



Serine Hydroxymethyltransferase ShrA (PA2444) Controls Rugose Small-Colony Variant Formation in *Pseudomonas aeruginosa*

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Pseudomonas aeruginosa causes many biofilm infections, and the rugose small-colony variants (RSCVs) of this bacterium are important for infection. We found here that inactivation of PA2444, which we determined to be a serine hydroxymethyltransferase (SHMT), leads to the RSCV phenotype of P. aeruginosa PA14. In addition, loss of PA2444 increases biofilm formation by two orders of magnitude, increases exopolysaccharide by 45-fold, and abolishes swarming. The RSCV phenotype is related to higher cyclic diguanylate concentrations due to increased activity of the Wsp chemosensory system, including diguanylate cyclase WspR. By characterizing the PA2444 enzyme in vitro, we determined the physiological function of PA2444 protein by relating it to S-adenosylmethionine (SAM) concentrations and methylation of a membrane bound methyl-accepting chemotaxis protein WspA. A whole transcriptome analysis also revealed PA2444 is related to the redox state of the cells, and the altered redox state was demonstrated by an increase in the intracellular NADH/NAD+ ratio. Hence, we provide a mechanism for how an enzyme of central metabolism controls the community behavior of the bacterium, and suggest the PA2444 protein should be named ShrA for serine hydroxymethyltransferase related to rugose colony formation.

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INTRODUCTION

Pseudomonas aeruginosa is an opportunistic pathogen that is responsible for many biofilm infections including those associated with ventilator-associated pneumonia, urinary and peritoneal dialysis catheters, bacterial keratitis, otitis externa, lungs (Macé et al., 2008), and burn wounds (Gjødsbøl et al., 2006). Persistence of this bacterium is linked to its ability to form biofilms (Ryder et al., 2007) and rugose small-colony variants (RSCVs) (Drenkard and Ausubel, 2002); for example, approximately 30% of the antibiotic-resistant colonies of *P. aeruginosa* clinical isolate PA14 were RSCVs (Drenkard and Ausubel, 2002). RSCVs have a wrinkled colony morphology, and these cells have elevated aggregation, attachment, and exopolysaccharide (EPS) production (Starkey et al., 2009). Previously, our lab discovered that tyrosine phosphatase TpbA controls the RSCV phenotype in *P. aeruginosa* through diguanylate cyclase (DGC) TpbB (Ueda and Wood, 2009; Pu and Wood, 2010). Our original findings were verified by an independent group which corroborated that TpbB is important for persistence related to cystic fibrosis through RSCVs (Malone et al., 2010). RSCVs are important for infection but their regulation is poorly characterized so it is an important

challenge to identify the molecular mechanisms behind the formation of these variants (Häussler, 2010).

Serine hydroxymethyltransferase (SHMT) is a pyridoxal 5'-phosphate (PLP)-dependent enzyme which has been extensively studied from different species as it is one of the few PLP-dependent enzymes that can be found in all living organisms (Florio et al., 2010). SHMT catalyzes the reversible conversion of glycine and (6S)-5,10-methylenetetrahydrofolate [(6S)-5,10-CH2-THF] to L-serine and (6S)-tetrahydrofolate [(6S)-THF] (Schirch and Szebenyi, 2005; Figure 1A); SHMT is utilized by the cell for generating one-carbon fragments for the synthesis of diverse metabolites such as quorum-sensing (QS) signals, nucleotides, methionine, thymidylate, and choline (Rao et al., 2003). Bacterial SHMTs from Escherichia coli (Scarsdale et al., 2000), Bacillus species (Bhatt et al., 2010) and Mycobacterium tuberculosis (Chaturvedi and Bhakuni, 2003) have been characterized, and usually these studies focused on enzyme mechanism and sometimes the role of SHMT in the regulation of central metabolism. However, a SHMT from Pseudomonas sp. has not been characterized and the physiological function is not understood.

In this study, we demonstrate that the SHMT ShrA (serine hydroxymethyltransferase related to rugose formation) from P. aeruginosa, which is encoded by shrA (PA2444, previously annotated as glyA2 based on bioinformatics), controls rugose colony morphology by regulating the second messenger cyclic diguanylate (c-di-GMP) through the Wsp system. We characterized the physiological function of this reversible enzyme by assaying in vitro enzyme activity and found that ShrA catalyzes the reaction to form Ser and (6S)-THF faster than it catalyzes the reaction to form Gly and (6S)-5,10-CH2-THF. When inactivated, the loss of ShrA should cause a build-up of (6S)-5,10-CH2-THF which would drive methylation of WspA which results in an increase c-di-GMP levels and biofilm formation. Our results provide one of the first links between central metabolism and cell community behavior.

RESULTS

Inactivation of ShrA Leads to RSCV Morphology

Previously, by screening 5,850 transposon mutants for altered biofilm formation, we identified 137 transposon mutants of *P. aeruginosa* PA14 with over 3-fold enhanced biofilm formation (Ueda et al., 2009). We then tested the colony morphology on Congo-red plates for 28 strains with the highest biofilm formation and identified a strain, the *shrA* mutant, with the RSCV phenotype in addition to the previously identified *tpbA* mutant (Ueda and Wood, 2009; **Figure 2A**). These colonies were smaller in size than the PA14 wild-type strain (**Figure 2A**) and developed a distinctive red and wrinkled morphology at both 37° and 25° C within $1\sim2$ days whereas the wild-type was smooth and non-wrinkled at 37° C and only slightly rough at 25° C (**Figure 2B**). Therefore,



FIGURE 1 | Mechanism of how ShrA controls c-di-GMP concentrations. (A) Proposed mechanism of how ShrA controls rugose morphology. The red box highlights the reaction scheme for SHMT ShrA. (B) Organization of the genes encoding ShrA and the glycine cleavage system. (C) Scheme for the Wsp chemosensory system. For the metabolites, THF is tetrahydrofolate, SAM is S-adenosylmethionine, SAH is S-adenosylhomocysteine and c-di-GMP is cyclic diguanylate; for the proteins, Gcv is the glycine cleavage system, MetF is a 5,10-methylenetetrahydrofolate reductase, MetH is a methionine synthase, MetK is a methionine methyltransferase, WspA is a membrane bound methyl-accepting chemotaxis protein, WspC is a methyltransferase, WspF is a methylesterase, WspD is a scaffold protein, WspE is a histidine kinase, and WspR is a diguanylate cyclase.

the *shrA* mutation converts wild-type cells into the RSCV phenotype.

Inactivation of ShrA Increases Biofilm Formation

The RSCV phenotype of the *shrA* mutant was associated with dramatically increased biofilm formation (**Figure 3**). In shake flasks, the *shrA* mutant developed biofilms at the air/liquid interface (**Figure 3A**) starting from an early growth stage (turbidity at 600 nm of 0.2). In polystyrene 96-well plates, the *shrA* mutant increased biofilm formation at the air/liquid interface as well as at the liquid/solid interface (**Figures 3B,C**). The increase was 76 ± 2-fold compared to PA14-WT after 4 h of incubation at 37°C and 119 ± 13-fold after 8 h incubation. This increase in biofilm formation occurred even though the *shrA* mutant grows at nearly the same rate $(1.10 \pm 0.08/h)$ as the wild-type $(1.13 \pm 0.00/h)$ in rich medium.



To corroborate the 96-well plate results, biofilm formation was tested using a flow cell assay with continuous flow of fresh medium (**Figure 4**). The biofilms formed by the *shrA* mutant have a 17-fold increase in biomass ($2 \pm 2 \ \mu m^3/\mu m^2$ for PA14 vs. $33 \pm 8 \ \mu m^3/\mu m^2$ for the *shrA* mutant), a 7-fold increase in substratum coverage ($13 \pm 13\%$ for PA14 vs. $96 \pm 2\%$ for the *shrA* mutant), and a 16-fold increase in thickness ($2 \pm 2 \ \mu m$ for PA14 vs. $32 \pm 8 \ \mu m$ for the *shrA* mutant). In addition, the biofilm architecture of the *shrA* mutant was smoother and flatter than that of PA14 (roughness coefficient 1.6 ± 0.3 for PA14 vs. 0.06 ± 0.01 for the *shrA* mutant). Therefore, the *shrA* mutation dramatically increases biofilm formation.

Inactivation of ShrA Increases EPS While Decreasing Motility

The red colony phenotype shown on the Congo-red plates of *shrA* mutant (**Figure 2B**) is due to Congo-red binding to EPS; hence, we quantified the amount of EPS bound to cells of PA14 and the *shrA* mutant at both 37° C and 25° C using a Congo-red binding assay and found the *shrA* mutant produces 45-and 7-fold more EPS than wild-type PA14 at 37° and 25° C,



FIGURE 3 | Inactivation of *shrA* increases biofilm formation in shake flasks and in microtitre plates. (A) The *shrA* mutant formed biofilms at the glass surface as indicated by the yellow arrow after overnight incubation in LB at 37°C with shaking at 250 rpm. Total biofilm formation (at the liquid/solid and air-liquid interfaces) (B), and bottom biofilm formation on the polystyrene plates (C) by PA14 and the *shrA* mutant after incubation in LB at 37°C without shaking. Six wells were used for each culture. Error bars indicate standard deviations from three independent cultures.

respectively (**Figure 5A**). The *pelA* mutant (negative control) did not form EPS at both temperatures tested. Considering that the Congo-red may not be an EPS-specific stain, we also used an anthrone-H₂SO₄ assay to test the EPS production. Corroborating the Congo-red assay results, the *shrA* mutation increased EPS production by 6.2 ± 0.4 compared to the wild-type strains at 37° C.

Reduced motility has been reported for other RSCVs (Ueda and Wood, 2009); hence, we examined motility for





FIGURE 5 | Inactivation of *shrA* increases EPS production and reduces motility. **(A)** EPS production by the Congo-red assay after incubation in LB for 24 h at 25°C or 16 h at 37°C. Error bars represent standard deviations from two independent cultures. **(B)** Swarming motility and **(C)** swimming motility of *P. aeruginosa* PA14 and the *shrA* mutant at 37°C after 24 h. Three plates were used for each culture and two independent cultures were used for each strain.

the *shrA* mutant; the *rhlR* (Köhler et al., 2000) and *flgK* (O'Toole and Kolter, 1998) mutants were used as negative controls for swarming and swimming motility, respectively.

Although PA14 swarmed on the surface of plates at 24 h, the *shrA* mutation abolished swarming, like the *rhlR* mutation (**Figure 5B**). In addition, swimming motility was also reduced for the *shrA* mutant compared to the PA14 wild-type strain (**Figure 5C**). Therefore, the *shrA* mutation dramatically increases EPS, abolishes swarming, and decreases swimming.

Inactivation of ShrA Increases Cellular c-di-GMP Concentrations

RSCV morphology and the related phenotypes of increased biofilm, increased EPS, and reduced motility have been associated with increased concentrations of the second messenger c-di-GMP (Ueda and Wood, 2009); therefore, we measured the cellular c-di-GMP concentrations of PA14 and shrA. The shrA mutation increased c-di-GMP concentration 18 \pm 3-fold (Figure 6). The cellular c-di-GMP level of the shrA mutant was 1.6 \pm 0.2 pmol/mg cells. This is comparable to the c-di-GMP production of a small colony variant (around 2 pmol/mg cells) (Meissner et al., 2007). The c-di-GMP concentration of the wild-type strain was 0.09 ± 0.03 pmol/mg cells. This value is lower than the reported value (around 0.6 pmol/mg) for P. aeruginosa wild-type 20265 (Simm et al., 2009). However, the difference could be due to the strain difference as well as differences in growth conditions (we harvested the cells at a turbidity of 2.0 while the strain used for comparison was incubated in Luria-Bertani (LB) medium for 48 h). Therefore, the increased RSCV morphology, EPS, biofilm, and motility phenotypes of the shrA mutant may be explained by the increased c-di-GMP concentrations.

Complementation of Biofilm Formation with ShrA

To verify whether the phenotypes observed in the *shrA* mutant were caused by loss of function of ShrA, we confirmed the transposon insertion in *shrA* by PCR. Furthermore, complementation of the biofilm phenotype of the *shrA* mutation by producing ShrA production from a plasmid was investigated. As expected, *shrA* gene expression using plasmid pMQ70-*shrA* reduced total biofilm formation of the *shrA* mutant by 90%



(B) *P. aeruginosa* PA14, (C) the *shrA* mutant, and (D) the *shrA* mutant spiked with 7.5 μM c-di-GMP. (E-H) spectra of the corresponding c-di-GMP peak of (A–D), respectively.

at 24 h (**Figure 7A**) and reduced the bottom biofilm formation by 43% (**Figure 7B**); *P. aeruginosa* forms biofilms along the sides of the 96-well plates as well on the bottom. Total biofilm formation of the PA14 strain was also delayed by *shrA* gene overexpression (**Figure 7C**). This result was confirmed by using another expression plasmid, pMJT1-*shrA*, from which expression of *shrA* reduced biofilm formation of PA14 strain 4.8 \pm 2.9-fold after incubation in LB medium at 37°C for 24 h (**Figure 7D**). Therefore, the increased biofilm phenotype of the *shrA* mutation could be complemented.

Genetic Suppressor Screening

To investigate how ShrA regulates the RSCV phenotype, genetic screening was conducted using Tn5-*luxAB* transposon mutagenesis to find suppressive loci for the RSCV phenotype from the *shrA* mutation. The double mutant library (*shrA* plus random gene inactivation) was first screened for a reduction in aggregation and only the cells remaining in the supernatant that failed to aggregate like the *shrA* mutant were grown on

Congo-red plates with selective antibiotics; approximately 5,000 double mutant colonies were screened. After incubating at 37°C for 2 days, colonies displaying a white and smooth shape like the wild-type strain were chosen. Eleven of these colonies were sequenced from the transposon: four of these had the Tn5-*luxAB* insertion in the *pel* locus (**Table 1**), and one had the insertion in the PA0839 gene. Critically, there were six suppressive mutations in four genes of the *wsp* chemosensory system: *wspA*, *wspD*, *wspE*, and *wspR* (**Table 1**). Therefore, the *shrA* mutation appears to cause the RSCV phenotype through an alteration of the Wsp chemosensory system.

ShrA Is an Oligomer Containing One PLP per Dimer

The physiological role of SHMT is to catalyze the reversible conversion of glycine and (6S)-5,10-CH₂-THF to L-serine and (6S)-THF (Schirch and Szebenyi, 2005; **Figure 1A**). (6S)-5,10-CH₂-THF provides the largest part of the one-carbon units



FIGURE 7 | ShrA reduces biofilm formation. (A) Total biofilm formation and (B) bottom biofilm formation on polystyrene plates for production of ShrA in the *shrA* mutant via pMQ70-shrA, and (C) total biofilm formation on polystyrene plates for production of ShrA in *P. aeruginosa* PA14 via pMQ70-shrA. LB containing 0.05% arabinose at 37°C was used. (D) Total biofilm formation of *P. aeruginosa* PA14 on polystyrene plates for production of ShrA via pMJT1-shrA in LB containing 0.2% arabinose at 37°C for 24 h. Six wells were used for each culture and three independent cultures were used for each strain. Error bars represent the standard deviations.

TABLE 1	Suppressive loci for the shrA mutation.
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Strain	PAO1 ID	PA14 ID	Gene name	Gene function
Suppressor 23	PA3702	PA14_16500	wspR	WspR, diguanylate cyclase/response regulator
Suppressor 8	PA3704	PA14_16470	wspE	WspE, histidine kinase/response regulator
Suppressor 9	PA3704	PA14_16470	wspE	
Suppressor 6	PA3705	PA14_16460	wspD	WspD, scaffold protein, regulates WspE activity
Suppressor 10	PA3705	PA14_16460	wspD	
Suppressor 29	PA3708	PA14_16430	wspA	WspA, a membrane bound methyl-accepting chemotaxis protein, will activate WspE when methylated
Suppressor 3	PA3058	PA14_24560	pelG	Predicted membrane protein related to EPS production, PelG
Suppressor 24	PA3061	PA14_24510	pelD	L-lactate permease related to EPS production, PeID
Suppressor 4	PA3064	PA14_24480	pelA	Oligogalacturonide lyase related to EPS production, PelA
Suppressor 25	PA3064	PA14_24480	pelA	Oligogalacturonide lyase related to EPS production, PelA
Suppressor 11	PA0839	PA14_53410		Probable transcriptional regulator

Genetic suppressor screening identified additional mutations that mask the phenotypes of the shrA mutant.

available to the cell (Stover and Schirch, 1990) and is a precursor for S-adenosylmethionine (SAM) synthesis through the folate pathway (**Figure 1A**; Shoeman et al., 1985). Since the methyltransferase WspC (analog to CheR) uses SAM to methylate the receptor WspA (Springer and Koshland, 1977), we hypothesized that inactivation of ShrA may increase the availability of the one-carbon units which will drive the methylation of WspA.

To check the hypothesis, we characterized the *in vitro* activity of the purified enzyme. Recombinant ShrA with a 6X-His tag at the carboxy terminus was produced in *E. coli* BL21 (DE3),

and the protein was purified (>95%) (Figure S1A). For SHMT, the dimer is the minimum necessary structure for the catalytic activity (Bhatt, Bhakuni, Kumar, Khan and Siddiqi); SHMT from *E. coli* as well as that from several bacterial sources is dimeric whereas SHMT from mammalian sources is a homotetramer. We checked the oligomerization state of ShrA by glutaraldehyde cross-linking and found ShrA shifted from a monomer to an oligomer after cross-linking using glutaraldehyde as determined by SDS-PAGE (Figure S1B). The oligomer protein band was not a sharp band of dimer at 90 kDa but was broad, indicating possible crosslinking products from a dimer to a tetramer, or an oligomer

with intra-molecular cross-linking by glutaraldehyde. Therefore, ShrA from *P. aeruginosa* PA14 is present as an oligomer as are other bacterial SHMTs.

SHMT is a PLP-dependent enzyme in which PLP is covalently attached to the enzyme (Chaturvedi and Bhakuni, 2003). SHMT from *E. coli* as well as SHM2 from *M. tuberculosis* contain 2 PLP/(mol enzyme dimer) (Chaturvedi and Bhakuni, 2003); however, SHM1 from *M. tuberculosis* contains only 1 PLP/(mol enzyme dimer) (Chaturvedi and Bhakuni, 2003). Therefore, we checked the PLP content for ShrA. The molar ratio of PLP to ShrA monomer is 0.5 ± 0.1 as calculated from four samples. This ratio is similar to that of SHM1 from *M. tuberculosis*, but different from other bacterial SHMTs.

ShrA Primarily Produces Ser and (6S)-THF from Gly and (6S)-5,10-CH₂-THF

According to our hypothesis, ShrA should primarily catalyze the conversion of Gly and (6S)-5,10-CH2-THF to Ser and (6S)-THF instead of the reverse reaction (Figure 1A) for the production of one-carbon units as its pivotal role (Rao et al., 2003; Florio et al., 2010). To confirm this, we characterized the enzyme activity in vitro toward both Ser/(6S)-THF and Gly/(6S)-5,10-CH₂-THF using purified ShrA; Figure 8A shows the chromatogram of each amino acid separated by highperformance liquid chromatography (HPLC). For the forward reaction (Figure 1A), the substrates Gly (20 mM) and (6R,S)-5,10-CH₂-THF (2 mM) were reacted with ShrA, and the product (6S)-THF was immediately recycled back to the substrate (6S)-5,10-CH₂-THF by using excess formaldehyde, so the substrate (6R,S)-5,10-CH₂-THF concentration was constant throughout the reaction. The product Ser was quantified (Figure 8B), and protein dialysis buffer instead of the protein was used as the negative control (Figure 8C). The enzyme specific activity was calculated to be 4.24 \pm 0.01 μ mol/min/mg. In order to confirm that the value represents V_{max}, we also increased the substrate concentrations to 30 mM Gly and 3 mM (6R,S)-5,10-CH₂-THF, and the activity was determined to be similar (4.01 μ mol/min/mg).

For the reverse reaction, the amount of Gly produced from the substrates Ser (4 mM) and (6*R*,*S*)-THF (2 mM) was analyzed (**Figure 8D**), and the specific activity was 1.6 \pm 0.3 μ mol/min/mg. Protein dialysis buffer instead of the protein was used as the negative control (**Figure 8E**). When the concentrations of the substrates were increased to 6 mM Ser and 3 mM (6*R*,*S*)-THF, the specific activity was nearly identical (1.5 \pm 0.3 μ mol/min/mg); hence, these values are equivalent to V_{max}. Therefore, ShrA is active as an SHMT, and the V_{max} is 2.8-fold higher for converting Gly/(6*S*)-5,10-CH₂-THF to Ser/(6*S*)-THF than for the reverse direction.

The *shrA* Operon Also Encodes Proteins to Decrease Gly and Produce (6S)-5,10-CH₂-THF

The *shrA* operon encodes genes for both Ser and Gly metabolism (**Figure 1B**), including those for the glycine cleavage system (GcvH2/P2/T2) and a serine dehydratase (SdaA). The GCV





system is responsible for the degradation of glycine and converts (6S)-THF to (6S)-5,10-CH₂-THF (**Figure 1A**), so the GCV system is an additional source for the one-carbon units.

Therefore, the proteins encoded by this operon should regulate the two amino-acid metabolism cooperatively. To investigate this, we checked the colony morphology of the transposon mutants (**Figure 2C**) and found that all of these mutants have the same morphology as the wild-type strains. Therefore, only ShrA inactivation leads to RSCV in the *shrA* operon since only this mutation should increase the one-carbon source (6*S*)-5,10-CH₂-THF.

Differentially Regulated Genes in Biofilm Cells of the *shrA* Mutant

To explore the mechanism for ShrA regulation and the impact of ShrA inactivation on the whole genome (in addition to the Wsp chemosensory system), a whole-transcriptome analysis was performed with biofilm cells on glass wool for the shrA mutant after incubating at 37°C for 7 h. There were 24 genes found to be induced for the shrA mutant biofilm more than 1.8-fold, 12 of which were related to phage. In contrast, 172 genes were found to be repressed in shrA biofilm cells more than 1.8-fold. Among them, 21 genes were related to denitrification (Table S1), such as the gene that encodes nitrite reductase nirS, genes encode nitric oxide reductase norBCD, and the gene encodes nitrous reductase nosZ. Another large group of affected genes is related to iron acquisition: there are total 65 genes in this group and 41 of these were related to siderophores, including the *pvd* gene clusters encoding for pyoverdine production, and the *pch* gene cluster encoding pyochelin. Another 13 genes were related to redox enzymes; for example, some of the FMNH₂/NAD dependent proteins such as SsuDE and some oxygenases were repressed. The common feature for the encoded proteins is that they are related to electron transport and might be regulated by cell redox homeostasis.

ShrA Decreases the NADH/NAD⁺ Ratio

Based on the transcriptome results, we examined the role of ShrA on redox homeostasis by investigating the NADH/NAD⁺ ratio since it is representative of the intracellular redox state (Price-Whelan et al., 2007). We utilized bacteria grown on agar surfaces since colonies have been used to study the development of cell communities (Ramos et al., 2010) and found that the *shrA* mutation increased the NADH/NAD⁺ ratio from 0.25 \pm 0.04 for the wild-type strain to 0.50 \pm 0.03. Therefore, the intracellular NADH/NAD⁺ ratio was increased 2.0 \pm 0.3-fold by the *shrA* mutation.

Nitrate Abolished the *shrA* Rugose Phenotype as an Electron Acceptor

To check whether the enhanced NADH/NAD⁺ ratio plays a role in the rugose phenotype of the *shrA* mutant, we examined the effect of nitrate (NO₃⁻) on *shrA* colony morphology. NO₃⁻ is a terminal electron acceptor (Williams et al., 1978), and it reduces cellular NADH/NAD⁺ levels (Price-Whelan et al., 2007) while it does not affect the total concentrations of NADH and NAD⁺ (Price-Whelan et al., 2007; **Figure 1A**). Addition of 30 mM KNO₃ (a physiologically-relevant concentration, Price-Whelan et al., 2007) completely converted the *shrA* rugose colony into the smooth colony morphology of the wild-type strain (**Figure 2B**). In comparison, the rugose phenotype of another RSCV mutant, *tpbA* (Ueda and Wood, 2009), was only partially reduced by nitrate (**Figure 2B**). Therefore, the increased NADH/NAD⁺ ratio caused by inactivating ShrA positively regulates the RSCV phenotype.

ShrA Increases Pyocyanin Levels

To further investigate the role of ShrA in redox homeostasis, we investigated its effect on pyocyanin levels; pyocyanin is secreted and generates reactive oxygen species (Bodelón et al., 2016). We found that inactivating ShrA led to a 2.4 ± 0.4 -fold increase in pyocyanin levels. As expected, the negative control *phzS* reduced pyocyanin 5.1 \pm 3.3-fold In agreement with our results, increased levels of c-di-GMP have been shown to increase pyocyanin levels (Lo et al., 2016). Note that reduced transcription of *shrA* via inactivation of PA2449 (GcsR), a positive regulator of the operon that includes *shrA*, has also been reported to reduce pyocyanin (Lundgren et al., 2013), but this was later shown to likely be due a *P. aeruginosa* PAO1 strain artifact (Sarwar et al., 2016).

DISCUSSION

Inactivation of the SHMT ShrA Leads to Increased c-di-GMP

In this study, we demonstrated that ShrA is an SHMT that regulates the RSCV phenotype of P. aeruginosa; this is the first biochemical characterization of this enzyme in P. aeruginosa. The RSCV phenotype for the shrA mutant and the related hyper-biofilm formation, increased EPS production, and reduced swarming motility may be explained by the 20fold increased intracellular c-di-GMP concentrations, that we measured (Figure 6). c-di-GMP is an ubiquitous intracellular second messenger that acts as a central regulator in bacterial physiology, especially in regulating the transition between motile and sessile states (Hengge, 2009), and there is ample evidence to correlate RSCV with increased c-di-GMP concentrations (Hickman et al., 2005; Ueda and Wood, 2009). For example, inactivation of protein tyrosine phosphatase TpbA caused RSCV formation through constitutive activation of a DGC TpbB, which results in elevated c-di-GMP concentrations (Ueda and Wood, 2009; Pu and Wood, 2010).

ShrA Controls c-di-GMP through the Wsp Chemosensory System

To examine how the SHMT ShrA controls c-di-GMP concentration, we screened rugose suppressor mutants (**Table 1**) and identified the *wsp* two-component chemosensory system (*wspADER*) is a necessary component downstream of ShrA for causing the RSCV phenotype. WspR is a DGC containing a CheY domain and a GGDEF domain, which catalyzes the production of c-di-GMP (Hickman et al., 2005). Deletion of *wspF* in *P. aeruginosa* causes the constitutive activation of WspR that leads to the rugose phenotype (Hickman et al., 2005). By comparing the Wsp system to the well-studied Che chemotaxis system, the signal cascade was predicted (**Figure 1C**; Bantinaki et al., 2007). Upon an external signal, the membrane bound methyl-accepting chemotaxis protein (MCP), WspA, is methylated

which activates autophosphorylation of the histidine kinase WspE. The methylation of WspA is controlled by the opposing activities of the methyltransferase WspC and the methylesterase WspF. The phosphorylated WspE relays the phosphate group to the CheY domain of WspR which activates its DGC activity. The predicted functions of WspC and WspF have been confirmed in *P. fluorescens* (Bantinaki et al., 2007) in that deletion of *wspF* or overexpression of *wspC* will convert a smooth colony into a rugose colony.

The suppressor mutants we identified in wspA, wspD, wspE, and *wspR* are consistent for the signal cascade (Figure 1C) in that WspA, WspD and WspE are all positive regulators of WspR as a DGC. Of significance is that it indicates ShrA controls cellular c-di-GMP concentrations through the DGC WspR, especially by influencing WspA, the most upstream component among these four genes. PA14 has 37 putative c-di-GMP related proteins, including 16 proteins with a GGDEF domain, 5 with a PDE domain, and 16 that contain both domains; however, not all of these proteins with DGC and PDE domains are active enzymes since some of the domains are used for regulation (Kulasakara et al., 2006). Hence, among the GGDEF proteins, only PA1107, TpbB, WspR, and PA5487 increased biofilm formation when overexpressed (Kulasakara et al., 2006). Previously, using the same phenotype suppressor screening for the *tpbA* RSCV mutant, only the DGC TpbB was found downstream (Ueda and Wood, 2009). In contrast, for this shrA mutant, only the WspR system was identified downstream rather than the other DGCs. Hence, ShrA controls c-di-GMP concentrations through the Wsp system rather than other DGCs. Perhaps these two means of controlling c-di-GMP for biofilm formation (Wsp and TpbB) are distinct and arise based on the use of two different appendages for surface sensing.

ShrA Reduces One Carbon Equivalents to Reduce Rugose Cells

The common concept for the physiological function of SHMT is that this enzyme is utilized by the cell for generating one-carbon fragments to (6S)-5,10-CH₂-THF for the synthesis of diverse metabolites (Rao et al., 2003; Florio et al., 2010). However, the *in vitro* ShrA enzyme activity indicates that ShrA prefers converting Gly/(6S)-5,10-CH₂-THF to Ser/(6S)-THF rather than the reverse reaction. Therefore, ShrA reduces the availability of one-carbon units.

According to this physiological function of ShrA, we propose the mechanism of how it controls the RSCV phenotype shown in **Figure 1A**. Inactivation of ShrA leads to accumulated (6*S*)-5,10-CH₂-THF, which provides the one-carbon building blocks. Delivering the one-carbon source to SAM via the folate pathway (Shoeman et al., 1985) (**Figure 1A**), including 5,10methylenetetrahydrofolate reductase MetF, methionine synthase MetH, and methionine methyltransferase MetK, allows SAM to transfer the methyl group to WspA by the methyltransferase WspC, which activates the DGC WspR.

The non-rugose phenotype of the gcv mutants (**Figure 2C**) also supports our hypothesis. The GCV system produces (6S)-5,10-CH₂-THF (**Figure 1B**). Thus, inactivation of the GCV

system should lead to the reduction of the one-carbon units rather than their accumulation as the shrA mutation did. Encoding the GCV system together with ShrA in the same operon seems to keep the supply of active one-carbon units balanced while inactivation of *shrA* alters this balance. Of course, the impact of Gly on cell physiology is complex; for example, we found that 200 mM Gly inhibited the growth of PA14 in tryptone medium at 25°C, and high concentrations of glycine inhibit the growth of many bacteria (Ratomahenina et al., 1979; Minami et al., 2004). A possible mechanism for this inhibition is that UDP-N-acetylmuramate-alanine ligase activity is reduced by glycine and thus cell wall component synthesis is impaired (Minami et al., 2004). It is also interesting to note that a glycine resistant mutant of Pseudomonas stutzeri entails increased activity of the serine hydroxymethyltransferase (Ratomahenina and Galzy, 1981) which would reduce Gly concentrations which is consistent with our mechanism.

Cell Redox State Plays a Role in the RSCV Phenotype via ShrA

The increase in the NADH/NAD⁺ ratio is consistent with our proposed mechanism (**Figure 1A**); inactivation of ShrA prevents the catabolism of Gly through the ShrA pathway which should enhance the GCV pathway and lead to increased NADH. The increased NADH levels then serve as a cofactor of MetF for the production of 5-methyltetrahydrofolate $[(6S)-5-CH_3-THF]$ and thus increase the availability of one-carbon units. The fact that addition of the electron acceptor NO₃⁻ abolished the rugose morphology of the *shrA* mutant but not the *tpbA* mutant (**Figure 2B**) indicates that the altered NADH/NAD⁺ ratio plays a key role for the rugose phenotype of the *shrA* mutant and that NADH is a required factor rather than a side effect. Hence, inactivation of ShrA alters the cell redox state, which cooperatively regulates colony morphology through the folate pathway and the Wsp system.

The redox state of *P. aeruginosa* has been linked to colony morphology before since a *phz* mutation enhances the NADH/NAD⁺ ratio (Price-Whelan et al., 2007) and results in large rugose colonies at 20° C (Dietrich et al., 2008). However, the regulation mechanism has not been elucidated yet. Our finding links the redox state and a downstream diguanylate cyclase, WspR, which further regulates the rugose morphology through the ubiquitous second messenger c-di-GMP.

Inactivation of ShrA Represses Iron Acquisition

Our whole transcriptome analysis of the *shrA* biofilm cells linked this mutation to denitrification, iron acquisition, and the redox state (Table S1). However, for the *wspF* mutant, which also results in RSCV and increased c-di-GMP due to increased WspR activity, these same genes were differently regulated (Hickman et al., 2005). For example, the *shrA* mutation repressed genes for denitrification (*nir, nor* and *nos* gene clusters) while these genes were not changed by the *wspF* mutation. Hence, the repression of

these genes in *shrA* mutant is not simply due to increased c-di-GMP levels; instead, they were regulated due to the redox change upstream of the Wsp system.

One common feature for the iron acquisition, denitrification and redox genes repressed in the transcriptome analysis is that they could all be directly or indirectly related to electron transport or the cell redox state. Indeed, an increased intracellular NADH/NAD+ ratio was observed for the shrA mutant, indicating a more reduced cellular environment. A large group of genes that were repressed upon inactivating ShrA were those related to iron acquisition, including siderophorerelated genes (Table S1). In P. aeruginosa, the expression of these iron acquisition genes is strictly regulated in response to the environmental iron concentrations through the ferric uptake regulator (Fur) protein (Ochsner and Vasil, 1996). At high iron concentrations, Fur binds to Fe²⁺ and the Fur-Fe²⁺ complex represses these genes. For the shrA mutant, various Fur-regulated genes were found to be repressed along with the *pvd* and *pch* genes, including pfeR, tonB, fumC, fpvA, phuR, fiuA, pirA, piuA, and fptA (Ochsner and Vasil, 1996; Cornelis et al., 2009). Since the *shrA* cells are in a more reduced state with higher NADH levels and since NADH (redox potential $E'_0 = -320 \text{ mV}$) is capable of reducing Fe^{3+} to Fe^{2+} ($\dot{E}'_0 = 770 \text{ mV}$) (Berg et al., 2002), it is reasonable to postulate that the Fe²⁺/Fe³⁺ ratio increased upon inactivating ShrA so that there is more Fe²⁺ accessible for Fur, which leads to the repression of these iron acquisition genes. Hence, the altered redox state may be the cause the repression of the iron acquisition genes.

Inactivation of ShrA Represses Denitrification

The transcriptome analysis with the *shrA* mutant also revealed the denitrification gene clusters (*nir*, *nor*, and *nos* genes) were repressed. The expression of the denitrification machinery of *P. aeruginosa* is affected by redox signaling through the transcription factors ANR (anaerobic regulation of arginine deiminase and nitrate reduction) and DNR (dissimilative nitrate respiration regulator) (Giardina et al., 2008). In *anr* or *dnr* mutants, *nirS* and *norCB* transcription is repressed (Arai et al., 1995). ANR is regulated by the redox state through the [4Fe-4S]²⁺ cluster oxygen sensor while DNR is an NO sensor through the cofactor ferrous heme (Giardina et al., 2008). The redox state of the cofactors [4Fe-4S]²⁺ and ferrous heme may be altered by the increased NADH levels in the *shrA* mutant. Hence, the altered redox state may explain the repression of the denitrification genes.

ShrA Regulation

SHMT could be regarded as the initial enzyme of a branched pathway, and is likely to involve complex regulation (Dev and Harvey, 1984b). Using *E. coli* auxotrophic cultures, Harvey et al. found that SHMT synthesis is subject to active control mechanisms which respond to the requirements for various end products of the folate pathway, for purine biosynthesis, and for methylation reactions, as well as to serine limitation (Dev and Harvey, 1984b). In *E. coli*, the rate of SHMT synthesis is a hyperbolic function of (homo-Cys/SAM) (Dev and Harvey,

1984a), thus it is postulated that homo-Cys acts as an inducer of SHMT and SAM acts as a corepressor, suggesting that it is the requirements of methionine for methylation reactions which controls the SHMT synthesis (Dev and Harvey, 1984a).

The regulation of the SHMT ShrA in *P. aeruginosa* is not clear yet, but it is also likely to be complicated. *shrA* was identified to be a QS induced gene in PAO1 by Schuster et al. (Schuster et al., 2003) by comparing the transcriptome of the wild-type strain with a *lasRI rhIR* mutant whose QS was repressed. The *gcv* genes in the same operon were also found to be induced by QS. *shrA* is also repressed 955-fold in *ppyR* (PA2663) biofilm cells compared to the PAO1 wild-type (Attila et al., 2008). Since the *ppyR* mutant also repressed QS genes, the repression of *shrA* could be further linked to QS. For the microarray data of the *shrA* mutant biofilm cells, the *shrA* gene as well as two other genes in the same operon, *gcvT2* and *sdaA*, were all repressed comparing with the PA14 wild-type. Hence, the *shrA* gene in *P. aeruginosa* should be highly regulated.

Overall, we have discovered insights into how central metabolism; i.e., the conversion of the amino acids glycine to serine by hydroxymethyltransferase ShrA, impact the sessile lifestyle and virulence of the opportunistic pathogen *P. aeruginosa*. Furthermore, we demonstrate that the mechanism by which ShrA impacts biofilm formation is via reduction of c-di-GMP through its reduction of the intermediates glycine, (6S)-5,10-CH₂-THF, methionine, and *S*-adenosylmethionine (**Figure 1A**).

MATERIALS AND METHODS

Strains and Growth Conditions

All strains and plasmids used in this study are listed in Table 2. P. aeruginosa PA14 (wild-type) and its isogenic mutants were obtained from the Harvard Medical School (Liberati et al., 2006). P. aeruginosa and Escherichia coli were grown in LB medium (Sambrooke et al., 1989) at 37°C unless noted. Gentamicin ($15 \mu g/mL$) was used for growth of the *P. aeruginosa* transposon mutants, carbenicillin (300 µg/mL) was used to maintain P. aeruginosa plasmid pMQ70 and pMJT1, and kanamycin (50 µg/mL) was used to maintain E. coli plasmid pET28b. Transposon insertion of the shrA mutant was verified as described previously (Ueda and Wood, 2009). Briefly, the PCR product amplified using primers PA14_33010-VF and PA14_33010-VR (Table 3) from the chromosomal DNA of shrA mutant was 1 kb larger than that from the wild-type, which corresponds to the size of the transposon. In addition, the DNA fragment corresponding to the end of the transposon and shrA gene was amplified with shrA chromosomal DNA using primers PA14_33010-VF and GB-3a (Table 3) and PA14_33010-VR and R1 (Table 3) but these pairs of primers did not amplify PA14 wild-type chromosomal DNA.

Complementation of P. aeruginosa Mutants

For complementation of the *shrA* mutations, *shrA* was expressed under the control of the pBAD promoter in pMQ70 (Shanks et al., 2006) and in pMJT1 (Olsen et al., 1982). *shrA* was

TABLE 2 | Strains used in this study.

Strain	Genotype or description	Reference
P. aeruginosa		
PA14	Wild-type strain	Liberati et al., 2006
PA14_33010 (PA2444, shrA)	PA14_33010 Ω <i>Mar2xT7</i> , Gm ^R	Liberati et al., 2006
PA14_19120 (PA3477, <i>rhIR</i>)	PA14_19120 Ω <i>Mar2xT7</i> , Gm ^R	Liberati et al., 2006
PA14_50360 (PA1086, <i>flgK</i>)	PA14_50360 Ω <i>Mar2xT7</i> , Gm ^R	Liberati et al., 2006
PA14_24480 (PA3064, <i>pelA</i>)	PA14_24480	Liberati et al., 2006
PA14_13660 (PA3885, <i>tpbA</i>)	PA14_13660 Ω <i>Mar2xT7</i> , Gm ^R	Liberati et al., 2006
PA14_33040 (PA2442, <i>gcvT2</i>)	PA14_33040	Liberati et al., 2006
PA14_33030 (PA2443, <i>sdaA</i>)	PA14_33030	Liberati et al., 2006
PA14_33000 (PA2445, <i>gcvP2</i>)	PA14_33000	Liberati et al., 2006
E. coli		
BL21(DE3)	F ⁻ ompT hsdS _B ($r_B^- m_B^-$) gal dcm λ(DE3) Ω placUV5::T7 polymerase	Novagen
HB101	pro leu thi lacY Str ^r endol ⁻ $recA^- r^- m^-$	Ditta et al., 1980
S17-1(λ <i>pir</i>)/pUT-miniTn5 <i>-luxAB</i>	Tc ^R Sm ^R Tp ^R mod ⁺ res thi pro recA hsdR17 ${\it \Omega}$ RP4-TC::Mu-Km::Tn7 with pUT-miniTn5-luxAB	Simon et al., 1983; Ramsey and Whiteley, 2004
TG1	K12, lac–pro supE thi hsdD5 (F' traD36 proA ⁺ B ⁺ lacl ^q lacZ M15)	Sambrooke et al., 1989
PLASMIDS		
pMQ70	Car ^R , Ap ^R , P _{BAD} , expression vector	Shanks et al., 2006
pMQ70-shrA	Car ^R , Ap ^R , P _{BAD} ::shrA, complementation plasmid	This study
pMJT1	Car ^R , Ap ^R , P _{BAD} , expression vector	Olsen et al., 1982
pMJT1-shrA	Car ^R , Ap ^R , P _{BAD} ::shrA, complementation plasmid	This study
pRK2013	Mobilizing conjugation plasmid	Figurski and Helinski, 1979
pET28b	Km ^R , P _{T7} expression vector	Novagen
pET28b-PA2444cHis	Km ^R , P _{T7} :: <i>shrA-cHis</i> ⁺ , expression vector for ShrA-cHis	This study

Gm^R, Tc^R, Km^R, Car^R, Cm^R, and Ap^R indicate gentamicin, tetracycline, kanamycin, carbenicillin, chloramphenicol, and ampicillin resistance, respectively.

amplified using a Pfu DNA polymerase with primers PA2444-F-NheI and PA2444-R-HindIII (**Table 3**) for pMQ70-*shrA*, and primers PA2444-F-NheI and PA2444-R-XbaI for pMJT1-*shrA*. PCR products were cloned into the restriction sites of pMQ70 or pMJT1. The pMQ70-*shrA* and pMJT1-*shrA* plasmids were confirmed by DNA sequencing with the pMQ70-F/R primers or pMJT1-F/R primers (**Table 3**). pMQ70-*shrA* was transformed from the donor strain TG1/pMQ70-*shrA* into PA14 strains by conjugation using helper strain HB101/pRK2013 as described previously (Ueda and Wood, 2009). The plasmid pMJT1-*shrA* was transferred into the PA14 strains by electroporation (Choi et al., 2006).

Colony Morphology

Colony morphology was checked on Congo-red plates (10 g/L tryptone, 40 μ g/mL Congo-red, 20 μ g/mL Coomassie brilliant blue and 15 g/L agar) as described previously (Ueda and Wood, 2009) or with Congo-red plates supplemented with 30 mM potassium nitrate. Plates were incubated at 37°C or room temperature.

Static Biofilm Assay

Biofilm formation was examined in 96-well polystyrene plates using crystal violet staining (Fletcher, 1977). Overnight cultures of *P. aeruginosa* were diluted to a turbidity of 0.05 at 600 nm with fresh LB medium, and then 200 μ L of diluted bacterial culture was incubated in 96-well polystyrene plates for 4, 8, 24, and 50 h. Six wells were used for each strain, and at least two independent cultures were used for each experiment.

Flow-Cell Biofilm Assay

The flow cell experiments were performed as previously described (Ma et al., 2011) with modifications. At 37°C, the flow cells were inoculated with cultures in 5% LB medium at an initial turbidity at 600 nm of 0.05 and at a flow rate of 10 mL/h for 2 h, then fresh 5% LB medium was added at 10 mL/h for 3 days. After 72 h of incubation, biofilms were stained with SYTO9 for 20 min in the dark. Biofilm images from nine random positions were visualized with IMARIS confocal software (Bitplane, Zurich, Switzerland) and analyzed by COMSTAT confocal software (Heydorn et al., 2000).

Motility Assay

Swimming motility was examined using 0.3% agar plates with 1% tryptone and 0.25% NaCl as described previously (Sperandio et al., 2002), and swarming motility was examined with BM-2 plates (62 mM potassium phosphate, 2 mM MgSO₄, 10 μ M FeSO₄, 0.1% casamino acid, 0.4% glucose, and 0.5% Bacto agar) (Overhage et al., 2008). Motility was measured after 24 h. Three plates were tested for each culture, and two independent cultures were used. The *flgK* (Liberati et al., 2006) and *rhlR*

TABLE 3 | Primers used in this study.

Name	Sequence (5'-3')	Description Verification of <i>shrA</i> tn insertion	
PA14_33010-VF	TGACACCGACACGACAAGAG		
PA14_33010-VR	GATAGCCCTCGGCATACTTG	Verification of shrA tn insertion	
GB-3a	TACAGTTTACGAACCGAACAGGC	Verification of shrA tn insertion	
R1	ATCGACCCAAGTACCGCCAC	Verification of shrA tn insertion	
PA2444-F-Nhel	GCCCCCGCTAGCAAGAAGGAGATATACCATGTTCAGCAA GCACGACCAGCTCCAGG	Construction of pMQ70-shrA and pMJT1-shrA	
PA2444-R-HindIII	GCCCCCAAGCTTCTAGTGATGGTGATGGTGATGTCAATCAGCG TAGACCGGGAAGTGC	Construction of pMQ70- <i>shrA</i> and pET28b-PA2444cHis	
PA2444-R-Xbal	GCCCCCTCTAGATCAATCAGCGTAGACCGGGAAGTGC	Construction of pMJT1-shrA	
pMJT1-F	CTCTTCTCGCTAACCAAACC	Sequencing of pMJT1 derivatives	
pMJT1-R	GTGGAATTGTGAGCGGAT	Sequencing of pMJT1 derivatives	
PA2444-F-Xbal	GCCCCCTCTAGAAAGAAGGAGATATACCATGTTCAGCAAGC ACGACCAGCTCCAGG	Construction of pET28b-PA2444cHis	
pMQ70-F	GCGTCACACTTTGCTATGCCATAGC	Sequencing pMQ70 derivatives	
pMQ70-R	CTACTGCCGCCAGGCAAATTCTGTTT	Sequencing pMQ70 derivatives	
LuxAB inside	CTTTCAATTTCCGCTTTCAAGC	Identification of transposon insertion	
LuxAB outside	CGATGGTGAGTTGTTCAAAATC	Identification of transposon insertion	
Arb1	GGCCACGCGTCGACTAGTACNNNNNNNNNGATAT	Identification of transposon insertion	
Arb2	GGCCACGCGTCGACTAGTAC	Identification of transposon insertion	
T7 promoter	TAATACGACTCACTATAGGG	Sequencing pET28b derivatives	
T7 terminator	GCTAGTTATTGCTCAGCGG	Sequencing pET28b derivatives	

(Liberati et al., 2006) mutants were used as a negative controls for swimming and swarming, respectively.

EPS and Pyocyanin Assay

EPS production was quantified by two methods. The first method (Ueda and Wood, 2009) was based on the amount of Congored that binds to the EPS in T-broth. PA14 *pelA* mutant (Liberati et al., 2006) was used as the negative control. Due to aggregative phenotype of the *shrA* mutant, cell pellets for all strains were sonicated at 3 W for 10 s three times to dissociate the cells as described previously (Ueda and Wood, 2009). Cell viability after sonication was tested by CFU counting using the drop plating method to make sure the mild sonication method did not cause cell lysis. The second method used anthrone-H₂SO₄ (Zhang et al., 2008) for quantification of the glucose equivalents in EPS. Normalized pyocyanin levels were assayed spectrophotometrically after overnight growth in LB medium as described previously after chloroform extraction (Wood and Wood, 2016).

Total RNA Isolation and Microarray Analysis

The *P. aeruginosa* genome array (Affymetrix, P/N 510596) was used to investigate differential gene expression in biofilm cells between PA14 and the *shrA* mutant. Biofilm cells were harvested from 10 g of glass wool (Ren et al., 2004a) after incubation for 7 h in LB with shaking at 250 rpm, and RNA was extracted with the RNeasy Mini Kit (Qiagen, Valencia, CA) using a bead beater (Biospec, Bartlesville, OK) (Ren et al., 2004a) with RNA*later* buffer (Applied Biosystems, Foster City, CA) to stabilize the

RNA. cDNA synthesis, fragmentation, hybridizations, and data analysis were as described previously (González Barrios et al., 2006). For each binary microarray comparison of differential genes expression, if the gene with the larger transcription rate did not have a consistent transcription rate based on the 13 probe pairs (P < 0.05), these genes were discarded. A gene was considered differentially expressed when the P-value for comparing two chips was lower than 0.05 (to assure that the change in gene expression was statistically significant and that false positives arise <5%) and when the expression ratio was higher than the standard deviation for the whole microarrays (1.8 for PA14 wild-type and shrA mutant) (Ren et al., 2004b). The microarray raw data are deposited at the Gene Expression Omnibus (GSE29879) of the National Center for Biotechnology Information; all data are MIAME compliant and have been deposited in a MIAME compliant database as detailed on the Functional Genomics Data Society website http://fged.org/ projects/miame/.

Genetic Suppressor Screening

To isolate the suppressive loci for RSCV formation due to the *shrA* mutation, a double mutant library was generated using the Tn5-*luxAB* transposon with the *shrA* strain as described previously (Ramsey and Whiteley, 2004). Screening cells with mutations in addition to *shrA* was performed in two steps as described previously (Ueda and Wood, 2009) with the first step used to discard cells with the aggregative phenotype and the second step to pick *P. aeruginosa* double mutants with smooth surfaces (*shrA* was red and wrinkled). The insertion position of Tn5-*luxAB* transposon was determined by two-step PCR as described previously (Ramsey and Whiteley, 2004) with primers LuxAB inside and Arb1 for the first round PCR and LuxAB outside and Arb2 for the second round PCR (**Table 3**). The PCR product was sequenced using primer LuxAB outside and a BigDye Terminator Cycle Sequencing kit (Applied Biosystems, Foster City, CA).

c-di-GMP Assay

c-di-GMP was quantified using HPLC as described previously (Ueda and Wood, 2009) with slight modifications. Strains were grown in 1 LLB medium for 16 h at 250 rpm. HPLC was conducted using C18 reverse-phase column (150*3.9 mm, 4 µm, Nova-Pak, Waters) at a flow rate of 1 ml/min. Solvent A was 0.15 M TEAA buffer (pH 5.0). Solvent B was acetonitrile. The gradient was as follows: t = 0, 0% solvent B; $t = 35 \min, 12\%$ solvent B; t = 36 min, 80% solvent B; t = 41 min, 80% solvent B; $t = 42 \min$, 0% solvent B; $t = 55 \min$, 0% solvent B. Each sample had a running time of 55 min. A photodiode array detector (Waters, Milford, MA) was used to detect nucleotides at 254 nm after the HPLC separation step. Synthetic c-di-GMP (BIOLOG Life Science Institute, Bremen, Germany) was used as a standard. The peak corresponding to c-di-GMP from the extract of the *shrA* mutant was verified by co-elution with standard c-di-GMP. This experiment was performed with two independent cultures.

Plasmid Construction of pET28b-PA2444cHis and Purification of Recombinant ShrA-cHis

shrA was amplified with Pfu DNA polymerase using primers PA2444-F-XbaI and PA2444-R-HindIII (Table 3). The PCR product was digested with XbaI and HindIII and was cloned into the XbaI and HindIII sites of the pET28b vector. The resulting plasmid, pET28b-PA2444cHis has shrA fused to a 6 \times His tag at the C-terminus (ShrA-cHis) and under control of the T7 promoter. The pET28b-PA2444cHis plasmid was confirmed by DNA sequencing with the T7 promoter and T7 terminator primers (Table 3). Production of ShrA-cHis was induced in E. coli BL21(DE3) cells with 1 mM IPTG at a turbidity of 1.0 at 600 nm overnight at room temperature. Cells were resuspended in 20 mL lysis buffer (50 mM potassium phosphate buffer, pH 7.6, 400 mM NaCl, 1 mM dithiothreitol (DTT), and $50\,\mu\text{M}$ PLP) and disrupted twice by a French Press (Thermo Electron Corporation, Waltham, MA). ShrA-cHis was purified using a Ni-NTA resin (Qiagen, Valencia, CA) as described by the manufacturer's protocol. Purified ShrA-cHis was dialyzed against buffer (50 mM potassium phosphate buffer, pH 7.6, 50 mM NaCl, 8% glycerol, and 1 mM DTT) for three times at 4°C overnight to remove free PLP. Then the protein was concentrated using a 10 kDa cut-off centrifugal concentrator (Millipore, Billerica, MA), and the protein concentration was measured by using a Pierce BCA assay kit (Pierce, Rockford, IL).

ShrA Enzyme Activity

The forward and backward SHMT activities were determined using an HPLC-based fluorometric assay using purified recombinant protein. For the backward reaction toward the

production of glycine and (6S)-5,10-CH₂-THF (Figure 1A), the reaction mixture was prepared as described previously (Chaturvedi and Bhakuni, 2003) with slight modification. Briefly, 100 µL of assay mixture contained 50 mM sodium phosphate buffer, pH 7.6, 4 or 6 mM L-Ser, 2 or 3 mM (6R,S)-THF, 2 mM DTT, 1 mM EDTA, 50 µM PLP and approximately $0.5\,\mu M$ of purified ShrA. After incubation at $37^{\circ}C$ for 20 min, the reaction was stopped by boiling for 10 min and centrifuged at 13,000 rpm for 20 min to remove protein. The amount of Ser and Gly was quantified using the amino acid analysis by HPLC equipped with a fluorescence detector as described previously in (Wu and Meininger, 2008). Generally, 50 µl supernatant was treated with 50 µl of 1.5 M HClO₄ for 3 min, and then neutralized using 25 µl of 2 M K₂CO₃ and 1.125 ml H₂O. The sample was vortexed and centrifuged to obtain the supernatant for HPLC analysis. For HPLC, the sample was mixed with o-phthaldialdehyde (OPA) reagent solution for 1 min in a reaction loop and immediately delivered into the C18 column without delay. Amino acids react with OPA in the presence of β -mercaptoethanol to form highly fluorescent products.

For the forward reaction toward the production of serine and (6S)-THF (Figure 1A), the reaction mixture was prepared as described previously (Wei and Roje, 2011) with slight modifications. Briefly, (6R,S)-5,10-CH2-THF was obtained by incubating (6R,S)-THF with 13 mM formaldehyde in 50 mM sodium phosphate buffer (pH 7.6) with 1 mM DETA. For the reaction, 100 µL assay mixture contained 50 mM sodium phosphate buffer, pH 7.6, 20 or 30 mM Gly, 2 or 3 mM (6R,S)-5,10-CH₂-THF, 7.8 mM formaldehyde, 2 mM DTT, 1 mM EDTA, 50 µM PLP and approximately 0.5 µM of purified ShrA. The excess amount of formaldehyde ensures that the product (6S)-THF was immediately recycled back the substrate (6S)-5,10-CH2-THF, thereby the substrate (6*R*,*S*)-5,10-CH₂-THF concentration was constant throughout the reaction. After incubating at 37°C for 20 min, the reaction was stopped by boiling for 10 min and centrifuged at 13,000 rpm for 20 min to remove protein.

Oligomerization State of ShrA by Cross-Linking Using Glutaraldehyde

ShrA cross-linking was performed using a mild approach which employs the glutaraldehyde vapor as described previously (Fadouloglou et al., 2008). Briefly, ShrA protein was prepared as 0.5 mg/mL in 50 mM sodium phosphate buffer (pH 7.5) with 50 mM NaCl and 1 mM DTT. The hanging drop protein solution in a cover slip was incubated with the vapor from the 25% glutaraldehyde solution in a sealed well for various time points, and the size of the cross-linked products was examined by SDS-PAGE.

PLP Content for ShrA

The amount of PLP bound to the enzyme was determined for the recombinant enzyme as described previously (Ulevitch and Kallen, 1977) after extensive dialysis of the purified recombinant enzyme. Briefly, PLP will react with L-cysteine to form thiazolidine, and the amount of thiazolidine produced was determined from the absorbance at 335 nm.

Intracellular NADH and NAD⁺

Overnight cultures were diluted into fresh LB medium with an initial turbidity at 600 nm of 0.005. The culture (100 μ L) was spread on LB plates and incubated at 37°C for 24 h. Cells on plates were collected by rapidly resuspending in 5 mL of LB and centrifuging. Extraction of NAD⁺ and NADH was carried out using an acid and alkaline extraction method (Price-Whelan et al., 2007), and the NAD⁺ and NADH concentrations were quantified using an enzyme cycling assay with alcohol dehydrogenase (Price-Whelan et al., 2007) with slight modifications. Room temperature was used for the cycling assay conditions, and NADH and NAD⁺ standard samples (1~10 μ M) were used for calibration. Three independent cultures were used for each strain.

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AUTHOR CONTRIBUTIONS

TW conceived the project. MP, LS, SS, and TG conducted the experiments. MP and TW wrote the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb. 2018.00315/full#supplementary-material

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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