Feathermoss and epiphytic *Nostoc* cooperate differently: expanding the spectrum of plant–cyanobacteria symbiosis

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Dinitrogen (N\(_2\))-fixation by cyanobacteria in symbiosis with feathermosses is the primary pathway of biological nitrogen (N) input into boreal forests. Despite its significance, little is known about the cyanobacterial gene repertoire and regulatory rewiring needed for the establishment and maintenance of the symbiosis. To determine gene acquisitions and regulatory changes allowing cyanobacteria to form and maintain this symbiosis, we compared genomically closely related symbiotic-competent and -incompetent *Nostoc* strains using a proteogenomics approach and an experimental set up allowing for controlled chemical and physical contact between partners. Thirty-two gene families were found only in the genomes of symbiotic strains, including some never before associated with cyanobacterial symbiosis. We identified conserved orthologs that were differentially expressed in symbiotic strains, including protein families involved in chemotaxis and motility, NO regulation, sulfate/phosphate transport, and glycosyl-modifying and oxidative stress-mediating exoenzymes. The physical moss–cyanobacteria epiphytic symbiosis is distinct from other cyanobacteria–plant symbioses, with *Nostoc* retaining motility, and lacking modulation of N\(_2\)-fixation, photosynthesis, GS-GOGAT cycle and heterocyst formation. The results expand our knowledge base of plant–cyanobacterial symbioses, provide a model of information and material exchange in this ecologically significant symbiosis, and suggest new currencies, namely nitric oxide and aliphatic sulfonates, may be involved in establishing and maintaining the cyanobacteria–feathermoss symbiosis.

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Introduction

Dinitrogen (N\(_2\))-fixation by filamentous heterocystous cyanobacteria living in symbiosis with pleurocarpous feathermosses (for example, *Pleurozium schreberi* and *Hylocomium splendens*) and acrocarpous mosses (for example, *Sphagnum fuscum*) serves as the primary input of nitrogen (N) into boreal forests and can account for 50% of total N input (DeLuca *et al.*, 2002, 2008; Turetsky *et al.*, 2012; Rousk *et al.*, 2015). The boreal forest ecosystem occupies 11% of the Earth’s terrestrial surface (Bonan and Shugart, 1989), where feathermoss often account for >65% of the forest floor (DeLuca *et al.*, 2008). This partnership has a vital role in primary productivity of the boreal forest (Turetsky, 2003; Lindo and Gonzalez, 2010). High-resolution secondary ion mass spectrometry verified transfer of fixed N from the cyanobacteria to the moss host, demonstrating that the symbiosis governs the main N entrance to boreal forests (Bay *et al.*, 2013). Despite the significance of the symbiosis, knowledge of the interaction between the two partners is rudimentary and, in particular, little is known about the gene repertoire and regulation needed to form the symbiosis. No studies have examined global gene or protein expression of the moss or the cyanobacterial partner.

Current knowledge on colonization steps and maintenance of cyanobacterial–plant symbioses is primarily based on studies of endophytes,
specifically the liverwort Blasia, and hornworts Anthoceros and Phaeoceros, where the cyanobacteria, mainly Nostoc sp., are extracellular but enclosed in a specialized cavity, and with the angiosperm Gunnera, where the cyanobacteria are located intracellularly (reviewed in Santi et al., 2013). During the establishment of these symbioses, there are two phases of interaction, an early phase, which includes chemical signalling between partners, followed by a later phase where the cyanobacteria are physically associated with the host.

In the early phase, the cyanobacteria differentiate from vegetative filaments into a transient motile stage termed hormogonia. The differentiation is induced by a variety of environmental stimuli and host-produced chemical signals of unknown structure, known as hormogonia-inducing factor, HIF (Rasmussen et al., 1994; Meeks and Elhai, 2002; Meeks, 2003; Adams and Duggan, 2008). The feathermoss secretes HIF and possibly other chemotactants to recruit and direct the hormogonia toward the host, according to the only study addressing feathermoss colonization (Bay et al., 2013). Cyanobacterial genes responding to plant HIF have been identified, and many appear to be involved in signal transduction and transcriptional activation (Campbell et al., 2007; Duggan et al., 2013; Risser and Meeks, 2013; Campbell et al., 2015). A recent study has identified che-family genes as important for both motility and symbiotic competence (Duggan et al., 2013), whereas genes involved in phototaxis, chemotaxis and motility (ptx, hmp, hps and pil genes) are thought to be implicated in the colonization process (Campbell et al., 2003, 2008; Risser and Meeks, 2013; Risser et al., 2014).

The initial signaling and motility stage is followed by a physical interaction phase between host and cyanobacterium. In the symbiosis with hosts such as Gunnera, Blasia and Anthoceros, the cyanobacteria exhibit shared morphological and physical changes (reviewed in Adams and Duggan, 2012). Specific changes include repression of hormogonia development regulated by the hrm locus, increases in heterocyst frequency (hetR gene) and N₂-fixation activity (nif genes), repression of GS-GOGAT activity, and reduced photosynthetic CO₂ fixation (reviewed in Adams and Duggan, 2012). These changes imply that the cyanobacterium shifts from photo-autotrophy to heterotrophic carbon metabolism, with organic C provided by host to the cyanobacteria to sustain N₂-fixation (Meeks and Elhai, 2002). The carbohydrate exported from the host is not identified, but in Gunnera it was suggested to be glucose and/or fructose (Khamar et al., 2010). In exchange, the cyanobacteria release 80–90% of their fixed N, mainly as ammonia for Gunnera, liverworts and hornworts symbioses (Meeks, 2009).

The aim of this study is to determine what gene acquisitions and regulatory rewiring allow cyanobacteria to form symbioses with feathermosses. To identify gene acquisitions, we sequenced and compared the genomes of novel Nostoc isolates to target putative symbiotic-competent gene clusters. To target regulatory rewiring, we then selected closely related symbiotic-competent and -incompetent Nostoc strains to investigate gene, protein and exoprotein expression during contact with the feathermoss. To distinguish effects of different phases of colonization on gene and protein expression, we used a novel experimental system to isolate physical versus chemical contact conditions with the host (Bay et al., 2013). We mined the gene expression of shared orthologs using random forest analysis, a supervised machine learning analysis based on decision tree predictors to identify expression signatures of genes that best discriminate symbiotic cyanobacteria from the non-symbiotic. The cellular and extracellular proteomes were also examined for patterns associated with cyanobacterial adaptation to the moss contact. We hypothesized that (1) sets of genes are absent from the genome of symbiotic-competent Nostoc and that these genes are of importance for initiating and maintaining a symbiotic association with feathermosses, (2) symbiotic-competent Nostoc strains exhibit a distinct expression profile in response to contact with the moss, as compared with incompetent strains, (3) competent Nostoc foster a distinct extracellular environment when in chemical contact with the host and (4) the epiphytic location of the symbiosis on feathermoss involves different physiological and metabolic processes during colonization relative to the endophytic symbioses in the liverwort/hornwort and Gunnera.

Materials and methods

Sampling, cyanobacteria isolation and growth conditions

Two feathermosses, P. schreberi and H. splendens, were randomly collected in June and September 2013 at the site of Rutjeheden in Reivo boreal forest nature reserve in northern Sweden (65°80′N–19°10′E). Mosses were maintained on a thin layer of vermiculite in the greenhouse at 10 °C and light 35 μmol photons m⁻² s⁻¹ (see Supplementary Methods for description of the isolation procedure). Individual cyanobacteria isolates were grown in liquid BG11o media (Rippka et al., 1979) with constant shaking under continuous light at 35 μmol photons m⁻² s⁻¹ and 20°C. Nostoc punctiforme ATCC 29133 and Nostoc sp. CALU 996 (N. 996), a free-living symbiotic-incompetent soil isolate (culture collection of St Petersburg State University, St Petersburg, Russia), which is phylogenetically similar to symbiotic Nostoc (Papaefthimiou et al., 2008) were also grown as liquid cultures in BG11o.

DNA extraction, genome sequencing and analysis

DNA was extracted from N. 996 and five Nostoc strains isolated from mosses (Supplementary Table S1, Supplementary Methods for description)
and sequenced using PacBio technology at the Joint Genome Institute (JGI) (see Supplementary Methods and Supplementary Figures S1 and S2 for description of genome sequencing, annotation, assembly and genome closure of N. 996). After kmer binning to remove low abundance contaminant contigs (Laczný et al., 2015), the chromosome and plasmid assemblies for the newly sequenced cyanobacteria were used to build a concatenated core gene phylogeny across all cyanobacteria. The existing phylogeny from the cyanoGEBA project (Shih et al., 2013) was constructed using a concatenated alignment set of proteins from 126 cyanobacterial genomes (Wu and Eisen, 2008) and provided a validated starting alignment. The existing cyanoGEBA alignment was used to construct HMMs for each core protein (HMMER3) (Eddy, 2011), with subsequent searches of the new genomes and retrieval of the correct proteins. The new genomes were added to the alignment as concatenated core protein strings, with a reconstruction of the whole genome phylogeny (Dupont et al., 2012) using RaxML (Stamatakis, 2014). All alignments and subsequent trees were manually examined with subsequent re-generation. The HMMER-based protein family annotations were used to compare the presence, absence and relative abundance of protein domains between all of the cyanobacterial genomes. To define orthologous protein families (OPFs) shared between Nostoc sp. Moss2 (N. Moss2), N. punctiforme and N. 996 (OPF_all) and between symbiotic strains (OPF_symb), we used reciprocal best BLAST hit clustering with predicted amino-acid protein sequences (see Supplementary Methods).

Experimental set up for chemical and physical contact
To mimic the chemical and physical contact of the cyanobacteria–moss symbiosis, experiments between P. schreberi and N. Moss2, N. 996 or N. punctiforme were performed as described by Bay et al. (2013) with slight modifications. For each cyanobacteria strain, triplicate cultures were set up with 1 and 8 μm membrane pore size filters (Corning, Inc., Corning, NY, USA) positioned above the cyanobacteria (for chemical and physical contact, respectively). In addition, triplicates were set up for moss alone (without cyanobacteria) and for each of the cyanobacteria (no mosses) (see Supplementary Methods). The chemical contact experiment was repeated to gain enough material for exoproteomic analyses, with a merging of media from both experiments before analyses. After 24 h, cyanobacteria and filtered spent media from the 1 μm experiment were harvested by centrifugation at 4 °C and promptly frozen in liquid N, and stored at -80 °C until RNA extraction. The mosses from the 8 μm experiment were harvested when colonization was noticed (4–5 weeks), frozen in liquid N and stored at -80 °C until RNA extraction (see Supplementary Methods for RNA and protein extraction procedure).

Transcriptome analysis
The moss–cyanobacteria physical contact transcriptome was treated as a mini-metatranscriptome (Supplementary Methods for description of transcriptomes sequencing, annotation and assembly). For all Nostoc-specific mapping reads, raw transcriptomic read mappings were normalized using TMM (Robinson and Oshlack, 2010) and analyzed for differentially expressed genes using the Bioconductor package EdgeR (Robinson et al., 2010). For symbiotic-competent cyanobacteria, there were two pairwise comparisons: (i) cyanobacteria in isolation versus chemical contact with P. schreberi and (ii) cyanobacteria in isolation versus physical contact with P. schreberi. For N. 996, the only possible pairwise comparison was the cyanobacteria in isolation versus chemical contact with P. schreberi.

A random forest analysis of the gene expression data was also used to identify putative gene expression patterns statistically associated with chemical or physical contact (see Supplementary Methods). The differential gene expression of profile-identified putative symbiotic genes defined by the random forest classification, hereafter referred to as PIPS genes, was visualized using AutoSOME software and co-expression modules were identified as described by Newman and Cooper (2010). Genomic locations of OPF_all+symb orthologs were manually curated and spatially colocalized regulons (SCRs) were defined as sets of orthologs genomically localized with a given PIPS gene; we examined the transcription profile of flanking genes in either direction in the genome of N. punctiforme, N. Moss2 and N. 996, cut-off distance of a flanking region was decided based on manual curation. SCRs were compared between the three strains, and spatially conserved and co-regulated SCRs (based on transcriptomics and/or proteomics) in the symbiotic cyanobacteria were selected for further analysis.

Proteome and exoproteome analysis
Extracted proteins from cyanobacteria alone (that is, N. 996, N. punctiforme, N. Moss2) and cyanobacteria–moss in chemical contact (that is, N. 996, N. punctiforme, N. Moss2) were prepared for proteome analyses as described in Supplementary Methods. Raw mass spectrometry data were converted to peak lists (DTA files) using DeconMSn (Mayampurath et al., 2008) and searched with MS-GF+ (Kim and Pevzner, 2014) against predicted proteins from the JGI-annotated genome for each species as the search database, except for N. punctiforme, which utilized the Uniprot N. punctiforme ATCC 29133 genome. The identified spectra were filtered based on their MSGF+ scores.
and only the proteins with two proteo-specific peptides were conserved resulting in a protein and peptide false discovery rate <1%. For the quantitative analysis, the tandem mass tag reporter ion intensities were extracted with MASIC (Monroe et al., 2008). Tandem mass tag reporter intensities were summed from the different peptides belonging to the same proteins.

For exoprotein extraction, 1.5–2 ml spent media from five replicate incubations of moss alone and moss-cyanobacteria in chemical contact for 24 h were analyzed for each N. Moss2, N. 996 and N. punctiforme (30 samples total). Scaffold (Proteome Software Inc., Portland, OR, USA) was used to validate peptide and protein identified by tandem mass spectrometry (MS/MS) and to calculate normalized weighted spectra values (NSAF). NSAF values were then averaged between replicates and log₂ fold change values calculated between the two treatments. A Student’s t-test was used to determine differences between the treatments (P < 0.05). (For a detailed description of extraction methods, search database and peptide identification thresholds see Supplementary Methods).

Results
Comparative genomic analysis
To determine if symbiotic-competent strains have unique genome contents, we sequenced five Nostoc strains that were isolated from mosses. For comparative genomics, we also sequenced one free-living Nostoc strain (N. 996), which we have experimentally verified as lacking symbiotic competence with P. schreberi, Blasia pusilla and Gunnera manicata. To establish the evolutionary relatedness of the six newly sequenced cyanobacteria, the sequences were included with 31 other Nostocales genomes in a phylogenomic inference built from the concatenated alignments of 15 conserved proteins (Figure 1a). The sequenced cyanobacteria belong to one clade, which consists of 12 strains currently classified as Nostocaceae, two as Microchaetaceae (Microchaete sp.

Figure 1. Nostocales species tree and distribution of HMM occurrences missing in Nostoc sp. CALU 996 (N. 996). (a) Maximum-likelihood phylogeny of Nostocales based on 15 conserved protein comparisons of the five moss symbiotic-competent strains sequenced in our study (in blue), the one non-symbiotic competent (in red) and 31 other Nostocales sequenced during the cyanoGEBA project (Shih et al., 2013). The genomes of Prochlorococcus marinus MED4 and Synechococcus sp. WH8102 were included as outgroups. Bootstrap values (out of 100) are shown at each node. Red boxes show subclades A and B, which include the new symbiotic-competent strains. (b) Distribution of the HMM occurrences missing in N. 996. Columns in the heat map represent individual gene families (Pfams or TIGRfams) grouped into eight functional protein categories. Increasing representation of each gene family in a given cyanobacteria strain is shown by increasing protein domains counts and depth color.
PCC 71269 and *Toiyophryx* sp. PCC 7601) and one as Rivulariaceaee (*Calothrix* sp. PCC 7507). The five moss-derived isolates were divided into two sub-clades within the Nostocales. The first subclade (A on tree) contained two of the isolates, *Nostoc* sp. Moss6 and *Nostoc* sp. Moss5 and were closely related to the free-living cyanobacteria *Anabaena variabilis* ATCC 29413 and *Nostoc* sp. PCC 7120. A second subclade (B) contained the other three isolates, *Nostoc* sp. Moss3, *Nostoc* sp. Moss4 and N. Moss2, which grouped together with the symbiotic strain *N. punctiforme* ATCC 29133. The symbiotic-incompetent strain N. 996 is a weakly supported (50%) basal outgroup for subclade B (Figure 1a).

We performed protein family profiling to identify gene families that were retained in the symbiotic strains but absent from the symbiotic-incompetent N. 996. We identified 32 gene families that lacked an HMM hit in N. 996 but were found in all feathermoss symbiotic isolates (Figure 1b, Supplementary Table S2). Many of the 32 gene families have non-redundant functional roles in gas vesicle formation, gene regulation and signaling, and sulfur metabolism. Three of these gene families have predicted functions in gas vesicle formation (PF00741, PF05121 and PF06386). Among the seven regulatory domains are a heme-nitric oxide (NO; PF07700) binding protein, and a heme NO-binding associated protein (PF07701). Domains involved in sulfur metabolism include putative aliphatic sulfonate transporter substrate bindin protein (TIGR01728), and a putative aliphatic sulfonate monoxygenase (TIGR03565). The aliphatic sulfonate transporter substrate binding protein was the most abundant differentially retained gene family found in symbiotic cyanobacteria, ranging from four protein domain counts in the N. Moss2 genome to 11 counts in *Nostoc* sp. Moss6 (Figure 1b, Supplementary Table S2).

**Transcriptional and proteomic regulation by *Nostoc* in physical or chemical contact with *P. schreberi***

To better understand the changes in gene regulation occurring during different phases of the cyanobacterial association with the moss, we performed transcriptomic, proteomic and exoproteomic analyses on triplicate cultures of *N. Moss2, N. punctiforme* and N. 996 under three conditions: cyanobacteria in isolation, in chemical contact with the moss and in physical contact with the moss (Supplementary Dataset S1 and Supplementary Table S3). To compare gene and protein expression between different strains and genomes, it is necessary to know the orthology of the different genome contents. Therefore, the predicted proteins from the genomes of *N. punctiforme*, *N. Moss2* and N. 996 were clustered into orthologous groups (Supplementary Table S4). OPFs shared between all three strains (OPF\(_{\text{all}}\)) consisted of 3117 orthologs, and 953 orthologs were only shared between N. 996 and *N. Moss2* (OPF\(_{\text{symb}}\)) (Figure 2, Supplementary Table S4). For the transcriptomics in chemical contact, all of the OPF\(_{\text{all}}\) and OPF\(_{\text{symb}}\) orthologs were detected. Despite the overwhelming representation of moss transcripts in the physical contact treatment data set, we identified reads mapping to 97–85% of OPF\(_{\text{all}}\) and OPF\(_{\text{symb}}\) orthologs during physical contact with *P. schreberi* (Figure 2). Proteomics identified 75% and 57% of OPF\(_{\text{all}}\) and OPF\(_{\text{symb}}\), respectively, and 5.8% and 3.5% of the orthologs were detected in the exoproteome (Figure 2). The depth of transcriptomic and proteomic coverage allows for maximal data input to analyses.

**Most abundant orthologous transcripts and proteins**

We identified the 10% highest expressed OPF\(_{\text{all}}\) and OPF\(_{\text{symb}}\) at the transcript, protein and exoprotein levels for each strain, based on average normalized counts in chemical and physical contact with the moss. This included 1179 transcripts across all cyanobacteria in chemical contact, 549 proteins and 112 exoproteins (Figures 3a–c). Figures 3a–c show the orthologs that are present in OPF\(_{\text{all}}\) and OPF\(_{\text{symb}}\). *N. Moss2* and *N. punctiforme* shared a unique complement of abundant transcripts and proteins belonging to OPF\(_{\text{symb}}\) and also a set included in OPF\(_{\text{all}}\) that are more highly expressed in symbiotic-competent cyanobacteria than N. 996 (Figures 3a–c). The OPF\(_{\text{symb}}\) orthologs are involved in pigment biosynthesis, exopolysaccharide production, extracellular proteins and RNA-binding proteins (Figures 3a and b). Three specific gene families missing from N. 996 were highly expressed, a protein of unknown function (PF13448), a secreted protein (metallopeptidase PF01447, PF02868), and one gas vesicle protein (PF00741) (Figure 3a). Only the PF13448 containing protein was found as highly produced in the proteome and exoproteome of symbiotic cyanobacteria (Figures 3b and c).
To investigate how cyanobacteria adapt their lifestyle post-colonization, we compared the 10% most abundant OPFall and OPFsymb orthologs when in isolation (that is, free-living N-fixing state) with the most abundant orthologs identified in physical contact (that is, symbiotic N-fixing state). In both symbiotic strains, most of the highly expressed orthologs in isolation are also significantly highly expressed in physical contact (Figures 3d–f).
were involved in CO₂ fixation (rbcLXS), photosynthesis (photosystem I psaABCDEFKL, photosystem II psbABCD), and cytochrome b₆-f complex (petABCD). GS-GOGAT (the glutamine synthetase gltA and the glutamate synthase gltB), phosphorus assimilation (pts), oxidative stress (peroxiredoxin and glutaredoxin synthesis), heterocyst differentiation (hetR), and N₂-fixation (nifBHKL). This represents a major departure from endophytic cyanobacterial-plant symbioses.

Symbiotic-competent exprotein composition

Of the 214 orthologs detected in the exoproteomes of symbiotic-competent strains, 33 are unique to OPFₕₜᵦₘ (Figure 2, Supplementary Dataset S1, Supplementary Table S5). OPFₕₜᵦₘ orthologs included a putative pectate lyase (polysaccharide lyase family 9 CAZy domain, cluster #669), and hemolysin type C-binding protein (cluster #7058; also identified as PIPS gene, see below). Both proteins were upregulated in the exoproteome when in contact with the host, and were in the top 10% abundance in the cellular proteome. Of the OPFₜₜᵦₘ exoproteins, several involved in oxidative stress response were differentially expressed in the competent strain exoproteomes, including a catalase protein (cluster #2351; PIPS gene), upregulated under chemical contact in the N. punctiforme exoproteome and not detected in N. 996. This catalase was also transcriptionally upregulated in both N. Moss2 and N. punctiforme under physical contact (Supplementary Dataset S1, Supplementary Table S5). In addition, a ferritin protein (cluster #5454; PIPS gene) and a flavorubredoxin (cluster #2363) were detected in all three exoproteomes, but only upregulated in the presence of the moss in the competent strains.

Machine learning identification of symbiosis-specific expression patterns

A supervised machine learning approach, random forest analysis with decision trees, was used to identify expression patterns of the subset of OPFₜₜᵦₘ+symb orthologs that best differentiated the symbiotic cyanobacteria N. Moss2 and N. punctiforme (4070 orthologs, Supplementary Dataset S1) from the non-symbiotic N. 996 transcriptome across physical and chemical contact conditions. We identified 405 PIPS genes (282 OPFₜₜᵦₘ and 123 OPFₜₜᵦₘ orthologs) that together predicted the cyanobacterial symbiotic state with 98.3% confidence (Supplementary Table S6).

We conducted unsupervised clustering of the expression profile of PIPS genes using self-organizing maps (Newman and Cooper, 2010), which revealed 21 colonization phase-dependent gene co-expression modules (Supplementary Figure 3). The second large module (35 OPFₜₜᵦₘ and 70 OPFₜₜᵦₘ) was upregulated PIPS genes during chemical contact with the moss, and was particularly enriched in genes involved in gas vesicle organization and chemotaxis (Supplementary Figure 3a). The largest module (164 OPFₜₜᵦₘ and 17 OPFₜₜᵦₘ orthologs) comprised PIPS upregulated genes during the physical phase and was enriched with GO terms involved in reactive oxygen species homeostasis, photosynthesis, c-di-GMP biosynthetic process and sugar, phosphorus and sulfur metabolisms (Supplementary Figure 3c). The other 19 modules were composed of genes similarly regulated during both phases and merged into one cluster, which contained 83 OPFₜₜᵦₘ and 36 OPFₜₜᵦₘ orthologs with genes enriched in GO terms associated with chemotaxis, type II secretion system and pili genes (Supplementary Figure 3b).

SCRs shared between the strains

We examined the genomic spatial organization of the 4070 OPFₜₜᵦₘ+symb orthologs and combined it with the differential gene expression analyses to identify spatially colocalized and co-regulated sets of genes (SCRs) shared in N. Moss2 and N. punctiforme and genes differentially expressed between the symbiotic-competent and -incompetent strains (Table 1, Figure 4, Supplementary Table S7). For a

**Table 1** SCRs identified in N. Moss2, N. punctiforme and N. 996

| SCR function | Interaction phase | SCR name | SCR statistics |
|--------------|------------------|----------|----------------|
| Chemotaxis and motility | Chemical contact | gvp and ptx | 22 genes; 20 OPFₕₜᵦₘ; 18 PIPS genes |
| | Chemical contact | hrm | 6 genes; 4 OPFₕₜᵦₘ; 2 PIPS genes |
| | Chemical and physical contact | pix | 13 genes; 6 OPFₕₜᵦₘ; 6 PIPS genes |
| | Chemical and physical contact | hmp | 8 genes; 7 OPFₜₜᵦₘ; 1 PIPS gene |
| | | hps | 15 genes; 15 OPFₜₜᵦₘ; 9 PIPS genes |
| | | pil I | 15 genes; 13 OPFₜₜᵦₘ; 4 PIPS genes |
| | | pil II | 31 genes; 30 OPFₜₜᵦₘ; 8 PIPS genes |
| Sensing and signal transduction | Chemical and physical contact | Heme NO-binding | 12 genes; 6 OPFₜₜᵦₘ; 4 PIPS genes |
| | Chemical and physical contact | FG-GAP | 6 genes; 5 OPFₜₜᵦₘ; 1 PIPS gene |
| | | Flavonohydrolase | 10 genes; 10 OPFₜₜᵦₘ; 0 PIPS genes |
| | | Glycosyltransferase | 11 genes; 11 OPFₜₜᵦₘ; 2 PIPS genes |
| | | Fructose/mannose metabolism | 13 genes; 13 OPFₜₜᵦₘ; 2 PIPS genes |
| Transport | Physical contact | Aliphatic sulfonate transporter | 5 genes; 4 OPFₜₜᵦₘ; 1 PIPS gene |
| | | Sulfate/phosphate transport | 12 genes; 8 OPFₜₜᵦₘ; 1 PIPS gene |

Abbreviations: FG-GAP, FG-GAP repeat-containing protein; NO, nitric oxide; OPF, orthologous protein family; PIPS, profile-identified putative symbiotic; SCRs, spatially colocalized regulons.
detailed description of the SCRs, see Supplementary Results. Overall, we identified 14 SCRs, with two that were regulated in chemical contact with *P. schreberi*, nine SCRs differentially expressed during both chemical and physical contact and three differentially expressed only during physical contact (Table 1, Supplementary Table S7). The SCRs were mostly mono-functional, as expected given the genomic colocalization, and separately involved in chemotaxis and motility, sensing and signal transduction and macromolecule biosynthesis and transport. We found four SCRs composed of OPF orthologs, which were differentially expressed in N. 996 compared with symbiotic strains. Those SCRs were associated with hormogonia motility (*hmp*, *hps*, *pil I*, *pil II* and glycosyltransferase) (Table 1, Supplementary Table S7). Six SCRs were composed of gene families missing from N. 996, involved in chemotaxis (*gvp* and *ptx*, and *pix*), motility (*pil II*), sensing of NO (heme NO-binding) and other functions (FG-GAP repeat-containing protein and aliphatic sulfonate transport) (Figure 4, Table 1, Supplementary Table S7).

**Discussion**

**Overview and hypotheses restatement**

With any data-rich study, experimental design and explicit hypotheses focus a robust analysis. Here, the genomes of closely related *Nostoc* strains, five competent and one incompetent for symbioses with feathermosses, were sequenced and compared. From our phylogenetic analysis, the feathermoss *Nostoc* isolates are divided in two subclades suggesting the taxonomy of *Nostoc* and related genus need further revisions (Komárek, 2010). The feathermoss isolate most closely related to the model symbiotic strain, *N. punctiforme*, and the closely related symbiotic-incompetent strain were chosen for further global gene and protein expression experiments. The samples for these analyses originated from a unique culturing framework allowing for controlled and replicated sampling of cyanobacteria under three conditions: in isolation, in chemical contact with *P. schreberi* (early colonization phase), and in physical contact with *P. schreberi* (late colonization phase). Our hypotheses framing the data analyses...
were that (1) symbiotic-competent Nostoc strains contain symbiosis-related gene families not found in incompetent strains, (2) symbiotic-competent Nostoc have a shared expression profile in response to contact with P. schreberi, distinct from incompetent strains, (3) symbiotic-competent strains foster a different extracellular environment when in chemical contact with the mosses and (4) the epiphytic nature of the cyanobacteria–moss symbiosis is unique because of novel regulation of metabolism and phenotype, as well as molecular currency exchange. The first hypothesis entails fundamental evolutionary adaptations shaped by gene acquisition or selective gene retention, whereas the second and third hypotheses require regulatory reprogramming and likely involve host-based signaling. The final hypothesis challenges singularity in cyanobacterial host adaptations, thereby refining existing theory to create a gradient of molecular adaptations based upon defined criteria of physical contact, namely endophyte, both intra and extracellular, and epiphyte. Each of these hypotheses will be addressed with regards to the experimental results below, and based on interpretations of integrated results, we will detail a model of new molecular currency exchanges between host and symbiont.

**Gene families specific to symbiotic strains**

We found that symbiotic-competent cyanobacteria genomes contain 32 gene families not found in closely related strains including the symbiotic-incompetent N. 996. These families were significantly expressed during the early colonization phase and are enriched in functional attributes such as chemotaxis and motility, signaling, and aliphatic sulfonate transport. This includes known molecular participants in cyanobacteria–plant symbioses. For example, seven upregulated gas vesicle genes (gvp), here predicted as PIPS genes, previously showed transcriptional upregulation in N. punctiforme upon addition of HIF, indicating a role in cyanobacterial–plant symbioses establishment (Campbell et al., 2008; Christman et al., 2011; Risser et al., 2014). Deletion of gvpA showed that this gene is essential to N. punctiforme motility on a solid substrate (Risser et al., 2014). The upregulation of gvp and ptx genes specifically during the early phase indicate that P. schreberi activates a signal transduction pathway for hormogonia motility unique to this phase. By contrast, another taxis locus pix, which is missing in N. 996, was upregulated during both colonization phases, suggesting phase-dependent signaling pathways. A FG-GAP repeat-containing protein (Npun_R6586) was also only present in the genomes of the symbiotic-competent strains, and found among the 10% most abundant transcripts, cellular proteins, and exoproteins during chemical contact with P. schreberi. The FG-GAP repeat-containing protein may have a role in the establishment of symbiosis, and is clearly secreted, given its abundance in the exoproteome. Two more gene families missing in N. 996 genome are involved in NO signaling. The heme-NO binding and a NO-associated (annotated as a histidine kinase) genes were identified as PIPS genes and were upregulated during chemical and physical contact, suggesting a role of NO in signaling during establishment and maintenance of the symbiosis. Increased NO concentration may lead to upregulation of the heme-NO binding gene and c-di-GMP synthesis genes inducing biofilm and adhesion of cyanobacteria to the host. Rao et al. (2015) showed that Silicibacter sp. TrichCh4B and the marine cyanobacterium Trichodesmium erythraeum have a NO-mediated mechanism for symbiosis, in which T. erythraeum produces a chemical signal, inducing NO formation in the symbiont. NO activates the heme-NO binding protein signaling pathway of Silicibacter sp. TrichCh4B, leading to higher levels of cellular c-di-GMP controlling adhesion and biofilm formation. As NO synthase genes were not identified in our Nostoc strains, the host would be the likely source of NO. There is evidence that P. schreberi NO synthase gene was upregulated when in physical contact with P. schreberi and N. punctiforme and accumulation of NO was observed by fluorescence probing with 4,5-diaminofluorescein-2 diacetate (unpublished work). Biofilm formation is often critical to host–symbiont communication, especially when it is mediated by diffusible signaling molecules (Norsworthy and Visick, 2013), so biofilm formation may favor host–symbiont NO-mediated communication and enhance currency exchange. NO-mediated signaling may also allow the host to guide cyanobacteria migration and colonization of new plant tissue as NO is highly produced in protonema cells during the growth of the moss Physcomitrella patens (Medina-Andres et al., 2015).

In support of our second hypothesis, we found that OPFs shared by both symbiotic-competent and -incompetent cyanobacteria are expressed differently in contact with the moss. Our random forest analysis shows that more than half of the 405 PIPS genes, those with expression profiles diagnostic of the phenotype, are OPF	extsubscript{all} orthologs. These are 282 genes present in the symbiotic-incompetent strain that have been transcriptionally reprogrammed. In particular, genes involved in hormogonia motility (hmp, hps, pil I, pil II and glycosyltransferase), further classified into SCRs, were upregulated in the two competent strains but not in N. 996. N. punctiforme motility involves co-ordinated expression of the pil genes encoding a type IV pilus that acts as a nanomotor, driving motility on a secreted polysaccharide layer encoded by the hps genes, with hormogonium gliding regulated by the hmp locus (Duggan et al., 2007; Risser and Meeks, 2013; Risser et al., 2014; Khayatan et al., 2015). The glycosyltransferase SRC may also be involved in the motility process by producing an excreted polysaccharide. The motility loci were also
upregulated during physical contact with \textit{P. schreberi}, which we propose are because of the cyanobacteria being epiphytic, retaining motility and chemotaxis allowing for colonization of new parts of the moss. Along with upregulation of the motility loci, the \textit{hrm} locus, which controls hormogonia repression (Cohen and Meeks, 1997; Meeks, 1998; Campbell \textit{et al.}, 2003), was downregulated or not detected in physical contact with \textit{P. schreberi}. Our results suggest that although \textit{N. 996} shares many of the genes coding for motility, chemotaxis and cellular differentiation, these are insensitive to the chemical signaling derived from \textit{P. schreberi}. This differential expression of a shared gene set is a concrete indication of moss-derived chemical signaling of regulatory reprogramming in the symbiotic-competent strains. Such organism-specific information exchange is one hallmark of symbiotic interactions.

Exoprotein composition provides support for our third hypothesis, which the symbiotic-competent strains foster a distinct extracellular environment when in chemical contact with their host, specifically through glycosyl modifications and oxidative stress mediation. In cyanobacteria, hydrolytic exoproteins in biofilms support heterotrophic acquisition of complex polymers, whereas in a free-living cyanobacterium diverse oxoenzymes, such as hemo-lisin, virulent factor-like metalloproteases and chitinases, may indicate competitive interactions dominate (Christie-Oleza \textit{et al.}, 2015; Stuart \textit{et al.}, 2016). Our data provide evidence for specific oxoenzymes produced by the symbiotic-competent cyanobacteria in response to the moss. For example, microbial pectate lyases have been implicated in plant pathogenesis (reviewed in Payasi \textit{et al.}, 2009), and legume-produced pectate lyases allow symbiont-rhizobia infection (Xie \textit{et al.}, 2012). Our symbiont-specific putative pectate lyase exoprotein may perform a similar function, assisting in colonization of the feathermoss surface through cell wall modifications. We also found that \textit{hpsJ}, involved in polysaccharide synthesis, is downregulated in the exproteome of \textit{N. punctiforme} providing evidence for a modification of the exopolysaccharide composition during interaction with the moss. Deletion of \textit{hpsJ} affects polysaccharide composition, indicating the role of this gene in defining the nature of hormogonium polysaccharide (Risser \textit{et al.}, 2014; Khayatan \textit{et al.}, 2015). Among the 10% most abundant exoproteins expressed in chemical contact, we also found several linked to oxidative stress. Specifically, we found two flavorubredoxins upregulated in the exproteome of both symbiotic-competent cyanobacteria and downregulated in \textit{N. 996}. There were also several splay-competent exoproteins that may be involved in oxidative stress mediation, including a catalase and ferritin, supporting the importance of mediating reactive nitrogen species and reactive oxygen species in establishing and maintaining the symbiosis.

In support to our last hypothesis, we found that the cyanobacteria in symbiosis with feathermoss have a distinct morphological and metabolic regulation relative to endophytic cyanobacteria–plant symbioses. In endophytic systems, cyanobacterial heterocyst frequency and the \textit{N2}-fixation activity increases when in contact with the host (Adams and Duggan, 2012). We did not observe any statistically significant upregulation of \textit{nifH} or \textit{hetR} genes when associated with \textit{P. schreberi} relative to \textit{N}-fixing cells in isolation, indicating that the cyanobacteria do not increase \textit{N2}-fixation activity when on the moss. Unequivocally, cyanobacterial fixed \textit{N2} is transferred to the moss (Bay \textit{et al.}, 2013), and the continued expression of \textit{nifH} here is consistent with that. Despite lack of differential expression when comparing isolation and physical contact treatments, both \textit{hetR} and \textit{nif} genes were among the 10% most abundant transcripts when on the moss. In other cyanobacterial symbioses, except the cycad association, ammonium is released by \textit{Nostoc}, achieved partly by a modulation of glutamine synthetase activity, with a reduction from 70% to 15% of \textit{in vitro} activity of the enzyme during symbiosis with \textit{Gunnera} and \textit{Anthoceros}, respectively (reviewed in Meeks, 2009). In our study, \textit{glnA} (glutamine synthetase, GS) and \textit{glkB} (glutamate synthase, GOGAT) transcripts are highly abundant post-colonization and statistically indistinguishable from the free-living condition. The results could indicate that the cyanobacteria may not transfer ammonium to the moss, but rather amino acids, as in the cycad–cyanobacteria symbiosis (Costa and Lindblad, 2002). Finally, in symbiosis with \textit{Gunnera} and \textit{Blasia}, cyanobacterial light-dependent \textit{CO2} fixation is reduced by 90–100%, in return for combined \textit{N}, the cyanobacteria is compensated by organic C from their host (Meeks and Elhai, 2002; Khamar \textit{et al.}, 2010). By contrast, our data indicate that light-dependent \textit{CO2} and photosynthesis-related genes are highly and equally abundant in association with \textit{P. schreberi} as in isolation, suggesting that cyanobacteria remain photo-autotrophic when in symbiosis with the moss.

Nitrogenase activity of feathermoss associated cyanobacteria is limited by phosphorus availability (Rousk \textit{et al.}, 2016), and sulfur is an essential component of abundant Fe-S clusters in a mature nitrogenase. Neither seems logically available in sufficient quantities on a plant surface. During physical contact, we observed transcriptional upregulation of phosphate and sulfate transporters in the symbiotic-competent strains. Interestingly, the protein families of aliphatic sulfonate transporters and monoxygenases in the genomes of moss symbiotic-competent cyanobacteria are highly upregulated during physical contact. The alkane sulfonate sulfur-specific catabolism pathways were identified using phylogenetic profiling and hypothesized to catalyze the anabolic incorporation of sulfonate-derived sulfur into homocysteine, a direct input to
cysteine synthesis. We propose aliphatic sulfonates may act as a ‘currency’ in the symbiosis, which has been suggested in marine diatom-bacteria interactions (Durham et al., 2015). In diatom–bacteria interactions, the bacteria utilizes carbon from the sulfonate, whereas in our study the genomically colocalized genes for sulfur anabolism suggest the sulfur is the desired component of the sulfonate. These findings indicate that cyanobacteria epiphytically located on the moss do not obligately depend on the host for fixed carbon, and instead the host may ‘reward’ their associated symbiotic cyanobacteria with an aliphatic sulfonate compound.

Our study defined a repertoire of important functional genes for the moss–cyanobacteria symbiosis composed of previously described OPFs as well as OPFs never before associated with cyanobacteria–plant symbioses. The latter include genes involved in gas vesicles formation, NO-sensing and aliphatic sulfonate transport. We also show that the competent strains differentially express a unique suite of genes related to motility, NO regulation, sulfate/phosphate transport compared with the non-competent strain. Based on these findings, we propose a model for the establishment and maintenance of the symbiosis, schematically presented in Figure 5. We demonstrate that the Nostoc symbiosis with feathermoss is distinct from that with other plants, most likely due to its epiphytic nature. The differences include the retention of motility and chemotaxis post-colonization, as well as constitutive regulation of N fixation, photosynthesis, GS-GOGAT cycle, and heterocyst formation. The epiphytic cyanobacteria is spatially mobile and chemotaxic, therefore predictably redeployed, while still exchanging needed metabolites with the host. Along with describing essential genes to the Nostoc feathermoss symbioses, and newly described pathways that either shared among cyanobacterial symbionts, or specific to epiphytic symbioses, our conceptual

Figure 5 Conceptual model for establishment and maintenance of cyanobacteria on feathermoss. A putative time-line includes the following steps: (1) moss releases NO into the environment, which is sensed by the heme NO-binding protein (H-NOX) fused with associated histidine kinase (HK), (2) cyanobacteria respond with reactive nitrogen species mediation by the secretion of flavohemoglobin (F) and FG-GAP repeat-containing protein (G) as chemical cues to the host and other cyanobacteria, (3) NO signaling pathway triggers hormogonia differentiation (hrm locus) and gas vesicle formation (gvp locus), (4) chemotaxis and motility (pxz, pix and hmp loci and pil and hsp loci, respectively), (5) cyanobacteria colonize the moss and differentiate into vegetative stage, (6) the host signals to the cyanobacteria through NO for spatial localization and re-direction of the hormogonia, (7) increased NO signaling induces biofilm formation to favor host–symbiont NO-mediated communication and enhance currency exchange, (8) production of aliphatic sulfonate (AS) compound and transfer from the host to the cyanobacteria. The cyanobacteria provide N as amino acid (AA) to the moss. Newly described loci/function are in bold.
model and newly described gene candidates provide targets for verification by gene knockout and testing on feathermosses and other hosts.

Data availability
The data have been deposited in https://gold.jgi.doe.gov/studies?id=Gs0110198.

Conflict of Interest
The authors declare no conflict of interest.

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