Cyanobacterial diversity and activity in modern conical microbialites


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ABSTRACT

Modern conical microbialites are similar to some ancient conical stromatolites, but growth, behavior and diversity of cyanobacteria in modern conical microbialites remain poorly characterized. Here, we analyze the diversity of cyanobacterial 16S rRNA gene sequences in conical microbialites from 14 ponds fed by four thermal sources in Yellowstone National Park and compare cyanobacterial activity in the tips of cones and in the surrounding topographic lows (mats), respectively, by high-resolution mapping of labeled carbon. Cones and adjacent mats contain similar 16S rRNA gene sequences from genetically distinct clusters of filamentous, non-heterocystous cyanobacteria from Subsection III and unicellular cyanobacteria from Subsection I. These sequences vary among different ponds and between two sampling years, suggesting that coniform mats through time and space contain a number of cyanobacteria capable of vertical aggregation, filamentous cyanobacteria incapable of initiating cone formation and unicellular cyanobacteria. Unicellular cyanobacteria are more diverse in topographic lows, where some of these organisms respond to nutrient pulses more rapidly than thin filamentous cyanobacteria. The densest active cyanobacteria are found below the upper 50 μm of the cone tip, whereas cyanobacterial cells in mats are less dense, and are more commonly degraded or encrusted by silica. These spatial differences in cellular activity and density within macroscopic coniform mats imply a strong role for diffusion limitation in the development and the persistence of the conical shape. Similar mechanisms may have controlled the growth, morphology and persistence of small coniform stromatolites in shallow, quiet environments throughout geologic history.

INTRODUCTION

The macroscopic record of attached, laminated sedimentary rocks ‘accretionary from a point or limited surface’ (Semi-khatov et al., 1979) called stromatolites may record microbial interactions with sediments throughout Earth’s sedimentary history (Grotzinger & Knoll, 1999). Stromatolites are conspicuous features in many Archean and Proterozoic carbonate strata, but the relationships between microscopic processes that build individual laminae and the preservable macroscopic shapes of stromatolites are not well understood. Evidence that microbial growth, movement and orientation influenced the formation of stromatolite laminae is particularly elusive in progressively older stromatolites owing to the abundance of precipitated cements and recrystallization (e.g., Hofmann & Jackson,
Small conical stromatolites and tufts as old as 3.0 billion years stand out as a biologically influenced end-member of precipitated stromatolites with modern, distinctly biogenic analogs that grow only in the presence of cyanobacteria (Fig. 1, Walter et al., 1972, 1976; Grey, 1980; Beukes & Lowe, 1989; Buick, 1992; Jones et al., 2002; Bosak et al., 2009; Petroff et al., 2010). Given these morphological similarities, one may ask whether similar factors have been responsible for the growth and persistence of coniform mats throughout geologic history. Here, we explore these factors by examining cyanobacterial diversity, biomass density and temporal changes in cyanobacterial communities in modern coniform microbialites from Yellowstone National Park (YNP, Figs 1 and 2).

Microscopic examinations of modern conical mats focus primarily on thin filamentous cyanobacteria. Walter et al. (1976) uses microscopy and laboratory enrichment cultures to identify filamentous cyanobacterium Phormidium tenue var. granuliferum as the principal cone-building organism. Variably lithified filaments morphologically similar to P. tenue also can be seen in cones from YNP and New Zealand (Cady & Farmer, 1996; Jones et al., 2002), whereas unicellular organisms and Pseudoanabaena are more commonly observed in the surrounding mats (Walter et al., 1976). The true molecular, morphological and functional diversity of cyanobacteria in coniform mats likely exceeds these microscopic estimates. Even before the advent of molecular techniques, Weller et al. (1975) reported the formation of cones in a number of axenic cyanobacterial cultures but did not describe any phenotypic or morphological differences among these cultures. Recent molecular studies show the presence of multiple genetically distinct cyanobacteria in photosynthetic mats from hot springs (Ferris et al., 2003; Lacap et al., 2005, 2007; Lau et al., 2005; Ward et al., 2006; McGregor & Rasmussen, 2008; Boomer et al., 2009; Foster et al., 2009). A single study (Lau et al., 2005) analyzed the cyanobacterial diversity in cones, ridged cones and flat-topped cones from one of the outflows of Black Sands Pool in YNP and reported six cyanobacterial sequences in all cones: one Phormidium-like sequence, a sequence similar to unicellular Synechococcus, and four sequences with the closest similarity to uncultured cyanobacteria. Because the mats around cones contained only Synechococcus-like sequences that were distinct from those in cones, these authors suggested that cones and mats, as distinct components of coniform mats, contained different microbial assemblages (Lau et al., 2005).

It remains unclear how to marry the reported molecular diversity of cyanobacteria in coniform mats with microscopic diversity.

Fig. 1 Coniform mats in alkaline hot springs in Yellowstone National Park (YNP) (A, B) Photographs of coniform mats in small depressions and quiet ponds. (A) Dense cones. (B) Coniform mat consisting of cones (c) and surrounding mats (m). (C) Transmitted light micrograph of a vertical 30-μm-thick section through a cone showing thin dark laminae (arrow) and thick, porous laminae. Arrow points to the region magnified in (E). (D) Photograph of a YNP cone and the adjacent mat used in the 30-min labeling experiment. A similar cone was used in the 60-min incubation. Small divisions on the scale are 1 mm long. (E) Transmitted light micrograph of the vertical section of a cone showing aligned filaments in a thin lamina (arrow) and randomly oriented filaments in the porous laminae pl. (F) Transmitted light micrograph of the vertical section through the mat showing heavily lithified thin laminae (dark, Si grains) and empty regions. The presence of Si was confirmed by energy-dispersive X-ray spectroscopy.

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observations. Namely, filamentous cyanobacteria are commonly seen in mats (e.g., Walter et al., 1976), but only sequences similar to unicellular *Synechococcus* are reported from the mats around cones (Lau et al., 2005). It also remains unclear whether only one organism drives the formation of coniform mats in YNP, or whether these structures contain multiple genetically distinct organisms capable of vertical aggregation. Here, we address these questions by:

- analyzing the diversity of cyanobacteria in cones and mats in multiple ponds around four different hot springs
- characterizing the ability of different cyanobacteria to form vertical structures in laboratory enrichment cultures
- measuring the activities and the densities of cells at the topographic highs and lows of natural coniform mats in YNP.

Our findings reveal that modern coniform mats contain multiple, genetically distinct filamentous and unicellular cyanobacteria and that these cyanobacterial communities vary in space and time, contributing to the interannual persistence of coniform structures. At any given point in time, the growth, degradation and lithification of microbes in coniform mats depend on the position within the three-dimensional topography.

**MATERIALS AND METHODS**

**Sample collection**

Samples of conical stromatolites used in laboratory enrichments, field incubations with labeled carbon, microscopic analyses and a subset of molecular analyses were collected in one pond of Mound Spring in YNP in September 2008. Cones that were incubated with labeled carbon were not used in molecular analyses, but adjacent cones from the same pond were sampled and analyzed instead. Most samples for comparative molecular analyses were collected in August 2009 as pairs of cones and adjacent mats (Fig. 1B, D) from 14 ponds fed by the outflow channels of four different hot springs in the Sentinel Meadows thermal area: Mound Spring, Flat Cone Spring, Bison Pool, and the Buggers (Table 1, Fig. 2). Some of these samples were also analyzed by transmitted electron microscopy (TEM, Fig. 3D). All field incubations and sample collections were performed under permit (#YELL-2008-SCI-5758 for the sampling of coniform mats in Sentinel Meadows and Fairy Creek). All samples were excised in duplicates by a sterile surgical blade and placed in sterile Eppendorf tubes, stored on dry ice during field collection and transport, and stored at $-80^\circ$ C in the laboratory. Flow in ponds in 2009 was monitored by placing the floating lid of a 15-mL BD FalconTM tube (BD Biosciences, San Jose, CA, USA) on the surface of the liquid and measuring the distance that the lid travelled in time.

**Light and epifluorescence microscopy**

Samples were fixed with 2.5% glutaraldehyde in 0.1 m sodium cacodylate buffer (pH 7.4) for 2 h. Buffer-washed samples were embedded in Sakura TissueTek O.C.T. compound (VWR, West Chester, PA, USA), frozen for at least 2 h and sectioned to a thickness of 10–30 μm using a cryo-stat (Leica, Wetzlar, Germany). Thin sections were washed with buffer and imaged using an Axio Imager M1 epifluorescence microscope (Carl Zeiss, Thornwood, NY, USA) with differential interference contrast and phase contrast modules. LSM510 confocal microscope with quasar spectral detection (Carl Zeiss) was used to distinguish
between chlorophyll autofluorescence (CY3 filter, excitation 558 nm and emission 568 nm) and the orange autofluorescence of phycoerythrin (FITC filter, excitation 490 nm and emission 520 nm).

### Genomic DNA extraction

Genomic DNA from approximately 0.5 g of all samples was extracted using UltraClean soil DNA isolation kit...
DENITRIFYING MIXTURES CONTAINED (PER 50 L)
72 CYCLES WITH 2 MIN EXTENSION TIME FOLLOWED BY NINE CYCLES
15 CM
9°
DEL MAR, CA, USA) ON A GEL (0.75 MM THICK, GEL ELECTOPHORESIS (DGGE) SYSTEM (C.B.S. SCIENTIFIC, VALENCIA, CA, USA) AND RE-AMPLIFIED IN A NESTED PCR USING PRIMERS SPECIFIC TO CYANOBIOTIA: FORWARD PRIMER 359F (GGG GAA TYT TCC GCA ATG GG) WITH A GC CLAMP (CCG CCG CCG CCC GCC GCG CGG GTC CCG CCG CCG CCC CCG GGG G) AND AN EQUIMOLAR MIXTURE OF REVERSE PRIMER 781R(a) (GAC TAC TGG GGT ATC TAA TCC CAT T) AND 781R(b) USING THE AMPLIFICATION CONDITIONS DESCRIBED PREVIOUSLY. PCR PRODUCTS WERE RE-ANALYZED BY DGGE TO CONFIRM SINGLE-BAND RECOVERY, EXCISED FROM THE GEL AND PURIFIED WITH THE QIAQUICK PCR PURIFICATION KIT (QIAGEN) PRIOR TO SEQUENCING.

PHYLLOGENETIC AND OCCURRENCE DATA ANALYSES


TO ANALYZE THE OCCURRENCE OF VARIOUS OTUS IN SAMPLES FROM DIFFERENT PONDS AND HOT SPRINGS, UPGMA HIERARCHICAL CLUSTERING ANALYSIS WAS PERFORMED WITH AVERAGE LINKAGE, AND JACCARD DISTANCE (LEGENDRE & LEGENDRE, 1998) USING THE HIERARCHY MODULE OF THE SCIPY PACKAGE (V. 0.9.0) OF THE PYTHON PROGRAMMING LANGUAGE (JONES ET AL., 2001). TO
identify cone- or mat-specific OTUs, the statistical significance of the association of OTUs with morphology or temperature was assessed separately for each OTU using a two-sided Fisher's exact test (Agresti, 1990), as implemented in the statistical package of the R programming language (v. 2.11.0), www.r-project.org. In this analysis, temperature values were discretized to ‘hot’ (≥ 30 °C) or ‘cold’ (<30 °C). Additionally, the dependence of the distribution of OTUs on continuous temperature values was assessed using logistic regression. Linear logistic regression was also used to test the dependence of distribution of OTUs on temperature. Dendrograms were generated using UPGMA with average linkage.

Measurements of photosynthetic rates

Oxygen concentrations and photosynthetic rates were determined using a Clark-type oxygen microsensor (OX-25 and OX-50; Unisense, Aarhus, Denmark) with a guard cathode (Revsbech, 1989). Linear calibration of microsensors was performed using air-saturated and anoxic pond water. The diffusive flux of oxygen was calculated from Fick’s first law of one-dimensional diffusion (Jørgensen & Revsbech, 1985):

\[ J_o = \frac{D_e \cdot \frac{dC_o}{dz}}{C_o} \]

where \( D_e \) is an effective diffusion coefficient and \( \frac{dC_o}{dz} \) is the slope of the microprofile at depth \( z \). 
De in the bacterial colony was assumed to be 2.0 \( \times 10^{-6} \) cm² s⁻¹ (Lassen et al., 1998; Grotzschel & de Beer, 2002). Net photosynthetic rates in individual layers were determined from the balance between oxygen fluxes through the overlying and underlying areas. The gross photosynthetic rate was measured using oxygen microelectrodes and the light/dark shift method (Revsbech & Jørgensen, 1983).

Enrichment cultures

Castenholz-D (Castenholz, 1988) agar plates (2%) were scored with parallel lines using dry calcium alginate swabs (Burton & Lee, 1978; Vaara et al., 1979), inocula from Mound Spring 2008 were placed at the beginning of each scored line and incubated for approximately 24 h at 45 °C in a light gradient (from 13.9 to 12.8 µm photons m⁻² s⁻¹) to enrich for phototactic, gliding cyanobacteria. The migration of filaments was monitored by epifluorescence microscopy as described previously. Single filaments that had glided along the scored path were excised by a sterile surgical blade and cultured in modified liquid Castenholz-D medium at pH 8 (Bosak et al., 2009) containing autoclaved silica sand (VWR International, Radnor, PA, USA). This process was repeated 25 times. The liquid cultures were grown with a fluorescent cold light source with a 12-h day, 12-h night cycle and the medium was exchanged periodically. The purity of cultures was tested by plating the culture onto 100% LB, 10% LB, and glucose/fructose (5 mM final concentration) agar plates, microscopically, and by the amplification of a single 16S rRNA gene from biofilms grown in the liquid medium using universal bacterial primers 27F (5’-AGA GTT TGA TGG CTG CTC AG-3’) and 1492R (5’-TAC GGY TCT TAC CCP TAC GAC ACT Y-3’) (Lane, 1991). The resulting enrichments were characterized by microscopy and by a series of growth experiments at various light intensities that encompassed measurements of chlorophyll A and visual characterization of cyanobacterial aggregation. Cyanobacteria enriched by this process are described in Figs 5 and 6 as XAN-1, 8, 12, and FYG.

Carbon isotope labeling

Individual cones and approximately 1 cm wide surrounding mats (as determined in Fig. 1D) were labeled in YNP using a modified Castenholz-D medium (Castenholz, 1988; Bosak et al., 2009) amended with sodium bicarbonate-\(^{13}\)C (98 atom%, Isotec, Inc., Miamisburg, Ohio). The total \(^{13}\)CO₂ was 2.58 mmol per labeling in 100 mL of the incubation medium. All samples were incubated in YNP from 1 to 2 pm at ambient light intensity of approximately 1500 µE m⁻² s⁻¹, in the host spring at ambient pond temperature (42 °C).

Transmission electron microscopy (TEM)

Isotopically labeled and unlabeled control samples were fixed by 2.5% glutaraldehyde in 0.1 m sodium cacodylate buffer (pH 7.4), washed with buffer, and post-fixed in 1% osmium tetroxide and 1.5% potassium ferrocyanide. The fixed samples were dehydrated through a gradient series of acetone/water solutions and embedded in epoxy resin (Epon). Ultrathin sections (70–100 nm) and semi-thin sections (approximately 500 nm) were cut from the resin blocks by using a Reichert-Jung Ultracut E ultramicrotome (Reichert-Jung AG, Vienna, Austria) with a Diatome diamond knife (Diatome AG, Biel, Switzerland). The ultrathin sections were transferred onto 200-mesh Cu TEM grids with Formvar-carbon support film for NanoSIMS-TEM correlation imaging. TEM was performed by a FEI Tecnai 12 TEM (FEI Company, Eindhoven, the Netherlands) equipped with a Gatan Bicoscan CCD camera Model 792 (Gatan, Inc., Pleasanton, CA, USA). An accelerating voltage of 120 kV was applied in bright-field mode.

Secondary ion mass spectrometry (NanoSIMS) mapping

Secondary ion mass spectroscopy (SIMS) imaging was performed using a NanoSIMS-50 Ion Microprobe ( Cameca,
Gennevillieres, France) operating in scanning mode. The semi-thin sections (approximately 500 nm) were cut at the same time as the ultrathin sections for observation in TEM, providing complementary chemical information by SIMS to the high spatial resolution structural information by TEM.

A tightly focused, high-energy primary ion beam of Cs+ bombards the sample surface and generates negative secondary ions during the collision cascade. These secondary ions are directed to a magnetic mass spectrometer that ensures high transmission with a high mass resolution. The parallel detection system allows the simultaneous recording of up to five ion species, permitting reliable comparisons between different ions, especially between isotopic ions from the same microvolume. The primary beam, operated in probe scanning mode, steps over the surface of the sample to create images of the selected ion species. In our study, survey mosaic secondary ion images were generated as 256 × 256 pixel rasters of 60 µm × 60 µm pixels, with the probe intensity of 5 pA and a typical probe size of 150 nm. To obtain high-resolution images of intracellular details, the raster was reduced to 15–30 µm, with the probe size smaller than 100 nm and an intensity of about 1 pA. Images were created for the following: 12C, 13C, 12C–14N, 13C–14N, and 32S. Figures S2 and S3 provide examples of survey mosaic images and position-dependent high-resolution scans. The dwell time per pixel was 15 ms for survey images and 30 ms for high-resolution images. High mass resolution separated 13C–14N– from the isotopic 12C–15N– species (Guerquin-Kern et al., 2005) and prevented the mass interference from 11B–16O– species. Compared with ion species 12C–, 13C–, 12C–14N– and 13C–14N– have higher intensities and are therefore both more statistically reliable and better able to indicate carbon associated with biomass. Thus, the ratio (%) of 13C–14N–/12C–14N– was used to track the relative carbon uptake (enrichment).}

Color ratio images were generated in OpenMIMS, an ImMaGi (Rasband, 1997–2006) plugin developed by Claude Lechene’s Laboratory. OpenMIMS allows the visualization of ratio value in a HSI (Hue-Saturation-Intensity) image (Benson, 2003; Poczatek et al., 2009).

To calculate the average 13C–14N–/12C–14N– ratio in the cells in each 60 µm × 60 µm survey image, the information pertaining to the microbial biomass (enriched in 13C) has to be distinguished from the background resin (natural 13C abundance). The biomass has a much higher CN– ion signal than the resin, allowing the generation of a mask by adjusting the threshold level on the sum image of 13CN– and 12CN–. The pixels in the resulting binary mask image have values of 1 or 0, corresponding to areas with and without cellular biomass, respectively. The total number of pixels with the value of 1 in the binary mask image was used as a measure of total cellular biomass. This binary mask image was multiplied by the measured 13CN–/12CN– ratio image to generate a 13CN–/12CN– ratio map, which eliminated the resin background. This biomass ratio map was used to calculate the net carbon uptake: the ratio characteristic for the unlabeled controls (N = 3, numerical value 1.25%) was first subtracted from each pixel value, and pixel values in thus corrected ratio of 13CN–/12CN– were integrated over the entire 13CN–/12CN– ratio map. To calculate the 13C enrichment in individual filaments, we manually outlined individual filaments in the biomass ratio map and integrated the pixel values within the outlined area (ROI, region of interest) after the subtraction of the unlabeled control value (1.25%). Only distinctly filamentous cells were selected for this analysis to avoid bias caused by the random truncation during sectioning.

RESULTS

Environmental parameters at sampling sites

Coniform mats in YNP form in quiet ponds and other surface depressions where the flow is low (such as buffalo steps) and the pH is alkaline (pH > 8) (Fig. 1A,B, Table 1). The water level in the outflow channels in 2009 was low, periodically exposing some mats and covering others by only a thin film of water (Table 1). The temperature of the water around sampled coniform mats varied from 22.0 to 51.3 °C (Tables 1 and 2) and the flow on the surface was detectable only in two ponds in Mound Spring (44.6 and 55.3 °C). Coniform mats around Mound Spring in September 2008 were submerged under 2–4 cm of warm water (40–45 °C).

Morphology and ultrastructure of cyanobacteria in cones and mats

Filaments in the dense thin laminae of cones were nearly parallel to the growth surface (Fig. 1C,E), whereas those in the porous thick laminae were sparsely distributed and randomly oriented with respect to the growth surface (Fig. 1C,E). Thick, porous laminae were rare or absent from the more heavily lithified adjacent horizontal mats (Fig. 1F), where many degraded cells and cells encrusted by silica were present (Figs S1–S3).

Cones sampled in 2008 contained various morphologically distinct cells that exhibited chlorophyll a autofluorescence (Fig. 3A), as well as <0.8-µm-thick filaments similar to those of Roseiflexus- and Chloroflexus-like organisms (Figs 3A and S4; Pierson & Castenholz, 1974; Pierson et al., 1985; Hanada et al., 2002). The most abundant cyanobacteria in the thin and thick laminae of cones were >100-µm-long filaments composed of 0.8–1 µm wide and 2–3-µm-long rectangular cells. These cells contain 3–4
peripherally arranged thylakoids, approximately 150–750-nm-wide subterminal granules (Nash, 1938) and cell walls that were commonly surrounded by fibrous mucilage. The terminal cells in some filaments were spear shaped (Figs 3C and 4A). These cells were less common in the tips of cones from Mound Spring that were sampled in 2009. YNP cones from 2008 and 2009 also contained sausage-shaped 0.9–1.2-μm-wide and 3–5-μm-long cells with 3–4 peripherally arranged thylakoids, intracellular granules (150–300 nm in diameter) and cell walls surrounded by fibrous mucilage (Fig. 3D). These sausage-shaped cyanobacteria, similar to Synechococcus sp. (e.g., Jones et al., 1998; Cannon et al., 2001; Castenholz, 2001), were very abundant in the intermittently exposed tips of cones from 2009 (Fig. 3D), and rare in the tips of permanently submerged cones from 2008. The silicification of the sticky matrix between these cells in cone tips (Fig. 3D) was likely promoted by the evaporation in the thin film of water or during periodic exposure (Table 1), but was still less heavy than the silicification in mats sampled in 2008. The approximately 0.8-μm-wide filaments and the 1 × 3 μm, sausage-shaped cells actively took up inorganic carbon (Fig. 4) and lacked the orange autofluorescence of phycocyanin. This autofluorescence was instead observed within some relatively rare, 2-μm-wide oval cyanobacteria morphologically similar to Synechococcus (Fig. 3E). These oval or round cyanobacteria occurred only individually in cones, primarily on the very surface, but were more common in the adjacent topographic lows (Figs 4C, D and S2). Numerous ghosts of lysed cells and silica-encrusted cells were present in the topographic lows (Figs 1 and S1–S3). More extensive degradation of cells in mats also was reported by Walter et al. (1976).

### Molecular diversity of cyanobacteria in cones and mats

Sequences of cyanobacterial 16S rRNA genes in field samples of coniform mats from four different thermal features and from two different years attested to the contribution of a number of genetically distinct cyanobacteria to the construction and the interannual persistence of modern conical stromatolites (Fig. 5). The 33 samples from 2009, each consisting of a cone and the adjacent mat, contained a total of 18 different operational taxonomic units (OTUs) forming two clusters similar to filamentous Leptolyngbya (Fig. 5). Also present were 13 OTUs similar to the 16S rRNA genes of cyanobacteria from cultured representatives of Synechocystis and Synechococcus genera (Fig. 5). The PCR amplification method using primers specific to cyanobacteria also detected an unaffiliated OTU (OTU 6) similar to candidate division SR1 (Fig. 5, Davis et al., 2009) and Acidobacteria (not included in the phylogenetic tree).

Table 2 summarizes the observed distribution of OTUs in the sampled ponds.

<table>
<thead>
<tr>
<th>OTU cluster</th>
<th>OTU</th>
<th>Number of ponds in each spring where OTU was present</th>
<th>OTU temperature range (°C)</th>
</tr>
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<tr>
<td>Subsection III filamentous, non-heterocystous</td>
<td>1</td>
<td>MS(5), BG(3), BS(1)</td>
<td>30.0–44.6</td>
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<tr>
<td></td>
<td>2</td>
<td>BG(5)</td>
<td>30.0–42.1</td>
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<tr>
<td></td>
<td>3</td>
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<td></td>
<td>4, 9, 29</td>
<td>FC(1)</td>
<td>22.2</td>
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<td></td>
<td>5</td>
<td>BS(1)</td>
<td>35.4</td>
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<td>18</td>
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<td>22.2–51.3</td>
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<td>32</td>
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<td>22.2–51.3</td>
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<td></td>
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<td>28</td>
<td>BS(2)</td>
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<td></td>
<td>31</td>
<td>MS(3)</td>
<td>32.5–51.3</td>
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<td>7</td>
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<td></td>
<td>8</td>
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<tr>
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<td>12</td>
<td>MS(5), BG(3), BS(2), FC(1)</td>
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<td>14</td>
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</table>
S5 and S6. Cones contained 2–10 OTUs from cyanobacterial Subsection III (‘filamentous, non-heterocystous’; Castenholz, 2001) and 0–3 of the ‘unicellular’ Synechococcus- or Synechocystis-like OTUs from Subsection I (Castenholz, 2001), whereas the adjacent mats contained 0–5 OTUs from Subsection III and 0–5 unicellular OTUs (Fig. S6). Eight OTUs from the larger Leptolyngbya-like Subsection III cluster, (OTU 18–24, 32), occurred in samples from all hot springs (Table 2). Two of these ubiquitous OTUs (19, 22) occurred primarily in cones ($P = 0.0001$, Fisher’s exact test, Fig. S6) and four (20, 21, 23, 24) almost always occurred together in cones and mats (Fig. S5, Jaccard similarity indices higher than 0.7). In fact, the presence or absence of these four OTUs defined the two major spatial clusters of cyanobacterial diversity (Fig. S5). The larger Subsection III cluster also contained sequences that were detected only around individual hot springs (Table 2). Various motile filamentous cyanobacteria from our laboratory enrichment cultures grouped with this cluster. Some cyanobacteria in these laboratory enrichment cultures (XAN-1 and XAN-12) formed cones under all growth conditions, while others (XAN-8) did not (Fig. 6). Also present in nearly all ponds was a sequence (OTU 1) from the smaller Subsection III cluster (Fig. 5, Table 2) that occurred primarily in cones ($P = 0.001$, Fig. S6). This sequence was similar to a filamentous cyanobacterium forming porous tubes and films in our laboratory enrichment cultures (FYG, Figs 5 and 6), and to the only OTU from Subsection III that was detected in coniform mats around Black Sands Pool (Lau et al., 2005). Other sequences from this cluster (OTU 2, 3, 9) were detected both in mats and in cones, but only in individual hot springs (Table 2, Fig. S5).

Operational taxonomic units similar to 16S rRNA gene sequences of unicellular cyanobacteria were absent from more than 50% of the cones from 2009, but were detectable in 75% of the mats (all data shown in Fig. S6). These OTUs varied spatially, with only two detected in more than one spring (OTU 7, 11) and eight detected only in single samples of mats or cones (Table 2, Fig. S5). OTU 11, exhibiting the highest similarity to Synechococcus C9 and to a sequence previously reported in the mats surrounding YNP cones (Lau et al., 2005; Fig. 5), was detected primarily in mats ($P = 0.0001$, Fig. S6). OTUs 12–17 are most similar to Synechococcus P1 and to sequences previously detected only in mats surrounding the YNP cones (Lau et al., 2005). In contrast to the previous report, we detected at least some of these OTUs in cones (e.g., OTU 12, 13). Other unicellular OTUs (25–27) were similar to Synechococcus sp. clone JulG-B6 and Synechocystis (OTU 7, 8), and were present in both mats and cones.

All major cyanobacterial clusters from 2009 were also represented by 16S rRNA sequences detected in five
samples of cones and mats from one pond fed by Mound Spring in 2008 (Fig. 5). The samples from 2008 contained sequences from the larger Subsection III cluster, one sequence from the smaller Subsection III cluster and three sequences similar to those of unicellular cyanobacteria from Subsection I (Fig. 5). However, none of the field samples from 2008 contained the co-occurring OTU 20, 21, 23, 24, although these OTUs were present in all examined ponds in 2009 (including Mound Spring). Furthermore, unicellular sequences were detected only in one of five mats sampled in 2008. Therefore, similar core cyanobacterial clusters contributed to the growth of cones in 2008 and 2009, but individual sequences representing these clusters in Mound Spring varied between the 2 years. Factors responsible for these variations remain unknown and may include both lower temperatures of some ponds in Mound Spring in 2009 (Tables 1 and 2), and different hydrological conditions in Mound Spring between 2008 and 2009 (Table 1).

Cyanobacterial activity in cones and surrounding mats

To quantify the differential accumulation of biomass in mats and cones, we examined the photosynthetic activity at the approximately 100-μm scale using microelectrodes. The highest net photosynthetic rates in coniform mats occurred in the upper 0.5 mm of the cones (Fig. 7). The ratio of respiration to gross photosynthesis was

Fig. 5 Operational taxonomic units (OTUs) of cyanobacteria and environmental samples from coniform mats in Yellowstone National Park (YNP) were inserted into pre-established phylogenetic tree of full-length 16S rRNA gene sequences in Greengenes database in ARB. Numbers on branches indicate bootstrap values obtained by RAxML. Sequences from denaturing gradient gel electrophoresis (DGGE) bands were grouped at 97% sequence similarity threshold. Sequences from enrichment cultures were left as unique sequences. Green and blue text represents sequences of cyanobacteria that do or do not form cones, respectively, in laboratory enrichment cultures. Boldface OTU 5 represents cyanobacterial sequences from samples collected in 2009, whereas 2008_MS and DGGE sequences represent sequences from samples collected in 2008. Italicized text (gel band) are cyanobacterial sequences previously identified in coniform mats (Lau et al., 2005). Accession numbers of GenBank entries are listed after the names of the corresponding organisms, gel bands or clones. Yellow panel denotes Synechococcus-like sequences, green panel denotes filamentous Leptolyngbya-like cluster, and blue panel denotes unicellular sequences similar to Synechocystis. Only bootstrap values larger than 50% are shown.
that likely represent some form of carbon concentrating granules (Fig. 3C,D). Indeed, after a 30-min incubation with $^{13}\text{C}$ bicarbonate, these granules were the most prominent loci containing detectable labeled carbon in the uppermost 120 μm of the cone (Fig. 4A). Some label was also present in the filament sheaths at 130 μm below the cone tip (Fig. S7A), but was absent from most other filamentous and sausage-shaped cells in the cone and in the mat. The less abundant, round and oval cyanobacterial cells in the mat exhibited a faster response to the influx of carbon than the immediately adjacent filamentous and sausage-shaped cells, because they contained uniformly distributed $^{13}\text{CN}^{-}$ even after 30 min (Fig. 4C). The rather slow response of the abundant filamentous and sausage-shaped cells to the influx of labeled carbon did not reflect the time required to synthesize the carbon concentrating granules, because abundant, morphologically identical granules were present in the cells from untreated and unlabeled samples (Fig. 3C,D).

After a 60-min incubation, the label was distributed throughout filamentous and elongated cells in cones and mats (Fig. 4B). Most new inorganic carbon was present in the 100-μm-thick region below the uppermost 90-μm layer of the cone (Fig. 8A). The uptake of $^{13}\text{C}$ between 90 and 150 μm in the cone was 1.5 times higher than the uptake at the very surface, and more than four times higher than the uptake below the top 0.1 mm of the mat (Fig. 8A). This was owing primarily to the large density of active cells, rather than to the higher activity of individual cells in the tip of the cone (Fig. 8B). The uptake of labeled carbon below the surface of the cone decreased with depth (Fig. 8A). This profile arose from the decreasing density of active cells within the upper 500 μm and not from the decreasing activity of individual cells with depth (Fig. 8B, C). Cells in the uppermost 90 μm of the cone departed from this trend, because individual filaments incorporated lower amounts of label and the overall cell density was lower than in the underlying region (Fig. 8B, C).
All parts of mat contained less cellular biomass than the cone and the amount of labeled carbon in cells in the mat decreased with depth, with the possible exception of the very surface (Fig. 8A,B). Active round cells (Fig. 4D) accounted for the threefold increase in the amount of labeled carbon and biomass at approximately 100 μm below the surface (Fig. 8A,B), but similar cells were rare in another isotopically examined mat from the same pond incubated for 30 min. Thus, the topographic lows were laterally heterogeneous, supported smaller active populations of cyanobacteria and exhibited visual and chemical evidence of organic degradation and heavy silicification (Figs 1, 7 and S1–S3).

**DISCUSSION**

**Microbial communities in cones and mats**

The shapes and the internal textures of modern conical microbialites ultimately preserve little information of the instantaneous molecular diversity and the complexity of microbial interactions. These interactions, however, ensure the persistence of stromatolite-forming communities over timescales necessary to attain macroscopic sizes and turn to rock. Our study reveals a previously unrecognized and spatiotemporally variable composition of cyanobacterial communities in coniform mats in YNP, demonstrates different cellular activities in topographic highs and lows, and hints at factors responsible for the persistence of coniform mats over seasonal and yearly timescales. The molecular diversity of these cyanobacteria varies among cones and mats within the same pond, among different ponds and hot springs (Table 2), and between two successive sampling years (Fig. 5). In the broadest sense, these findings are consistent with the previous reports of morphologically similar filamentous cyanobacteria in cones and mats (Walter et al., 1976; Brock, 1978; Cady & Farmer, 1996; Jones et al., 1998, 2002), and of morphologically more diverse cyanobacteria in the mats (Walter et al., 1976; Brock, 1978).

Unlike previous studies, the findings presented here suggest that cyanobacteria responsible for the formation of cones in YNP cannot be described as a single species (Walter et al., 1976), characterized by a single 16S rRNA sequence from Subsection III (Lau et al., 2005), or by an unchanging group of cyanobacterial 16S rRNA sequences (Lau et al., 2005).

Cyanobacterial 16S rRNA sequences from coniform mats around four different hot springs in Sentinel Meadows in 2009, a single hot spring sampled in 2008 and 2009, and from another thermal area (Lau et al., 2005) form clusters. As a whole, these clusters are well represented across space and time. However, the diversity of sequences within individual clusters and samples may, in part, depend on the hydrological conditions. These
conditions would stress communities in ponds with oscillating or rather low temperatures (Tables 1 and 2) and communities in temporarily exposed mats (Rothrock & García-Pichel, 2005). Coniform mats are generally absent from ponds colder than approximately 30 °C and warmer than approximately 55 °C (e.g., Walter et al., 1976; Jones et al., 2002) and Synechococcus-like unicells are the dominant cyanobacterial ecotypes above approximately 48 °C (Ferris & Ward, 1997). Therefore, one may expect different cyanobacterial populations in coniform mats exposed to temperatures below 30 °C and above 45 °C than in coniform mats growing permanently within this temperature range. Our current dataset does not support statistically significant relationships of any OTUs with temperature within the observed range (22–51 °C). Some possible differences among samples at the lower and the higher end of this temperature range include the presence of three unique OTUs from Subsection III in the coldest pond (Table 2), the absence of cone-specific and ubiquitous OTU1 at the lowest and the highest temperatures, respectively, and the absence of otherwise ubiquitous OTUs 20, 21, 23, 24 from the warmest pond (Table 2, Fig. S5). At present, these differences remain speculative because the dataset contains samples from only one pond below 30 °C and from only one pond above 47 °C, respectively. Analyses of more samples from temperatures higher than 48 °C and lower than 30 °C, and from more hydrologically stable hot springs elsewhere in YNP could test these suggested trends in a statistically meaningful manner.

The sampling and processing methods used here average the cyanobacterial diversity over entire cones and entire mats, painting a picture in which cyanobacteria with similar or identical 16S rRNA sequences exhibit measurable topography-dependent differences in growth, activity and aggregation (Figs 8 and S6). This picture is consistent with laboratory observations of cone formation in the cultures of filamentous cyanobacteria that were enriched from field cones (XAN-1 and XAN-12 in Fig. 6). These cyanobacteria grow at different rates in cone tips, between mm- to cm-tall cones and on cone sides (Petroff et al., 2011), giving rise to laminated pinnacles in highly enriched laboratory cultures. Other thin and motile filamentous cyanobacteria enriched from field cones appear to be incapable of initiating cone formation under identical growth conditions (XAN-8 in Fig. 6), although their 16S rRNA sequences are very similar to those of known cone formers (XAN-1) and various field OTUs (Fig. 5). Therefore, the diversity of field OTUs likely reflects a number of cyanobacteria capable of cone formation, as well as some that cannot initiate this process but can aggregate into rather flat, tightly woven mats (XAN-8 in Fig. 6), or into thin tubular structures and finely woven floating films (FYG in Fig. 6, after ‘filamentous yellow glider’, Bosak et al., 2009, 2010). These organisms may contribute to the complex morphologies of coniform mats in the field, which often include filmy, draping, laminated bridges.

Three ‘filamentous’ cyanobacterial OTUs from YNP cones are cone-specific and one ‘unicellular’ OTU is mat-specific, suggesting that some cyanobacteria are uniquely adapted to microenvironments within macroscopic cones and mats. Adaptations to the varying intensities and qualities of light and different aggregation behaviors also may underpin the diversity of the four co-occurring OTUs found in many, but not all, cones and mats (OTU 20, 21, 23, 24). Alternatively, this diversity may arise from random fluctuations within the rapidly growing thin layers (Figs 7 and 8) that maintain sectors populated by like individuals even when these individuals carry neutral mutations (Hallatschek & Nelson, 2010; Mitri et al., 2011). Given that 16S rRNA sequences are but a limited measure of the underlying genetic and the physiological diversity (e.g., Moore et al., 1998; Ward et al., 2006), the differences among cyanobacteria from Subsection III in coniform mats may be better resolved by examining longer 16S rRNA sequences, amplifying various functional genes, and analyzing the distribution of OTUs at finer spatial scales within cones and mats.

Thin filamentous cyanobacteria are commonly thought to be condicio sine qua non for the formation of cones, but these structures also contain metabolically active and abundant unicellular cyanobacteria (Figs 4 and 5) and thin filaments that may belong to Chloroflexus- and Roseiflexus-like photosynthetic bacteria (Figs 4E and S4). Our analyses do not identify any of the unicellular cyanobacterial OTUs as cone-specific, in contrast to previous suggestions (Lau et al., 2005). Instead, these OTUs vary among hot springs, ponds, among samples from individual ponds and in time (Figs 5 and S6). This variation may arise as a result of the occasional colonization of coniform surfaces by cyanobacteria from other areas in hot springs. These colonizers may form patchily distributed small populations (Papke & Ward, 2004). The common occurrence of sausage-shaped cyanobacteria in old, decaying coniform mats in the laboratory further points to the ability of these organisms to grow when cone-forming filamentous cyanobacteria do not. Different abundances of individual unicellular OTUs are thus to be expected in actively growing, lithifying, or decaying coniform mats. Samples from more points in time and space would also help distinguish organisms that actively shape coniform mats from organisms that colonize previously formed topography, but are unable to either initiate or maintain cones and the depressions between cones over longer timescales. Such colonization may account for the abundance of sausage-shaped cells in the tips of lightly submerged cones sampled in Mound Spring in 2009 (Fig. 3D), given that mainly thin filamentous cyanobacteria could be identified in the tips of fully

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submerged cones from Mound Spring sampled in 2008 (Fig. 3B).

Thin filaments of Chloroflexus or Roseiflexus-like organisms are commonly seen in field cones (Fig. 4E), but are generally absent from laboratory cultures. When growing in cyanobacterial mats, these organisms can be phototrophic or photoheterotrophic (e.g., van der Meer et al., 2003, 2005) and can contribute to the formation of laminae (Doemel & Brock, 1974). Their contribution to the uptake of inorganic carbon in YNP cones is thought to be small (Walter et al., 1976; Brock, 1978). Indeed, NanoSIMS maps and the accompanying TEM images reveal little labeled carbon in the thin filaments in field cones, even when these filaments are surrounded by highly enriched cyanobacterial cells (Fig. S4). Thin filamentous phototrophs are thus not the main loci of primary production during the main photosynthetic period, but our current data cannot distinguish between their slow autotrophy and photoheterotrophic uptake of labeled cyanobacterial exudates produced during the 60-min incubation. Metabolic potential and function of these thin filaments in coniform mats can be probed in more detail by tracking the distribution of labeled carbon over longer times through high-resolution ultrastructural and NanoSIMS maps (Fig. S4B,C). These measurements should be accompanied by measurements of bulk dissolved organic carbon and 13C-labeled glycolate and should be conducted at different times of day, including periods of low light (van der Meer et al., 2005).

Carbon uptake and cone growth

Measurements of 13CN− allow comparisons of the uptake of inorganic carbon among different cells within the same layer of the mat. This method underestimates the rate of the uptake of inorganic carbon because it does not account for the supply of 13CN− produced by the remineralization of unlabeled organic material. However, the recycling of carbon should not affect the observed relationship between cell densities and the position in the mat (Petroff et al., 2011). Field cones (Fig. 8) and laboratory-grown cones (Petroff et al., 2011) exhibit similar profiles of the uptake of carbon with depth. These profiles match the model predictions for biofilms whose growth is limited by the diffusion of nutrients from the overlying stratified solution (Petroff et al., 2011). This model predicts the highest density of active cells in cone tips, because these areas are the closest to the overlying source of nutrients (or light) and are curved. More filaments thus compete for nutrients (or light) in areas of high cell density (cone tips). The YNP cones, which grow at high light intensities (>1500 μE m−2 s−1), depart from the light-limited or nutrient-limited model in the uppermost approximately 100-μm-thick layer (Fig. 8A). This region contains fewer cells and individual filaments there take up less carbon than those in the dense active layer below the upper 100 μm. In contrast, the uptake of labeled carbon in laboratory cones grown at 200 μE m−2 s−1 is the highest at the very surface (Petroff et al., 2011). The somewhat lower activity of filaments at the surface of field cones may be attributed to photorespiration and the faster recycling of organic carbon, but it does not support the migration of light-limited growth phototactic cyanobacteria toward topographic highs (Walter et al., 1976). Because the very tip contains few degraded cells and ghosts, the faster recycling of carbon in this region should stem primarily from the microbial degradation of extracellular substances and photorespiration (Bateson & Ward, 1988), rather than from the lysis of inactive cells.

Small-scale isotopic labeling and mapping reveals differences in cyanobacterial responses to the influx of inorganic carbon and nitrogen, potentially limiting nutrients within YNP cones (Wiegert & Fraleigh, 1972; Jahinke et al., 2004; Lolaco, 2012). Not only are filamentous cyanobacteria in cones able to extend across diffusive gradients and access nutrients in the overlying solution, but these organisms in cones and mats also concentrate carbon and nitrogen in approximately 300–750 nm large granules (e.g., polysaccharides or cyanophycin). The ability to concentrate carbon and nitrogen may confer an advantage during growth under nutrient-limiting conditions (Shibata et al., 2001; McGinn et al., 2003). The large round photosynthetic cells below the surface of the horizontal mats respond to the influx of carbon and nutrients faster than the filamentous and sausage-shaped cyanobacteria, have a laterally discontinuous distribution, and occur below the upper 150 μm of the mat. These observations suggest that the oval cells grow sporadically during nutrient pulses and are adapted to lower light intensities. Our current experiments track the initial incorporation and distribution of labeled inorganic carbon in the cells of primary producers, but do not determine the longer-term fate of the newly synthesized organic carbon. The latter may critically influence the preservation of coniform mats and their robustness because the abundance of excreted substances (i.e., extracellular biomass) may distinguish the sturdy, cohesive cones in the field from the often filmy, internally porous, transparent structures in the highly enriched laboratory cultures (Fig. 6). The extracellular compounds contain less nitrogen and sulfur relative to cellular biomass and, with the possible exception of cyanobacterial sheaths (Fig. S7), may not be detectable as CN− ions by NanoSIMS (Figs 4 and S2). Future studies can investigate these processes by following the production and distribution of new cellular biomass and exopolymeric substances, the flow of carbon from primary producers to other organisms, and total microbial diversity in coniform mats.
Growth and preservation of cones by silicification

The topography-dependent and temporally variable growth of coniform mats in modern hot springs leads to the formation of porous and dense laminae, textural differences between cones and mats, and distinct macroscopic topography. Cones contain more porous laminae formed by filaments that are oriented randomly or are perpendicular with respect to the growth surface. Recent studies report that these orientations occur primarily in actively photosynthesizing, diffusion-limited mats (Petroff et al., 2010). Structures that form under light-limited conditions, where diffusion limitation should be less pronounced, contain thin, dense laminae of mutually aligned prostrate filaments (Bosak et al., 2009). Thus, the higher availability of nutrients at cone tips in stratified solutions (Jorgensen & DesMarais, 1990) and the depletion of these nutrients within the mat (Petroff et al., 2011) may promote the growth of cone tips and lead to the formation of porous laminae characterized by less abundant cells and more abundant exopolymeric substances (Fig. S7). Textural record of these topography-dependent and time-dependent processes can be preserved by precipitating silica. In some flat laminated microbialites, actively growing microbes outpace silicification, giving rise to porous laminae encrusted by silica, whereas biomass-poor and silica-rich laminae form in autumn/winter when the growth is slow and microbes are unable to outpace silicification (Konhauser et al., 2001, 2004). In keeping with this model, the sparse and more heavily encrusted cells in the topographic lows of coniform mats may be attributed to the slower accumulation of biomass in topographic lows relative to mineral precipitation. Conversely, the fast accumulation of biomass in actively growing cone tips outpaces mineral precipitation and results in unmineralized or only lightly mineralized cells. These preservational mechanisms in coniform mats can give rise not only to the alternating cell-rich and cell-poor laminae driven by seasonal variations in growth, as is the case in stratiform microbialites (Konhauser et al., 2001), but also to the laminae produced by the more rapidly varying orientation of cyanobacterial filaments in cones (e.g., Walter et al., 1976; Petroff et al., 2010). All cones shown and analyzed in this study were, at best, lightly mineralized, and their fine organic laminae may not be preserved by silicification. This does not diminish the value of these coniform mats as model systems in which to study biological aspects of stromatolite morphogenesis, as faster silicification has the potential to preserve small-scale textures in cones from some less accessible localities in YNP (e.g., Fig S3A–C in Bosak et al., 2009). A more critical difference between modern coniform mats and ancient conical stromatolites is mineralogical: most modern small cones are preserved by silicification, but ancient cones were preserved primarily by calcium carbonate. Therefore, direct insights into stromatolites as records of past chemical conditions and possible influences of microbial metabolisms on carbonate precipitation require more studies of the growth and preservation of coniform mats in carbonate-precipitating environments.

CONCLUSIONS

Filamentous cyanobacteria build the vertical scaffolding of conical microbialites that grow in the shallow pools of modern alkaline hot springs. These organisms are present throughout the cones and the adjacent horizontal mats, and most of their 16S rRNA gene sequences are not specific to either cones or mats. Field sequences of filamentous cyanobacteria in coniform mats belong to two principal clusters and have cultured representatives that form vertical structures in highly enriched laboratory cultures. Cyanobacterial sequences vary among different hot springs, among ponds in the same hot spring, and even in one location between two sampling years. Therefore, the growth of modern conical microbialites depends on a number of genetically distinct filamentous cyanobacteria rather than a single organism or an unchanging group of organisms. unicellular sequences in coniform mats vary even among samples from the same pond and are often undetectable, although photosynthetically active Synechococcus-like cyanobacteria are present in cones. Most new cellular biomass in cones grows in the tips, but not at the very surface. In mats, the very surface is both the densest and the most active growth layer, while organic degradation is prevalent in the mat interior. There, cyanobacteria in hot spring coniform mats morphologically diverse and photosynthetically active unicellular cyanobacteria quickly respond to nutrient pulses, but are unable to extend across diffusive gradients and form cohesive porous laminae.

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REFERENCES


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Cyanobacteria in hot spring coniform mats


SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Fig. S1 Lithification in cones and mats.

Fig. S2 Cone-mat sample (photograph, scale in mm) incubated with labeled carbon for 60 minutes showing the distribution of 60 µm × 60 µm NanoSIMS maps and high resolution maps showing different cell morphologies.

Fig. S3 NanoSIMS analysis of a cone (A) and a mat (B) incubated for 60 min in labeled inorganic carbon in 2008.

Fig. S4 Distribution of unlabeled and labeled carbon in thin filaments in the cone incubated with labeled inorganic carbon for 60 min.

Fig. S5 Spatial distribution of OTUs in the samples of cones and mats from four different springs.

Fig. S6 Distribution of OTUs in cones and mats from four different springs in 2009.

Fig. S7 NanoSIMS maps of YNP cones showing possible cyanobacterial sheaths.

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