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## Deletion Mutant Library for Investigation of Functional Outputs of Cyclic Diguanylate Metabolism in *Pseudomonas aeruginosa* PA14

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We constructed a library of in-frame deletion mutants targeting each gene in *Pseudomonas aeruginosa* PA14 predicted to participate in cyclic di-GMP (c-di-GMP) metabolism (biosynthesis or degradation) to provide a toolkit to assist investigators studying c-di-GMP-mediated regulation by this microbe. We present phenotypic assessments of each mutant, including biofilm formation, exopolysaccharide (EPS) production, swimming motility, swarming motility, and twitch motility, as a means to initially characterize these mutants and to demonstrate the potential utility of this library.

**P**seudomonas aeruginosa is a ubiquitous Gram-negative bacterium found in a diverse array of environments, ranging from soil to the mammalian host. As an opportunistic bacterial pathogen, individuals with a compromised immune system, epithelial damage, or cystic fibrosis (CF) can succumb to its colonization and infection (1–3). In addition to its arsenal of virulence factors, including exotoxin A, exoenzyme S, phenazines, rhamnolipids, and lipopolysaccharides, *P. aeruginosa* is also capable of switching its life-style from a planktonic growth mode to a surface-associated life-style in response to environmental stimuli. Biofilm-associated infections have garnered significant attention in the medical context due to the increased antibiotic tolerance of these communities (4–6) and their potential as a reservoir of infection in the host (7, 8).

Cyclic diguanylate (c-di-GMP) is a ubiquitous second-messenger molecule capable of regulating a myriad of cellular functions in bacteria. Studies to date have demonstrated numerous functions regulated by this cyclic dinucleotide, including, but not limited to, bacterial physiology (9–14), group behavior (15–21), and virulence (21, 22). The level of c-di-GMP in the cell is contingent on the activity of two classes of enzymes: diguanylate cyclase (DGC) and phosphodiesterase (PDE). DGCs, with a canonical GG(D/E)EF motif, synthesize c-di-GMP from two molecules of GTP (23, 24), while PDEs, with a canonical E(A/E/V)L motif, degrade c-di-GMP to pGpG (24). More recently, proteins with an HD-GYP domain were also shown to demonstrate c-di-GMP phosphodiesterase activity (25). Current models correlate high intracellular c-di-GMP levels to a sessile life-style, or a biofilm state, while low levels of this signal promote motility and/or planktonic growth. Such a simple model likely underestimates the complexity of the c-di-GMP signaling network, as implied by 40 DGCs and PDEs encoded in the genome of P. aeruginosa PA14.

A surging interest in c-di-GMP and its functional outputs in *P. aeruginosa* has resulted in a large number of publications focused primarily on forward-genetics approaches (9, 13, 16–18, 26–28). For many, the emergence of a publicly available, nonredundant transposon insertion library in *P. aeruginosa* PA14 (29) proved very useful in advancing studies of this microbe. For example, Kulasakara et al. benefited from this library in their genome-wide analysis of DGC and PDE mutants in *P. aeruginosa* PA14 (30). Although useful, drawbacks of transposon insertion mutants are 2-fold: (i) the potential for partial loss of function at the insertion

site and (ii) the possibility of polarity, causing either loss or increased expression of downstream genes.

Here, we constructed a library of in-frame deletion mutants targeting each gene in *P. aeruginosa* PA14 predicted to participate in c-di-GMP synthesis or degradation. We present phenotypic assessments of each mutant, including biofilm formation, exopolysaccharide (EPS) production, swimming motility, swarming motility, and twitch motility, as a means to initially characterize these mutants as well as to demonstrate the potential value of this library to investigators in the field.

#### MATERIALS AND METHODS

Bacterial strains, media, and chemicals. Strains, plasmids, and primers used in this study are listed in Tables S1 to S3 in the supplemental material. *Pseudomonas aeruginosa* strain UCBPP-PA14 (abbreviated *P. aeruginosa* PA14) was used in this study. *P. aeruginosa* PA14 and *Escherichia coli* were routinely cultured in lysogeny broth (LB) (31) at 37°C and, when appropriate, solidified with 1.5% agar and/or supplemented with gentamicin (Gm) at 10  $\mu$ g ml<sup>-1</sup> (*E. coli*) and 50  $\mu$ g ml<sup>-1</sup> (*P. aeruginosa*). Minimal M63 salts (32) supplemented with MgSO<sub>4</sub> (1 mM) plus glucose (0.2%) and Casamino Acids (0.5%) or supplemented with MgSO<sub>4</sub> (1 mM) plus arginine (0.4%) were used for biofilm assays. Swarming, swimming, and twitching motility medium contained M8 salts (33) with glucose (0.2%), Casamino Acids (0.5%), and MgSO<sub>4</sub> (1 mM). Congo red (CR) agar (1.5% agar) medium was the same as that used for swarming, swimming, and twitching motility assays but further supplemented with Congo red (40  $\mu$ g ml<sup>-1</sup>) and Coomassie brilliant blue dye (20  $\mu$ g ml<sup>-1</sup>), as reported previously (34).

**Molecular techniques.** Plasmids constructed during the course of this study were prepared by using homologous recombination in *Saccharomyces cerevisiae* (35). All restriction enzymes were obtained from New England BioLabs (Ipswich, MA). Plasmids constructed in yeast were subsequently extracted by a modified "smash-and-grab" method (36) and

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electroporated into *E. coli* S-17 for confirmation by colony PCR (37). PCR-verified constructs were conjugated into *P. aeruginosa* PA14, as previously reported (15), and exconjugants were selected and counterselected by gentamicin and 5% sucrose, respectively. PCR amplification and subsequent DNA sequencing using primers flanking the site of deletion were performed to verify all resulting mutant candidates.

**Biofilm assays.** Biofilm formation assays on polyvinylchloride (PVC) plates (VWR Scientific, Waltham, MA) were performed as previously reported (38), with the following minor modifications. Bacterial cultures grown overnight in LB were diluted 1:100 in minimal M63 salts supplemented with  $MgSO_4$  (1 mM) and either glucose (0.2%) and Casamino Acids (0.5%) or arginine (0.4%). Plates were incubated overnight at 37°C, followed by staining with 0.1% crystal violet and subsequent solubilization with 30% acetic acid. Solubilized crystal violet was transferred onto a new polystyrene microtiter dish (VWR Scientific, Waltham, MA) and quantified by measuring its optical density at 550 nm. The resulting values were normalized to wild-type (WT) *P. aeruginosa* strain PA14 (positive control) biofilm levels to simplify comparisons across numerous strains.

**Congo red assays.** Congo red assays were performed as previously described (18, 39). Qualitative assessment of Congo red staining was performed considering coloration (gradation from white to red) compared to WT *P. aeruginosa* strain PA14 (positive control) as well as colony morphology (wrinkly versus smooth). Each strain was tested on at least two different days with 3 to 4 technical replicates per day.

Swimming motility assays. Swimming motility assays were performed as previously reported (40), with the following minor modifications. Swim agar (0.3%) plates were poured and left to dry at room temperature ( $\sim$ 25°C) for  $\sim$ 4 h prior to inoculation, with 3 to 4 replicates per strain. Plates were used on the day that they were poured. Each strain was tested on at least three separate days. All swim plates were incubated upright at 37°C for 24 h prior to measurements using ImageJ software (NIH). Swim zones of a strain, measured as the percentage of plate coverage, were normalized to the WT *P. aeruginosa* strain PA14 (positivecontrol) swim zone on the same plate. The resulting values from each set of replicates were averaged to yield a final swim zone coverage value for the particular strain.

Swarming motility assays. Swarming motility assays were performed as previously reported (40), with the following minor modifications. Swarm agar (0.5%) plates were poured and left to dry at room temperature ( $\sim$ 25°C) for  $\sim$ 4 h prior to inoculation. Five to six replicates per strain per trial were tested, and all studies were repeated at least twice on different days. All swarm plates were incubated upright at 37°C for 16 h, followed by 24 h at room temperature. Swarming motility was quantified by using ImageJ software (NIH). Surface coverage was calculated by dividing the area of the swarm by the area of the entire plate, and the values from each replicate were averaged. These values were normalized to values for WT *P. aeruginosa* strain PA14 (positive-control) swarm coverage from the same batch of plates to facilitate comparisons among strains.

It is important to note that even the nonswarming  $\Delta flgK$  mutant is capable of ~10% surface coverage, resulting from the colony formed at the site of inoculation expanding as bacterial cell numbers increase. Thus, strains with swarm coverage of 10 to 20% may indicate no significant change from the negative control.

**Twitch motility assays.** Twitch motility assays were performed as previously reported (41), with minor modifications, as follows. Twitch motility agar (1.5%) plates contained minimal M8 salts supplemented with  $MgSO_4$  (1 mM), glucose (0.2%), and Casamino Acids (0.5%). Cells were stabbed into the bottom of the agar plate by using a toothpick and incubated upright at 37°C overnight, followed by 48 h of incubation at room temperature (~25°C). Each strain was tested on three separate days. After incubation, agar was carefully removed, and the basal surface was stained with 0.1% crystal violet (Sigma, St. Louis, MO) to facilitate visualization of the twitch zones. Twitch zones were determined by using ImageJ software (NIH), by measuring the area covered by the twitch area of WT *P*.

*aeruginosa* PA14 (positive control). The resulting twitch zone values were averaged to yield a final twitch zone value for the particular strain.

**Statistical analysis.** The two-tailed Student *t* test was performed pairwise, between the wild type and each strain, by using GraphPad Prism software (GraphPad, La Jolla, CA).

#### **RESULTS AND DISCUSSION**

**Complete library of strains carrying diguanylate cyclase and phosphodiesterase deletion mutations.** Analysis of the genome of *P. aeruginosa* PA14 revealed 40 genes with predicted GGDEF, EAL, and HD-GYP domains either singly or in combination (30). These c-di-GMP metabolism proteins can be divided into four subclasses: 16 GGDEF-only, 5 EAL-only, 16 dual-domain GGDEF-EAL, and 3 HD-GYP-only proteins (Fig. 1). It is important to note that while alignments of primary amino acid sequences have been largely successful in determining whether a GGDEF/EAL/HD-GYP domain is either functional or degenerate, definitive conclusions require empirical data. For example, *in vitro* analyses showed minimal DGC activity in a seemingly degenerate GDSIF motif in the PA14\_65540 (*fimX*) gene (13) as well as for the PA14\_53140 (*rbdA*) gene, which has the canonical GGDEF motif (28).

The presence of other domains in proteins carrying GGDEF/ EAL/HD-GYP domains can provide some functional context for a particular protein. Therefore, we first performed an analysis of all 40 c-di-GMP metabolism proteins for putative domains using the publicly available SMART algorithm (42, 43). Of the 40 sequences queried, 19 are predicted to be either exported or membrane bound based on the presence of either a transmembrane (TM) or signal domain(s) at the N-terminal region of the protein. These include 8 GGDEF-only (PA14\_20820, PA14\_26970, PA14\_40570, PA14\_49890, PA14\_50060 [roeA], PA14\_53310, PA14\_56280 [sadC], and PA14\_65090), 2 EAL-only (PA14\_14530 and PA14\_36260), and 9 dual-domain GGDEF-EAL (PA14\_03720, PA14\_07500, PA14\_21190, PA14\_37690, PA14\_42220 [mucR], PA14\_53140 [rbdA], PA14\_56790 [bifA], PA14\_60870 [morA], and PA14\_71850) proteins (Fig. 1). We are aware of alternative predictive algorithms, and while such predictions are useful, we emphasize that they are sometimes inconsistent with experimental studies. For example, PA14\_66320 (*dipA*) was shown to be an inner membrane or inner membrane-associated protein (26) despite its lack of obvious transmembrane or signal domains (Fig. 1).

The PAS domain was the most prevalent domain identified in our analysis. Observed in 12 different proteins, the number of PAS domains can range from 1 (e.g., PA14\_03790) to 4 (e.g., PA14\_07500) within a single protein. PAS domains are generally regarded as signal sensors, including signals such as oxygen and redox potential (44, 45). Moreover, PAS domains can also mediate protein-protein interactions (46, 47), perhaps allowing these proteins to participate in larger complexes. Each PAS domain was frequently associated with a downstream PAC domain, with the few exceptions to this organization including PA14\_60870 (*morA*), PA14\_65540 (*fimX*), and PA14\_66320 (*dipA*) (Fig. 1). The PAS-PAC organization is thought to provide structural stability to the PAS domain.

Other domains with putative or known sensor or signal transduction functions are also predicted with high frequency. These domains include REC, HAMP, and CHASE domains. REC domains, typically found in response regulators in two-component signaling systems, are predicted in three GGDEF-only proteins (PA14\_16500 [*wspR*], PA14\_57140, and PA14\_64050), two EAL-

	GGDEF only				EAL only			
PA#	PA14#	F	Predicted domains	PA#	PA14#	Predicted domains		
0169 (siaD)	02110		+	3947 (rocR)	12810			
0290	03790			3825	14530	+		
0338	04420	-		2200	36260	+		
3702 (wspR)	16500			2133	36990			
3343	20820			- (pvrR)	59790			agand
3177	23130				HD-GYP onl	У	L	
2870	26970			PA#	PA14#	Predicted domains	Symbol	Domain
1851	40570			4108	10820			CBS
1120 (yfiN)	49890			2572	30830			
1107 (roeA)	50060			4781	63210			GAF
0847	53310	+					+	Unknown function
4332 (saac)	57140							сс
4843	64050	_						PSB
4929	65090	Ŧ						Signal
5487	72420						1	ТМ
GGDEF + EAL								
PA# PA1		4#		Predict	ted domains			CHASE
0285 0372		20	++-					НАМР
0575	075	00						HD
3311	211	90 70	-+-					EAL
3258	218	70 20				-		GGDEF
2072	315	90 90						РАС
1727 (mucl	R) 422	20						PAS
1433	459	30	_					7ТМ
1181	491	60						REC
0861 ( <i>rbdA</i>	A) 531	40						Repeat
4367 (bifA	) 567	90	+	+				I
4601 (mor	4) 608	70						
4959 (fimX	() 655	j40 <u> </u>						
5017 (dipA	663	20		-0-4				
5295	699	00	-					
5442	718	50		-				

FIG 1 Predicted domains in GGDEF-only, EAL-only, HD-GYP-only, and dual-domain GGDEF-EAL proteins. Based on the SMART algorithm, putative functional domains were identified in each protein (42, 43). Note that the cartoon depiction of each protein is not drawn to scale. The CBS (cystathionine beta-synthase) domain is predicted to serve as a binding site for adenosine and derivatives, as well as playing a regulatory role in regulating enzyme activity; the GAF (cGMP-specific phosphodiesterases, adenylyl cyclases, and EhlA) domain is present in cGMP-specific phosphodiesterases, adenylyl cyclases, and phytochromes as well as FhlA, a regulator of nitrogen fixation in bacteria; the CC (coiled-coil) domain is a structural motif in which  $\alpha$ -helices intertwine; the PSB (periplasmic substrate binding) domain participates in ligand binding; Signal indicates the protein secretion signal; the TM (transmembrane) domain is the inner membrane transmembrane domain; the CHASE domain (extracellular sensory domain) is a predicted ligand binding domain; the HAMP (histidine kinases, adenylyl cyclases, methyl binding proteins, phosphatases) domain is a signal transduction domain; HD indicates the HD-GYP domain, a domain found in c-di-GMP degrading phosphodiesterases; the EAL (phosphodiesterase) domain is found in c-di-GMP-degrading phosphodiesterases; the GGDEF (diguany-late cyclase) domain is a signal transduction domain; the PAS domain fold; the PAS domain is a signal transduction domain is a signal transduction domain; the REC (receiver) domain is a signal transduction domain; the REC (receiver) domain is a signal transduction domain is a signal transduction domain; the REC (receiver) domain is conserved in response regulators; and Repeat indicates repeat sequences.

only proteins (PA14\_12810 [*rocR*] and PA14\_59790 [*pvrR*]), two HD-GYP-only proteins (PA14\_30830 and PA14\_63210), and one dual-domain GGDEF-EAL protein (PA14\_65540 [*fimX*]). HAMP and CHASE domains are less common, predicted in three (two GGDEF-only and one dual-domain GGDEF-EAL) and two (one GGDEF-only and one dual-domain GGDEF-EAL) proteins, respectively. Another domain, 7TM, is also thought to be involved in signal transduction. Described as "7 transmembrane receptors with diverse intracellular signaling modules," it is predicted in just one GGDEF-only protein, PA14\_65090 (Fig. 1).

The GAF domain (regulatory domain in cyclic <u>G</u>MP-specific phosphodiesterases, <u>a</u>denylyl cyclases, and <u>FhlA</u>) is predicted in two dual-domain GGDEF-EAL proteins (PA14\_31330 and PA14\_66320 [*dipA*]). For PA14\_66320 (*dipA*), binding of cyclic AMP (cAMP) to the GAF domain induced its phosphodiesterase activity (26). Whether the same function applies to PA14\_31330 has not yet been explored. Other domains, including CBS (cystathionine beta-synthase) and PSB (periplasmic substrate binding) domains, were predicted in the dual-domain GGDEF-EAL proteins PA14\_21870 and PA14\_07500, respectively. Functions of CC (coiled-coil) (PA14\_63210 and PA14\_60870 [*morA*]) and repeat (PA14\_02110) domains remain unknown for these proteins (Fig. 1).

Medium-specific impact of DGCs and PDEs on biofilm formation. Given the known role of c-di-GMP in biofilm formation, we tested the biofilm formation of each mutant under two growth conditions used routinely in our laboratory. The first, glucose supplemented with Casamino Acids (glucose+CAA), is the standard assay medium used in most of our previously reported studies (16, 17), and glucose is a common growth substrate used by many groups studying biofilm formation (48–51). The second substrate analyzed is arginine, one of the few carbon sources which *P. aeruginosa* can ferment and which we have shown previously boosts the levels of c-di-GMP by ~4-fold compared to growth on glucose+CAA (52). In all biofilms assays, the wild-type strain and the biofilm-deficient, hyperswarming  $\Delta sadC \Delta roeA$  double mutant (18) served as controls.

Among the mutants in the GGDEF-only subclass, the majority of mutants formed biofilms at levels ~60 to 70% of those of the wild type, whereas wild-type levels of biofilm formation were observed for strains carrying the  $\Delta$ PA14\_20820,  $\Delta$ PA14\_40570,  $\Delta$ PA14\_53310,  $\Delta$ PA14\_57140, and  $\Delta$ PA14\_65090 mutations. Apart from  $\Delta$ PA14\_20820, these phenotypes are in agreement with previous findings (30). Two mutants,  $\Delta$ PA14\_50060 (*roeA*) and  $\Delta$ PA14\_56280 (*sadC*), formed the weakest biofilms among this group, at roughly 20 to 50% of wild-type levels (Fig. 2A); these observations are in agreement with previous reports of these two well-characterized mutants (17, 18).

Most of the PDE mutants, including both EAL-only and HD-GYP-only subclasses, showed no significant difference from the wild-type strain when biofilm formation was assessed on glucose+CAA medium. In fact, the  $\Delta$ PA14\_30830 and  $\Delta$ PA14\_63210 HD-GYP mutants formed significantly reduced biofilms, at ~50% of wild-type levels (Fig. 2A). Based on the currently accepted model wherein elevated c-di-GMP levels promote biofilm formation, the fact that we observed reduced biofilm formation when we mutated genes coding for putative c-di-GMP-degrading phosphodiesterases is somewhat surprising. The lack of any biofilm-related phenotype among strains mutated for the production of EAL-only proteins (30) and the various phenotypes among

strains lacking HD-GYP-only proteins (21) have been reported previously.

The last subclass, strains mutated for dual-domain GGDEF-EAL proteins, consisted of strains that were both weaker and stronger biofilm producers than the wild type. In contrast to P. aeruginosa PAO1 carrying a  $\Delta mucR$  mutation, which formed a biofilm equivalent to that formed by the wild type (12), the  $\Delta PA14_{42220}$  (*mucR*) mutant formed the weakest biofilm among mutants in this class, at  $\sim$ 60% of wild-type levels. Strains carrying the  $\Delta$ PA14\_21190,  $\Delta$ PA14\_56790 (*bifA*), and  $\Delta$ PA14\_66320 (dipA) mutations, as expected (30), were all hyperbiofilm producers relative to the wild type, ranging from 160 to 300% of wild-type strain levels (Fig. 2A). Strains carrying  $\Delta$ PA14\_56790 (*bifA*) and  $\Delta PA14_{66320}$  (*dipA*) mutations, in particular, were previously reported to demonstrate enhanced biofilm formation (16, 26) and thus serve to validate our findings. Interestingly, mutation of the homolog of the PA14\_53140 (rbdA) gene was shown to cause a hyperbiofilm phenotype in P. aeruginosa PAO1 (28, 30), but mutation of this gene showed wild-type levels of biofilm in the P. aeruginosa PA14 background (Fig. 2A). Thus, while there are similarities between strains PAO1 and PA14 of P. aeruginosa, these data suggest strain-specific differences in the roles of these proteins in biofilm formation.

As mentioned above, our group has shown that a biofilm medium supplemented with L-arginine (0.4%) promotes c-di-GMP production and, as a consequence, results in more pronounced biofilm formation by *P. aeruginosa* PA14 (52). Under these conditions, L-arginine is utilized as the sole carbon source in the presence of oxygen. Thus, to facilitate observations of minor changes in biofilm formation and potential medium-specific effects of the loss of the DGCs and/or PDEs, we tested the same library of mutants in minimal medium supplemented with arginine.

In general, the GGDEF-only subclass mutants showed negligible effects on biofilm formation in arginine medium. However, two mutants, the previously reported  $\Delta$ PA14\_50060 (*roeA*) and  $\Delta$ PA14\_56280 (*sadC*) mutants (17, 18), showed impaired biofilms, at approximately 40 to 50% of wild-type levels, while the  $\Delta$ PA14\_04420,  $\Delta$ PA14\_20820,  $\Delta$ PA14\_26970, and  $\Delta$ PA14\_64050 mutants produced hyperbiofilms, reaching 110 to 140% of wild-type levels (Fig. 2B).

Similarly, most strains carrying mutations in genes coding for the EAL-only and HD-GYP-only subclasses showed wild-type phenotypes, as shown by  $\Delta$ PA14\_36260,  $\Delta$ PA14\_36990,  $\Delta$ PA14\_ 59790 (*pvrR*), and  $\Delta$ PA14\_30830 mutant strains. However, noticeably weaker biofilm ( $\Delta$ PA14\_12810 [*rocR*] and  $\Delta$ PA14\_ 10820) and hyperbiofilm ( $\Delta$ PA14\_14530 and  $\Delta$ PA14\_63210) phenotypes were also observed (Fig. 2B).

The mutations in genes with dual GGDEF-EAL domains also showed various degrees of biofilm formation, with the  $\Delta$ PA14\_42220 (*mucR*) and  $\Delta$ PA14\_60870 (*morA*) mutants showing the weakest biofilm formation in this class of mutants, at 50 to 60% of wild-type levels. In contrast, the  $\Delta$ PA14\_21190,  $\Delta$ PA14\_ 56790 (*bifA*),  $\Delta$ PA14\_65540 (*fimX*),  $\Delta$ PA14\_66320 (*dipA*), and  $\Delta$ PA14\_71850 mutants produced hyperbiofilms at levels ranging from 110 to 150% of wild-type levels. Other mutants were not significantly different from the wild type, with a few showing a small, but significant, decrease, as seen, for example, with the  $\Delta$ PA14\_31330 mutant (Fig. 2B).

In sum, a total of 16 mutants showed medium-specific changes in biofilm formation. Of these, minimal medium supplemented



FIG 2 Biofilm formation. Shown is an analysis of biofilm formation by *P. aeruginosa* PA14 strains carrying mutations in genes encoding GGDEF-only, EAL-only, HD-GYP-only, and dual-domain GGDEF-EAL proteins in glucose+CAA (A)- and arginine (B)-supplemented media. The biomass of the attached cells on PVC plates was measured after 24 h of static incubation at 37°C. Final values corresponding to each mutant were normalized to the value for wild-type *P. aeruginosa* strain PA14, which was set to a value of 1, for ease of comparison. OD<sub>550</sub>, optical density at 550 nm. \*, P < 0.05.

with arginine increased biofilm formation in 14 of the 16 mutants. The net increase occurred in both hyperbiofilm mutants (e.g.,  $\Delta$ PA14\_66320 [*dipA*]) as well as those with weak biofilms (e.g.,  $\Delta$ PA14\_03790 and  $\Delta$ PA14\_72420), which is consistent with the previously reported arginine-dependent increase in intracellular c-di-GMP levels (52). Surprisingly, two mutants ( $\Delta$ PA14\_60870 [*morA*] and  $\Delta$ PA14\_10820) showed an arginine-dependent decrease in biofilm formation. It is important to note that the phenotypic differences highlighted in this work are unlikely to be attributed to growth defects, as all strains except the  $\Delta$ *bifA* strain, which is known to be a hyperbiofilm former, showed no growth defect in glucose minimal medium (see Fig. S1 in the supplemental material).

Effects of c-di-GMP on exopolysaccharide production in *P. aeruginosa*. Extracellular matrix production is typically associated with biofilm formation; exopolysaccharides (EPS) are a critical component of this matrix. In *P. aeruginosa* PA14, two different types of EPS, Pel and alginate, are synthesized, of which Pel has been reported to be a critical factor for biofilm formation *in vitro* (39). Requiring the activities of the PelA to PelG proteins (53, 54), the level of Pel EPS production is typically positively associated with the intracellular pool of c-di-GMP (16, 18). Thus, investigat-

ing the impact of mutating genes encoding GGDEF-only, EALonly, HD-GYP-only, and dual-domain GGDEF-EAL proteins on EPS production is of particular interest.

The qualitative assay used here exploits the ability of Congo red (CR) dye to bind a range of macromolecules, including its high affinity for cellulose in plants and fungi (55, 56). This dye can also bind amyloids (57). CR has been used to identify mutants defective in the production of Pel in the context of pellicle production and biofilm formation in P. aeruginosa PA14 (16-18, 39). While acknowledging the potential for CR to stain other macromolecules, a Pel-deficient *P. aeruginosa* PA14 mutant ( $\Delta pelA$ ) fails to bind CR (58), resulting in the production of smooth, white colonies, and thus, this mutant serves as a negative control (Fig. 3). In these assays, all strains are scored on a scale from 0 to 3, with the WT being scored as 2, the hyper-CR-binding  $\Delta$ PA14\_56790 (*bifA*) mutant being given a score of 3, and the Pel-deficient  $\Delta pelA$  mutant being assigned a score of 0. The  $\Delta wspR$  mutant, scored as 1, shows an intermediate phenotype between the wild type and the  $\Delta pelA$  mutant (Fig. 3 and Table 1).

There were four mutants that scored "0" ( $\Delta$ PA14\_50060 [*roeA*],  $\Delta$ PA14\_07500,  $\Delta$ PA14\_60870 [*morA*], and  $\Delta$ PA14\_10820) across the four mutant subclasses (Table 1). These mutants



FIG 3 Congo red assays. Shown are representative images of Congo red binding assays for *P. aeruginosa* PA14 and strains carrying mutations in the *pelA*, PA14\_16500 (*wspR*), and PA14\_56790 (*bifA*) genes. As cited in text, the  $\Delta pelA$ mutant, the  $\Delta PA14_16500$  (*wspR*) mutant, *P. aeruginosa* PA14 (wild type), and the  $\Delta PA14_56790$  (*bifA*) mutant are representative of scores of 0, 1, 2, and 3 on the score spectrum, respectively (see Table 1 for a complete list). The  $\Delta PA14_56790$  (*bifA*) mutant also forms a wrinkly colony morphology, which is indicated by "W" in Table 1.

failed to bind any CR following overnight incubation at 37°C and a lengthy incubation ( $\sim 2$  days) at room temperature. The  $\Delta$ PA14\_50060 (*roeA*) mutant was previously reported to show no CR binding (18). The next class, scored as "1," included eight mutants (ΔPA14\_16500 [wspR], ΔPA14\_20820, ΔPA14\_23130, ΔΡΑ14\_53310, ΔΡΑ14\_36990, ΔΡΑ14\_37690, ΔΡΑ14\_30830, and  $\Delta PA14_{63210}$  across the subclasses. In particular, diminished CR binding of  $\Delta$ PA14 16500 (*wspR*) was expected, given its role as a DGC in P. aeruginosa (59), and thus, this mutant served as a representative of this class of mutants. We observed 17 mutants with the wild-type phenotype and thus scored them as "2." Finally, there were 11 mutants that bound CR to a greater extent than the wild type (generally after overnight incubation at 37°C) and/or had a wrinkly colony morphology, which we interpret as excessive EPS production (Table 1). The previously reported  $\Delta$ PA14\_56790 (bifA) mutant (16) showed strong CR binding and a wrinkly colony morphology and served as the representative of this class of mutants, scored as "3" (Fig. 3 and Table 1). Similarly, the  $\Delta$ PA14\_21190 mutant also formed a wrinkly colony with strong CR binding. In addition to the CR binding score, mutants demonstrating wrinkly colony morphology are indicated with a "W" in Table 1.

Despite the strong association between biofilm formation and EPS production, there was surprisingly little correlation between these two phenotypes. For example, some mutants with low-level CR binding (e.g.,  $\Delta$ PA14\_16500 [*wspR*] and  $\Delta$ PA14\_07500) demonstrated minor defects in biofilm formation under both medium conditions (Fig. 2 and Table 1). Conversely, some mutants with high-level CR binding (e.g.,  $\Delta$ PA14\_31330 and  $\Delta$ PA14\_69900) formed wild-type biofilms (Fig. 2 and Table 1). However, some strains with high-level CR binding and/or wrinkly colony morphology did yield hyperbiofilm phenotypes, for example, the  $\Delta$ PA14\_21190,  $\Delta$ PA14\_56790 (*bifA*), and  $\Delta$ PA14\_66320 (*dipA*) mutants, as reported previously (16, 26). Thus, EPS production, as measured by CR binding, may not be indicative of the mutant's biofilm formation phenotype.

**c-di-GMP impacts flagellar-based swimming motility.** The link between c-di-GMP and motility is well documented (60). Thus, we analyzed each mutant for its impact on swimming motility. Motility was assessed by using a standard soft agar (0.3%) plate assay. It is important to note that plate-based motility assays also measure bacterial chemotaxis in addition to motility.

Of the 16 GGDEF-only mutants, 4 showed an increase  $(\Delta PA14\_16500 [wspR], \Delta PA14\_23130, \Delta PA14\_49890 [yfiN], and$ 

 $\Delta$ PA14\_56280 [*sadC*]) and 4 showed a decrease ( $\Delta$ PA14\_02110 [*siaD*],  $\Delta$ PA14\_03790,  $\Delta$ PA14\_26970, and  $\Delta$ PA14\_72420) in swimming motility compared to the wild type. The remaining mutants showed wild-type swimming motility (e.g.,  $\Delta$ PA14\_0420) (Fig. 4).

Wild-type swimming phenotypes were dominant among mutants of the EAL- and HD-GYP-only subclasses (e.g.,  $\Delta$ PA14\_12810 [*rocR*] and  $\Delta$ PA14\_36990). However, two mutants, carrying mutations in the PA14\_14530 and PA14\_10820 genes, showed significant decreases (~30% of wild-type levels) and increases (~140% of wild-type levels) in swimming motility, respectively (Fig. 4).

TABLE 1 Congo red phenotypes of DGC and PDE mutants<sup>a</sup>

	Congo red
Strain	binding score
WT	2
$\Delta pelA$	0
$\Delta PA14 02110 (siaD)$	3
ΔΡΑ14 03790	3
ΔΡΑ14_04420	3
$\Delta PA14\_16500 (wspR)$	1
ΔΡΑ14_20820	1
ΔΡΑ14_23130	1
ΔPA14_26970	3
ΔΡΑ14_40570	2
$\Delta PA14_{49890}$ (yfiN)	2
$\Delta$ PA14_50060 ( <i>roeA</i> )	0
ΔPA14_53310	1
$\Delta PA14_56280 (sadC)$	3
ΔPA14_57140	2
ΔPA14_64050	2
ΔPA14_65090	2
ΔPA14_72420	2
$\Delta$ PA14_12810 (rocR)	2
ΔPA14_14530	3
ΔPA14_36260	2
ΔPA14_36990	1
$\Delta PA14_59790 (pvrR)$	2
ΔPA14_03720	2
ΔPA14_07500	0
ΔPA14_21190	3 <sup>W</sup>
ΔPA14_21870	2
ΔPA14_31330	3
ΔPA14_37690	1
$\Delta$ PA14_42220 ( <i>mucR</i> )	2
ΔPA14_45930	2
ΔPA14_49160	2
ΔPA14_53140	2
$\Delta PA14_56790 \ (bifA)$	3 <sup>W</sup>
$\Delta$ PA14_60870 (morA)	0
$\Delta PA14_{65540}$ (fimX)	2
$\Delta PA14_{66320} (dipA)$	3
ΔΡΑ14_69900	3
ΔPA14_71850	2
ΔΡΑ14_10820	0
ΔPA14_30830	1
APA14 63210	1

<sup>*a*</sup> The scale ranges from 0 to 3, where 0 is equivalent to the negative control ( $\Delta pelA$  mutant), 1 is represented by the  $\Delta wspR$  mutant, 2 is equivalent to the wild type, and 3 represents hyperbinding of Congo red (*bifA* mutant). Samples of each phenotype are shown in Fig. 3. A superscript "W" indicates a mutant that showed a wrinkly colony phenotype.



FIG 4 Swimming motility. Shown is an analysis of swimming motility of wild-type *P. aeruginosa* strain PA14 as well as strains carrying mutations in genes encoding GGDEF-only, EAL-only, HD-GYP-only, and dual-domain GGDEF-EAL proteins. The area covered by the swimming motility zone was normalized to that of the wild-type strain, which was set to a value of 1, for ease of comparison. The nonmotile  $\Delta flgK$  mutant served as a negative control. \*, P < 0.05.

Finally, 11 of the 16 dual-domain GGDEF-EAL subclass mutants showed reduced swimming motility compared to the wild type. Of these, the motility of six mutants ( $\Delta$ PA14\_07500,  $\Delta$ PA14\_21190,  $\Delta$ PA14\_56790 [*bifA*],  $\Delta$ PA14\_60870 [*morA*],  $\Delta$ PA14\_66320 [*dipA*], and  $\Delta$ PA14\_71850) was significantly reduced (<50%) compared to the motility of the wild type. Two mutants ( $\Delta$ PA14\_03720 and  $\Delta$ PA14\_31330) showed ~70% of wild-type levels of motility. In contrast, three mutants ( $\Delta$ PA14\_2220 [*mucR*], and  $\Delta$ PA14\_65540 [*fimX*]) were hypermotile, at 110 to 120% of wild-type levels (Fig. 4).

As described above, we observed three mutants with a hyperbiofilm phenotype ( $\Delta$ PA14\_21190,  $\Delta$ PA14\_56790 [*bifA*], and  $\Delta$ PA14\_66320 [*dipA*]). These same mutants were impaired in swimming motility (Fig. 4), which is consistent with the general model that high intracellular c-di-GMP levels favor biofilm formation, while low intracellular c-di-GMP levels favor motility. Interestingly, there were two other mutants with similarly impaired swimming motility (e.g.,  $\Delta$ PA14\_72420 and  $\Delta$ PA14\_ 14530) that showed reduced biofilm formation. The complexity underlying c-di-GMP-dependent phenotypes and exceptions to the general model were demonstrated previously (18).

**c-di-GMP regulation of swarming motility.** *P. aeruginosa* PA14 is capable of a second form of flagellum-dependent motility: swarming. Swarming motility also utilizes rhamnolipid surfactants to move across a semisolid surface and is routinely assayed on 0.5% agar plates. Swarming has also been linked to c-di-GMP in *P. aeruginosa* PA14 (16–18, 61).

Among the 16 GGDEF-only mutants, 5 mutants ( $\Delta$ PA14\_02110 [*siaD*],  $\Delta$ PA14\_03790,  $\Delta$ PA14\_04420,  $\Delta$ PA14\_26970, and  $\Delta$ PA14\_72420) swarmed at  $\leq$ 50% of wild-type levels (Fig. 5A). Additionally, two mutants ( $\Delta$ PA14\_23130 and  $\Delta$ PA14\_57140) demonstrated a smaller but significant decrease in swarm coverage. In contrast, five other mutants ( $\Delta$ PA14\_16500 [*wspR*],  $\Delta$ PA14\_20820,  $\Delta$ PA14\_49890 [*yfiN*],  $\Delta$ PA14\_50060 [*roeA*],  $\Delta$ PA14\_53310, and  $\Delta$ PA14\_64050) were hyperswarmer strains, showing surface coverages that were 110 to 150% of wild-type levels (Fig. 5A).

Reduction in swarm coverage was observed for three mutants

( $\Delta$ PA14\_14530,  $\Delta$ PA14\_30830, and  $\Delta$ PA14\_63210) of the EALonly and HD-GYP-only subclasses, with the  $\Delta$ PA14\_30830 mutant showing the highest level of coverage, at 60% of wild-type levels. Three mutants of this class were hyperswarmers ( $\Delta$ PA14\_ 12810 [*rocR*],  $\Delta$ PA14\_36260, and  $\Delta$ PA14\_10820), with 110% of wild-type swarm coverage. The product of the PA14\_12810 (*rocR*) gene was previously linked to Cup fimbria expression, but its contribution to swarming motility was unknown (62, 63).

Mutants with reduced swarm coverage outnumbered hyperswarmers within the dual-domain GGDEF-EAL subclass. In fact, only two mutants in this group ( $\Delta$ PA14\_42220 [*mucR*] and  $\Delta$ PA14\_65540 [*fimX*]) were hyperswarmers, surpassing wild-type surface coverage at 150%. The hyperswarming phenotype of the  $\Delta$ PA14\_42220 (*mucR*) mutant of *P. aeruginosa* PA14 is in contrast to a previous report by Hay et al., wherein  $\Delta$ *mucR* mutant swarming motility was on par with that of wild-type *P. aeruginosa* strain PAO1 (12). Differences in the strains used and/or medium compositions may explain this discrepancy. Seven mutants of the GGDEF-EAL subclass ( $\Delta$ PA14\_07500,  $\Delta$ PA14\_21190,  $\Delta$ PA14\_ 53140 [*rbdA*],  $\Delta$ PA14\_56790 [*bifA*],  $\Delta$ PA14\_60870 [*morA*], and  $\Delta$ PA14\_66320 [*dipA*]) showed reduced swarming. The decreased swarming of the  $\Delta$ PA14\_56790 (*bifA*) and  $\Delta$ PA14\_53140 (*rbdA*) mutants is in line with previous reports (16, 28).

In general, weak swarmers (e.g.,  $\Delta$ PA14\_02110 [*siaD*] and  $\Delta$ PA14\_07500) are also weak swimmers (Fig. 3). Interestingly, the hyperswarming phenotype (e.g.,  $\Delta$ PA14\_50060 [*roeA*] and  $\Delta$ PA14\_53310) did not always translate into a hyperswimming phenotype (Fig. 3). These findings indicate a complex relationship between swimming and swarming motility.

Impact of mutations on pilus-mediated twitching motility. *P. aeruginosa* can also utilize type IV pili (TFP) to move across hard surfaces ( $\geq$ 1% agar) in a process known as twitch motility (41, 64), which is distinct from the two flagellar-dependent modes of motility, swimming and swarming, described above. We investigated the effects of the DGC and PDE mutants on twitching motility. The  $\Delta$ *pilA* mutant, lacking the type IV pilin (65), served as the negative control (Fig. 6).

A small subset of GGDEF-only mutants ( $\Delta$ PA14\_03790,



FIG 5 Swarming motility. Shown is an analysis of swarming motility of wild-type *P. aeruginosa* PA14 as well as strains carrying mutations in genes encoding GGDEF-only, EAL-only, HD-GYP-only, and dual-domain GGDEF-EAL proteins. (A) The area covered by the swarm and its tendrils was measured and normalized to that of wild-type *P. aeruginosa* strain PA14, which was set to a value of 1, for ease of comparison. (B) Representative images of swarming motility. Reduced swarm coverage includes reduced tendrils ( $\Delta$ PA14\_60870 [*morA*]) or tendril-less growth of the colony ( $\Delta$ PA14\_56790 [*bifA*]). The nonmotile  $\Delta$ *flgK* mutant served as a negative control. \*, *P* < 0.05.

 $\Delta$ PA14\_26970, and  $\Delta$ PA14\_72420) demonstrated reduced twitch zones, at <50% of wild-type levels. Six other mutants ( $\Delta$ PA14\_ 04420,  $\Delta$ PA14\_23130,  $\Delta$ PA14\_40570,  $\Delta$ PA14\_56280 [*sadC*],  $\Delta$ PA14\_57140, and  $\Delta$ PA14\_64050) also showed statistically significant decreases in their respective twitch zones but to a lesser degree than the previous group of mutants. In comparison, only one mutant ( $\Delta$ PA14\_20820) had a hypertwitch phenotype, with its twitch zone being ~120% of that of the wild type (Fig. 6).

Several mutations of EAL-only and HD-GYP-only genes conferred a reduction in twitch motility. A total of four mutants ( $\Delta$ PA14\_14530,  $\Delta$ PA14\_36260,  $\Delta$ PA14\_30830, and  $\Delta$ PA14\_ 63210) showed significant decreases in their respective twitch



**FIG 6** Twitch motility. Shown is an analysis of twitching motility by wild-type *P. aeruginosa* strain PA14 as well as strains carrying mutations in genes encoding GGDEF-only, EAL-only, HD-GYP-only, and dual-domain GGDEF-EAL proteins. Twitch zones were measured and normalized to that of wild-type *P. aeruginosa* strain PA14, which was set to a value of 1, for ease of comparison. The  $\Delta pilA$  mutant served as a negative control. \*, P < 0.05.

zones compared to the wild type. In particular,  $\Delta$ PA14\_14530 and  $\Delta$ PA14\_63210 showed twitch zones that were <50% of that of the wild type (Fig. 6).

Of 16 mutants of the dual-domain GGDEF-EAL class, 8 mutants ( $\Delta$ PA14\_21190,  $\Delta$ PA14\_21870,  $\Delta$ PA14\_31330,  $\Delta$ PA14\_ 49160,  $\Delta$ PA14\_56790 [*bifA*],  $\Delta$ PA14\_65540 [*fimX*],  $\Delta$ PA14\_ 69900, and  $\Delta$ PA14\_71850) were statistically significantly impaired for twitch motility, and only 2 ( $\Delta$ PA14\_03720 and  $\Delta$ PA14\_37690) showed larger twitch zones than that of the wild type (Fig. 6). It is also worth noting that our data reproduced a twitch-negative phenotype for a strain carrying PA14\_65540 (*fimX*), a known c-di-GMP-dependent regulator of TFP biogenesis in *P. aeruginosa* (13).

**Conclusions.** Here, we present a library of in-frame deletion mutations targeting the 40 c-di-GMP metabolism proteins identified in the genome of *P. aeruginosa* PA14. Assessment of biofilm formation, swimming motility, swarming motility, twitch motility, and EPS production (via CR binding) was performed by using this mutant library (summarized in Table S4 in the supplemental material). Analysis of these mutants revealed complex relationships among these phenotypes, consistent with the complex nature of the c-di-GMP signaling system. Going forward, this mutant library should serve as a helpful tool for elucidating the role of c-di-GMP metabolism proteins and their regulated pathways.

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