# c-di-GMP Turn-Over in *Clostridium difficile* Is Controlled by a Plethora of Diguanylate Cyclases and Phosphodiesterases

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# Abstract

Clostridium difficile infections have become a major healthcare concern in the last decade during which the emergence of new strains has underscored this bacterium's capacity to cause persistent epidemics. c-di-GMP is a bacterial second messenger regulating diverse bacterial phenotypes, notably motility and biofilm formation, in proteobacteria such as Vibrio cholerae, Pseudomonas aeruginosa, and Salmonella. c-di-GMP is synthesized by diguanylate cyclases (DGCs) that contain a conserved GGDEF domain. It is degraded by phosphodiesterases (PDEs) that contain either an EAL or an HD-GYP conserved domain. Very little is known about the role of c-di-GMP in the regulation of phenotypes of Gram-positive or fastidious bacteria. Herein, we exposed the main components of c-di-GMP signalling in 20 genomes of C. difficile, revealed their prevalence, and predicted their enzymatic activity. Ectopic expression of 31 of these conserved genes was carried out in V. cholerae to evaluate their effect on motility and biofilm formation, two well-characterized phenotype alterations associated with intracellular c-di-GMP variation in this bacterium. Most of the predicted DGCs and PDEs were found to be active in the V. cholerae model. Expression of truncated versions of CD0522, a protein with two GGDEF domains and one EAL domain, suggests that it can act alternatively as a DGC or a PDE. The activity of one purified DGC (CD1420) and one purified PDE (CD0757) was confirmed by in vitro enzymatic assays. GTP was shown to be important for the PDE activity of CD0757. Our results indicate that, in contrast to most Gram-positive bacteria including its closest relatives, C. difficile encodes a large assortment of functional DGCs and PDEs, revealing that c-di-GMP signalling is an important and well-conserved signal transduction system in this human pathogen.

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# Introduction

Clostridium difficile is a Gram-positive, anaerobic, spore-forming bacterium causing mild diarrhea to fulminant colitis in humans. Due to spreading of hypervirulent and high toxin-producing strains, C. difficile has caused in the last decade several epidemics in Europe and North America where it is now the leading cause of nosocomial diarrhea [1–3]. Its ability to sporulate allows this bacterium to remain dormant for years and survive to harsh conditions such as gastric acid after ingestion or the presence of oxygen in the environment. C. difficile-associated diseases are commonly associated with antibiotic usage, which creates a favorable niche for C. difficile to grow and cause infection in part by disrupting the gut microflora.

Bis-(3'-5')-cyclic dimeric guanosine monophosphate (c-di-GMP) is a bacterial second messenger controlling diverse bacterial phenotypes mostly known to be involved in the transition from free-living, motile to biofilm lifestyle in Gram-negative bacteria [4,5]. c-di-GMP has also been shown to be involved in the

development and cell cycle control in Caulobacter crescentus [6,7], and the modulation of virulence in several pathogens such as Vibrio cholerae, Vibrio vulnificus, Bordetella pertussis or Pseudomonas aeruginosa [8-11]. c-di-GMP is synthesized from 2 GTP molecules by enzymes named diguanylate cyclases (DGCs) that contain a GGDEF domain [12]. It is degraded respectively into pGpG or 2 GMP by phosphodiesterases (PDEs) that contain an EAL (PDEA) or a HD-GYP domain [13,14]. GGDEF domains were named by Hecht and Newton according to their conserved amino acid motif GG[D/E]EF [12]. EAL and HD-GYP domains are also named based on conserved amino motifs within these domains [13,14]. Whole genome analysis of a large number of bacterial species has revealed that the number of genes coding for enzymes involved in c-di-GMP turn-over varies widely between different species [15]. Genomes of proteobacteria generally encode a much wider array of such enzymes compared to those of Gram-positive bacteria. For instance, 66 genes coding for predicted enzymes involved in c-di-GMP turn-over are found in Shewanella oneidensis, 62 in V. cholerae, 41 in Pseudomonas aeruginosa, 29 in Escherichia coli, 6 in Bacillus subtilis

# **Author Summary**

c-di-GMP is a bacterial intracellular signalling molecule regulating motility, biofilm formation, cell cycle control, or virulence in Gram-negative bacteria. The function and importance of this molecule still remain unknown in Grampositive bacteria, even in important emerging pathogens such as Clostridium difficile, which causes from mild to deadly intestinal infections and has lately been on the rise in the healthcare setting and in the community. Here, we expose in the genomes of C. difficile strains a large number of conserved genes encoding proteins involved in the synthesis, degradation, or sensing of c-di-GMP, in contrast with most other Gram-positive bacteria including C. difficile's closest relatives. We confirmed the activity of most of these well-conserved proteins in a microorganism for which typical behavior alterations associated with variation of the intracellular c-di-GMP pools are known. We further confirmed the c-di-GMP synthesis and degradation activities of two purified C. difficile proteins. Our results indicate that c-di-GMP signalling is important in the lifecycle of this pathogen. This finding is particularly exciting because the c-di-GMP signalling network could serve as a target for the development of new drugs against C. difficile-associated diseases that are commonly associated with antibiotic usage.

and 1 in *Staphylococcus aureus*. In fact, very little is known about c-di-GMP's input in the regulation of phenotypes within Gram-positive bacteria. GdpS, the sole predicted staphylococcal GGDEF domain-containing protein, positively regulates biofilm formation in both *S. aureus* and *S. epidermidis*, and expression of protein A, a major virulence factor in *S. aureus*. However, GdpS does not appear to be an active DGC *in vitro* and its C-terminal GGDEF domain is not involved in these two phenotypes [16,17].

Although the enzymes producing and degrading c-di-GMP share customary domains in all bacteria, when known, the downstream effectors and pathways regulating the different phenotypical responses are usually different. c-di-GMP-sensing proteins containing the characterized c-di-GMP-binding PilZ domain [18] and some non-PilZ proteins are known c-di-GMP binding receptors. In E. coli and related bacteria, the PilZ domaincontaining protein YcgR was found to decrease motility by interacting with the flagellar motor to control its direction and rotation speed upon binding of c-di-GMP [19,20]. In V. cholerae, VpsT, a key transcription regulator that inversely regulates the expression of genes associated with motility and biofilm formation, was only recently found to be active following binding of c-di-GMP to its atypical receiver domain [21]. Recently, the first c-di-GMP-binding riboswitches (c-di-GMP-I) have been discovered in the genomes of V. cholerae, C. difficile and other bacteria [22]. Riboswitch Cd1 of C. difficile is located upstream of the large operon coding for the synthesis of the flagellum and was found to have an "off" switch action on transcription in an in vitro transcription assay and in  $\beta$ -galactosidase assays in *B. subtilis* [22]. Additionally, self-splicing of an unusual group I intron in C. difficile genome was found to be allosterically controlled by a c-di-GMP-II riboswitch aptamer, likely enabling the translation of a putative surface protein upon c-di-GMP binding [23]. Furthermore, 37 genes of C. difficile 630 that encode putative proteins containing GGDEF and/or EAL domains are available in the SignalCensus database [24]. Together, these observations suggest that c-di-GMP is a key signalling component in this emerging pathogen; yet studies on proteins regulating the intracellular c-di-GMP level, i.e. DGCs and PDEs, are still lacking.

The identification and characterization of enzymes producing or degrading c-di-GMP is a critical step to determine and understand the relevance of this second messenger in *C. difficile*'s lifecycle and virulence. In this study, we analyzed the prevalence and conservation of genes coding for putative DGCs and PDEAs in the genomes of 20 *C. difficile* isolates. Thirty-one conserved genes were assayed for their ability to encode functional DGC or PDEAs by evaluating the effect of their expression in *V. cholerae.* Most of these proteins conferred phenotypes that were consistent with their predicted function in our heterologous expression model. Our results indicate that, unlike the vast majority of Grampositive bacteria including the Clostridiaceae, *C. difficile* regulates its intracellular c-di-GMP pools via a plethora of functional DGCs and PDEAs.

# Results

# Domain composition and activity prediction of putative DGCs and PDEAs encoded by *C. difficile* 630

Initial examination of the Pfam database 24.0 [25] for proteins involved in c-di-GMP turn-over in C. difficile 630, correlated to data provided by the NCBI's SignalCensus database [24], reveals a total of 37 proteins containing a GGDEF (Pfam PF00990) and/ or an EAL (Pfam PF00563) domain: 18 proteins have a GGDEF domain, one protein contains an EAL domain and 18 proteins have both GGDEF and EAL domains. Proteins containing both GGDEF and EAL domains have been shown to act either as DGCs or as PDEAs, or to exhibit both activities [14,26,27]. Furthermore, not all proteins containing a GGDEF or an EAL domain have been shown to exhibit DGC or PDEA activity. Several conserved amino acids in the GGDEF and EAL domains are predicted to be important to confer enzymatic activity [28,29], among which the highly conserved motifs GG[D/E]EF and EXLR, respectively. These motifs and other conserved amino acid residues were sought for in the primary sequences of the 37 putative c-di-GMP-signalling proteins encoded by the genome of C. difficile 630 by multiple sequence alignment (Figure 1). Briefly, among those 37 proteins, 15 are most likely active DGCs, 18 could be active PDEAs, 1 protein (CD0522) could either be an active DGC and/or PDEA, and 3 contain a predicted catalytically inactive GGDEF domain (Table S1). Except for CD0522, all the proteins with both GGDEF and EAL domains were predicted to be PDEAs having an inactive GGDEF partner domain since these have a degenerated GG[D/E]EF motif. Most of the 37 proteins are predicted to have at least one sensor domain, transmembrane regions or a signal peptide region.

# Distribution of the putative c-di-GMP-signalling components in *C. difficile*

To assess whether c-di-GMP signalling components are conserved within the species *C. difficile*, orthologs of the proteins identified in strain 630 were exhaustively sought for in 19 other partially or completely sequenced *C. difficile* genomes using tblastn. Most genes found in *C. difficile* 630 are conserved in all the strains examined with 31 of the 37 GGDEF and/or EAL proteins having an ortholog in at least 15 strains and only CD2753 and CD2754 being unique to strain 630 (Figure 2A). The putative glycosyltransferase CD2545, the sole protein predicted to contain a PilZ domain (Pfam PF07238), has orthologs in all the other analyzed strains (Figure 2A). Furthermore, two RNA targets (named herein Cd1-a and 84Cd, respectively) that have been shown to bind c-di-GMP, the riboswitch Cd1 located upstream of the flagellum synthesis operon, and the self-splicing group I intron (tandem riboswitch-ribozyme) are conserved in a subset of strains [22,23].

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**Figure 1. Domain composition and organization of the 37 putative c-di-GMP-signalling proteins encoded by** *C. difficile* **630.** Proteins names are listed on the left and predicted sizes in amino acids are indicated on the right. GGDEF domains are shown in green and EAL domains are shown in red. The DGC PleD from *C. crescentus* (YP\_002517919) and the PDEA RocR from *P. aeruginosa* (NP\_252636) are shown as references to highlight amino acids important for enzymatic activity. The putative functions and positions of important amino acid residues [28,29] are listed above each domain in black (conserved) or grey (non-conserved) and their position in PleD and RocR are indicated. Blue boxes are predicted signal peptide regions. Black boxes represent predicted transmembrane regions. Grey boxes represent predicted coiled-coil motifs. Additional sensor domains predicted by Pfam 24.0 are also shown in white. Proteins are not drawn to scale. Cache 1, Calcium channels and chemotaxis receptor family 1; PAS 3, PAS fold family 3; IP, primary inhibitory site; I<sub>5</sub>, secondary inhibitory site. doi:10.1371/journal.pgen.1002039.q001

The conservation pattern of the c-di-GMP regulatory proteins and downstream effectors clearly follows the phylogenetic distribution of the strains (Figure 2A and 2B). Interestingly, the cluster of 10 hypervirulent NAP1/BI/027 strains (cluster CD196/2007855 in Figure 2B) regroups the strains encoding the lowest number of c-di-GMP signalling components.

Like CD2753 and CD2754 that are specific to C. difficile 630, we assumed that other strains could also encode c-di-GMP signalling proteins that were absent from all of the other strains. The Microbial Signal Transduction database (MiST<sub>2</sub>) [30], which currently provides data for 14 out of our 19 strains, contains 3 putative c-di-GMP turn-over proteins that are absent from C. difficile 630's proteome. Notably, CdifA\_020200002673, is unique to strain ATCC43255 (Figure 2A), and is the only HD-GYP domain-containing protein detected in C. difficile. Based on analysis of amino acid conservation, it is predicted to have both DGC and PDE enzymatic activities (Figure S1). To identify additional strainspecific genes encoding c-di-GMP signalling proteins, we performed profile hidden-Markov model (profile HMM) searches with the Pfam HMMs for GGDEF, EAL, HD and PilZ conserved domains using HMMER3 software against the proteomes of strains 2007855, BI1, BI9, CF5, M120 and M68 as predicted by GeneMark.hmm (Table S2). No additional c-di-GMP-signalling proteins were detected. Besides CD2545, no other PilZ domaincontaining protein was found. Finally, no GGDEF, EAL, HD-GYP or PilZ domain-containing proteins were found to be encoded by C. difficile plasmids pCD630, pCD6, pCDBI1 or the 300 kb putative phage or plasmid from BI1 (Table S2)

The remarkable prevalence of c-di-GMP-signalling components among *C. difficile* strains suggests that c-di-GMP is an important second messenger in this bacterium. The functionality of the 31 most conserved GGDEF and/or EAL domain proteins among *C. difficile* strains (Figure 2A) was therefore assessed to confirm their biological activity.

# Heterologous expression of DGC and PDEA encoding genes in *V. cholerae*

To the best of our knowledge, no model of Gram-positive bacteria is currently available to efficiently and reliably evaluate in vivo the enzymatic activity of proteins regulating the intracellular levels of c-di-GMP. Moreover, to circumvent the tedious laboratory procedures associated with working with C. difficile and the lack of information regarding the phenotypes regulated by c-di-GMP in this bacterium, assessment of the enzymatic activity of the 31 putative DGCs and PDEAs (cloned from C. difficile 630) was carried out by heterologous expression in V. cholerae. Characteristic phenotype alterations associated with variations of the intracellular c-di-GMP concentration of V. cholerae are easily observable and measurable. High levels of intracellular c-di-GMP concentrations increase biofilm formation and decrease motility whereas lower c-di-GMP concentrations cause the opposite effects [31]. As expected, ectopic expression of most of the predicted DGCs containing the canonical GG[D/E]EF motif decreased cell motility (Figure 3A) and increased biofilm formation (Figure 3C),

supporting the hypothesis that these C. difficile proteins are genuine and functional DGCs (Table S1). Replacement of the second glycine of the GGDEF motif by site-directed mutagenesis has been shown to greatly reduce the enzymatic activity of other DGCs [32,33]. Such a mutation in CD1420 (mutant G204E) abolished alterations of both biofilm and motility phenotypes (Figure 4A, 4B and 4C) confirming that the phenotypical alterations observed in V. cholerae are linked to the enzymatic activity of this functional C. difficile DGC. Moreoever, the expression of CD1420 in V. cholerae N16961 led to a dramatic increase of the c-di-GMP level compared to the same strain expressing LacZ (Figure S2B and S2D, Text S1). Furthermore, we observed that CD1015, CD1185 and CD1420, the DGCs causing the strongest phenotypical shifts (Figure 3A and 3C), lacked the amino acid motif RXXD that is part of the retro-inhibition site (I-site) of the GGDEF domain (Figure 1) [34]. The putative DGC CD2887 significantly enhanced biofilm formation without affecting the motility of V. cholerae. Interestingly, CD2887 contains a GGEEY motif instead of the canonical GG[D/E]EF motif suggesting that it might not be a functional DGC. However, Malone and colleagues [35] showed that upon substitution of the F amino acid residue for a Y residue in the A-site, the DGC WspR of Pseudomonas fluorescens retains its activity. Unlike the putative DGCs presented above, several predicted DGCs that appear to possess all the conserved amino acid residues necessary for c-di-GMP synthesis (e.g. CD2385 and CD3365) did not modulate biofilm formation or motility of V. cholerae in our experimental conditions. Yet we cannot conclude that these putative DGCs are not functional since they might not have been produced, may have been unstable or may lack the appropriate activating signal in the heterologous host. Unexpectedly, CD0537, which also has a canonical A-site, did not enhance biofilm formation but enhanced cell motility by  $\sim 60\%$ . While this increase is notable compared to other DGCs, it remained very modest compared to the increase promoted by PDEAs (see below) suggesting that in our experimental setting, this putative DGC was not functional. The unexpected result on motility could result from partial sequestration of intracellular c-di-GMP by CD0537's I-site upon overexpression of the protein. Additionally, the lack of apparent DGC activity of CD0537's might simply be due to a lack of phosphorylation of its phosphoreceptor REC domain, a modification that could be catalyzed by the putative kinase *cheA* (CD0539) that was absent in our assay. Other DGCs have been shown to be activated by phosporylation of their phosphoreceptor REC domain [29,32,36]. As expected, CD1028 and CD2384, which respectively possess the degenerated QKDMI and GGEEI motifs, did not alter V. cholerae's phenotypes. Consistent with our observation, the substitution of the F residue of the A-site for an I residue alone is known to eliminate the activity of WspR of P. fluorescens [35].

Expression of several putative PDEAs (CD0757, CD1515, CD1616, CD1840 and CD2663) significantly enhanced cell motility on soft agar by 2 to 4 fold (Figure 3B and Table S1). While five putative PDEAs exhibited significant activity (CD0204, CD1421, CD1515, CD2134 and CD2663), in most cases, the



Figure 2. Conservation of c-di-GMP-signalling components of *C. difficile* **630** in other strains of *C. difficile*. (A) Protein orthologs exhibiting  $\geq$ 90% identity are indicated by dark grey boxes. More divergent proteins sharing between 85 and 89% identity or containing

major gaps are indicated by light grey boxes. Proteins that are absent are indicated by empty boxes. The c-di-GMP-sensing riboswitch Cd1 ((RS) Cd1) and 84Cd aptamer from the tandem riboswitch-ribozyme ((RZ) 84Cd) are indicated by a dark gray box when they shared  $\geq$ 90% and were in the same genetic context as in strain 630. CdifQ2682, CdifA2673, and CdifA6827 were used to refer to proteins CdifQCD-2\_02020002682 (ZP\_05400016) from *C. difficile* QCD-23m63, CdifA\_020200002673 (ZP\_05349638) and CdifA\_020200006827 (ZP\_05350459) from *C. difficile* aTCC 43255, respectively. (B) Phylogenetic tree of *C. difficile* isolates based on *rpoB* nucleotide sequences. Bootstrap values are indicated at branch point in percent. Only values  $\geq$ 70% are shown.

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impact of ectopic expression of these proteins on biofilm formation was modest and not significant (Figure 3D). The weak response of *V. cholerae* N16961 to PDEA activity could be due to its low basal level of biofilm formation in our assays. However, we indirectly confirmed that the *V. cholerae* motility response to overexpression of these proteins was linked to PDEA activity by mutating the glutamic acid residue of the EVLxR motif of CD0757, which is critical for enzymatic activity. Indeed, unlike the wild-type protein, overexpression of CD0757-E339A did not enhance the motility of *V. cholerae* N16961 on soft agar (Figure 4A and 4C).

## CD0522 has dual enzymatic activity

Since CD0522 has a particular combination of GGDEF and EAL domains suggesting that it could have both DGC and PDEA activities, it was analyzed apart from those in Figure 3. CD0522 contains two predicted N-terminal GGDEF domains and one predicted C-terminal EAL domain (Figure 1 and Figure 4D). Our analysis indicates that the first GGDEF domain and the EAL domain should be catalytically active due to the conservation of the A-sites. However, the second GGDEF domain contains the strongly degenerated motif YADVF suggesting that it is catalytically inactive (Figure 1 and Figure 4D). CD0522 or its individual domains were tested in our V. cholerae heterologous expression model to verify these predictions (Figure 4C, 4E and 4F). Since the variations of cell motility and biofilm formation we observed upon ectopic expression of CD0522 were not statistically significant, we could not clearly establish a DGC activity for the complete CD0522. On the other hand, expression of the N-terminal fragment N-448, which encompasses the first GGDEF domain, reduced motility by more than half and increased biofilm by  $\sim 7$ fold, suggesting that this fragment of CD0522 acts as a functional DGC (Figure 4E and 4F). The observed phenotypes correlate with a marked increase of intracellular c-di-GMP upon expression of N-448 in V. cholerae N16961 compared to the same strain expressing LacZ (Figure S2B and S2C, Text S1). DGC catalytic activity of CD0522 was also indirectly confirmed by mutating the glutamic acid residue of the EAL motif to abolish any possible PDEA activity (Figure 4D). Expression of CD0522-E814A altered phenotypes as expected for a DGC, as observed for N-448 (Figure 4C, 4E and 4F). Expression of the C-terminal fragment C-307 led to a modest but significant increase of motility. As expected, expression of the degenerated GGDEF-containing central fragment M-305 did not lead to any significant change of phenotype compared to the control. The degenerated GGDEF domain of M-305 could be involved in the regulation of the PDEA activity of the EAL domain of CD0522. We observed that motility of cells expressing MC-591, which contains the central and Cterminal fragments, increased by  $\sim 50\%$  (Figure 4E), which is consistent with a diminution of intracellular c-di-GMP. Increased motility was also observed when we overexpressed CD0522-G366E, a protein containing a substitution of the second glycine



**Figure 3. Phenotypic changes triggered by the expression of** *C. difficile* **630 genes encoding predicted DGCs or PDEAs in** *V. cholerae* **N16961.** Relative motility of cells over-expressing (A) predicted DGCs or (B) predicted PDEAs. Relative biofilm formation of mutant cells over-expressing (C) predicted DGCs or (D) predicted PDEAs. *V. cholerae* over-expressing *lacZ* was used as a reference control. The means and standard deviations obtained from at least three independent assays are shown. One-way ANOVA with a Dunnett multiple comparison post-test was used to compare the strains over-expressing putative c-di-GMP signalling proteins and the control strain (LacZ) (\*, P<0.05; \*\*, P<0.01; \*\*\*, P<0.001). doi:10.1371/journal.pgen.1002039.g003

residue of the first GGDEF to abolish any DGC activity (Figure 4C and 4E). Complex proteins such as CD0522 that are composed of several GGDEF and EAL domains suggest a possible two-way c-di-GMP control and must be studied in detail to reveal what stimuli switches their enzymatic activity between the DGC or PDEA state.

### In vitro enzymatic assays

CD1420 and CD0757 enzymatic activities were further assessed in vitro to corroborate the results obtained in the V. cholerae expression model and confirm that the C. difficile proteins are genuine DGC and PDEA, respectively. These proteins were chosen for their strong activity in V. cholerae and the simplicity of the structure of the N-terminal sensor domain that suggested little requirements for *in vitro* assays (Figure 1). Purified CD1420 in its native form was able to produce c-di-GMP from GTP as substrate and the accumulation of the product increased with time (Figure 5A). Conversely, purified CD0757 did not produce c-di-GMP from GTP even after 1 h incubation. Therefore, we confirmed that CD1420 is a functional DGC. The absence of DGC activity of CD0757 suggests that it contains an inactive GGDEF domain and acts as a PDEA only. CD0757 was then assessed for PDEA activity on c-di-GMP. cdi-GMP hydrolysis by PDEAs is known to yield the linear diguanylate pGpG [37]. We incubated purified CD0757 with radiolabeled c-di-GMP, yielding small amounts of pGpG. This characteristic PDEA activity was abolished by denaturing the protein prior to the assay (Figure 5B). Inactive GGDEF domains have been shown to enhance PDEA activity of an adjacent EAL domain by binding GTP [26]. Addition of GTP to the enzymatic reaction increased noticeably the PDEA activity of CD0757 presumably through binding to the GGDEF domain like for PdeA (CC3396) from *C. crescentus*. After a 30-min incubation period, virtually all the c-di-GMP was converted to pGpG. Marginal degradation of c-di-GMP to GMP by CD0757 was detected as previously shown to occur with another PDEA [37].

#### Discussion

Studies on c-di-GMP have addressed with some depth many aspects regarding the proteins involved in its synthesis (DGCs and PDEAs) and the molecular targets of c-di-GMP such as proteins and riboswitches in several bacteria. While c-di-GMP signalling has been extensively studied in many Gram-negative bacteria like



Figure 4. Alteration of *V. cholerae* N16961 motility and biofilm formation upon expression of wild-type or mutated putative DGCs and PDEAs from *C. difficile* 630. (A) Relative motility upon expression of wild-type or mutated CD0757 or CD1420. (B) Relative biofilm expression upon expression of wild-type or mutated CD0757 or CD1420. (C) Motility of the strain expressing the wild-type (WT) and mutated proteins on LB soft agar plates at 30°C. The picture shown is representative of four independent experiments. (D) Schematic representation of CD0522 protein fragments expressed. The amino acids in the signature motifs of the GGDEF or EAL domains are indicated above each domain. Positions of the mutations in the GGDEF or EAL domains are indicated by asterisks. (E) Relative motility upon expression of full-length, mutated or truncated CD0522. (F) Relative biofilm formation upon expression of full-length, mutated or truncated CD0522. (F) Relative biofilm formation upon expression of full-length, mutated or truncated CD0522. (F) Relative biofilm formation upon expression of full-length, mutated or truncated CD0522. (F) Relative biofilm formation upon expression of full-length, mutated or truncated CD0522. (F) Relative biofilm formation upon expression of full-length, mutated or truncated CD0522. (F) Relative biofilm formation upon expression of full-length, mutated or truncated CD0522. (F) Relative biofilm formation upon expression of full-length, mutated or truncated CD0522. The means and standard deviations obtained from four independent assays are shown. One-way ANOVA with a Dunnett multiple comparison post-test was used to compare the cells over-expressing putative c-di-GMP signalling proteins and the control strain (LacZ) (\*, P<0.05; \*\*, P<0.01; \*\*\*, P<0.001). doi:10.1371/journal.pgen.1002039.g004

*C. crescentus, E. coli, V. cholerae, Salmonella* and *Pseudomonas,* very few studies have been carried out on Gram-positive bacteria. To the best of our knowledge, the staphylococcal GGDEF domain protein GdpS has been the only c-di-GMP regulatory protein studied to date in low G+C Gram-positive bacteria. GdpS does not seem to have any measurable DGC activity [16]. The recent discovery of a functional c-di-GMP binding riboswitch in *C. difficile* and *Bacillus cereus,* as well as the prediction of several other similar riboswitches in other Gram-positive bacteria has revived the interest in studying c-di-GMP metabolism in these microorganisms. The recent characterization of a c-di-GMP-dependent self-splicing group I ribozyme in *C. difficile* further reinforces the role of c-di-GMP in Gram-positive bacteria.

In this work, we have shown that many of the genes encoding putative DGCs and PDEAs of *C. difficile* behave like genuine DGCs and PDEAs in heterologous expression experiments (Figure 3 and Figure 4, Table S1). This number of c-di-GMP regulatory proteins encoded by *C. difficile* is high compared to what is found in its closest relatives (Table S3), and also among the Firmicutes in general (median = 1) [38]. Analysis of the genomes of 49 strains of Clostridiaceae representing 27 species revealed that most contain less than 20 of such genes (Table S3). Only 2 species of *Clostridium* were found to encode more putative DGCs/PDEAs than *C. difficile, Clostridium asparagiforme* DSM15981 and *Clostridium bolteae* ATCC BAA-613, two newly characterized yet barely

studied species isolated from human fecal samples [39,40]. The disparity in the occurrence of c-di-GMP signalling proteins is remarkable. The two species coding for the lowest number of c-di-GMP regulatory proteins, Clostridium hiranonis and Clostridium bartlettii, are the closest phylogenetically related species to C. difficile (Figure S3 and Table S3). On the opposite, the two species coding for the highest number of GGDEF, EAL or HD-GYP protein, C. asparagiforme and C. bolteae, are among the most distant species from C. difficile. Additionally, C. difficile, which encodes with one exception no HD-GYP domain proteins, seems to be an exception among the Clostridiaceae and contrasts with Clostridium beijerinckii which encodes 14 HD-GYP domain proteins (Table S3), while retaining the same number of c-di-GMP regulatory proteins as C. difficile. In addition, C. difficile does not seem to carry any gene encoding c-di-GMP-signalling proteins that could have been recently exchanged by horizontal transfer with the 3 other Clostridium species containing the highest number of such proteins (Table S3 and data not shown). The Clostridiaceae seem to have a high number of c-di-GMP-signalling proteins among the Firmicutes in general, but it remains similar to other Firmicutes of comparable size (3000-4000 genes, median = 10) [38].

The high number of c-di-GMP turn-over proteins in *C. difficile* is likely indicative of the importance of this second messenger in the bacterium's lifecycle and suggests a major role in regulating different phenotypes. The diversity of N-terminal structures



Figure 5. TLC analysis of the enzymatic activities of CD1420 and CD0757. (A) DGC activity of purified CD1420. (B) PDEA activity of purified CD0757. Snake venom phosphodiesterase (SVPD) was used as a positive control. doi:10.1371/journal.pgen.1002039.g005

suggests that their function is not redundant. Instead, these proteins could individually act in a functionally or spatially sequestered way, in addition to being temporally regulated through differential expression. It has been shown that DGCs usually are not interchangeable and can contribute to very specific and distinct phenotypes for a unique microorganism. For example, while the DGC YddV of E. coli impacts poly-N-acetylglucosamine production, other DGCs like AdrA do not [41]. Instead AdrA controls the production of cellulose, another exopolysaccharide, in E. coli and Salmonella [42,43]. Furthermore, the prevalence of DGCs and PDEAs in C. difficile could also indicate the importance of these proteins in sensing and relaying a diversified array of environmental conditions through their sensor domains or their eventual differential expression. In V. cholerae, the PDEA CdpA is not expressed in vivo until the late stage of infection in a mouse colonization model [44]. C. difficile vegetative cells encounter various environmental conditions during their journey through the gastrointestinal tract, during which c-di-GMP signalling might play a role in regulating diverse phenotypes.

Despite the current lack of experimental data, it is reasonable to assume that c-di-GMP regulates at least two phenotypes in C. difficile: flagella synthesis/motility and polysaccharide synthesis. A putative c-di-GMP-binding PilZ domain is located in the putative glycosyltransferase CD2545, which is predicted to be a cellulose synthase. Interestingly, although motility is commonly controlled by c-di-GMP in bacteria, the c-di-GMP-responsive effectors that likely regulate this phenotype in C. difficile appear to differ from those found in V. cholerae, E. coli and related bacteria. The c-di-GMP-sensing riboswitch Cd1 appears to control the transcription of the large operon of genes essential for assembling the flagellum apparatus [22]. Bacterial flagella are obviously important for motility but can also be involved in adhesion. Adhesion to mouse cecal mucus of the flagellin FliC and of the flagellum cap protein FliD of C. difficile has been demonstrated in vitro [45]. Moreover, FliD has been shown to specifically adhere to cultured cells [45]. Therefore, c-di-GMP signalling might impact both cell motility and adhesion of *C. difficile* to mucosal surface through the regulation of flagellum assembly. Additionally, c-di-GMP signalling may play a significant role in the excessive inflammation caused by *C. difficile* infection since flagellin is a very potent immunogenic protein recognized as a proinflammatory ligand by toll-like receptor 5 (TLR-5) located at the baso-lateral surface of intestinal cells (reviewed in [46]).

Except for one EAL protein (CD3650), all of C. difficile's PDEAs contain a GGDEF domain predicted to be non-catalytically active as shown in vitro for CD0757 (Figure 1 and Figure 5). Composite proteins containing both GGDEF and EAL domains are relatively frequent, representing approximately one third of proteins with such domains [47]. Some of these composite proteins act as DGCs, PDEAs, or have both activities (reviewed in [47]). Inactive GGDEF or EAL domains can act as sensor domains rather than catalytic domains. GGDEF domain proteins with degenerated active sites have been reported to bind c-di-GMP at their conserved I-site, as for the C. crescentus protein PopA [48], or to retain the ability to bind GTP at their degenerated active site, as for the C. crescentus protein PdeA [26]. Christen and colleagues [26] have demonstrated that binding of GTP to the inactive GGDEF domain of PdeA of C. crescentus, strongly enhanced the PDEA activity of the C-terminal EAL domain. The authors formulated two hypotheses to explain why the PDEA activity is linked to GTP intracellular concentrations: (i) to prevent GTP pools to drop by the uncontrolled successive activities of DGCs and PDEAs and (ii) to sense physiological changes. Intracellular GTP levels have been shown to impact the activity of CodY, a major transcriptional regulator in many low G+C Gram-positive bacteria such as C. difficile, S. aureus, Streptococcus pneumoniae, Streptococcus pyogenes, Streptococcus mutans, Listeria monocytogenes, B. cereus and Bacillus anthracis in which it affects virulence gene expression ([49] and references therein). CodY is known to have greater affinity to target DNA promoter regions upon binding of two synergistic

effectors, GTP and branched-chain amino acids [50,51]. *C. difficile* CodY has been shown to repress the expression of the toxin A and B genes (*tcdA* and *tcdB*), through binding to the promoter region of the positive transcriptional regulator TcdR [52]. A recent study aimed at identifying all DNA promoter regions targeted by *C. difficile* CodY as well as genes differentially expressed in a *codY* null mutant [49]. Interestingly, among the 165 genes identified with altered expression, PDEA genes CD0757 and CD1476 were highly derepressed. Additionally, DNA regions containing CD1476, CD2385, CD2873, CD2965 and CD3650 were identified as CodY binding-sites. These data suggest a probable interplay between the c-di-GMP and CodY signalling pathways, known to be important in the regulation of many metabolic genes and of the major virulence factors, toxins A and B [49,52].

To the best of our knowledge, no model of Gram-positive bacteria is currently available to efficiently and reliably evaluate the enzymatic activity of proteins regulating the intracellular levels of c-di-GMP. With the recent availability of molecular tools for *C. difficile* genetic manipulation [53–55], it will finally be possible to study in detail the many genes involved in c-di-GMP signalling and turn-over in this bacterium and to identify the phenotypes associated with the variation of intracellular c-di-GMP pools. The need to decipher the regulatory mechanisms underlying *C. difficile*'s behaviors is imperative to the development of new therapeutics and treatment strategies. Particularly, the bacterial signalling pathways and phenotypes involved at the colon mucosal interface ought to be addressed.

## **Materials and Methods**

#### Bioinformatics

Proteins containing the c-di-GMP-associated conserved domains (GGDEF, EAL, HD for HD-GYP and PilZ) were searched for in the Clostridiaceae proteomes on the Pfam 24.0 server [25]. HD domains were further analyzed to identify HD-GYP domains by looking for the HD-GYP amino acid motif by multiple alignment with the HD-GYP domain of Rpfg from Xanthomonas campestris 8004 (Accession number AAY49388) using ClustalW version 2.0.12 [56]. Other conserved domains, signal peptides, transmembrane regions, and coiled-coil motifs annotations are as determined on Pfam 24.0 [25]. Identification of proteins with c-di-GMP-associated conserved domains in C. difficile strains other than 630 was achieved with the hmmsearch program of the HMMER 3.0 software (http://hmmer. org/). C. difficile annotated protein sequences were retrieved for the 13 other strains and 2 plasmids available in the NCBI Refseq database (Table S2) [57]. Protein sequences from C. difficile 2007855, BI1, BI9, CF5, M120 and M68 genomes and extrachromosomal sequences (Table S2) were predicted using Gene-Mark.hmm for Prokaryotes version 2.4 [58]. Profile hidden Markov models (profile-HMMs) of c-di-GMP-associated conserved domains were downloaded from Pfam 24.0. The bit score threshold values used in every search were the "trusted cutoff" values for the Pfam profile-HMMs. Proteins containing the c-di-GMP-related conserved domains identified using HMMER 3.0 software were further analyzed to identify other conserved domains (Pfam 24.0), signal peptides and transmembrane regions (Phobius [59]), and coiled-coil motifs (ncoils [60]). Nucleotide and amino acid conservation of selected C. difficile 630 genes and proteins were assessed with the appropriate BLAST algorithms [61]. Since most of the genomes are drafts, pseudogenes were ignored and assumed to be the results of sequencing errors. As a matter of fact, pseudogenes are found in many of these strains even for important, unique and well-conserved genes such as the gene encoding DNA polymerase I (data not shown).

Phylogenetic trees were generated using the neighbor-joining method as implemented by ClustalX version 2.012 [56] from gapless alignments of nucleotide sequences. Nucleotide sequences were aligned using ClustalW version 2.0.12 [56] and gap columns were removed using Jalview version 2.5 multiple alignment editor [62]. The reliability of each tree was subjected to a bootstrap test with 1000 replications. Trees were edited using FigTree version 1.3.1 (http://tree.bio.ed.ac.uk/software/figtree/).

#### Growth conditions

Bacterial strains were routinely grown in Luria–Bertani (LB) broth at 37°C in an orbital shaker and maintained at -80°C in LB broth containing 15% (v/v) glycerol. Ampicillin (Ap) was used at 100 µg ml<sup>-1</sup> when needed. For induction of gene expression in the strains carrying arabinose-inducible vectors (pBAD series), L-arabinose was added to the growth medium at a final concentration of 0.02% (w/v).

## Bacterial strains and plasmid construction

The bacterial strains and plasmids used in this study are described in Table S4. The oligonucleotides used for plasmid constructions are described in Table S5. For expression of putative DGCs and PDEAs in *V. cholerae*, genes cloned in pBAD-TOPO were amplified by PCR with their native Shine-Dalgarno sequence using *C. difficile* 630 genomic DNA as a template. Truncated versions of CD0522 were cloned to include the native Shine-Dalgarno sequence of CD0522. DNA was amplified to express CD0522 protein fragments N-448, M-305, C-307 and MC-591 respectively containing the N-terminal 448 amino acids (aa), 305 aa encompassing the middle domain, the 307 aa in C-terminal and 591 aa encompassing the middle and C-terminal domains (Figure 4D).

Plasmids pCD0522-G366E, pCD0522-E814A, pCD0757-E339A and pCD1420-G204E, which accordingly contain amino acid substitutions in their respective conserved GG[D/E]EF or EXL A-sites, were created by site-directed mutagenesis of pCD0522, pCD0757 and pCD1420 using the QuickChange Lightning Site-directed Mutagenesis Kit (Stratagene) using primer pairs listed in Table S5. The mutations introduced were designed to create new EarI or PvuII restriction sites for initial screening of the mutated plasmids. Mutated genes were verified by sequencing.

For CD0757 and CD1420 proteins purification, the corresponding genes were amplified by PCR from *C. difficile* 630 genomic DNA and cloned into BamHI/SalI-digested pGEX6P-1 in frame with the glutathione S-transferase (GST) coding sequence.

#### Molecular biology methods

All the enzymes used in this study were obtained from New England BioLabs and were used according to the manufacturer's instructions. Plasmid DNA was prepared with a Qiaprep Spin miniprep kit (Qiagen). Genomic DNA of C. difficile 630 was extracted using the illustra bacteria GenomicPrep mini spin kit (GE Healthcare). PCR assays were performed with the primers described in Table S4 in 50 µl of PCR mixtures with 1 U of Pfu Ultra DNA polymerase (Agilent). PCR conditions were as follows: (i) 3 min at 94°C, (ii) 30 cycles of 30 s at 94°C, 30 s at suitable annealing temperature, and 30-300 s at  $72^{\circ}$ C, and (iii) 5 min at 72°C. When needed PCR products were purified using a QIAquick PCR Purification Kit (Qiagen) according to the manufacturer's instructions. E. coli was transformed by electroporation according to Dower and colleagues [63]. V. cholerae was transformed by electroporation according to Occhino and colleagues [64]. In both cases, transformation was carried out in

0.1 cm electroporation cuvettes using a Bio-Rad GenePulser Xcell apparatus set at 25  $\mu F,$  200  $\Omega$  and 1.8 kV.

## Motility and biofilm formation assays

Motility and biofilm assays were performed as described before [32]. Briefly, a semi-solid medium composed of 1% tryptone, 0.5% NaCl, 0.3% agar supplemented with ampicillin and L-arabinose was used to evaluate motility of V. cholerae mutant strains during over-expression assays at 30°C. Motility was assessed from the comparison of the surface area  $(mm^2)$  of the colonies from plate images captured and analyzed using a Gel Doc XR system and Quantity One software (Bio-Rad). The capacity of V. cholerae mutant strains to form biofilm was determined after 6 h static growth in LB broth containing ampicillin and L-arabinose at 30°C. Bound crystal violet was solubilized with 200 µl of 95% ethanol and quantified by absorbance at 595 nm in a Model 680 microplate reader (Bio-Rad). Motility and biofilm formation assays were carried in triplicate and data were normalized as fold expression compared with the control LacZ over-expressing bacteria. Data from at least three independent experiments were combined.

#### Production of recombinant proteins

Overnight-grown cultures of E. coli BL21 bearing pGCD0757 or pGCD1420 were diluted 1:100 in fresh  $2 \times$  YTA broth and incubated at 37°C with agitation. Protein expression was induced with 0.1 mM IPTG (isopropyl 1-thio-β-D-galactopyranoside) at mid-exponential phase ( $OD_{600}$  of 0.6) for CD0757 or at lateexponential phase ( $OD_{600}$  of 1.2) for CD1420. The cultures were grown for an additional 4 h at 37°C for CD0757 or 2 h at 25°C for CD0757. Cells were collected by centrifugation, re-suspended in PBS containing 1% Triton X-100 and protease inhibitors (Protease Inhibitor Cocktail, Sigma), and lysed by sonication. CD0757 and CD1420 were recovered by affinity chromatography using the GST purification module (GE Healthcare) with the PreScission protease (GE Healthcare) according to the manufacturer's instructions. After elution, proteins samples were dialyzed against the conservation buffer (50 mM Tris-HCl pH 7.8, 250 mM NaCl, 25 mM KCl, 10 mM MgCl<sub>2</sub>, 30% glycerol) for 18 h in D-Tube Dialyzer Maxi (MWCO 12-14, Novagen), concentrated by centrifugation on Amicon Ultra-15 columns (MWCO 10, Millipore), and stored at  $-20^{\circ}$ C. Protein concentration was estimated using a BCA Protein Assay Kit (Thermo-Scientific) and purity was determined by SDS-PAGE analysis.

#### Assays for enzymatic activity and TLC analysis

Diguanylate cyclase and phosphodiesterase activities were measured according to previously described procedures [32,37] with the following modifications. Diguanylate cyclase assays were performed with approximately  $1-2 \mu g$  of purified proteins in a final volume of 50 µl. Reaction mixtures were pre-incubated for 5 min at 30°C in the reaction buffer (50 mM Tris-HCl pH 7.8, 250 mM NaCl, 25 mM KCl, 10 mM MgCl<sub>2</sub>). DGC reactions were initiated by adding 33.3 nM [ $\alpha$ -<sup>33</sup>P]-GTP (0.1  $\mu$ Ci  $\mu$ l<sup>-1</sup>) and incubated at 30°C. Samples were taken at various times, and the reactions were stopped by addition of one volume 0.5 M EDTA. Radiolabeled c-di-GMP for phosphodiesterase activity assays was synthesized using purified DgcK [32]. Purified DgcK (30 µg) was incubated 8 h at 30°C in the reaction buffer to completely convert  $[\alpha^{-33}P]$ -GTP into c-di-GMP. Reactions were stopped by denaturing at 99°C for 15 min, centrifuged for 2 min at 16,000 g to elimate DgcK and recover the supernatant containing the radiolabeled c-di-GMP. Phosphodiesterase assays were performed with approximately 1-2 µg of purified proteins in a final volume of 50 µl of reaction buffer containing 20 nM prepared radiolabeled c-di-GMP ( $0.1 \ \mu$ Ci  $\mu$ l<sup>-1</sup>) with or without 100  $\mu$ M GTP. One unit of snake venom phosphodiesterase (Phosphodiesterase I, Worthington) suspended in SVPD conservation buffer (100 mM Tris-HCl pH 8.0, 100 mM NaCl, 14 mM MgCl<sub>2</sub>, 50% glycerol) was used as a positive control in PDEA assays. Proteins denatured at 99°C for 15 min were used as negative controls in both DGC and PDEA assays. Reaction products were analyzed by TLC as described before [32]. Briefly aliquots (2–4  $\mu$ l) were spotted on polyethyleneimine-cellulose TLC plates (Sigma) previously washed in 0.5 M LiCl and air dried. Plates were then soaked for 5 min in methanol, dried, and developed in 2:3 (v/v) saturated (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>/ 1.5 M KH<sub>2</sub>PO<sub>4</sub> (pH 3.5). Plates were allowed to dry prior to exposition to a phosphor imaging screen (Molecular Dynamics). Data were collected and analyzed using a FX molecular imager and the Quantity One software (Bio-Rad).

### Supporting Information

**Figure S1** Domain composition and organization of the 3 putative c-di-GMP-signalling proteins encoded by *C. difficile* strains other than strain 630. Domain composition and organization of the proteins are depicted essentially as in Figure 1. The HD-GYP domain is colored in orange.



**Figure S2** Analysis of intracellular c-di-GMP by HPLC. 10  $\mu$ M synthetic c-di-GMP (A), nucleotide extracts from *V. cholerae* N16961 expressing *lacZ* from pBAD-TOPO/lacZ/V5-His (negative control) (B), the DGC N-448 from pCD0522-(N-448) (C), or the DGC CD1420 from pCD1420 (D). Spectra of synthetic c-di-GMP (E), and of peaks with corresponding retention times in the nucleotide extracts from *V. cholerae* N16961 expressing the DGC N-448 (F) or the DGC CD1420 (G). 100 mg of cells were used for all nucleotide extractions.

(TIF)

**Figure S3** Phylogenetic tree of selected Clostridiaceae species. The tree was generated by alignment of the *rpoB* sequences using the neighbor-joining method. The support for each branch is indicated by the value at each node (in percent) as determined by 1,000 bootstrap samples. Only values  $\geq$ 70% are shown. *Staphylococcus aureus* COL and *Bacillus subtilis* 168 were taken as outgroup strains. Colored boxes highlight the *Clostridium difficile* strains (cyan), the two Clostridiaceae strains with the most c-di-GMP regulatory proteins (yellow) and the two Clostridiaceae strains with the least c-di-GMP regulatory proteins (magenta). Total c-di-GMP regulatory proteins per species are shown in parenthesis next to colored boxes.

(TIF)

**Table S1** Predicted and observed activities of the 31 mostconserved putative c-di-GMP turn-over proteins.(DOC)

**Table S2** Accession numbers of genomes of *C. difficile* strains used in this study.

 (DOC)

**Table S3**Clostridiaceae-encoded proteins containing GGDEF,EAL, or HD-GYP domains.(DOC)

 Table S4
 Bacterial strains and plasmids used in this study.

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 Table S5
 DNA sequences of the oligonucleotides used in this study.

 (DOC)

**Text S1** Supplemental Materials and Methods. (DOC)

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## **Author Contributions**

Conceived and designed the experiments: EB VB. Performed the experiments: EB VB. Analyzed the data: EB VB. Contributed reagents/ materials/analysis tools: VB L-CF FM. Wrote the paper: EB L-CF FM VB.

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