

Mechanosensing of shear by Pseudomonas aeruginosa leads to increased levels of the cyclic-di-GMP signal initiating biofilm development

Christopher A. Rodesney^a, Brian Roman^a, Numa Dhamani^a, Benjamin J. Cooley^{a, 1}, Parag Katira^b, Ahmed Touhami^c, and Vernita D. Gordon^{a,d,2}

^aDepartment of Physics, Center for Nonlinear Dynamics, The University of Texas at Austin, Austin, TX 78712; ^bMechanical Engineering Department, San Diego State University, San Diego, CA 92182-1323; 'Department of Physics, The University of Texas Rio Grande Valley, Brownsville, TX 78520; and ^dInstitute for Cellular and Molecular Biology, The University of Texas at Austin, Austin, TX 78712

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Biofilms are communities of sessile microbes that are phenotypically distinct from their genetically identical, free-swimming counterparts. Biofilms initiate when bacteria attach to a solid surface. Attachment triggers intracellular signaling to change gene expression from the planktonic to the biofilm phenotype. For Pseudomonas aeruginosa, it has long been known that intracellular levels of the signal cyclic-di-GMP increase upon surface adhesion and that this is required to begin biofilm development. However, what cue is sensed to notify bacteria that they are attached to the surface has not been known. Here, we show that mechanical shear acts as a cue for surface adhesion and activates cyclic-di-GMP signaling. The magnitude of the shear force, and thereby the corresponding activation of cyclic-di-GMP signaling, can be adjusted both by varying the strength of the adhesion that binds bacteria to the surface and by varying the rate of fluid flow over surface-bound bacteria. We show that the envelope protein PilY1 and functional type IV pili are required mechanosensory elements. An analytic model that accounts for the feedback between mechanosensors, cyclic-di-GMP signaling, and production of adhesive polysaccharides describes our data well.

mechanosensing | biofilm | cyclic-di-GMP | Pseudomonas aeruginosa | mechanobiology

In higher eukaryotes, mechanosensing and mechanotransduction
are well established to be of widespread importance (1, 2). In are well established to be of widespread importance (1, 2). In contrast, very little is known about how prokaryotes respond to mechanical inputs. However, bacteria adapt to a wide range of mechanically differentiated and changing environments (3). An important example is biofilm development, in which the transition from suspension in a fluid environment to adhesion to a solid substrate is associated with radical changes in intracellular signaling and gene expression. What cue is sensed to notify bacteria that they have become attached to a surface is not known. Notably, transitioning from fluid suspension to surface attachment can result in an abrupt increase in the shear experienced by bacteria. Increased shear can arise from fluid flow and from bacterial motility (Fig. $S1A$ and B). In both these cases, adhesive forces binding one side of a bacterium to the surface will resist displacement and thereby result in shear. It was recently found that surface-attached Escherichia coli can respond to shear by increasing virulence (4). In this work, we investigate the role of shear as a cue to activate signaling to trigger biofilm development.

Pseudomonas aeruginosa is a biofilm-forming, opportunistic human pathogen (5). When planktonic *P. aeruginosa* cells attach to a solid surface, intracellular levels of cyclic-di-GMP (c-di-GMP) increase (6–8). c-di-GMP is a second messenger used by P. aeruginosa and many other bacteria to regulate the expression of genes associated with biofilm initiation (9). What specific cue is sensed upon surface adhesion and leads to increased c-di-GMP signaling is not known. This is a significant gap in our understanding of a fundamental microbiological process.

Recent work has suggested that P. aeruginosa and other bacteria may be able to sense mechanical changes in their environment through mechanosensitive proteins in their cell envelopes (10–13).

The envelope protein PilY1 has been identified as a key mediator of surface-associated behaviors and has been suggested as a possible mechanosensor (14). PilY1 has a region with weak similarity to the mechanoresponsive von Willebrand factor A domain (12), widely found in higher eukaryotes (15). In addition, type IV pili have recently been suggested as possible force sensors in a proposed mechanochemical model for virulence regulation in P. aeruginosa by a different intracellular signal, cyclic AMP (14, 16). Type IV pili extend, attach, and retract to exert forces that allow P. aeruginosa to move laterally across surfaces in a motility mode known as "twitching" (17, 18). Type IV pili have been associated with c-di-GMP signaling and biofilm development in Clostridium difficile (19) and P. aeruginosa (11).

Here, we show that shear cues P . *aeruginosa* that it is adhered to a surface and thereby results in increased c-di-GMP signaling. We demonstrate this in two ways: (i) modulating the shear force arising from bacterial motility on surfaces by modulating the polymermediated adhesion to the surface, and (ii) imposing variable external shear by varying fluid flow (Fig. $S1 \land A$ and B). By using fluid flow to impose shear with a magnitude and time-dependence that we control, we can distinguish the effects of twitching motility from those of environmental forces, and we can distinguish the effects of mechanical coupling from those of mechanosensing. We propose an analytic model for bacterial surface sensing that incorporates inputs from both surface adhesion per se and external shear stress, as well as a biological feedback loop between c-di-GMP and adhesion.

Significance

It is well established that mechanical inputs, such as strain and elasticity, can be sensed by eukaryotic cells and can impact phenotype and behavior. In contrast, very little is known about how prokaryotes may respond to mechanical inputs. Here, we show that bacteria can sense shear and can respond by initiating biofilms. This is an important advance in fundamental microbiology and mechanobiology. Biofilms are difficult to prevent using extant approaches. Our knowledge points the way to a hitherto-undeveloped type of antibiofilm surface that thwarts mechanosensing by not sustaining sufficiently high shear. This would prevent bacteria from sensing surface attachment, activating cyclic-di-GMP signaling, and forming a biofilm.

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¹Present address: Department of Physics and Astronomy, University of Georgia, Athens, GA 30602.

²To whom correspondence should be addressed. Email: [gordon@chaos.utexas.edu.](mailto:gordon@chaos.utexas.edu)

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Results

The Pel Polysaccharide Tightens Coupling to the Surface and Increases c-di-GMP Levels in Surface-Adhered Bacteria. Planktonic P. aeruginosa are coated in sticky extracellular polysaccharides (EPSs) that promote surface adhesion (20–22). The P. aeruginosa wild-type (WT) laboratory strain PAO1 in vitro makes two EPS materials, Psl and Pel; the isogenic transposon mutant Δpel does not produce Pel. We have previously shown that Pel modulates the geometry of surface attachment by making it more likely that cells will lie down flat, maximizing the area in contact with the surface (Fig. $S1\,C$ and $D(21)$ $D(21)$. We have also used atomic force microscopy (AFM) to pull bacteria off a surface and found that, near the surface, force increases more steeply with displacement for WT than for Δpel ([Fig. S2](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF2)) (21). Thus, Pel production increases both geometric and mechanical coupling to the surface.

Here, we investigate how Pel production is linked to surface sensing and c-di-GMP production. Conventional c-di-GMP studies have relied on discrete measurements of c-di-GMP levels before and after some specific event, with measurements typically separated on the timescale of hours (8, 11, 12, 23, 24). Therefore, the dynamics of c-di-GMP activation have been largely unknown. We determine c-di-GMP dynamics by using time-lapse confocal fluorescence microscopy of bacteria containing the plasmid pCdrA::gfp, which is a verified reporter for c-di-GMP; cdrA, and thus green fluorescent protein (GFP), is upregulated by c-di-GMP (25). We cycle between subpopulations on the surface, so that each subpopulation is imaged at 30-min intervals.

In the first 30 min following attachment to the surface, reporter populations of WT and Δpel both have a sharply peaked GFP intensity distribution (Fig. $1 \nA$ and E). We interpret this as the footprint of residual c-di-GMP levels from planktonic culture. The finding that WT and Δpel populations have the same reporter brightness distribution indicates that loss of pel functionality has not intrinsically disrupted the c-di-GMP production machinery.

Beginning 30 min after attachment, the brightness distribution for WT populations broadens before slowly narrowing (Fig. 1 B–D); the broadening indicates that a large subpopulation of WT cells achieves high production of c-di-GMP compared with the planktonic state. The 30-min delay likely reflects the timescales necessary for adhesion-triggered increase in c-di-GMP plus the timescales for GFP production and folding (26). The brightness distribution for Δ *pel* populations broadens much less upon surface adhesion than does the brightness distribution for WT populations (Fig. 1 F–H). WT and Δpel distributions remain significantly different for at least 240 min (Fig. 1I). For a threshold brightness of 550 a.u., for the first 30 min after attachment, less than 15% of either population is above threshold. After 90 min, the above-threshold fraction more than doubles for Δpel and more than triples for WT (Fig. 1J). After 2 h, the above-threshold fraction of the Δpel population falls back to its initial level (below 15%). The above-threshold fraction of the WT population is still about 30% even after 4 h.

c-di-GMP levels averaged over the entire population peak at about 60 min after surface attachment and then decrease toward a base level (Fig. 2A). Thirty minutes after adhesion, the average intensity of WT reporters has increased by 61%, whereas the average intensity of Δpel reporters has increased by only 30%. Moreover, WT achieve a peak increase in brightness of 74% compared with 50% for Δpel, and subsequently the WT drop toward a baseline level approximately three times more slowly than do the Δpel.

Our results show that Pel production enhances the magnitude and lifetime of the response to the surface without intrinsically changing c-di-GMP production for bacteria that are not attached to a surface. Moreover, our results show that the time course of c-di-GMP signaling is nonmonotonic.

Slower Surface Motility, Indicative of Greater Friction, Is Associated with Stronger Surface Sensing. The pili-driven twitching motility of P. aeruginosa can take on two distinct modes: flat "crawling," where

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Fig. 1. Histograms of average fluorescence intensity per cell for WT (A-D) and Δpel (E–H) populations during 30-min intervals that begin at the times indicated. For each histogram, $n \geq 100$ cells from at least five independent experiments. Less than 30 min after surface attachment, WT (A) and Δpel (E) histograms are indistinguishable (two-sample Kolmogorov–Smirnov test, $P =$ 0.87). After 30 min, both WT and Δpel populations have broadened toward higher intensities, with the WT showing a larger change. (Insets) Sample confocal micrographs taken during each time window. This time series of micrographs tracks the brightness of one WT bacterium and one Δpel bacterium and their daughter cells. Purple, lower intensity; yellow, higher intensity. (I) P values from comparing WT and Δpel populations using a two-sample Kolmogorov– Smirnov test. (J) The percentage of cells in each population that have a mean intensity brighter than 550 a.u.

the cell is lying flat on the surface, and upright "walking," where the cell is in contact with the surface at only one pole (Fig. $S1 C$ and D, and Fig. 3 A and B) (27). We see that both WT and Δpel retain surface motility, demonstrating that pilus function is not lost with the loss of pel. Because the force-generating capabilities of pili are intrinsic to the proteins that they are made of, we expect WT and Δpel to have the same pilus machinery and function. We further note that WT, on average, move about 25% more slowly on the surface than do Δpel (Fig. 3C). This discrepancy is even greater for cells that spend most of their lifetime lying flat. In contrast, for cells that spend most of their lifetime attached by only one end, WT and Δ *pel* do not have a significant difference in speed (Fig. 3*D*). From these findings, we conclude that there is a friction-like force

Fig. 2. Pel production and mechanical shear both increase intracellular c-di-GMP levels. (A) Average intensity vs. time for the first 250 min after surface attachment. Both WT and Δpel (data points, left axis) start at the same level, sharply increase during the first hour, and then decrease. Corresponding curves are from our model (solid lines, right axis). (B) Peak intensity vs. applied shear stress. Each data point is the average of all cells imaged in the interval 30–120 min after surface attachment, corresponding to the peaks in A (gray shaded region). $n \ge 100$ cells, from at least two independent experiments. Error bars are SEM.

between the cells and the surface that originates from the polymers coating the cell body. The magnitude of this dissipative force increases with the adhesion area, which also increases the number of polymer contacts involved. We expected a priori that disruption of pel function should have no impact on the force generated by the pilus motor. This is supported by our finding that WT and Δpel have indistinguishable motility when average adhesion area, and therefore polymer-mediated friction, is low. Therefore, we infer that WT experience greater dissipative interactions with the surface, and therefore greater shear stress, than do Δpel.

Individual type IV pili exert forces of ∼100 pN (28); the collective activity of several pili, plus the polymer-mediated force opposing motion, plus hydrodynamic resistance, sum to give the net motility-associated force on a bacterium (29, 30). The net force is a shear force because the polymer-mediated force acts only where the bacterium is attached to the surface.

Mechanical Shear Increases c-di-GMP Levels. The association between loss of Pel, reduced c-di-GMP response to surface attachment, and reduced motility-associated shear, leads to the hypothesis that P. aeruginosa may be sensing mechanical shear as a cue that it is attached to the surface. Therefore, we apply external shear by varying the flow rate of liquid media over surface-attached bacteria. We measure c-di-GMP levels over the time window from 30 to 120 min after surface attachment, because this corresponds to the peak of the c-di-GMP time course for both WT and Δpel (Fig. 2A). We find that externally applied shear increases the peak intracellular levels of c-di-GMP for both WT and Δpel, in lockstep with each other (Fig. 2B). This serves as further evidence that the cellular machineries for surface sensing and for c-di-GMP production are not intrinsically perturbed by loss of pel function. This experiment also allows us to distinguish mechanical coupling, which is impacted by loss of Pel, from mechanosensing, which is not.

The maximum shear force on cells for which we measure c-di-GMP reporter intensity is only 0.002 pN, and we have seen cells adhered strongly at shear forces up to 0.02 pN. This is four orders of magnitude lower than pilus-generated forces. Thus, it is not surprising that we find that flow rarely removes cells from the surface $(Fig. S4)$ $(Fig. S4)$ and that bacteria translate freely even at high flow rates. The finding that even our highest flow rates do not remove bacteria from the surface also shows that the surface responses we measure do not arise from selective removal of weakly adhering or low-signaling subpopulations. However, other factors, such as rapid turnover of the medium and removal of bacterial products, may become important at high flow rates. This may be the cause of the decrease in reporter intensity at very high flow rates (Fig. 3B).

Shear Further Elevates c-di-GMP Levels in Populations Already Responding to Surface Adhesion. The finding that shear acts as a cue for surface adhesion raises the possibility that bacteria may be able to respond differentially to surface adhesion and to changes in external shear. To disentangle the effects of surface adhesion alone from surface adhesion coupled with externally applied shear stress, we allow WT cells to attach to a surface in the absence of flow. These bacteria experience an initial c-di-GMP response that peaks ∼60 min after surface attachment. Then, 150 min after surface attachment, we turn on flow to apply a shear stress of 0.013 pN/ μ m². This begins a second increase in c-di-GMP that we attribute directly to the mechanical input provided by shear flow (Fig. 4, hollow black circles). The second increase does not exactly reproduce the response profile seen when constant flow is applied from the beginning. Rather, the second increase has an intensity that is close to that of the response under constant flow at the observed time (hollow blue circles). This suggests that a timedependent response function may be activated upon initial adhesion that acts as an envelope to limit the magnitude of the response to external shear. Thus, the response under time-varying flow is fundamentally different from the response curves for no flow (filled red circles) and constant flow.

Functional Type IV Pili and PilY1 Are Required Mechanosensory **Elements.** We find that $\Delta p \, i$ bacteria are nonmotile on surfaces and do not increase c-di-GMP in response to surface attachment in the absence of external shear [\(Fig. S6](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF6)A). We also find that ΔpilY1 bacteria do not respond to external shear with any increase in c-di-GMP. Thus, we find that PilY1 is a required component for the mechanosensing system that is required for the c-di-GMP response to surface adhesion and shear.

ΔpilA is deficient in production of PilA, the major subunit for type IV pili (31), and thus cannot form type IV pili. Therefore, Δp *ilA* bacteria are nonmotile on surfaces. We found that Δp *ilA* cells adhere to the surface, in line with our previous findings (21). However, ΔpilA do not show a significant increase in c-di-GMP levels upon surface adhesion and also do not increase c-di-GMP in response to external shear ([Fig. S6](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF6)B).

Type IV pili are involved in the transport of PilY1 to the outer cell envelope (11). Therefore, our data showing that $\Delta p i A$ do not respond to surface adhesion with increased c-di-GMP are not an unambiguous indication that type IV pili are a requisite mechanosensor. Therefore, we repeated these experiments with ΔpilT bacteria, which are hyperpiliated but cannot produce the pilus retraction motor PilT and therefore are also nonmotile on

Fig. 3. Without Pel, cells move more quickly on surfaces. (A) Phase-contrast micrograph of PAO1 Δpel cells. The cell on the Left is fully in focus, lying flat on the surface. The circled cell on the Right is partially out of focus, tilting up off of the surface. (B) Schematic showing a side view of flat-lying and tilting cells. (C) Average instantaneous speed of WT and Δpel cells. n ≥ 80,000 data points, from three experiments. ***P < 0.001 from two-sample t test; error bars are SEM. (D) Scatter plot showing the mean speed of cells during an entire trajectory as a function of the fraction of the cell's tracked lifetime that was spent tilting. Linear fits are shown. Below a lifetime tilting fraction of 0.67, the fits lie outside of the 95% confidence intervals of each other, and Δpel are faster than WT. Detail for fit lines is shown in [Fig. S3](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF3).

the surface. If pilus functionality were necessary for force generation but not for force sensing, then Δp ilT cells would be unable to respond to surface adhesion per se, but would show a response to external force. However, PAO1 ΔpilT also failed to respond both to surface adhesion and to externally applied shear ([Fig. S6](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF6)C). This suggests that mechanosensing requires either pilus retraction functionality, or a structural feature disrupted in ΔpilT such as that required for surface deposition of PilY1.

Therefore, we conclude that both PilY1 and fully functional pili are necessary for a c-di-GMP surface-sensing response.

Modeling. Thus far, we have found that mechanical shear is a cue that increases c-di-GMP production and that the magnitude of the shear will depend on the EPS coating the bacteria. This implies a feedback loop, because increased c-di-GMP levels increase EPS production (23). To assess the degree to which a few-parameter model incorporating such a feedback loop can describe our experimental results, we model c-di-GMP time course using coupled differential equations for force on activated mechanosensors increasing c-di-GMP production (Eq. 1), EPS increasing sensor activation (by increasing coupling to the surface) (Eq. 2), and c-di-GMP increasing EPS production (Eq. 3):

$$
\frac{\partial C}{\partial t} = \alpha_1 Y_1 - \gamma_1 C + \alpha_5,
$$
 [1]

$$
\frac{\partial Y_1}{\partial t} = \alpha_3 E - \gamma_2 Y_1,\tag{2}
$$

$$
\frac{\partial E}{\partial t} = \alpha_4 C - \gamma_3 E. \tag{3}
$$

The model incorporates three biological factors: c-di-GMP (C), EPS (E) , and force sensors (Y_1) (Table 1). Parameters describe the coupling between biological factors (α values) and the constitutive rates of production and degradation (γ values) [\(Table S1\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=ST1).

The initial number of c-di-GMP molecules was estimated from ref. 32, and the number of force sensors and EPS were estimated using geometrical considerations. Initial parameter values were an ansatz to achieve qualitatively similar curves to our experimental data. Due to the simplified, black-box nature of our model, it is difficult to ascribe the actions of a particular protein or

gene to a given parameter. Rather, each parameter is likely a combination of the actions of multiple, possibly interconnected, genes and their products. The model posits that WT cells will activate more force sensors than Δpel cells upon adhesion to the surface because WT have stronger coupling to the surface than do Δ *pel*. The parameter α_1 describes the rate at which an activated force sensor triggers production of c-di-GMP. In our model, this includes the effects of variable force, so that α_1 is lower for cells in static culture and increases with increasing shear.

This model has a large number of free parameters, but it is not underdetermined because we can vary EPS production and the magnitude and time-dependence of shear to generate an arbitrarily large number of distinct c-di-GMP time courses. Keeping initial values for C and E constant, we fit Y_1 for both WT

Fig. 4. Shear stress can "restart" c-di-GMP response in surface-sensitive cells. c-di-GMP response for WT PAO1 as a function of time. Cells in static (filled red) and constant shear stress (hollow blue) conditions show a characteristic rise and monotonic decrease in time. Cells in semifilled black circles were grown in static culture for 150 min, after which point a constant shear stress was applied and fluorescence sharply increases. $n \geq 300$ cells, from at least three independent experiments. Error bars are SEM. Corresponding curves are from our model (solid and dashed lines, right axis). The c-di-GMP increase is not due to the influx of fresh media, as refreshment of media without a sustained shear stress does not elicit a response [\(Fig. S5\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF5).

Table 1. Biological factors in our model for mechano-activation of c-di-GMP

Biological factor	Initial value	Description
ϵ	120	No. of c-di-GMP molecules
	1.939 (WT)	No. of activated surface sensors
	$1,060 \ (\Delta \text{pel})$	
	12	Amount of EPS

and Δ *pel*. WT data from constant-flow experiments [\(Fig. S7\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF7) was then used to fit new α_1 values for each flow rate [\(Table S2](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=ST2)). These values were used to plot curves for both WT and Δpel [\(Fig. S7](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF7)). For WT at all external-shear values and for Δ*pel* at zero external shear and high external shear, the model curves are in excellent agreement with the data (Fig. 2 and [Fig. S7\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF7). For Δpel at intermediate-shear values, the agreement is less exact, but the trends are correct and the models in this case are pure predictions, using no free parameters. Furthermore, our model also describes the shape of the curve when external shear flow is activated 150 min after attachment to the surface (Fig. 4).

Thus, this basic model predictively describes the most salient features of our experimental results.

Discussion

Our primary finding is that *P. aeruginosa* uses mechanosensing to respond to surface attachment by increasing the production of c-di-GMP, an intracellular signal that initiates the transition from the planktonic to the biofilm phenotype. The EPS Pel enhances surface sensing by increasing friction-like interaction with the surface, and thereby increasing the shear resulting from twitching motility driven by type IV pili. Pel decreases the net mechanical work required to detach individual bacteria from a surface ([Fig. S8\)](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF8) (21). Strengthening the surface-sensing response of newly adhered cells might overcome any disadvantage of easier detachment, because strong c-di-GMP signaling should enhance transition to the biofilm phenotype. The difference in c-di-GMP signaling between WT and Δ *pel* may cause the reported difference in WT and Δ *pel* biofilm phenotype (33). We have shown that PAO1 can respond to shear stresses on the order of millipascals. The shear stresses and rates we apply are 2–100 times smaller than stresses commonly studied for biofilm formation or found in host physiology (34–36). This suggests that PAO1 and other bacteria may be remarkably sensitive to mechanical stresses in their environment.

Our results also show that the time course of the c-di-GMP signaling response to surface adhesion is nonmonotonic (Fig. 2A). This suggests that c-di-GMP may act as a toggle switch, wherein transient high signal levels set gene expression patterns to the biofilm state, which is then maintained at lower signal levels (37). Higher eukaryotes have mechanoresponsive toggle switches (38). Genetic toggle switches have been analyzed theoretically in systems biology, with models verified using switches constructed in E. coli and Bacillus subtilis (39–41). These switches settle into one of two bistable states, with a single spike in chemical or thermal cues serving to flip from one state to the other. Unlike a toggle switch, a pressure switch would require sustained high levels of c-di-GMP throughout the lifetime of the biofilm. Thus, use of a toggle switch instead of a pressure switch would reduce the metabolic burden of continually producing c-di-GMP, freeing biofilm cells to expend energy in other pursuits.

We find that PilY1 is a required mechanosensory element for c-di-GMP response to surface adhesion and shear. This agrees with Luo et al. (11) who found that PilY1 provides a signal to the diguanylate cyclase SadC to activate c-di-GMP production in surface-attached cells, but apparently contradicts Kuchma et al. (12), who found that deleting *pilY1* had no effect on c-di-GMP pools within bacteria. However, Kuchma et al. conducted their experiment on planktonic cells; we also see no difference in c-di-GMP levels for just-attached

populations of WT and ΔpilY1. Cell wall deformation has been suggested as a means of sensing surface adhesion. However, this deformation and subsequent triggering of mechanosensitive channels takes longer than an hour (42), too long to be responsible for the fast responses we see. This further implicates PilY1 as a surface-sensing element, not intrinsic to the c-di-GMP production machinery.

We see that $\Delta p i/T$ mutants, lacking pilus retraction, are deficient in a c-di-GMP response. Luo et al. find that knocking out pilus retraction via a $\Delta p i l U$ mutant does not completely block surface sensing (11). However, they use cAMP for their measure of surface response. They further propose a signaling cascade where cAMP is produced upstream of PilY1 and c-di-GMP. It is consistent with our findings that pilus retraction is not required for cAMP production, but may be necessary for downstream c-di-GMP production. In addition, our results also show that $\Delta p \mathcal{A}$, which altogether lack type IV pili, have constitutively higher levels of c-di-GMP production than do WT [\(Fig. S6](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF6)A). This hints at another potential purpose for the PilA protein and/or the pilus structure in c-di-GMP production.

Identification of mechanical stress as the cue leading to biofilm development points to the possibility of a new type of approach for engineering antibiofilm surfaces. Extant approaches to preventing biofilms focus on developing surfaces that resist bacterial attachment or are bactericidal (43–45). Without c-di-GMP, P. aeruginosa that otherwise form robust biofilms within 12 h instead form no biofilm whatsoever over at least 72 h (24). Our work here suggests that surfaces that do not sustain mechanical stress (that behave as 2D fluids or very soft solids) should hinder transition to the biofilm phenotype, and should do so by targeting highly conserved mechanisms for which evolutionary escape is likely to be difficult.

Our work also suggests that the mechanical stresses from the environment, such as shear stress by external flow, to which bacteria are subjected in early biofilm initiation, may impact properties of the mature biofilm. For several bacterial species, biofilms that are grown under high shear are more elastic and denser in matrix proteins and EPS than biofilms grown under low shear (34, 35, 46). Thus, early enhancement in c-di-GMP time course might allow biofilm-forming bacteria to adapt the strength and resilience of the biofilms formed to better resist mechanical removal.

Materials and Methods

Bacteria and Media. We used WT P. aeruginosa strain PAO1 and four singlegene transposon mutants in the PAO1 background, Δpel, ΔpilA, ΔpilT, and ΔpilY1 (47). To be able to measure c-di-GMP levels, we transformed strains with the reporter plasmid pCdrA:: gfp (25). This plasmid causes production of GFP to be increased when transcription of the cdrA gene increases in response to high intracellular levels of c-di-GMP. Strains transformed with a promotorless control plasmid pMH487 were used to measure background GFP expression. Bacteria were grown as described earlier, and time-lapse phase-contrast microscopy data acquired earlier were analyzed in this work to determine the relationship between EPS expression, tilting, and cell motility (21). See [SI Materials and Methods](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=STXT).

Laser-Scanning Confocal Fluorescence Microscopy. For measurements of c-di-GMP signaling, we volumetrically diluted the culture by either 500× or 50,000 \times . Samples for static experiments were prepared in the same way as for phase-contrast microscopy. For flow experiments, 500x diluted cultures were inoculated into flow cells ($2 \times 4 \times 40$ mm) and allowed to adhere to the surface of an untreated glass coverslip for 30–40 min, after which a peristaltic pump (Watson Marlow) began flow. Volumetric flow rates varied from 0.065 to 7.7 mL/min. Bacteria were imaged using an Olympus FV1000 motorized inverted IX81 microscope suite; the stage was enclosed by an incubation chamber heated to 37 °C. We use a 100 \times oil-immersion objective and bacteria were illuminated with a 488-nm laser using standard GFP filter sets. Image capture was controlled by FV10-ASW, version 3.1, software. Confocal Z stacks were taken with a depth of ∼4 μm and a spacing of 340 nm. For 50,000x dilutions and flow experiments, Z stacks were taken every 30 min for 4 h. The 500x dilutions were used to increase the ease with which cells could be found and imaged, so for static experiments Z stacks were taken every minute for the first 90 min, and for flow experiments, Z stacks were taken every 30 min for 4 h.

Confocal Z stacks were projected in the vertical direction to create images in the x–y plane. The projected length and the mean intensity of each bacterium were measured using the Fiji distribution of ImageJ software (48). The WT and Δpel populations have very similar distributions of projected lengths, which verifies that images of dimmer Δpel cells are not being artifactually truncated by brightness thresholding in a way that would magnify differences between the two populations ([Fig. S9](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=SF9)). The average intensity of each individual cell was computed and recorded along with the time elapsed from initial surface adhesion. Intensities were normalized by subtracting the average per-cell intensity of an ensemble of cells carrying a control plasmid for producing GFP that lacks the cdrA promoter (pMH874) to correct for natural variations in fluorescence between strains.

Calculation of Shear Stress in Flow Experiments. Flow was controlled using a Watson Marlow peristaltic pump, allowing us to set the 3D volumetric flow rate $(Q*_{3D})$. From this, we can calculate the pressure drop per unit length down our channel (49). See [SI Materials and Methods](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=STXT).

- 1. Jaalouk DE, Lammerding J (2009) Mechanotransduction gone awry. Nat Rev Mol Cell Biol 10:63–73.
- 2. Iskratsch T, Wolfenson H, Sheetz MP (2014) Appreciating force and shape—the rise of mechanotransduction in cell biology. Nat Rev Mol Cell Biol 15:825–833.
- 3. Persat A, et al. (2015) The mechanical world of bacteria. Cell 161:988–997.
- 4. Alsharif G, et al. (2015) Host attachment and fluid shear are integrated into a mechanical signal regulating virulence in Escherichia coli O157:H7. Proc Natl Acad Sci USA 112:5503–5508.
- 5. Wolcott R, Dowd S (2011) The role of biofilms: Are we hitting the right target? Plast Reconstr Surg 127:28S–35S.
- 6. Cotter PA, Stibitz S (2007) c-di-GMP-mediated regulation of virulence and biofilm formation. Curr Opin Microbiol 10:17–23.
- 7. Hengge R (2009) Principles of c-di-GMP signalling in bacteria. Nat Rev Microbiol 7: 263–273.
- 8. Gupta K, Liao J, Petrova OE, Cherny KE, Sauer K (2014) Elevated levels of the second messenger c-di-GMP contribute to antimicrobial resistance of Pseudomonas aeruginosa. Mol Microbiol 92:488–506.
- 9. Valentini M, Filloux A (2016) Biofilms and cyclic di-GMP (c-di-GMP) signaling: Lessons from Pseudomonas aeruginosa and other bacteria. J Biol Chem 291:12547–12555.
- 10. Otto K, Silhavy TJ (2002) Surface sensing and adhesion of Escherichia coli controlled by the Cpx-signaling pathway. Proc Natl Acad Sci USA 99:2287–2292.
- 11. Luo Y, et al. (2015) A hierarchical cascade of second messengers regulates Pseudomonas aeruginosa surface behaviors. mBio 6:e02456-14.
- 12. Kuchma SL, et al. cyclic di-GMP-mediated repression of swarming motility by Pseudomonas aeruginosa: The pilY1 gene and its impact on surface-associated behaviors. J Bacteriol 192:2950–2964.
- 13. Blanka A, et al. (2015) Constitutive production of c-di-GMP is associated with mutations in a variant of Pseudomonas aeruginosa with altered membrane composition. Sci Signal 8:ra36.
- 14. Siryaporn A, Kuchma SL, O'Toole GA, Gitai Z (2014) Surface attachment induces Pseudomonas aeruginosa virulence. Proc Natl Acad Sci USA 111:16860–16865.
- 15. Whittaker CA, Hynes RO (2002) Distribution and evolution of von Willebrand/integrin A domains: Widely dispersed domains with roles in cell adhesion and elsewhere. Mol Biol Cell 13:3369–3387.
- 16. Persat A, Inclan YF, Engel JN, Stone HA, Gitai Z (2015) Type IV pili mechanochemically regulate virulence factors in Pseudomonas aeruginosa. Proc Natl Acad Sci USA 112: 7563–7568.
- 17. Jin F, Conrad JC, Gibiansky ML, Wong GCL (2011) Bacteria use type-IV pili to slingshot on surfaces. Proc Natl Acad Sci USA 108:12617–12622.
- 18. Skerker JM, Berg HC (2001) Direct observation of extension and retraction of type IV pili. Proc Natl Acad Sci USA 98:6901–6904.
- 19. Purcell EB, McKee RW, Bordeleau E, Burrus V, Tamayo R (2016) Regulation of type IV pili contributes to surface behaviors of historical and epidemic strains of Clostridium difficile. J Bacteriol 198:565–577.
- 20. Cooley B, et al. (2016) Asymmetry and inequity in the inheritance of a bacterial adhesive. New J Phys 18:045019.
- 21. Cooley BJ, et al. (2013) The extracellular polysaccharide Pel makes the attachment of P. aeruginosa to surfaces symmetric and short-ranged. Soft Matter 9:3871–3876.
- 22. Ma L, et al. (2009) Assembly and development of the Pseudomonas aeruginosa biofilm matrix. PLoS Pathog 5:e1000354.
- 23. Irie Y, et al. (2012) Self-produced exopolysaccharide is a signal that stimulates biofilm formation in Pseudomonas aeruginosa. Proc Natl Acad Sci USA 109:20632–20636.
- 24. Hickman JW, Tifrea DF, Harwood CS (2005) A chemosensory system that regulates biofilm formation through modulation of cyclic diguanylate levels. Proc Natl Acad Sci USA 102:14422–14427.

Model. The model for mechanoactivation of c-di-GMP signaling was implemented and fit in COPASI (50) and iterated deterministically for 250 min using the LSODA (51) method.

AFM Experiments and Analysis. AFM data were taken as described in ref. 21; see [SI Materials and Methods](http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1703255114/-/DCSupplemental/pnas.201703255SI.pdf?targetid=nameddest=STXT). To determine the work of detachment, we numerically integrated the force versus separation curves from zero separation to the separation at which the force first returned to zero.

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- 25. Rybtke MT, et al. (2012) Fluorescence-based reporter for gauging cyclic di-GMP levels in Pseudomonas aeruginosa. Appl Environ Microbiol 78:5060–5069.
- 26. Golding I, Paulsson J, Zawilski SM, Cox EC (2005) Real-time kinetics of gene activity in individual bacteria. Cell 123:1025–1036.
- 27. Gibiansky ML, et al. (2010) Bacteria use type IV pili to walk upright and detach from surfaces. Science 330:197.
- 28. Maier B, et al. (2002) Single pilus motor forces exceed 100 pN. Proc Natl Acad Sci USA 99:16012–16017.
- 29. Touhami A, Jericho MH, Boyd JM, Beveridge TJ (2006) Nanoscale characterization and determination of adhesion forces of Pseudomonas aeruginosa pili by using atomic force microscopy. J Bacteriol 188:370–377.
- 30. Sabass B, Stone HA, Shaevitz JW (2017) Collective force generation by groups of migrating bacteria. arXiv:1701.00524.
- 31. Chang Y-W, et al. (2016) Architecture of the type IVa pilus machine. Science 351: aad2001.
- 32. Christen M, et al. (2010) Asymmetrical distribution of the second messenger c-di-GMP upon bacterial cell division. Science 328:1295–1297.
- 33. Yang L, et al. (2011) Distinct roles of extracellular polymeric substances in Pseudomonas aeruginosa biofilm development. Environ Microbiol 13:1705–1717.
- 34. Araujo P, et al. (2016) Influence of flow velocity on the characteristics of Pseudomonas fluorescens biofilms. J Environ Eng 142:04016031.
- 35. Lemos M, Mergulhao F, Melo L, Simoes M (2015) The effect of shear stress on the formation and removal of Bacillus cereus biofilms. Food Bioprod Process 93:242–248.
- 36. Stewart PS, Franklin MJ (2008) Physiological heterogeneity in biofilms. Nat Rev Microbiol 6:199–210.
- 37. Smits WK, Kuipers OP, Veening J-W (2006) Phenotypic variation in bacteria: The role of feedback regulation. Nat Rev Microbiol 4:259–271.
- 38. Whitfield JF (2008) The solitary (primary) cilium—a mechanosensory toggle switch in bone and cartilage cells. Cell Signal 20:1019–1024.
- 39. Gardner TS, Cantor CR, Collins JJ (2000) Construction of a genetic toggle switch in Escherichia coli. Nature 403:339–342.
- 40. Espinar L, Dies M, Çağatay T, Süel GM, Garcia-Ojalvo J (2013) Circuit-level input integration in bacterial gene regulation. Proc Natl Acad Sci USA 110:7091–7096.
- 41. Tian T, Burrage K (2006) Stochastic models for regulatory networks of the genetic toggle switch. Proc Natl Acad Sci USA 103:8372–8377.
- 42. Li J, et al. (2014) Residence-time dependent cell wall deformation of different Staphylococcus aureus strains on gold measured using surface-enhanced-fluorescence. Soft Matter 10:7638–7646.
- 43. Salwiczek M, et al. (2014) Emerging rules for effective antimicrobial coatings. Trends Biotechnol 32:82–90.
- 44. Graham M, Cady N (2014) Nano and microscale topographies for the prevention of bacterial surface fouling. Coatings 4:37–59.
- 45. MacCallum N, et al. (2015) Liquid-infused silicone as a biofouling-free medical material. ACS Biomater Sci Eng 1:43–51.
- 46. Herbert-Guillou D, Tribollet B, Festy D (2001) Influence of the hydrodynamics on the biofilm formation by mass transport analysis. Bioelectrochemistry 53:119–125.
- 47. Jacobs MA, et al. (2003) Comprehensive transposon mutant library of Pseudomonas aeruginosa. Proc Natl Acad Sci USA 100:14339–14344.
- 48. Schindelin J, et al. (2012) Fiji: An open-source platform for biological-image analysis. Nat Methods 9:676–682.
- 49. Gaver DP III, Kute SM (1998) A theoretical model study of the influence of fluid stresses on a cell adhering to a microchannel wall. Biophys J 75:721-733.
- 50. Hoops S, et al. (2006) COPASI—a COmplex PAthway SImulator. Bioinformatics 22: 3067–3074.
- 51. Petzold L (1983) Automatic selection of methods for solving stiff and nonstiff systems of ordinary differential equations. SIAM J Sci Statist Comput 4:136–148.

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