Novel Metabolic Attributes of the Genus Cyanothece, Comprising a Group of Unicellular Nitrogen-Fixing Cyanobacteria

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ABSTRACT The genus Cyanothece comprises unicellular cyanobacteria that are morphologically diverse and ecologically versatile. Studies over the last decade have established members of this genus to be important components of the marine ecosystem, contributing significantly to the nitrogen and carbon cycle. System-level studies of Cyanothece sp. ATCC 51142, a prototypic member of this group, revealed many interesting metabolic attributes. To identify the metabolic traits that define this class of cyanobacteria, five additional Cyanothece strains were sequenced to completion. The presence of a large, contiguous nitrogenase gene cluster and the ability to carry out aerobic nitrogen fixation distinguish Cyanothece as a genus of unicellular, aerobic nitrogen-fixing cyanobacteria. Cyanothece cells can create an anoxic intracellular environment at night, allowing oxygen-sensitive processes to take place in these oxygenic organisms. Large carbohydrate reserves accumulate in the cells during the day, ensuring sufficient energy for the processes that require the anoxic phase of the cells. Our study indicates that this genus maintains a plastic genome, incorporating new metabolic capabilities while simultaneously retaining archaic metabolic traits, a unique combination which provides the flexibility to adapt to various ecological and environmental conditions. Rearrangement of the nitrogenase cluster in Cyanothece sp. strain 7425 and the concomitant loss of its aerobic nitrogen-fixing ability suggest that a similar mechanism might have been at play in cyanobacterial strains that eventually lost their nitrogen-fixing ability.

IMPORTANCE The unicellular cyanobacterial genus Cyanothece has significant roles in the nitrogen cycle in aquatic and terrestrial environments. Cyanothece sp. ATCC 51142 was extensively studied over the last decade and has emerged as an important model photosynthetic microbe for bioenergy production. To expand our understanding of the distinctive metabolic capabilities of this cyanobacterial group, we analyzed the genome sequences of five additional Cyanothece strains from different geographical habitats, exhibiting diverse morphological and physiological attributes. These strains exhibit high rates of N₂ fixation and H₂ production under aerobic conditions. They can generate copious amounts of carbohydrates that are stored in large starch-like granules and facilitate energy-intensive processes during the dark, anoxic phase of the cells. The genomes of some Cyanothece strains are quite unique in that there are linear elements in addition to a large circular chromosome. Our study provides novel insights into the metabolism of this class of unicellular nitrogen-fixing cyanobacteria.

Cyanobacteria constitute a fascinating group of photosynthetic prokaryotes that have inhabited almost every sunlit ecosystem of the earth for ~3 billion years. The remarkable success of this group of microbes in adapting to a wide range of environmental and ecological conditions has largely been attributed to the presence of an extraordinarily flexible repertoire of metabolic pathways (1, 2). Cyanobacteria possessed an efficient cellular machinery to function in anaerobic environments that prevailed during the mid-/late Archaean era, and their metabolic activities are credited for the transitioning of the earth into the present-day oxygen-rich environment (3). In fact, many of the archaic metabolic traits have been retained in extant cyanobacterial species, enabling them to thrive in many diverse ecological niches.

The metabolic feats of cyanobacteria are exemplified by the ability of some strains to fix molecular nitrogen, a process sensitive to oxygen (4) and not found in any other known oxygenic organism (5, 6). Cyanobacteria have adapted various strategies to meet the cellular demands of nitrogen fixation, the most critical being the protection of the oxygen-sensitive nitrogenase enzyme (7). While some filamentous strains have developed specialized cells, called heterocysts, to accommodate this process, unicellular strains make use of the diurnal cycle to separate oxygen-evolving photosynthesis from oxygen-sensitive nitrogen fixation (8, 9). Recent studies have demonstrated the importance of unicellular nitrogen-fixing cyanobacteria in the marine nitrogen and carbon cycle (10, 11). The efficiency of nitrogen fixation exhibited by these microbes during the dark period of a day/night cycle suggests that they must have the ability to harvest and store sufficient
solar energy during the day, which in turn fuels the energy-intensive nitrogen-fixing process at night. The suboxic intracellular conditions that are maintained during the nitrogen fixation period also facilitate various fermentative processes that are commonly observed in many facultative and obligate anaerobes (12, 13).

*Cyanothece* is a genus of morphologically diverse, unicellular cyanobacteria that are known to inhabit a variety of ecological niches. This genus was created by Komárek (14) to include unicellular cyanobacteria with distinct morphological, ultrastructural, and genomic features (15–17). *Cyanothece* strains from various ecotypes have been studied in the past for their robust circadian rhythm, fermentative capabilities, and other biotechnological applications (12, 18, 19). *Cyanothece* sp. ATCC 51142 [hereinafter referred to as *Cyanothece* 51142], a prototypic member of this genus, was isolated from the Texas Gulf Coast and is one of the most potent diazotrophic strains yet characterized (20).

System-level studies with *Cyanothece* 51142 revealed many novel metabolic traits of this unicellular cyanobacterium which led to the determination of its genome sequence at the Washington University sequencing center (21). The studies revealed robust diurnal and circadian cycling of central metabolic processes in this strain, as well as a strong coordination of correlated processes at the transcriptional level (22). Interestingly, genome analysis of *Cyanothece* 51142 uncovered the presence of a 430-kb functional linear chromosomal element, the first such element to be identified in any photosynthetic bacterium. The arrangement of genes on this chromosome suggested a specific role for it in energy metabolism, and it was hypothesized that such linear elements with regulatory functions might be a distinctive trait of the genus *Cyanothece* (21). Also interesting from the genomic perspective is the finding that some atypical nitrogen-fixing strains, such as the endosymbiont spheroid body of the eukaryotic diatom *Rhopalodia gibba* and the unicellular marine cyanobacterium UCYN-A, which lacks photosystem II, have genomes closely related to those of *Cyanothece* spp. (23, 24). In particular, the nitrogenase gene clusters in both of these organisms is highly similar to that in *Cyanothece* 51142. It has been hypothesized that these organisms may have evolved as a result of targeted gene loss (loss of genes involved in photosynthesis while maintaining an elaborate gene cluster involved in nitrogen fixation) from a *Cyanothece*-like ancestor (23), thus suggesting a highly plastic nature of *Cyanothece* genomes as well as the robustness of their nitrogen-fixing machinery.

The most striking of the unique metabolic capabilities of *Cyanothece* 51142 is that cells can exhibit high rates of nitrogenase-mediated H₂ production under aerobic conditions, an unusual metabolic trait in oxygenic phototrophs (25). Furthermore, the metabolic versatility of this strain was demonstrated by its ability to switch between photoautotrophic and photoheterotrophic modes of metabolism depending on the availability of external carbon sources and the presence of an atypical alternative citramalate pathway for isoleucine biosynthesis (26).

In an effort to unravel the genomic basis of the observed metabolic traits of unicellular diazotrophic cyanobacteria, the genomes of five additional members of the genus *Cyanothece* (*Cyanothece* sp. strains PCC 7424, PCC 7425, PCC 7822, PCC 8801, and PCC 8802 [hereinafter referred to as *Cyanothece* 7424, 7425, 7822, 8801, and 8802]) were sequenced at the Joint Genome Institute, U.S. Department of Energy. The strains were collected from different geographical locations and exhibit considerable diversity with respect to cell size and pigment composition. A comparison of the genomes of the different *Cyanothece* strains revealed that members of this genus are metabolically versatile, each member having acquired unique metabolic capabilities. The capability of aerobic nitrogen fixation and the presence of a large, contiguous *nif* gene cluster distinguish this group of unicellular photosynthetic microbes. Analysis of the genes common and unique to five of the six *Cyanothece* strains revealed that the core *Cyanothece* genomes is an amalgamation of genes from strains associated with fermentative capabilities, such as *Microcystis* and *Microcoleus* strains, and from aerobic nitrogen-fixing filamentous strains. The key to the success of this group of organisms appears to lie in their ability to retain such useful metabolic traits as nitrogen fixation and anaerobic fermentation while simultaneously adapting and accommodating advanced cellular features of contemporary photosynthetic organisms.

**RESULTS**

General features of the *Cyanothece* genomes. Table 1 summarizes the general characteristics of the six sequenced *Cyanothece* genomes. *Cyanothece* 51142 is a marine (benthic) strain, whereas *Cyanothece* 7424, 7425, 7822, 8801, and 8802 were collected from different tropical and subtropical rice fields in Asia and Africa. The genome sizes of the six strains show considerable variation, ranging between ~4.8 and 7.8 Mbp. *Cyanothece* 7822 has the largest cell size (8 to 10 μm), as well as the largest genome, with ~6,600

| Characteristic of *Cyanothece* strains | ATCC 51142 | PCC 7424 | PCC 7425 | PCC 7822 | PCC 8801 | PCC 8802 |
|---------------------------------------|------------|----------|----------|----------|----------|----------|
| Cell size (μm)                        | 4–5        | 7–8      | 3–4      | 8–10     | 4–5      | 4–5      |
| Site of isolation                     | Port Aransas, TX | Rice field, Senegal | Rice field, Senegal | Rice field, India | Rice field, Taiwan | Rice field, Taiwan |
| Size (Mbp)                            | 5.46       | 6.55     | 5.79     | 7.84     | 4.79     | 4.80     |
| No. of coding sequences               | 5,304      | 5,710    | 5,327    | 6,642    | 4,367    | 4,444    |
| % G+C                                 | 37.1       | 38.0     | 49.9     | 39.6     | 39       | 39       |
| No. of plasmids                       | 4          | 6        | 3        | 3        | 3        | 4        |
| No. of linear chromosomal elements    | 1          | 0        | 0        | 3        | 0        | 0        |
| No. of pseudogenes                    | 6          | 177      | 131      | 341      | 200      | 206      |

a Size bar, 2 μm.
protein-coding genes, whereas *Cyanothece* 8801 and 8802 have the smallest genomes, with ~4,400 open reading frames. The genome of each strain consists of a circular chromosome and several (3 to 6) smaller plasmids. The plasmids range in size from ~10 kb (smallest plasmid in *Cyanothece* 51142) to ~330 kb (largest plasmid in *Cyanothece* 7424).

Like *Cyanothece* 51142, *Cyanothece* 7822 also carries linear chromosomal elements in its genomes. The single linear chromosome in *Cyanothece* 51142 did not exhibit significant synteny with any sequenced cyanobacterial genome except the partially sequenced genome of *Cyanothece* sp. strain CCY 0110. Based on this observation, it was suggested that the linear chromosome might be specific to the genus *Cyanothece* (21). Interestingly, our analyses showed that *Cyanothece* 7822 has 3 linear chromosomal elements in its genome. The largest of these is an 880-kb element with 595 coding sequences, followed by a 474-kb fragment with 422 coding sequences. The third linear element is 14 kb long with 13 coding sequences. About 50% of the coding genes in the largest linear chromosome are unique to *Cyanothece* 7822, and 229 of these are without any paralogs elsewhere in the genome. A large fraction of these genes encode ABC transporters, with two operons containing genes involved in phosphate and molybdenum transport (see Table S1 in the supplemental material). In addition, this linear element has a significant number of genes involved in carbohydrate metabolism, including a cluster of genes encoding carbohydrate degradation and glycosylation proteins. Several genes encoding proteins with regulatory functions, transposons, and CRISP-R-associated proteins, as well as proteins involved in aromatic compound degradation, are also present in this chromosome. A cluster of cytochrome oxidase (*cox*) genes involved in respiration is common to the linear chromosomes of both *Cyanothece* 51142 and 7822. About 65% of the genes on the second-largest element are unique to *Cyanothece* 7822, with 52% of the genes unique to this chromosomal element.

The GC content of five of the sequenced *Cyanothece* genomes is close to 40 percent. *Cyanothece* 7425 is an exception, with a GC content of 50%. The terrestrial *Cyanothece* strains have a high percentage of pseudogenes in their genome (Table 1). Although *Cyanothece* 51142 has only 6 pseudogenes, *Cyanothece* 7822 contains 341, whereas in *Cyanothece* 8801 and 8802, the strains with the smallest genomes, the ~200 pseudogenes account for ~4% of the genome.

**Phylogenetic analysis.** Phylogenetic analysis of 61 cyanobacterial genomes using 226 homolog protein groups (see Materials and Methods), revealed various novel aspects of the evolutionary history of the *Cyanothece* strains. Based on this analysis, five of the six *Cyanothece* strains (*Cyanothece* 51142, *Cyanothece* 8801, *Cyanothece* 8802, *Cyanothece* 7822, and *Cyanothece* 7424) branch into a single clade together with three other nitrogen-fixing unicellular cyanobacteria, *Cyanothece* CCY 0110, *Crocosphaera watsonii* WH 8501, and UCYN-A (Fig. 1). These eight strains are likely to have evolved from a common ancestor, from which the two nondiazotrophic *Microcystis* strains also seem to have branched. Two other nondiazotrophic strains, *Synechococcus* sp. strain 7002 and *Synechocystis* sp. strain 6803, form a distant branch within this unicellular nitrogen-fixing group (*Cyanothece* 51142 to *Synechococcus* sp. strain JA-2-3Ba). Striking in this analysis is the position of *Synechococcus* sp. strain 7002, which appears to have evolved separately and is phylogenetically close to *Acaryochloris marina* MBIC 11017 (a chlorophyll-d-containing strain), compared to any other *Cyanothece* strain. *Cyanothece* 7425 branched off earlier than most other nitrogen fixers except for three anaerobic nitrogen-fixing *Synechococcus* strains (*Synechococcus* sp. strain 7335, *Synechococcus* sp. strain JA-3-3AB [2–13], and *Synechococcus* JA-2-3Ba).

One of the two main branches in the phylogenetic tree contains only non-nitrogen-fixing cyanobacteria (consisting of *Synechococcus* sp. strain 6301, *Synechococcus* sp. strain 7942, and all alphacyanobacteria). The other branch (from *Nodularia spumigena* CCY 9414 to *Synechococcus* sp. JA-2-3Ba), although consisting predominantly of nitrogen fixers, is interspersed with non-nitrogen-fixing strains. As suggested by earlier phylogenetic studies of diazotrophic cyanobacteria (21, 27), it is likely that some of the cyanobacteria in the second branch lost their nitrogen-fixing capability in the course of evolution. The position of the newly sequenced *Cyanothece* 7425 (with a functional nitrogenase cluster), which branched off from a common ancestor with *A. marina* (a non-nitrogen-fixing strain), strengthens this premise.

**Shared and unique genes in *Cyanothece* genomes.** Based on NCBI protein BLAST analysis (see Materials and Methods for details), we identified 1,705 homologous gene groups that are shared by all of the six *Cyanothece* strains (see Table S2 in the supplemental material). Using the classification scheme in the CyanoBase database (28), 1,003 (59%) of these genes are associated with known functional categories. Genes related to nitrogen fixation, central carbon metabolism, photosynthesis, respiration, and most common amino acid biosynthetic pathways are included in this shared group of genes. When the protein sequences of these homologous genes were BLAST-aligned against all sequenced genomes (excluding those of three closely related strains; see Materials and Methods), >99.5% of the 1,705 groups had homologs in other cyanobacterial strains. These genes were distributed among several cyanobacterial strains (Table S2 in the supplemental material), with the highest number (more than 85%) of homologues in *Microcystis* sp. and filamentous nitrogen-fixing strains. In contrast, few of these genes (less than 50%) had a homolog in members of the alpha-cyanobacterial group.

**TABLE 2 Ortholog comparison between pairs of *Cyanothece* genomes**

| *Cyanothece* strain | ATCC 51142 | PCC 7424 | PCC 7425 | PCC 7822 | PCC 8801 | PCC 8802 |
|---------------------|------------|----------|----------|----------|----------|----------|
| ATCC 51142          | 4,864      | 2,791    | 2,059    | 2,739    | 2,823    | 2,838    |
| PCC 7424            | 2,791(54)  | 5,163    | 2,273(44)| 3,707(60)| 2,711(52)| 2,700(52)|
| PCC 7425            | 2,059(44)  | 2,273(48)| 4,729    | 2,725(47)| 2,055(43)| 2,064(44)|
| PCC 7822            | 2,739(47)  | 3,707(64)| 2,275(39)| 5,787    | 2,743(47)| 2,740(47)|
| PCC 8801            | 2,823(68)  | 2,711(66)| 2,055(50)| 2,743(66)| 4,132    | 3,844(93)|
| PCC 8802            | 2,838(68)  | 2,700(65)| 2,064(49)| 2,740(66)| 3,844(92)| 4,181    |

*Paralogs excluded.*
Cyanothece 7425 is the most disparate among the six strains, with more than 2,300 unique genes (Fig. 2) representing 43% of its protein coding genes. Cyanothece 8801 and 8802 have the same geographical origin and are more than 90% identical to each other at the genome level, with a single large region of inversion (Table 2; Fig. 3). Since most of the genes are shared between these two strains, they have very few unique genes (183 in Cyanothece 8801 and 263 in Cyanothece 8802). Sixty-eight percent of the genes in Cyanothece 8801 and 8802 have homologs in Cyanothece 51142, and their close phylogenetic association is also reflected in their proximity in the tree (Fig. 1). Also in accordance with their position in the tree, Cyanothece 7424 is closest to Cyanothece 7822, sharing more than 60% of the genes in their individual genomes. Although Cyanothece 7822 has the largest genome, only 27% of the genes are unique to this strain. Interestingly, even though Cyanothece 7424 and 7425 have a common ecological origin (Table 1), unlike Cyanothece 8801 and 8802, their genomes are very diverse.

A BLAST analysis of the individual protein sequences from each homologous group showed that the top hits for these sequences are spread among a number of cyanobacterial strains, indicating differences in the evolutionary pathways of the six Cyanothece strains. Interestingly, for all strains except Cyanothece 7425, the top hits for more than 70% of the group of 1,705 homologous genes were mostly from Microcystis aeruginosa, Microcoleus chthonoplastes, and Synechocystis 6803. In contrast, top BLAST hits for protein sequences of Cyanothece 7425 (67% of the homologous groups) were mostly from Acaryochloris marina, Thermosynechococcus elongatus, Microcoleus chthonoplastes, Oscillatoria sp. strain PCC 6506, and Nostoc punctiforme.

In contrast, BLAST results for the protein sequences unique to Cyanothece 51142, 7424, 7822, 8801, and 8802, against the entire sequence database, showed that more than 50% of them did not have any significant hits in any other organism. Furthermore, the top BLAST hits for the remaining unique proteins in these Cyanothece strains were spread among several organisms and could not be associated with any specific organism, as was seen with the shared genes. About 40% of the unique genes in Cyanothece 7425 did not show any significant hits with any other organism. More than 10% of the unique genes showed top hits in A. marina and N. punctiforme, with the remainder spread among several organisms. About 10% of the unique genes in the Cyanothece strains have homologs in several nonoxygenic bacteria, many of which are diazotrophic strains.

Due to the genomic diversity observed in Cyanothece 7425, its distant location in the phylogenetic tree compared to the other Cyanothece strains, and its proximity to the three anaerobic...
nitrogen-fixing *Synechococcus* strains, we assessed its relationship to all sequenced cyanobacterial strains. BLAST analysis of all protein sequences of *Cyanothece* 7425 against those of all other sequenced cyanobacteria revealed that 1,885 of its 5,327 genes had homologs in the other five *Cyanothece* strains. In contrast, only 1,335 genes are shared with the three *Synechococcus* strains. Considering all sequenced cyanobacterial strains, *Cyanothece* 7425 shares the highest number of genes with *Nostoc* sp. strain PCC 7120 (49%), *N. punctiforme* PCC 73102 (49%), *Cyanothece* 7822 (49%), *Anabaena variabilis* ATCC 29413 (48%), *Cyanothece* 7424, and *A. marina* MBIC 11017 (46%).

**Metabolic traits of Cyanothece.** (i) Nitrogen fixation and hydrogen production. Our analysis revealed the presence of a large nitrogenase (*nif*) gene cluster in each of the sequenced *Cyanothece* genomes, thereby establishing this genus as a group of unicellular diazotrophic cyanobacteria. Comparison of the *nif* gene clusters in all sequenced diazotrophic cyanobacterial strains showed that the largest contiguous cluster is present in *Cyanothece* 51142, consisting of 35 genes arranged in two adjacent regulons (Fig. 4). The *nif* clusters in *Cyanothece* 8801 and 8802 are identical to each other and closely resemble the cluster in *Cyanothece* 51142, with only three missing genes: the molybdate ABC transporter permease protein gene *modB*, the hypothetical gene *nifK*, and the hypothetical gene between the ferrous iron transport protein gene *feoA*2 and *modB*. The synteny of this *nif* cluster is also largely maintained in *Cyanothece* 7424 and 7822, although it is somewhat shortened by gene losses in these two strains. In contrast, the nitrogenase cluster in *Cyanothece* 7425 is interrupted by a 2.5-Mbp insertion in the middle of the cluster, separating *nif-HDK* from *nifE*. Also, in contrast to the other five *Cyanothece* strains, which possess the *hup* genes, encoding an uptake hydrogenase, an enzyme associated with nitrogenase activity and nitrogenase-mediated hydrogen production, the *Cyanothece* 7425 genome does not have genes for this enzyme.

We assessed the abilities of the six *Cyanothece* strains to fix nitrogen and produce hydrogen. *Cyanothece* 51142 showed the highest nitrogenase activity, as well as the highest rates of hydrogen production, followed by *Cyanothece* 8802 and 8801 (Table 3). All *Cyanothece* strains except *Cyanothece* 7425 exhibited nitrogenase activity and hydrogen production capacity under aerobic in-
cubation conditions (incubation with air in the headspace). In contrast, *Cyanothece* 7425 exhibited nitrogenase activity and hydrogen production ability only when an anaerobic environment was provided (incubation with argon in the headspace).

A comparison of the nitrogenase cluster of all the sequenced nitrogen-fixing cyanobacterial strains revealed an unusual arrangement of the *nifVZT* regulon in *Cyanothece* 7425, closely resembling the cluster in two other anaerobic nitrogen-fixing strains, *Synechococcus* JA-2-3Ba and *Synechococcus* JA-3-3Ab, which exhibit an inversion between *nifV* and *nifE*. While the organization of this cluster is known to be largely conserved among all nonheterocystous nitrogen-fixing strains (21), an alteration in the arrangement of the regulons or an inversion/insertion in the region was also observed in other anaerobic nitrogen-fixing cyanobacterial strains investigated in this study (*Synechococcus* 7335, *Oscillatoria* PCC 6506, and *M. chthonoplastes*) (Fig. 4). Interestingly, as with *Cyanothece* 7425, the genomes of these other anaerobic nitrogen-fixing strains also do not possess any gene for an uptake hydrogenase.

(ii) Other metabolic characteristics of the genus *Cyanothece*.

(a) Photosynthesis. As expected from their ability for oxygenic photosynthesis, the genomes of all the sequenced *Cyanothece* strains contain most of the genes encoding the core cyanobacterial proteins related to photosynthesis (29). However, a BLAST analysis of the *Cyanothece* genomes with all the annotated genes in the KEGG database showed that certain low-molecular-mass proteins associated with PSI and PSII are missing in some of the *Cyanothece* strains. While all six genomes encode genes for biosynthesis of the light-harvesting pigment phycocyanin, *Cyanothece* 7424, 7822, and 8801 also have genes encoding phycoerythrin, a pigment which imparts a brownish-green color to these strains. The core cyanobacterial genes encoding chlorophyll biosynthesis enzymes, Calvin cycle enzymes, and regulatory proteins (29) are present in all six *Cyanothece* genomes. Interestingly, *Cyanothece* 7424, *Cyanothece* 7425, and *Cyanothece* 7822 have two very similar copies of the *psaB* gene, a trait shared by *Synechococcus* 7335 and *Nostoc azollae* 0708 among the 61 sequenced cyanobacteria. *Cyanothece* 7822 was the only strain to have the second *psaB* gene contiguous to the *psaAB* operon.

(b) Carbon metabolism. Members of the genus *Cyanothece* have been documented to synthesize and store large amounts of carbohydrates in the form of glycogen granules (30) when grown under light/dark cycles. BLAST analysis of the *Cyanothece* proteins against the KEGG database showed that several genes involved in glycogen synthesis, degradation, and metabolism are present in the core group of *Cyanothece* genes (see Table S2 in the supple-

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**FIG 3** Alignment of the genomes of *Cyanothece* 8801 and *Cyanothece* 8802. The two genomes were aligned using the Mauve software program (54), using its default parameters. A high level of sequence similarity is observed between the two strains, as shown by the colored portions of the aligned regions. However, the two genomes differ from each other due to a 1.3-Mbp-long inverted region spanning from 2.5 × 10^6 to 3.8 × 10^6 locations of the genomes.

**FIG 4** Alignment of clusters of nitrogen fixation-related genes in the six *Cyanothece* strains and in five other sequenced anaerobic nitrogen-fixing strains. The cluster in *Cyanothece* 7425 is significantly different from the clusters in the other *Cyanothece* strains, with a 2.5-Mbp insertion between *nifK* and *nifN*. The *nifVZT* regulon also shows an inversion similar to sequences in two other *Synechococcus* strains (*Synechococcus* JA-3-3Ab and *Synechococcus* JA-2-3B). In *Oscillatoria* sp. 6506, the *nifE* and *nifN* genes are pseudogenes, shown in white. *Microcoleus* has a shorter cluster with *nifB* translocated next to *nifN*. *Synechococcus* 7335 has a 50-kbp insertion between *nifV* and *nifB* and has *nifZ* translocated between *nifK* and *nifE*. 
mental material). A cluster of four genes involved in the metabolism of polyhydroxylalkanoic acid is present in *Cyanothece* 7424, 7822, and 7425 (CyAn7424_0494-0497, CyAn7822_1326-1330, and CyAn7425_4054-4057) and in *M. aeruginosa*. In addition, the *Cyanothece* genome encodes several genes involved in the synthesis and utilization of diverse sugar molecules. All the *Cyanothece* strains have genes for cellulose synthesis and metabolism. *Cyanothece* 7424 and 7425 encode genes for sucrose synthase, whereas *Cyanothece* 51142, 7424, 7822, 8801, and 8802 have genes for trehalose metabolism.

All of the sequenced *Cyanothece* strains have genes encoding enzymes for complete glycolytic and pentose phosphate pathways and an incomplete tricarboxylic acid (TCA) cycle. Noteworthily is the presence of a gene encoding phosphoenolpyruvate carboxylase in five of the six sequenced *Cyanothece* strains. This enzyme, involved in the gluconeogenic conversion of oxaloacetate to phosphoenolpyruvate, is not very common among other cyanobacteria, occurring only in *Arthrospira*, *Microcystis*, and *Synechococcus*. Our analysis revealed two unusual genes in *Cyanothece* 7424 and 7822, encoding isocitrate lyase (PCC7424_4054 and CyAn7822_2461) and malate synthase (PCC7424_4055 and CyAn7822_2460), enzymes that are involved in the glyoxylate shunt of the TCA cycle. No other sequenced cyanobacterial genome has genes for these two enzymes.

(c) Nitrogen metabolism. The six *Cyanothece* strains exhibit diversity in various nitrogen metabolism pathways. Cyanophycin, a nitrogen reserve molecule in cyanobacteria, is a polymer of arginine and asparagine. Catabolism of L-arginine can serve as a source of nitrogen, carbon, and energy for the cells (31). Differences are observed in this catabolic pathway, suggesting that the pathway fulfills diverse roles in these *Cyanothece* strains. Although all six strains have an arginine decarboxylase that catalyzes the conversion of arginine to agmatine, the fate of agmatine appears to differ. Like most cyanobacterial strains, *Cyanothece* 7822 and 7424 possess an agmatinase enzyme that converts agmatine into putrescine and urea. The genomes of these two strains have an operon of seven genes (the largest among all sequenced cyanobacteria) encoding urease and its accessory proteins (CyAn7424_4411-4417 and CyAn7822_2693-2699), as well as genes involved in the conversion of putrescine into spermine and spermidine. *Cyanothece* 7425, 8801, and 8802, in contrast, convert agmatine to putrescine via an agmatine deaminase and N-carbamoylputrescine amidase. *Cyanothece* 8801 and 8802 can further process putrescine into spermidine and spermine. The *Cyanothece* strain 51142 genome does not have genes that can convert agmatine to putrescine, suggesting that agmatine is the preferred polyamine for this strain. It also does not contain any gene for urea metabolism. Carbamate kinase, an unusual cyanobacterial enzyme involved in the production of ATP from ADP and carbamoyl phosphate in the final step of the fermentative degradation of arginine (32), is present in *Cyanothece* 7822, 8801, and 8802 and in *Synechocystis* 6803.

Other interesting differences in amino acid metabolism pathways include the presence of the kynurenine pathway of tryptophan degradation in *Cyanothece* 7822, 7425, 8801, and 8802 and the methionine salvage pathway in *Cyanothece* 7424, 7822, 8801, and 8802.

(d) Anaerobic metabolic capabilities. Our earlier studies have shown that *Cyanothece* 51142 exhibits high levels of anaerobic metabolism capacity (22, 25). In fact, all of the *Cyanothece* strains show several biochemical pathways associated with anaerobic metabolism. *Cyanothece* 51142, 7424, and 7822 have genes for fermentative lactate production, and both *Cyanothece* 7424 and 7822 perform mixed acid fermentation with formate as the end product (13). *Cyanothece* 7822 has been shown to have a capacity for mixed acid fermentation (12), a pathway also observed in the genus *Microcystis*. Pathways for ethanol, acetate, and hydrogen production are found in most of the *Cyanothece* strains. Also, an anaerobic chlorophyll biosynthesis pathway involving protoporphyrin IX cyclase (BChE) is present in *Cyanothece* 7425 and 7822. This gene has a homolog in the filamentous cyanobacterial strain *Cyanidioschyzon merolae* and in noncyanobacterial strains like *Helio- bacillus mobilis* and *Rhodopseudomonas palustris*. While a gene for BChE has been identified in *Synechocystis* 6803 (29), this gene has little sequence similarity with the *Cyanothece* gene. Many of the genes in the five *Cyanothece* strains (except *Cyanothece* 7425) had top hits to *M. aeruginosa* (>700 genes) and *M. chthonoplastes* (>260 genes), both of which are associated with anaerobic environments and have been extensively studied for fermentative processes. Furthermore, a significant number of unique genes found in each of the *Cyanothece* strains have homologs in either facultative or obligate anaerobic bacteria.

(e) Other novel aspects of *Cyanothece* metabolism. *Cyanothece* strains have acquired or retained diverse metabolic traits that make them interesting model organisms for studying various biological processes. For example, *Cyanothece* 8801 and 8802 differ from the other *Cyanothece* strains and most other cyanobacterial strains in possessing genes that encode a V-type ATPase (a six-gene operon, CyAn8801_3221-3226 and CyAn8802_2894-2899). This operon is also present in *Cyanobium* species and *Synechococcus* sp. strain WH 5701. It is also important to note that *Cyanothece* 8801 and 8802 have a number of genes encoding proteins involved in phosphonate metabolism. These include a three-gene operon encoding phosphonate transporters. Part of this operon is an amidohydrolase gene that is involved in phosphonate metabolism. In addition, the C–P lyase system involved in phosphonate metabolism is also present in these *Cyanothece* strains. In some ecosystems, phosphonates comprise a significant proportion of the available phosphorous (33), and consequently some strains might have evolved the capability to metabolize them.

Cytochrome P450s in cyanobacteria have been implicated in several metabolic processes involved in natural product synthesis, and members of the *Cyanothece* genus have been shown to be particularly enriched for some of these heme oxygenases (34). Our analysis revealed several unique cytochrome P450s in the *Cyanothece* strains, some with homologs in *A. marina*. In particular, the *Cyanothece* 8801 and 8802 genomes have several genes (CyAn8801_2436, CyAn8801_1896, CyAn8802_3674, and CyAn8802_1920) encoding these proteins, and interestingly, these strains also have large operons encoding nonribosomal peptide synthetase modules and related proteins (CyAn8801_3021-3032 and CyAn8802_3090-3101).

Most of the *Cyanothece* strains (except *Cyanothece* 7424 and 7822) also possess an alkane biosynthetic pathway involving aldehyde decarboxylase and an acyl-ACP reductase (cce_0778 and cce_0779, CyAn7425_0398 and CyAn7425_0399, PCC8801_0455, and PCC8801_0872, and CyAn8802_0468 and CyAn8802_0898) (35). Pathways involved in the nonfermentative synthesis of higher alcohols have also been identified in all the *Cyanothece* strains.
(iii) Characteristics of unicellular nitrogen-fixing strains. In order to identify the genes that might influence the metabolism of unicellular nitrogen-fixing cyanobacteria, we BLAST-aligned the group of 1,705 homologous genes common to all *Cyanothece* strains against the entire sequenced cyanobacterial genome database. Among these, 59 genes were identified that are common to all unicellular and filamentous N$_2$-fixing strains but are lacking in most (~<6 out of 39 strains) non-N$_2$-fixing strains. These included most of the core N$_2$ fixation-related genes from the *nif* cluster, several genes encoding regulatory proteins, and genes involved in energy metabolism. In addition, several hypothetical and conserved hypothetical proteins (see Table S2 in the supplemental material) are found to be restricted to the N$_2$-fixing group. Among this group of 1,705 homologous genes, 51 genes were found to be present exclusively in the unicellular N$_2$-fixing strains. Most of these gene products are hypothetical or conserved hypothetical proteins with domains implicated in regulatory functions. Some transporters and regulatory proteins are also found to be present only in the unicellular strains.

Earlier studies have shown that many of the genes are differentially regulated under diazotrophic growth conditions in *Cyanothece* 51142. We BLAST-aligned the genes known to be diurnally regulated in *Cyanothece* 51142 (22, 36, 37) against all sequenced cyanobacterial genomes. Our results show that the core nitrogen fixation genes (~15 genes) are present in both unicellular and filamentous nitrogen-fixing strains (see Table S3). Another ~50 of these diurnally regulated genes are mostly restricted to the unicellular N$_2$-fixing strains. These include genes encoding proteins with regulatory functions (transcriptional and translational regulators and two component system proteins), transporters, signaling proteins, peroxiredoxins, and peroxidases and several hypothetical and conserved hypothetical proteins.

**DISCUSSION**

Our analyses of the six completely sequenced *Cyanothece* genomes revealed that many key metabolic features were conserved during evolution, while considerable diversity was also gained (Fig. 2). The metabolic traits common to the six *Cyanothece* strains are shared by many other cyanobacteria, suggesting that they must have been acquired from an ancient ancestor and retained in the extant strains. The plasticity of the *Cyanothece* genomes is evident from the fact that the strains have acquired many novel metabolic capabilities, which is reflected in their diverse genotypes and phenotypes (such as cell size, shape, and pigment composition). Two of the *Cyanothece* strains possess linear chromosomal elements, a feature not observed in any other photosynthetic bacteria studied to date. These chromosomal elements seem to accommodate specific adaptive features that might impart niche-specific advantages to the strains, as is suggested by the presence of a large number of genes encoding transposons and CRISP-R-associated proteins.

The significant difference observed in the numbers of predicted coding sequences between the *Cyanothece* strains suggests a substantial amount of loss or gain of genetic material over evolutionary time. *Cyanothece* 8801 and 8802 possess the smallest genomes and have a high percentage of pseudogenes, indicating that they might be undergoing a reductive genome evolution. A high percentage of pseudogenes is also observed in the genome of *N. azollae*, a strain that has undergone significant gene loss to adapt to a symbiotic lifestyle (38). Despite their small genomes, *Cyanothece* 8801 and 8802 possess many novel genes that are missing in the other *Cyanothece* strains and in most sequenced cyanobacteria, suggesting that they must have been acquired in response to some selective pressure. An outstanding example is the presence of the V-type ATPases, involved in numerous energy transduction pathways and known to be indispensable for plant growth, especially under different stress conditions (39, 40). Another plant-like feature in *Cyanothece* is the presence of a two-gene operon in *Cyanothece* 7424 and 7822 encoding enzymes involved in the glyoxylate cycle. In plants this cycle is implicated in the conversion of storage lipids into carbohydrates (41). This cycle is also known to impart metabolic versatility to some bacterial strains. However, no other cyanobacterial strain sequenced to date is known to possess the glyoxylate cycle.

Our phylogenetic analysis showed that *Cyanothece* 7425 separated from the other *Cyanothece* strains at an early stage of evolution. *Cyanothece* 7425 cells are smaller than those of the other *Cyanothece* strains and are more cylindrical. A GC content of ~40% is characteristic of the genus *Cyanothece*, and *Cyanothece* 7425 is an anomaly in this regard as well, with a GC content of ~50%. In contrast to the other five *Cyanothece* strains, *Cyanothece* 7425 fixes nitrogen only under anaerobic conditions and appears to share a common ancestor with three other anaerobic nitrogen-fixing *Synechococcus* strains. In contrast to the *nif* gene cluster in the five *Cyanothece* strains, the cluster in *Cyanothece* 7425 is disrupted by the insertion of a large fragment and exhibits inversions similar to those of the clusters in the *Synechococcus* strains. However, our protein BLAST analysis did not show significant homology of the *Cyanothece* 7425 genes with those of any *Synechococcus* strain. Also, unlike the other *Cyanothece* strains, very few of the *Cyanothece* 7425 genes had homologs in *Microcystis* and *Microcoleus*. These results indicate that *Cyanothece* sp. PCC 7425 represents a cyanobacterial strain that is losing its nitrogen-fixing ability and evolving independently of the other *Cyanothece* strains.

Another interesting observation in this study is the absence of an uptake hydrogenase in all the sequenced anaerobic nitrogen-fixing cyanobacteria, which suggests that this enzyme must be associated with aerobic nitrogen fixation in nonheterocystous cyanobacterial strains. *Raphidiopsis brookii*, a strain that has lost the ability to fix nitrogen and has eliminated most of the nitrogen fixation related genes (42), also does not have genes encoding the uptake hydrogenase. *Cyanothece* 7425 is phylogenetically closest to *A. marina* and shares a common ancestor with this strain, indicating that the latter lost its nitrogen-fixing ability in the course of evolution. Similarly, *T. elongatus*, an unicellular nitrogen-fixing strain, located between two anaerobic nitrogen fixers, appears to have lost its nitrogen-fixing ability. These evolutionary trends suggest that strains that have not adapted for functioning under aerobic conditions may not succeed in a predominantly oxygen-rich environment and therefore lose this ability with the simultaneous elimination of the nitrogenase cluster. Therefore, the nitrogenase cluster of *Cyanothece*, which appears to have evolved to function efficiently under ambient conditions, is evolutionarily selected for, as is seen in strains like UCYNA and the endosymbiont of *R. gibba*.

*Cyanothece* cells are unique in their ability to provide a platform for both aerobic and anaerobic metabolic processes at alternate phases of the diurnal cycle. While the unicellular *Microcystis* cells also have the capability to create an anoxic intracellular environment, they do not have genes required for nitrogen fixation. Five *Cyanothece* strains exhibit high rates of nitrogenase-mediated
hydrogen production under aerobic conditions, indicating that an anaerobic intracellular environment is created to protect the oxygen-sensitive nitrogenase enzyme. The *Cyanotheca* genomes also contain many genes for catalases and peroxidases, enzymes which protect oxygen-sensitive cellular constituents required for anaerobic metabolism. A large, contiguous *nif* gene cluster and the ability to perform aerobic nitrogen fixation distinguish the unicellular *Cyanotheca* cells from all other cyanobacteria. The presence of versatile metabolic pathways, such as nitrogen fixation and oxygen photosynthesis, and the ability to generate anoxic cellular environments under diazotrophic growth conditions make members of the genus *Cyanotheca* attractive model systems for studying various sunlight-driven biofuel-yielding pathways which entail microaerobic conditions.

**MATERIALS AND METHODS**

**Genome annotation.** The *Cyanotheca* 51142 genome was annotated at Washington University in St. Louis, MO (21), whereas the genomes of the other five *Cyanotheca* strains were annotated at the Joint Genome Institute, U.S. Department of Energy. In these five strains, genes were identified using the Prodigal software program (43) as part of the Oak Ridge National Laboratory genome annotation pipeline, followed by a round of manual curation using the JGI GenePRIMP pipeline (44). The predicted CDSs were translated and used to search the National Center for Biotechnology Information (NCBI) nonredundant database and the UniProt, TIGRfam, Pfam, PRIAM, KEGG, COG, and InterPro databases. These data sources were combined to assert a product description for each predicted protein. Noncoding genes and miscellaneous features were predicted using tRNAscan-SE (45), RNAMMER (46), Rfam (47), TMHMM (48), and signalP (49).

**Intergenome BLAST analysis.** Sequences of protein-coding genes of six completed *Cyanotheca* genomes (*Cyanotheca* 51142, 7424, 7425, 7822, 8801, and 8802) were downloaded from NCBI (http://www.ncbi.nlm.nih.gov) (as of 14 October 2010). Homolog genes between different strains were identified using NCBI protein-protein BLAST analysis (BLASTP 2.2.22 [50]). Two genes are defined to be homologs to each other if their reciprocal BLAST hits resulted in the following: (i) an E value of <1E−4, (ii) a ratio between length of the BLAST hit region and length of the complete protein >2/3, and (iii) a ratio between the raw score for two-protein BLAST and the raw score for “self-self” BLAST >1/3.5. All BLAST runs were conducted with the additional parameters “-num_descriptions 99999 -num_alignments 999999 -comp_based_stats F -seg No” in order to ensure all relevant alignments are analyzed.

Based on these analyses, all genes in the six *Cyanotheca* strains could be associated with 11,607 homolog groups. Among them, 1,705 homolog gene groups are shared by all six strains and are defined as the core genome (see Table S2 in the supplemental material). In addition, unique genes in individual strains were also identified.

In order to identify the evolutionary history of the *Cyanotheca* family, these common and unique genes were BLAST-aligned against the NCBI nonredundant protein database. These BLAST runs excluded three additional cyanobacterial strains, namely, *Crocosphaera* 8501, *Cyanotheca* CCY 0110, and the uncultured cyanobacterium UCYN-A. Among these, the *Crocosphaera* 8501 and *Cyanotheca* CCY 0110 genomes are complete. Further, the draft versions of genomes of the two strains reveal that 64% (3,799/5,958) and 69% (2,009/6,475), respectively, of the probable protein-coding genes in these strains share homologs with one or more of the other *Cyanotheca* strains.

**Phylogenetic tree construction.** Orthologous sets of proteins were identified across 61 cyanobacterial strains through an all-versus-all BLASTP v2.2.23 (50, 51) analysis of their respective proteomes. Orthology was defined as reciprocal best-match hits between proteomes, matching 66% of the length of the longer of the two proteins, with scores 1/10 of the higher of the self-self scores. Any highest-scoring protein with multiple identically scoring hits was discarded. Sets of orthologs were considered to be conserved if ≥ 75% of proteins within the set were orthologous to one another, resulting in 226 sets of genes with orthologs in all 61 proteomes. Each of the 226 sets of 61 proteins was individually aligned using the MAFFT v6.811b software program (52) and then concatenated into a single alignment, removing all columns containing gaps. The PHYLIP v3.64 (5) software package was used to generate the final consensus tree using the Fitch-Margoliash method with 100 bootstraps, global rearrangement, and 1 jumble per bootstrap. Distances were then back fit to the resulting consensus tree using maximum-likelihood estimates from the original concatenated alignment. The resultant tree was rendered using the Archaeopteryx v0.957b software program (53).

**Hydrogen production and nitrogenase activity measurement.** Hydrogen production and nitrogenase activity were measured following the protocol published in the work of Bandopadhyay et al. (25).**KEGG pathway mapping.** In order to identify genes that may be involved in different metabolic reactions, individual protein-coding genes were BLAST-aligned against the KEGG pathway database (http://www.genome.jp/kegg/pathway.html). For each reaction, protein sequences of all currently annotated genes from different organisms were BLAST-aligned against the *Cyanotheca* genomes. Following the same criteria utilized to identify the homolog genes in intergenome BLAST analysis, genes were assigned to relevant KEGG reactions if they were homolog to any of the currently annotated genes in the KEGG.

**Genome alignments.** Whole-genome alignments were performed using “ProgressiveMauve” (54) with default parameter values.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at http://mbio.asm.orglookup supp doi:10.1128/mBio.00214-11/-DCSupplemental. Table S1, XLSX file, 0.1 MB. Table S2, XLSX file, 0.7 MB. Table S3, XLSX file, 0.1 MB.

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