org/10.1016/j.jtbi.2009.09.009>, PMID: 19761779
- Rybtke MT**, Borlee BR, Murakami K, Irie Y, Hentzer M, Nielsen TE, Givskov M, Parsek MR, Tolker-Nielsen T. 2012. Fluorescence-based reporter for gauging cyclic di-GMP levels in *Pseudomonas aeruginosa*. *Applied and Environmental Microbiology* **78**:5060–5069. DOI: <https://doi.org/10.1128/AEM.00414-12>, PMID: 22582064
- Ryu MH**, Gomelsky M. 2014. Near-infrared light responsive synthetic c-di-GMP module for optogenetic applications. *ACS Synthetic Biology* **3**:802–810. DOI: <https://doi.org/10.1021/sb400182x>, PMID: 24926804
- Schreiber F**, Littmann S, Lavik G, Escrig S, Meibom A, Kuypers MM, Ackermann M. 2016. Phenotypic heterogeneity driven by nutrient limitation promotes growth in fluctuating environments. *Nature Microbiology* **1**:16055. DOI: <https://doi.org/10.1038/nmicrobiol.2016.55>, PMID: 27572840
- Serra DO**, Klauk G, Hengge R. 2015. Vertical stratification of matrix production is essential for physical integrity and architecture of macrocolony biofilms of *Escherichia coli*. *Environmental Microbiology* **17**:5073–5088. DOI: <https://doi.org/10.1111/1462-2920.12991>, PMID: 26234179
- Shrout JD**, Chopp DL, Just CL, Hentzer M, Givskov M, Parsek MR. 2006. The impact of quorum sensing and swarming motility on *Pseudomonas aeruginosa* biofilm formation is nutritionally conditional. *Molecular Microbiology* **62**:1264–1277. DOI: <https://doi.org/10.1111/j.1365-2958.2006.05421.x>, PMID: 17059568
- Starkey M**, Hickman JH, Ma L, Zhang N, De Long S, Hinz A, Palacios S, Manoïl C, Kirisits MJ, Starner TD, Wozniak DJ, Harwood CS, Parsek MR. 2009. *Pseudomonas aeruginosa* rugose small-colony variants have adaptations that likely promote persistence in the cystic fibrosis lung. *Journal of Bacteriology* **191**:3492–3503. DOI: <https://doi.org/10.1128/JB.00119-09>, PMID: 19329647
- Stewart PS**, Franklin MJ. 2008. Physiological heterogeneity in biofilms. *Nature Reviews Microbiology* **6**:199–210. DOI: <https://doi.org/10.1038/nrmicro1838>, PMID: 18264116
- van Gestel J**, Vlamakis H, Kolter R. 2015. Division of labor in biofilms: the ecology of cell differentiation. *Microbiology Spectrum* **3**. DOI: <https://doi.org/10.1128/microbiolspec.MB-0002-2014>, PMID: 26104716
- Vogel HJ**, Bonner DM. 1956. Acetylornithinase of *Escherichia coli*: partial purification and some properties. *The Journal of Biological Chemistry* **218**:97–106. PMID: 13278318
- Vroom JM**, De Grauw KJ, Gerritsen HC, Bradshaw DJ, Marsh PD, Watson GK, Birmingham JJ, Allison C. 1999. Depth penetration and detection of pH gradients in biofilms by two-photon excitation microscopy. *Applied and Environmental Microbiology* **65**:3502–3511. PMID: 10427041
- Wei Q**, Ma LZ. 2013. Biofilm matrix and its regulation in *Pseudomonas aeruginosa*. *International Journal of Molecular Sciences* **14**:20983–21005. DOI: <https://doi.org/10.3390/ijms141020983>, PMID: 24145749
- Wessel AK**, Arshad TA, Fitzpatrick M, Connell JL, Bonnacaze RT, Shear JB, Whiteley M. 2014. Oxygen limitation within a bacterial aggregate. *mBio* **5**:e00992. DOI: <https://doi.org/10.1128/mBio.00992-14>
- West SA**, Cooper GA. 2016. Division of labour in microorganisms: an evolutionary perspective. *Nature Reviews Microbiology* **14**:716–723. DOI: <https://doi.org/10.1038/nrmicro.2016.111>, PMID: 27640757
- Zhao K**, Tseng BS, Beckerman B, Jin F, Gibiansky ML, Harrison JJ, Luijten E, Parsek MR, Wong GCL. 2013. Psl trails guide exploration and microcolony formation in *Pseudomonas aeruginosa* biofilms. *Nature* **497**:388–391. DOI: <https://doi.org/10.1038/nature12155>, PMID: 23657259