

1	
2	
3	A Novel Capture Compound for the Identification and Analysis of Cyclic di-GMP
4	Binding Proteins
5	
6	Jutta Nesper ¹⁾ , Alberto Reinders ¹⁾ , Timo Glatter ^{*1)} , Alexander Schmidt [*] , and Urs Jenal
7	
8	
9	
10	
11	
12	Affiliations:
13	Biozentrum of the University of Basel, Klingelbergstrasse 50, CH-4054 Basel,
14	Switzerland
15	*Proteomics Core Facility, Biozentrum of the University of Basel, Klingelbergstrasse
16	50, CH-4054 Basel, Switzerland
17	
18	
19	1) These authors contributed equally to this work
20	
21	
22	
23	For correspondence: <u>urs.jenal@unibas.ch</u>
24	
25	
26	
27	Keywords: Caulobacter crescentus; Pseudomonas aeruginosa; Salmonella enterica
28	serovar typhimurium; c-di-GMP effector; EAL GGDEF; Capture Compound mass
29	spectrometry
30	
31	

1 Abstract

2 The second messenger cyclic di-GMP is a near-ubiquitous signaling molecule that 3 globally alters bacterial cell physiology to promote biofilm formation and community 4 behavior. Much progress was made in recent years towards the identification and 5 characterization of diguanylate cyclases and phosphodiersterases, enzymes involved 6 in the synthesis and degradation of this signaling compound. In contrast, our 7 knowledge of the nature and mechanistic details of c-di-GMP effector proteins lags 8 behind, primarily because effective tools for their specific enrichment and rapid 9 analysis are missing. In this report we demonstrate that a novel tri-functional c-di-10 GMP-specific Capture Compound (cdG-CC) can be effectively used to identify and validate c-di-GMP binding proteins. The cdG-CC was able to specifically and 11 12 efficiently pull-down bona fide c-di-GMP effector proteins. Furthermore, in 13 combination with mass spectrometry (CCMS), this technology robustly identified a 14 substantial fraction of the known c-di-GMP signaling components directly from cell 15 extracts of different model organisms. Finally, we applied the CCMS technique to 16 profile c-di-GMP binding proteins of Salmonella enterica serovar typhimurium. Our 17 studies establish CCMS as a powerful and versatile tool to identify and analyze 18 components of the cellular c-di-GMP pathway in a wide range of different organisms. 19

1 Cyclic di-GMP is a ubiquitous second messenger regulating growth and behavior of a 2 wide range of gram-positive and gram-negative bacteria. In particular, c-di-GMP 3 mediates the switch between single cell behavior and a community life style called 4 biofilm, which is often associated with chronic infections of bacterial pathogens [1]. 5 Major components of the regulatory network are the GGDEF and EAL domains that 6 are widespread in bacteria and catalyze c-di-GMP synthesis and degradation, respectively [2]. The list of cellular processes controlled by c-di-GMP is ever-7 8 increasing and includes the biosynthesis and secretion of surface adhesins and 9 exopolysacharide (EPS) matrix components, different forms of cellular motility, long-10 term survival and response to environmental stress, synthesis of secondary 11 metabolites, regulated proteolysis and cell cycle progression, delivery of anti-12 bacterial toxins, intracellular growth and the production of virulence factors in a 13 range of animal and plant pathogens [2-4]. Despite of this wide variety of cellular 14 functions that are modulated by c-di-GMP, the list of effector proteins has remained 15 relatively sparse [2-4]. These include PilZ, small switch-like domains that undergo 16 conformational change upon binding c-di-GMP. In addition, several members of the 17 CRP/FNR and response regulator superfamilies of transcription regulators were 18 shown to specifically bind c-di-GMP. Finally, a subgroup of GGDEF and EAL domains 19 was recognized as c-di-GMP effector proteins adopting their novel functionality 20 through the combined loss of catalytic activity and exploitation of their allosteric and 21 active site binding pockets, respectively. While most known effector proteins were 22 discovered through an "educated guess" approach that was based on their 23 functional linkage to c-di-GMP mediated cellular processes, unbiased screening for 24 novel effectors was hampered primarily by the lack of reliable and effective 25 biochemical tools for their enrichment and isolation. Only one global recent study 26 used a chemical proteomics approach to identify c-di-GMP binding proteins in P. 27 aeruginosa [5]. Here we introduce a novel tri-functional capture molecule (cdG-CC) 28 as an effective tool to identify specific c-di-GMP binding proteins directly from a 29 complex mixture of macromolecules. The compound is based on a chemical scaffold 30 harboring specificity, reactivity, and sorting properties (Fig. 1A) [6, 7].

To evaluate the potential of this compound for the specific enrichment and isolation
 of c-di-GMP binding proteins we first tested if the cdG-CC is able to pull down known

1 c-di-GMP effector molecules directly from a solution with purified protein. We used 2 the well-studied PilZ domain protein DgrA from C. crescentus [8]. The binding affinity 3 of wild-type DgrA for c-di-GMP is in the low nanomolar range. In contrast, the 4 binding mutant DgrAW75A has a Kd of 6.2 µM and a DgrA mutant lacking 5 coordinating Arg residues in the N-terminus (RR11AA) fails to bind c-di-GMP 6 completely [8]. 0.5 µM purified wild-type and mutant hexahistidine-tagged proteins 7 (Fig. 1B) were incubated with 10 μ M cdG-CC, UV cross-linked for 4 min and pulled 8 out with streptavidin coated magnetic beads. While DgrA was readily captured in the 9 absence of any other ligand, the addition of a 100x excess of c-di-GMP abolished 10 cdG-CC almost entirely (Fig. 1B). In contrast, an equal molar excess of GTP did not 11 interfere with the cdG-CC pull down (Fig. 1B) arguing that cdG-CC binding to DgrA is 12 highly specific. In agreement with this, the cdG-CC only inefficiently pulled-down the 13 DgrAW75A mutant form and completely failed to bind the DgrARR11AA mutant (Fig. 14 1B). These data are fully consistent with the binding affinities of the different DgrA 15 forms for c-di-GMP and strongly argue that cdG-CC is able to specifically capture 16 DgrA. Moreover, these experiments demonstrate that the cdG-CC is a valuable 17 diagnostic tool to verify candidate c-di-GMP binding proteins.

18 These results encouraged us to probe if the capture compound can be applied for 19 the selective enrichment of c-di-GMP binding proteins from a more complex mixture 20 of proteins. To test this, we captured soluble c-di-GMP binding proteins from cell 21 extracts and probed immunoblots with PopA specific antibodies. PopA is a GGDEF 22 effector protein that regulates cell cycle progression in *C. crescentus* in response to a 23 cellular upshift of c-di-GMP during the G1-S phase transition [9, 10]. To bind c-di-24 GMP PopA utilizes a conserved and well-defined binding pocket, which, in related 25 catalytic GGDEF domains, is used as an allosteric I-site for product inhibition of the 26 diguanylate cyclase (DGC) activity [11, 12]. As shown in Fig. 1C, PopA with a known 27 Kd for c-di-GMP of 2.5 μM, was readily captured from C. crescentus cell lysates using 28 3 μ M cdG-CC and a total of 400 μ g soluble protein. Pull down of PopA was inhibited 29 in the presence of a large excess of c-di-GMP (1mM), while GTP did not interfere 30 with cdG-CC binding (Fig. 1C), arguing that the cdG-CC interaction with PopA is highly 31 specific. Likewise, no PopA was bound to the cdG-CC when using a strain expressing 32 a PopA mutant that lacks the highly conserved Arg residue of the canonical RxxD I-

1 site binding motif (R357G) [9] (Fig. 1C). This indicated that the cdG-CC can enrich c-2 di-GMP binding proteins directly from whole cell extracts in a highly specific manner 3 and that this compound is suited for a global isolation procedure of c-di-GMP binding 4 proteins. Therefore we combined capture experiments with the analysis of isolated 5 proteins by LC-MS/MS (CCMS, [6, 7]). When applying 10, 5 or 2.5 μ M cdG-CC with 6 400 µg of soluble *C. crescentus* proteins, nine of eleven proteins predicted to contain 7 either a PilZ, GGDEF or EAL domain (Table 1A and Supplementary Table S1A) were significantly enriched as compared to the competition control based on the spectral 8 9 counts of the identified peptides. In addition to the analysis of soluble proteins we 10 also aimed at evaluating the efficiency of CCMS for the enrichment of c-di-GMP 11 binding proteins from membrane fractions. Although the numbers of spectral counts 12 were lower as compared to proteins from the soluble fraction, three of the five 13 integral membrane proteins predicted to bind c-di-GMP were identified when 400 14 μ g DDM solubilized membrane proteins and 10 μ M cdG-CC were used for CCMS (Table 1A and Supplementary Table S1A). Only four of the known components of the 15 16 C. crescentus c-di-GMP network were not identified by CCMS. Two of these are 17 integral membrane proteins with several predicted membrane spanning domains in 18 their N-terminal regions (CC0740, CC0896). It is possible that they were not 19 solubilzed by the detergent used or not detected by LC-MS/MS. Another possibility is 20 that they are not expressed, as it might be the case for CC3094 and CC3148.

21 To expand these studies to a different organism, we also tried to isolate known c-di-22 GMP binding proteins from P. aeruginosa PA01. With over 40 GGDEF and EAL 23 domain and eight PilZ domain proteins annotated, the complexity of the c-di-GMP 24 signaling network in this organism is much higher than in *C. crescentus*. Using 10 μ M, 25 7.5 µM, 5 µM, 2.5 µM and 1.25 µM cdG-CC and 350 µg of soluble protein extract, 26 several of these proteins were unambiguously identified by CCMS (Table 1B and 27 Supplementary Table S1B). This includes four PilZ domain proteins, four GGDEF and 28 two composite GGDEF-EAL domain proteins. The fraction of the P. aeruginosa 29 proteins predicted to bind c-di-GMP that were isolated by CCMS is substantially 30 lower (10 of 28 predicted soluble proteins) as compared to C. crescentus but is 31 comparable to a recent study using sepharose-coupled c-di-GMP [5]. The

1 discrepancy between C. crescentus and P. aeruginosa might be related to the fact 2 that many of these proteins are engaged in the transition between a motile, 3 planktonic and a sessile, surface attached life-style. Since C. crescentus has 4 integrated this developmental transition into its reproductive cycle [13], most of the 5 components of the c-di-GMP network orchestrating this switch, need to be 6 expressed in cells growing in liquid cultures [10]. In contrast, *P. aeruginosa* adapts its 7 expression program upon surface colonization in response to environmental cues 8 [14]. It is thus possible that several components of the *P. aeruginosa* core machinery 9 required to adapt to surface communities are simply not expressed in logarithmically 10 growing cultures in liquid media. To test this, it would be interesting to repeat theses 11 experiments with cell extracts harvested from surface grown P. aeruginosa 12 communities.

13 To evaluate whether cdG-CC based CCMS can be used to profile an entire cellular 14 network of c-di-GMP binding proteins we used S. typhimurium. This organism was 15 chosen to perform an unbiased CCMS experiment because its c-di-GMP network 16 appears to be of lower complexity as compared to other bacterial species [15]. In 17 contrast to the CCMS experiments with C. crescentus and P. aeruginosa, in which 18 spectral counts were extracted for known c-di-GMP binders, we tested whether 19 CCMS is capable to enrich for c-di-GMP binding proteins upon MS1 label-free 20 quantification. CCMS experiments were performed in triplicates with 350 µg soluble 21 whole cell proteins using 7.5 μ M cdG-CC and competition controls with an excess of 22 c-di-GMP (1 mM) (Supplementary Table S2A). Following mass spectrometry analysis 23 and label-free quantification (Supplementary Table S2B) significant differences in 24 protein enrichment between cdG-CC experiments and control samples with 25 competing c-di-GMP were visualized in a volcanoplot. The graph shows a significant 26 enrichment (>2 fold) of 36 proteins as compared to the control with a q-value <0.05 27 (Supplementary Table S2C and Fig. 2). Among the enriched proteins is the PilZ 28 domain protein YcgR (Fig. 2) [16]. Many of the identified components that were not 29 previously associated with the c-di-GMP network were metabolic proteins and 30 proteins involved in fatty acid and LPS biosynthesis (Fig. 2). Such proteins are of 31 great interest in the light of switching between a virulent planktonic and a surface 32 attached persistent lifestyle. However, since none of these proteins show homology

to known c-di-GMP binding proteins, they first need to be validated to specifically
bind c-di-GMP and be part of the c-di-GMP signaling network. In addition also some
background proteins are enriched. They are mostly binders of other nucleotides or
proteins that are known to be highly abundant, such as five tRNA-related proteins or
the chaperone GroEL (Supplementary Table S2C).

1 Acknowledgements

We thank Samuel Steiner for help in optimization of Capture Compound binding
assays. This work was supported by the Swiss National Science Foundation (SNF)
Sinergia grant CRSII3_127433.

5

1 **References:**

- 2 [1] Furukawa S, Kuchma SL, O'Toole GA. Keeping their options open: Acute versus
- 3 persistent infections. J Bacteriol. 2006;188:1211-7.
- 4 [2] Schirmer T, Jenal U. Structural and mechanistic determinants of c-di-GMP
 5 signalling. Nat Rev Microbiol. 2009;7:724-35.
- 6 [3] Hengge R. Cyclic-di-GMP reaches out into the bacterial RNA world. Sci Signal.7 2010;3:pe44.
- 8 [4] Sondermann H, Shikuma NJ, Yildiz FH. You've come a long way: C-di-GMP
 9 signaling. Curr Opin Microbiol. 2012.
- 10 [5] Duvel J, Bertinetti D, Moller S, Schwede F, Morr M, Wissing J, et al. A chemical
- proteomics approach to identify c-di-GMP binding proteins in *Pseudomonas aeruginosa*. J Microbiol Methods. 2012;88:229-36.
- [6] Koster H, Little DP, Luan P, Muller R, Siddiqi SM, Marappan S, et al. Capture
 compound mass spectrometry: A technology for the investigation of small molecule
 protein interactions. Assay Drug Dev Technol. 2007;5:381-90.
- [7] Lenz T, Fischer JJ, Dreger M. Probing small molecule-protein interactions: A new
 perspective for functional proteomics. J Proteomics. 2011.
- [8] Christen M, Christen B, Allan MG, Folcher M, Jeno P, Grzesiek S, et al. DgrA is a
 member of a new family of cyclic diguanosine monophosphate receptors and
 controls flagellar motor function in *Caulobacter crescentus*. Proc Natl Acad Sci U S A.
 2007;104:4112-7.
- [9] Duerig A, Abel S, Folcher M, Nicollier M, Schwede T, Amiot N, et al. Second
 messenger-mediated spatiotemporal control of protein degradation regulates
 bacterial cell cycle progression. Genes Dev. 2009;23:93-104.
- [10] Abel S, Chien P, Wassmann P, Schirmer T, Kaever V, Laub MT, et al. Regulatory
 cohesion of cell cycle and cell differentiation through interlinked phosphorylation
 and second messenger networks. Mol Cell. 2011;43:550-60.
- [11] Christen B, Christen M, Paul R, Schmid F, Folcher M, Jenoe P, et al. Allosteric
 control of cyclic di-GMP signaling. J Biol Chem. 2006;281:32015-24.
- [12] Wassmann P, Chan C, Paul R, Beck A, Heerklotz H, Jenal U, et al. Structure of
 BeF3- -modified response regulator PleD: Implications for diguanylate cyclase
 activation, catalysis, and feedback inhibition. Structure. 2007;15:915-27.

1 [13] Curtis PD, Brun YV. Getting in the Loop: Regulation of development in 2 Caulobacter crescentus. Microbiology and Molecular Biology Reviews. 2010;74:13-41. 3 [14] Kuchma SL, O'Toole GA. Surface-induced and biofilm-induced changes in gene 4 expression. Curr Opin Biotechnol. 2000;11:429-33. 5 [15] Solano C, Garcia B, Latasa C, Toledo-Arana A, Zorraquino V, Valle J, et al. Genetic 6 reductionist approach for dissecting individual roles of GGDEF proteins within the c-7 di-GMP signaling network in Salmonella. Proc Natl Acad Sci U S A. 2009;106:7997-8 8002. [16] Ryjenkov DA, Simm R, Romling U, Gomelsky M. The PilZ domain is a receptor for 9 10 the second messenger c-di-GMP: The PilZ domain protein YcgR controls motility in 11 enterobacteria. J Biol Chem. 2006;281:30310-4.

1 Figure legends:

2 Fig. 1 - The cdG-CC can specifically pull down bona fide c-di-GMP binding proteins 3 from purified and from crude cell extracts. A) Chemical structure of cdG-CC. C-di-4 GMP provides the selectivity for capturing proteins. The photo-reactivity group 5 allows irreversible nitren formation between the compound and the captured 6 protein. Biotin as sorting function allows the isolation of the compound by binding to 7 streptavidin coated magnetic beads. B) Immunoblot of purified and captured DgrA 8 (wt), DgrA W75A and DgrA RR11AA. In all capture experiments 10 µM cdG-CC was 9 present, in competition experiments the proteins were preincubated with a 100 x 10 excess of c-di-GMP (cdG) or GTP. All proteins were His-tagged and detected using 11 anti-His antibodies. C) Immunoblot of PopA. PopA (wt) or the PopA I-site mutant 12 R357G was expressed in NA1000∆popA (lanes marked with 'proteins'; note that this 13 I-site mutant is less abundant in the cell). PopA was captured in the presence of 3 14 μM cdG-CC. Addition of 1 mM c-di-GMP but not 1 mM GTP prevented the binding to 15 the cdG-CC. In contrast the PopA I-site mutant could not be captured at all. PopA 16 was detected using anti-PopA antibodies.

17

18 Fig. 2 - Volcanoplot based visualization of proteins significantly enriched by CCMS of 19 S. typhimurium. Following capturing, LC-MS/MS analysis and label-free quantification, 20 log2-intensity ratio of all detected peptide features between capturing and 21 competition experiment were calculated and plotted versus values derived from 22 significance analysis. Proteins within the significance thresholds for q-values < 0.05 23 and intensity ratios >2-fold are indicated in a box. Experiments in triplicate were 24 performed in the presence of 7.5 μ M cdG-CC and with 1 mM c-di-GMP added to the 25 competition reactions.

- 1 Table 1 Identified known c-di-GMP binding proteins.
- 2 A) C. cresentus

Protein Name	ID	Domain architecture	CCMS experiment/CCMS competition ¹⁾			
			No spectral counts of identified peptides			
Soluble fraction:						
Experiment ²⁾			1	2	3	
РорА	CCNA_01918	GGDEF	20/1	25/3	15/5	
PleD	CCNA_02546	Rec-Rec-GGDEF	18/0	21/2	13/5	
DgcA	CCNA_03394	GGDEF	5/0	3/0	1/0	
DgcB	CCNA_01926	GGDEF	12/1	16/3	10/2	
CC0655	CCNA_00692	PAC-GGDEF-EAL	8/0	18/0	19/0	
CC1086	CCNA_01140	PAS-EAL	8/0	5/0	4/0	
PdeA	CCNA_03507	GGDEF-EAL	3/0	5/0	6/0	
DgrA	CCNA_01671	PilZ	5/0	6/3	6/1	
DgrB	CCNA_03268	PilZ		2/0		

Membrane fraction:

Experiment³⁾

TipF	CCNA_00747	EAL	3/0	23/13	
PdeB	CCNA_00089	3x(MHYT)-PAS-GGDEF-EAL	3/0	6/0	
CC0857	CCNA_00900	CHASE4-GGDEF-EAL		2/0	

3

4 1) All competition experiments were performed in the presence of 1 mM c-di-GMP.

5 2) 3 independent experiments are indicated using 10, 5 or 2.5 μ M cdG-CC

6 respectively.

7 3) 2 independent experiments are indicated using 10 μ M cdG-CC.

1 B) Soluble proteins of *P. aeruginosa*

Protein Name	ID	Domain architecture	CCMS experiment/CCMS competition ¹⁾ No of spectral counts of identified peptides				
Experiment No ⁴⁾			1	2	3	4	5
FimX	PA4959	PAS-GDSIF-EVL	9/0	10/0	9/0	8/0	7/0
	PA3353	PilZ	5/0	6/0	6/0	10/0	7/0
WspR	PA3702	Rec-GGEEF	2/0	2/0	2/0		
	PA0012	PilZ			2/0	1/0	
	PA0169	GGEEF	2/0	2/0	2/0		
	PA4843	Rec-Rec-GGEEF		2/0	2/0	5/0	3/0
	PA2989	PilZ	1/0	1/0	1/0	1/0	1/0
	PA0290	PAS-Rec-GGEEF	1/0			2/0	2/0
	PA2567	GAF-SPTRF-EAL		1/0	1/0		1/0
	PA4608	PilZ	1/0	2/0	2/0	2/0	1/0

2 3

4) Eperiment 1 was performed with 10 μM cdG-CC, experiment 2 with 7.5 μM cdG-CC,

4 experiment 3 with 2.5 μ M cdG-CC, experiment 4 with 2.5 μ M cdG-CC and experiment 5 with 1.25 μ M cdG-CC.



Fig. 2 Click here to download high resolution image

Figure 2:



Materials & Methods Click here to download Supplementary material: SupplementaryMaMe.docx Table S1Click here to download Supplementary material: SupplementaryTable S1.xlsx

Table S2Click here to download Supplementary material: SuplementraryTable S2.xlsx