Mutation at Different Sites in the *Nostoc punctiforme* cyaC Gene, Encoding the Multiple-Domain Enzyme Adenylate Cyclase, Results in Different Levels of Infection of the Host Plant *Blasia pusilla*<sup>†</sup>

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The filamentous cyanobacterium *Nostoc punctiforme* forms symbioses with plants. Disruption of the catalytic domain of the *N. punctiforme* adenylate cyclase (CyaC) significantly increased symbiotic competence, whereas reduced infectivity was observed in a mutant with a disruption close to the N terminus of CyaC. The total cellular cyclic AMP levels were significantly reduced in both mutants.

The filamentous cyanobacterium *Nostoc punctiforme* is highly responsive to environmental changes and is able to regulate its cell type accordingly (29). The best-studied response is the patterned development of specialized dinitrogen-fixing cells, known as heterocysts (1, 43, 45). This cyanobacterium is also capable of differentiating motile filaments known as hormogonia (10, 30, 34), which act as the infective agents in symbiotic associations with plants. Potential host plants produce a hormogonium-inducing factor (3, 10, 13, 27, 34) and induce chemoattractive behavior in hormogonia (27, 31, 41).

Our knowledge of the molecular events involved in the differentiation of hormogonia and the factors affecting their behavior in response to plants is limited. In response to hormogonium-inducing factor, *N. punctiforme* mutants with disruptions in the *hrmU* and *hrmA* genes convert 80% of the filaments to hormogonia, compared with 30% of the filaments in the wild type (14). This increased capacity for hormogonium formation is associated with increased symbiotic competence in the host plant *Anthoceros punctatus*, a hornwort. Expression of *hrmU* and *hrmA* is induced by an aqueous extract of *Anthoceros* tissue, leading to the suggestion that a factor(s) in the extract prevents hormogonium formation, at least in part by its up-regulation of *hrmU* (13). Mutations in *sigH* and *tpRN* (encoding an alternative group 2 sigma factor and a tetratricopeptide repeat protein, respectively) are also associated with increased symbiotic competence, although in these cases this is not associated with increased hormogonium differentiation, suggesting that the response of the hormogonia to plant-derived signals has been modified (9, 28). Hormogonia are clearly important in the establishment of symbiotic associations, but their formation alone does not guarantee infection. Campbell and Meeks (10) observed that a strain of *Nostoc* failed repeatedly to infect *A. punctatus* despite converting more than 90% of its vegetative filaments to hormogonia. Similarly, Johansson and Bergman (20) noted several noninfective cyanobacteria that, despite being able to differentiate motile hormogonia and being found in abundance on the stems and glands of the angiosperm *Gunnera*, failed to establish symbioses (16, 34). Campbell et al. (11) showed that 1,827 genes are differentially transcribed in hormogonia of *Nostoc* (estimated 24 h after their induction), a number almost five times higher than the number in akinete-forming or N<sub>2</sub>-fixing cultures. The majority of the 944 upregulated genes were found to be involved in signal transduction and transcriptional activation, and this indicates that the hormogonium is a rather complex, metabolically active motile filament that is highly adapted to sense and respond to its environment.

Transposon mutant H1, originally identified in our laboratory during a screen of nitrogen starvation-responsive *N. punctiforme* mutants (21), exhibits a prolonged hormogonial phase in the presence of the liverwort *Blasia pusilla*, another host plant, and increased symbiotic competence (which is 2.5-fold higher than that of the wild type). Analysis revealed that the transposon had inserted within the putative catalytic domain of *cyaC* encoding an adenylate cyclase (AC) (NP01896 gene [http://genome.jgi-psf.org/finished_microbes/nospu/nospu_annotation.html]). ACs catalyze the formation of the intracellular messenger cyclic AMP (cAMP) (8, 12). Here we report reconstruction of the H1 *cyaC* mutant phenotype. In addition, we constructed a *cyaC* mutant in which the N terminus of CyaC was inactivated, which, by contrast, exhibited significantly reduced symbiotic competence. Total cAMP levels were dramatically reduced in both *cyaC* mutants, implying that reduced levels of cellular cAMP alone cannot explain the different levels of symbiotic competence observed in the two mutants.

Construction of *cyaC* mutants and phenotypic characterization of these mutants. Attempts to reconstruct the transposon mutant H1 by introducing the recovered transposon and flanking *Nostoc* DNA into wild-type *N. punctiforme* using the strategy of Black and Wolk (7) were unsuccessful; therefore, an omega neomycin phosphotransferase gene (Ω-<i>npt</i>) from pSCR9 (13) was ligated to the 3′ region of *cyaC* (nucleotide

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position 3212, which is close to position 3317, the site of the transposon interruption in H1) (Fig. 1) to generate mutant C3212. The $\Omega$-npt cassette was also introduced into the 5' end of cyaC (nucleotide position 1068), generating mutant C1068 (Fig. 1). DNA sequences containing the cyaC gene and flanking DNA were PCR amplified from N. punctiforme genomic DNA, and constructs were introduced into wild-type N. punctiforme as described previously (14). The carboxy-terminal and most conserved region in cyanobacterial ACs consists of a catalytic domain (Fig. 1). The N-terminal region of ACs is characteristically variable, and the differences in this region are believed to account for the differences in the regulatory properties of the enzyme activity (24; see reference 12 for a review of the domain structure of cyanobacterial ACs).

Upstream of the N. punctiforme CyaC catalytic domain are several distinct regulatory regions, which are also present in the CyaC proteins of the cyanobacterium Spirulina platensis (23) and Anabaena sp. strain PCC 7120 (24) and resemble the sensory kinase and response regulator proteins that consist the bacterial two-component signal transduction systems (37, 47). The regulatory domains are arranged sequentially from the N terminus and comprise a response regulator-like domain, two GAF domains (GAF is an acronym derived from the proteins of initial identification, mammalian cGMP-phosphodiesterases, Anabaena adenylate cyclases, and Escherichia coli transcription factor FhlA [formate hydrogen lyase transcriptional activator] [2]), a histidine kinase-like domain, and a second response regulator-like domain (Fig. 1). The highly conserved histidine residue of histidine kinases and the aspartate residue characteristic of response regulator proteins are present in the Nostoc CyaC protein (His-549, Asp-59 and Asp-869) (data not shown) and align with the phosphorylation sites (His-524, Asp-59, and Asp-847) previously reported for the Anabaena sp. strain PCC 7120 CyaC protein (24). In two-component signaling systems the conserved histidine residue of the histidine kinase is able to transfer a phosphoryl group to conserved aspartate residues in the response regulator domains. It has been shown that autophosphorylation of His-572 in the transmitter domain and phosphotransfer to Asp-895 in the receiver (R2) domain of CyaC of S. platensis regulate the activity of AC. Autophosphorylation of CyaC is probably regulated by a specific signal transferred from a primary signal sensor (22).

Mutants C1068 and C3212 grew at rates comparable to those of the wild type in BG11 media (35) both with and without a source of combined nitrogen. Also, no obvious differences were observed in the frequency of proheterocysts and heterocysts expressed by mutant and wild-type cultures 24 h after

FIG. 1. Schematic illustration of cyaC in the wild type (WT) and cyaC mutants C1068, C3212, and H1. The positions and orientations of npt (neomycin phosphotransferase gene) and the transposon (Tn5-1063) are indicated by triangles. Domain abbreviations: RR, response regulator-like domain; GAF, GAF-like domain; HK, histidine kinase-like domain; AC, adenylate cyclase-like domain. The figure is not drawn to scale. GAF domains are part of a large superfamily of proteins that bind diverse ligands and regulate the biochemical output of proteins in which they are found (2, 4, 38, 40). GAF domains appear to represent sensor domains which receive a signal that is subsequently transmitted to the catalytic domain via the response regulator domain.
The frequency of infected symbiotic cavities (auricles) was expressed as a percentage of the total number of auricles examined for each determination. At least eight cultures were examined for each strain, and at least 400 auricles were examined for each determination. The error bars indicate the standard errors of the means.

Coculture experiments with the symbiotic partner was as described by Wong and Meeks (44). The mutant strains formed hormogonia approximately 12 h earlier than the wild-type strain (data not shown), and the maximum mean frequencies of hormogonia observed between 18 and 60 h of coculture were 70, 92, and 95% for the wild type, C1068, and C3212, respectively. The numbers of symbiotic colonies detected in the auricles (dome-shaped symbiotic cavities located on the ventral surface of the Blasia tissue) were estimated over a 28-day coculture period (Fig. 2). Wild-type cultures exhibited a steady increase in the frequency of infection from approximately 15% after 7 days to a maximum mean value of approximately 26% (observed after both 21 and 28 days of coculture). Initially, mutant C3212 infected the Blasia symbiotic tissues slowly, and the percentages of symbiotically associated colonies were just 4.16 and 19.5% after 7 and 14 days of coculture, respectively. However, after 21 and 28 days of coculture the mean percentage of infected colonies had increased dramatically to 40.36 and 67.12%, respectively (Fig. 2).

FIG. 2. Mean frequencies of infection of B. pusilla tissue following coculture with N. punctiforme wild type (WT) and cyaC mutants C1068 and C3212. The frequency of infected symbiotic cavities (auricles) was estimated after 7, 14, 21, and 28 days of coculture with a host plant, the liverwort Blasia. The values are the mean number of infected auricles expressed as a percentage of the total number of auricles examined for each determination. At least eight cultures were examined for each strain, and at least 400 auricles were examined for each determination. The error bars indicate the standard errors of the means.

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Analysis of cellular cAMP levels. Since transfer of certain cyanobacteria from light to dark conditions results in an increase in cAMP levels (24, 32, 33, 36), we estimated cAMP levels at mid-exponential phase (optical density at 720 nm, 0.15 to 0.25) in the light and following transfer of the cells to dark conditions. Samples were collected after 1, 5, and 10 min of incubation in the dark (Fig. 3). The samples used for determination of cellular cAMP concentrations were prepared by the method of Terauchi and Ohmori (39), and the amount of cAMP in each lyophilized sample was determined using the cAMP Biotrak enzyme immunoassay system (Amersham Biosciences) according to the manufacturer’s instructions. When wild-type cells were incubated in the dark, the cAMP levels rapidly increased approximately 1.4-fold in 1 min and then decreased within 5 min (Fig. 3). This is in agreement with previous observations and is consistent with the role of cAMP as a signaling molecule (25, 33, 36). By contrast, the cyaC mutants did not show any corresponding increase in cellular cAMP levels following transfer from light to dark growth conditions (Fig. 3). The total cAMP levels were three- to fourfold higher in the wild type than in any of the three cyaC mutants, suggesting that cyaC function in relation to cAMP production was effectively disrupted in all three mutants (since cAMP levels in the H1 mutant were not determined in the study of Joseph [21], they are included here). The results support previous proposals that cyaC is responsible for maintaining steady-state cAMP levels within certain cyanobacteria (24) and imply that CyaC is probably the major AC in N. punctiforme. Anabaena sp. strain PCC 7120 has at least six different AC genes (cyaA, cyaB1, cyaB2, cyaC, cyaD, and cyaE) (24; references 12 and references therein), and searches of the N. puncti-
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N. punctiforme genome suggests that this organism also contains additional putative AC genes (including homologues of cyaA, cyaB1, cyaB2, cyaC, and cyaD), which may account for the small amounts of cAMP detected in all three of the mutant strains (Fig. 3). To our knowledge, this is the first account of the determination of cAMP levels in N. punctiforme. The mean concentration of cAMP (20 pmol cAMP per mg chlorophyll a) was estimated in wild-type cultures incubated in the light is almost fourfold higher than the concentration detected in Anabaena sp. strain PCC 7120 cells grown under similar conditions (25). By contrast, two other filamentous cyanobacteria, *S. platensis* and *Anabaena cylindrica*, grown in the light had more than-twofold-higher intracellular concentrations of cAMP, 51 and 48 pmol cAMP per mg chlorophyll a, respectively (36). However, the values increased to 179 and 189 pmol cAMP per mg chlorophyll a in *A. cylindrica* and *S. platensis*, respectively, after incubation in the dark, leading to the suggestion that light suppresses cAMP synthesis or perhaps stimulates the degradation of cAMP in cyanobacteria (36).

Mutant H1, containing the transposon Tn5 derivative Tn5-1063 (42) carrying the luxAB genes encoding the light-emitting protein complex luciferase, showed a higher level of luminescence under nitrogen-depleted conditions, and the luminescence was significantly higher in proheterocytes (21). Single filaments undergoing hormogonium formation under nitrogen-depleted conditions did not show any increase in luminescence (21).

Sequence analysis revealed that the transposon had inserted with luxAB oriented parallel to cyaC, implying that the increase in luminescence in response to nitrogen deprivation was driven by the cyaC promoter. Hood et al. (19) showed that nitrogen starvation increases intracellular cAMP levels fourfold in the filamentous cyanobacterium *Anabaena variabilis*. In *Anabaena flos-aquae*, transfer to nitrogen-depleted media causes an increase in cAMP levels (17). However, to a lesser degree, transfer to nitrogen-replete media also causes an increase in cAMP levels in this organism, raising doubts that nitrogen deprivation per se is a stimulus for cellular cAMP production (17). Similarly, mutational analysis of AC genes in another filamentous cyanobacterium, *Anabaena sp.* strain PCC 7120, did not reveal any nitrogen-dependent growth phenotypes (24), and cellular cAMP in *Synechocystis* sp. strain PCC 6803 appears to be unaffected by nitrogen deprivation (18).

It is not clear why the two cyaC mutants described here have very different phenotypes in terms of symbiotic competence, although distinct cyanobacterial phenotypes generated by mutations at different sites within the same gene (for example, taxA/Y1, which encodes a chemotaxis-related protein [6], and slr1443 encoding a protein kinase homologue required for motility in *Synechocystis* sp. strain PCC 6803 [26]), are not unusual. *N. punctiforme* CyaC is a large protein (1,177 amino acids) with multiple domains (Fig. 1). Mutant C1068 is inactivated in the second GAF domain, whereas C3212 is inactivated in the C terminus catalytic domain. It is not inconceivable that the different domains participate in different cellular activities and that the differences in symbiotic competence observed here may be a reflection of the different domain functions and the environmental signals that they respond to. Indeed, there is evidence that each domain (including a GAF domain, a kinase domain, and a receiver domain) of the tobacco plant ethylene receptor, NTHK1, may have specific roles in regulation of plant growth, the salt stress response, and gene expression (46). Therefore, the function with regard to the impact on symbiotic competence (if any) of the other domains within *Nostoc CyaC* requires further investigation.

Interestingly, mutants C1068 and C3212 (and H1) had low cAMP levels, implying that cAMP per se is not involved in symbiotic competence. There is evidence which suggests that pili expressed on the hormogonium surface are important in the establishment of *Nostoc-Blasia* symbioses (15). However, close examination, using shadow casting and transmission electron microscopy, revealed that the pili expressed on the hormogonium surface of cyaC mutants H1, C3212, and C1068 were, in terms of abundance, distribution, and structure, indistinguishable from the pili expressed on the surface of hormogonia differentiated by the wild type (data not shown), indicating that the explanation of the difference in the observed symbiotic competence phenotypes lies elsewhere. Recently, cAMP has been shown to be critical for the phototactic response of *Synechocystis* sp. strain PCC 6803 that is characterized by the formation of fingerlike projections from colonies on agar plates (5). The different symbiotic competence phenotypes reported here may imply that there are differences in the behavior of the mutant hormogonia in response to plant signals.

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