



Silica Biomineralization of *Calothrix*-Dominated Biofacies from Queen's Laundry Hot-Spring, Yellowstone National Park, USA

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Experiments on microorganisms capable of surviving silicification are often conducted to gain a better understanding of the process of silica biomineralization and to gain insights into microbially influenced rock formations and biofabrics like those found in ancient deposits such as the Early Archean Apex Chert formation (Schopf, 1993; House et al., 2000). An ideal microorganism for studying silicification is the large sheathed cyanobacterium *Calothrix*, which form distinctive organo-sedimentary structures in the low to moderate temperature regions of hydrothermal springs or columnar stromatolitic structures in aquatic systems. Our ability to identify and characterize microfossils from ancient deposits allows us to gain a better understanding of environmental conditions on early Earth. Here we characterized *Calothrix*-dominated biofacies along the outflow apron of Queen's Laundry Hot-Spring in Yellowstone National Park using microscopy and molecular techniques to examine biofacies morphology and phylogenetic diversity. We found that flow regime and temperature had a profound effect on community composition as identified by the observation of five distinct *Calothrix*-dominated communities and on biofacies architecture along the outflow apron.

Keywords: geomicrobiology, microfossils, mineral templating, hot-springs, biomineralization

INTRODUCTION

Precambrian cherts have been found to yield silicified microfossils suggesting that ancient microbial communities present in marine waters and hydrothermal ecosystems became embedded in colloidal amorphous silica and were subsequently entombed and preserved in the fossil record as distinct textures (Westall et al., 1995; House et al., 2000). Characterization of extant hot-spring environments provides us the opportunity to better understand and interpret the paleoenvironment in which ancient biofacies formed, allowing us to take a glimpse into Earth's early environments.

The geology of Yellowstone National Park (YNP) strongly influences the composition of hydrothermal fluids as meteoric waters interact at depth with silica rich acid-intermediate volcanic rock, thereby enriching hydrothermal fluids with dissolved silica (Channing and Butler, 2007). Silica deposits, referred to as sinter, form in and around the hot-spring once hydrothermal fluids

erupt at the surface. As hydrothermal fluids cool the solubility of amorphous silica is exceeded, e.g., due to changes in pH or temperature and supersaturation through evaporation, resulting in the polymerization of dissolved silica and subsequent precipitation (White et al., 1956; Rimstidt and Barnes, 1980; Benning et al., 2005; Hugo et al., 2011). Siliceous deposits in YNP can be studied as extant analogs of early Earth allowing for the study of biogenic silica deposition and microfossil formation (Channing and Butler, 2007).

In this study, we describe *Calothrix*-dominated biofacies composed of microbe-mineral assemblages in which mineral deposition exhibits a specific and unique suite of biological characteristics due to nucleation of minerals onto microbial biofilm/mat surfaces. Mineralized *Calothrix*-dominated mats heavily influence the fabrics of low- to mid-temperature siliceous sinters forming distinctive hot-spring-associated rock formations (Walter, 1972; Cady and Farmer, 1996; Jones et al., 2001; Konhauser et al., 2001). The distinctive *Calothrix* biofacies have been instrumental in the interpretation of hot-spring paleoenvironments (Walter et al., 1996; Campbell et al., 2001; Jones and Renaut, 2003; Hugo et al., 2011). Microbial populations in Queen's Laundry Hot-Spring are exposed to circumneutral geothermal groundwater fluids that are saturated with dissolved silica resulting in a series of abiotic silicification reactions (Cady and Farmer, 1996; Benning et al., 2004). Microfossils form as silica minerals nucleate on cell walls and in exopolysaccharides, completely impregnating organic material; this must occur rapidly for cell and mat morphology to remain intact (White et al., 1956).

Previous studies have identified *Calothrix*-like microstructures from sinter deposits of extinct hot-springs, suggesting that *Calothrix* biofilms directly influence the biofabric of sinter deposits in extant hot-springs (Walter, 1972; Cady and Farmer, 1996; Jones et al., 2001; Konhauser et al., 2001). Silicification of *Calothrix* is thought to be a passive process due to the rapid and extreme changes in fluid chemistry as fluids flow from the deep source vent(s) to the shallow outflow apron (Konhauser et al., 2003). Here we characterize microfossil formation from three distinct microenvironments along the outflow apron from Queen's Laundry Hot-spring and show that each of these microenvironments host distinct *Calothrix*-dominated communities.

Calothrix

Calothrix, constituents of the Rivulariaceae family, one of the most diverse cyanobacterial lineages, are present in marine, freshwater, and terrestrial environments. However, little is known about their genetic diversity from the natural environment due to the lack of data pertaining to the biodiversity of *Calothrix* present across habitats (Sihvonen et al., 2007). The filamentous cyanobacterium *Calothrix* has been observed to occupy the upper 1–2 mm of microbial mats formed in silica-dominated hot-springs across YNP (Walter, 1976). Rivulariaceae cyanobacteria such as *Calothrix* are morphologically complex, possessing specific characteristics that make them unique. These include tapering trichomes and terminal heterocysts which are enclosed in a thick fibrillar sheath of microfibrillar capillaceous

structures of exopolysaccharide (EPS) and lipopolysaccharides (LPS) forming large branched molecules that are insoluble in water making them resistant to dehydration (Westall et al., 1995; Hoiczkyk, 1998; Whitton, 2002). The multilayered exterior presents mechanical and permeability obstacles for larger molecules making it necessary for alternative transport mechanisms, such as porins, and junctional pore complexes (Hugo et al., 2011). These alternative transport mechanisms may contribute to the silicification of cyanobacteria in hot-spring environments, thereby allowing for the formation and preservation of microfossils (Hoiczkyk and Hansel, 2000). Monomeric dissolved silica is capable of passing through the outer sheath and accumulating in the extracellular space. Polymerization occurs as dissolved silica concentrations increase, trapping amorphous silica in the extracellular space which eventually leads to cell death and the formation of a microfossil. The growth pattern of these microorganisms directly influences the texture of rock formations in and around the hot-springs (Cady and Farmer, 1996).

Algal and bacterial biofacies composed of amorphous silica have been well-characterized and are known to occur around hot-spring environments in YNP forming structures reminiscent of stromatolites (Walter, 1972). However, we have characterized *Calothrix*-dominated biofacies that form not only textured sinters that are crudely laminated with wavy surface features similar to the commonly recognized columnar domes, but three distinct biofacies identified as: nodular mats, pustular mats (small domes), and terracette ridges (clusters of contiguous domes).

The actively growing surface of the *Calothrix* mat becomes encrusted with silica minerals so that filaments are completely entombed in a silica matrix deeper in the mat. The rate of mineralization is dependent on several environmental parameters such as pH, silica activity, temperature, and presence of water (Iler, 1979; Braunstein and Lowe, 1996; Walter et al., 1996). Filaments are oriented in the direction of water flow and become fossilized in their orientation due to rapid silica mineralization entombing filaments and EPS making them appear as bundles of filaments. Once completely entombed in silica the sheath is difficult to differentiate from the silica matrix. Due to their resilience to degradation silicified sheaths are more commonly preserved in the rock record than the internal trichome (Figure 1; Jones et al., 2001).

In this study, we characterize the biofacies and the microbial communities from three distinct *Calothrix*-dominated mats collected at Queen's Laundry Hot-Spring in YNP. These distinct biofacies were characterized from specimens collected along the temperature gradient of the outflow apron. This was done as a means to better understand the effects temperature and fluid dynamics have on community diversity and the morphology of each distinct biofacies, and to characterize the process of microfossil formation and preservation in the rock record.

Field Site Description

Queen's Laundry is a member of the Sentinel Meadows Group located in the Lower Geyser Basin of YNP, USA at 44°33'48" N, 110°52'13" W. The perimeter of the outflow apron is dominated by the sheathed filamentous cyanobacterium *Calothrix*, which

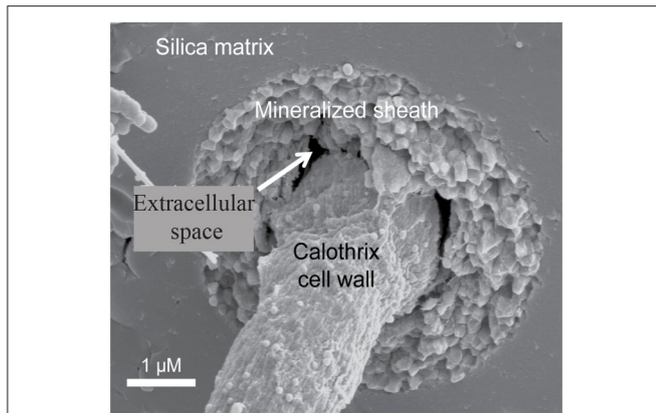


FIGURE 1 | Scanning electron micrograph of an entombed *Calothrix* filament, whose sheath is completely mineralized with silica allowing the cell to remain intact and preserved.

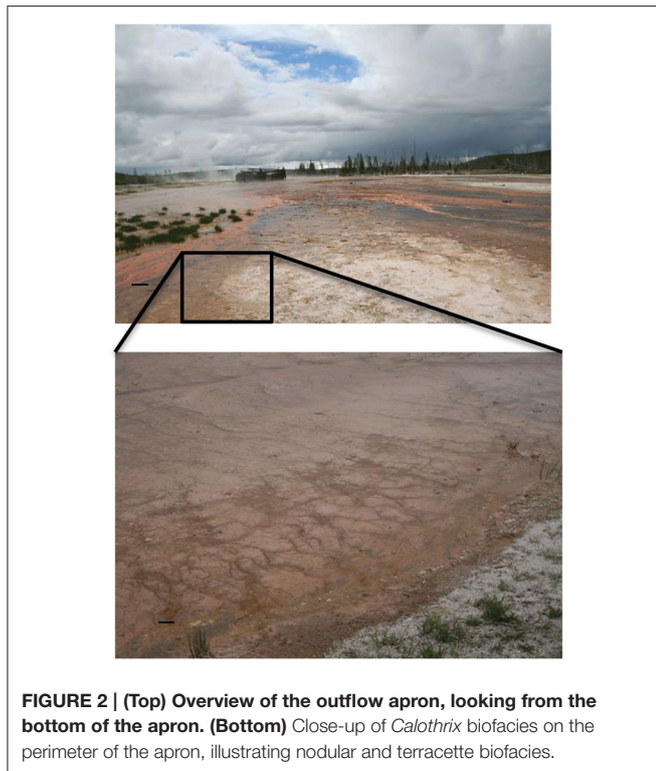


FIGURE 2 | (Top) Overview of the outflow apron, looking from the bottom of the apron. (Bottom) Close-up of *Calothrix* biofacies on the perimeter of the apron, illustrating nodular and terracette biofacies.

typically populates neutral to alkaline waters between 25 and 50°C. This gently boiling and surging hot-spring is comprised of a large deep pool with two source vents discharging $\sim 0.6 \text{ m}^3 \text{ h}^{-1}$ of silica-saturated hydrothermal (92°C) fluids into a narrow stream that meanders around the outflow apron, creating a thermal and pH gradient (Braunstein and Lowe, 1996; Hugo et al., 2011). The outflow apron broadens into a large braided channeled apron that gently slopes, draining into marshy grasslands (Figure 2). Geochemical data reported by the USGS show that the average concentration of silica from hot-springs in the Lower Geyser Basin is 3.1 mM (McCleniskey et al., 2002).

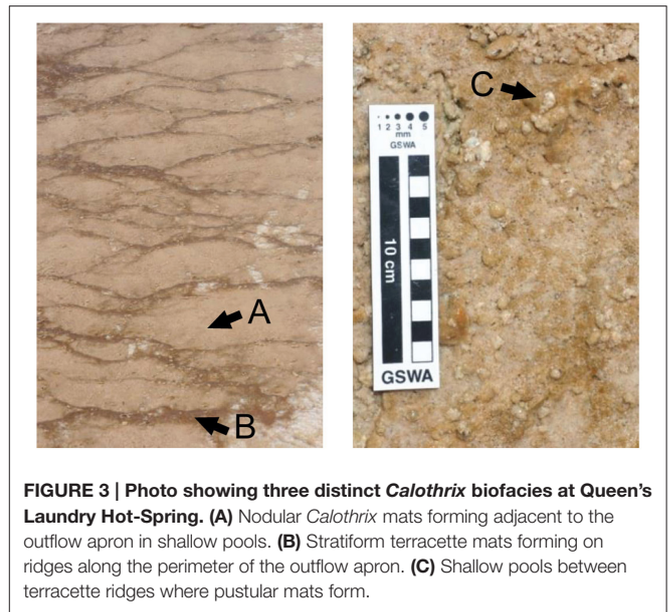


FIGURE 3 | Photo showing three distinct *Calothrix* biofacies at Queen's Laundry Hot-Spring. (A) Nodular *Calothrix* mats forming adjacent to the outflow apron in shallow pools. (B) Stratiform terracette mats forming on ridges along the perimeter of the outflow apron. (C) Shallow pools between terracette ridges where pustular mats form.

Biofacies characterized were: (1) nodular mats forming in shallow pools of thermal fluid located at the top and bottom of the outflow apron; temperatures $\sim 25^\circ\text{C}$ (Figure 3C), (2) stratiform terracette ridges forming along the length of the outflow apron and wetted by a thin sheet of flowing fluid; temperatures from 50°C at the top closest to the source vent to 25°C at the bottom (Figure 3B), and (3) pustular mats formed along the length of the outflow apron in shallow gently flowing thermal pools that form between stratiform terracette ridges; temperature of 25–50°C from top to bottom of the outflow apron (Figure 3A).

MATERIALS AND METHODS

Sample Collection and Preparation

Sinter and microbial mat specimens (1 × 1 cm) were collected and immediately fixed in a 3% glutaraldehyde solution of 0.2 μm-filtered spring water for microscopy. Samples collected for molecular analyses were fixed in RNALater (Life Technologies, Grand Island, NY). All samples were stored at 4°C while in the field. Prior to microscopic examination specimens were rinsed twice in 0.1 M cacodylate buffer to remove fixative. *Calothrix* biofacies were examined macroscopically and microscopically, characterizing gross morphology of intact mats and the morphology of individual cells.

Sonication Experiments

Silica encrusted *Calothrix* filaments and associated EPS were sonicated in order to remove mineral rinds. Prior to sonication, *Calothrix* mats exhibited a hirsute appearance from mineralization of the EPS around the filaments. Sonication at 30 s intervals were increasingly effective until 200 s, when cells began to lyse.

Stereomicroscope Analysis

Whole intact *Calothrix* mats were examined using a Nikon SMZ800 stereoscope; digital images were acquired using a Leica (Wetzlar, Germany) DFC digital camera. Specimens were first examined intact to characterize morphology, surface features, and growth orientation of filaments. Specimens were then sectioned lengthwise, with a scalpel and analyzed in cross-section to characterize the interior of the mat, the average specimen was 5 mm thick.

Optical Light Microscopy

Highly mineralized *Calothrix* mats required removal of mineral rinds in order to observe cellular characteristics using the optical light microscope (OLM). Specimens were prepared for observation by first removing the silica rind encasing filaments and then by removing the top actively growing portion of the *Calothrix* mat. Sonication of mineralized mats was an effective method of loosening and removing silica rinds that encrusted sheathed filaments. Densely packed filaments cemented in mineral matrices were loosened with the removal of minerals from the sheaths with increasing sonication time. After each

30 s sonication interval the overall mat structure was examined to characterize the integrity of *Calothrix* filaments using 20X objectives under phase contrast using a Leica (Wetzlar, Germany) DMRX OLM and images were acquired with an Apogee CCD camera.

Scanning Electron Microscopy

Calothrix mats from each biofacies were examined using a FEI Siron high-resolution scanning electron microscope (HR-SEM) equipped with an Energy Dispersive Spectroscopy (EDS) detector (FEI, Hillsboro, OR). Samples were prepared utilizing two dehydration techniques: (1) extensive chemical fixation with osmium tetroxide and dehydration with a graded ethanol series followed by critical point drying, or (2) chemically fixed with a 2% glutaraldehyde (Fisher, Pittsburg, PA) solution rinsed twice in 0.1 M cacodylate buffer and air-dried in a desiccator. Samples prepared using the first fixation technique were rinsed twice in 0.1 M cacodylate buffer soaking for 10 min each, post-fixed using a 1% osmium tetroxide (Fisher, Pittsburg, PA) solution prepared in 0.1 M cacodylate buffer (Fisher, Pittsburg, PA) for 1 h, after which samples were rinsed twice in 0.1 M cacodylate

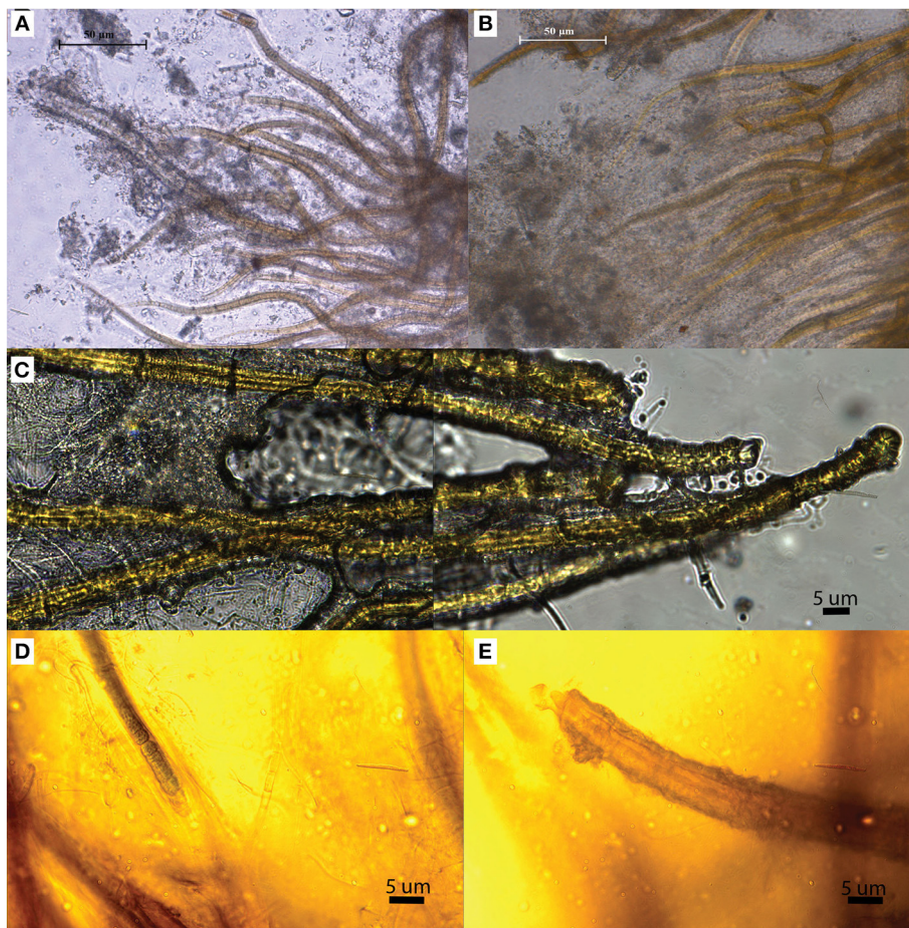
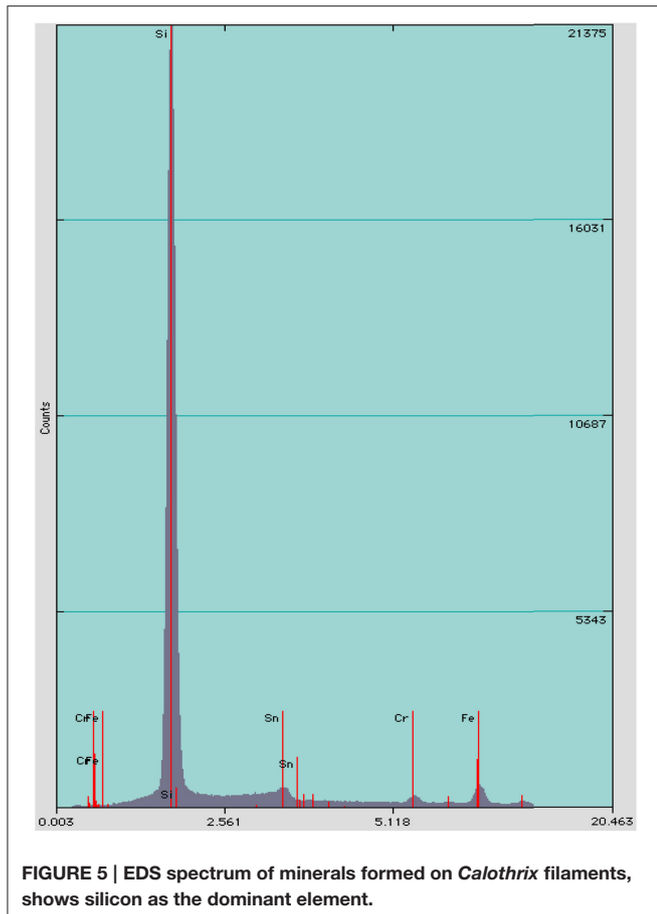


FIGURE 4 | OLM images illustrating the various stages of silicification. (A,B) Illustrate primary (A) and advanced (B) encrustation. (C) Filaments are cemented together in a solid silica matrix illustrating encasement. (D,E) Illustrate entombment, as filaments are not only encrusted in silica but entombed in a solid silica matrix.



buffer, dehydrated using a graded ethanol series (50, 70, 90, and 100%), and then critical point dried. Dehydrated specimens were mounted on aluminum pins and coated with 100Å Au-Pd.

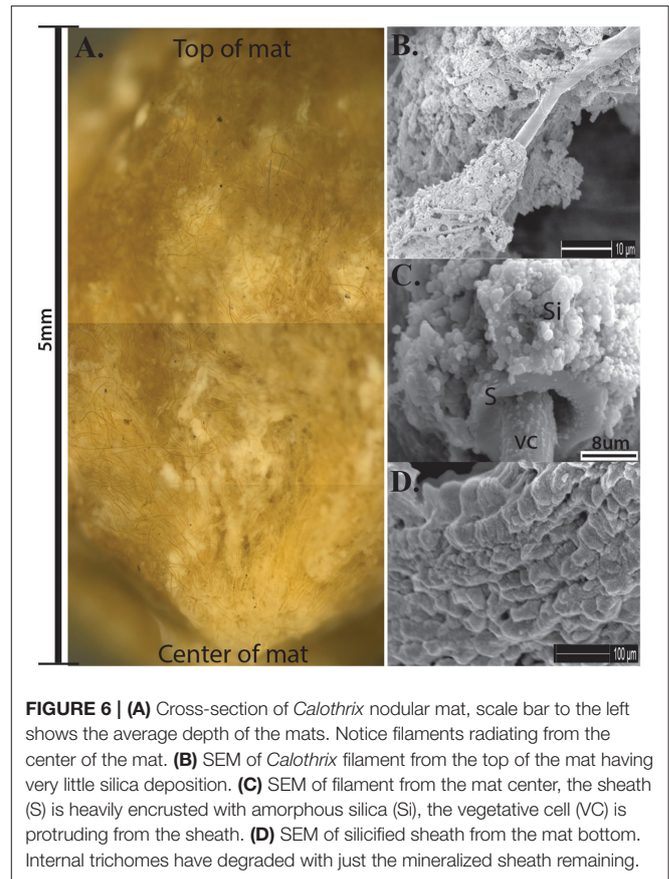
EDS analysis provides elemental and chemical analysis of a minerals associated with *Calothrix* filaments. Spectra were collected using 50 kV, and visualized using AZtecEnergy analysis software (Oxford Instruments).

Genomic DNA Extraction

Genomic DNA (gDNA) was extracted from samples using the FastDNA SPIN Kit for Soil (MP Biomedicals, Solon, OH) following the manufacturer's protocol with the modification that the gDNA was eluted into 10 mM Tris at pH 8. The purity and concentration of gDNA were determined with a Nanodrop ND-1000 spectrophotometer (Thermo Scientific, Wilmington, DE) and diluted to ~10 ng/μL for downstream molecular applications.

Community Diversity

Community diversity of *Calothrix* biofacies was analyzed using terminal restriction fragment length polymorphism (T-RFLP), providing a fingerprint of the community from each biofacies, based on the small subunit ribosomal RNA (SSU rRNA) gene. The methodology used for T-RFLP analysis



has been reported in detail in previous publications (Davis and Moyer, 2008; Rassa et al., 2009; Fleming et al., 2013). This technique accurately resolves populations in microbial communities of low to intermediate richness (Engebretson and Moyer, 2003). Electropherograms are imported into the program BioNumerics (Applied Maths, Austin, TX) where community fingerprints were compared using average Pearson product moment correlation (Häne et al., 1993). Community fingerprints were compared between 50 and 500 base pairs (bp) and the average Pearson product moment correlation coefficient was calculated for cluster analysis by unweighted pair group method with arithmetic mean (UPGMA) combining eight restriction digests (Davis and Moyer, 2008). The primer set used was 68F-FAM (5' 6-FAM - TdNA dNAC ATG CAA GTC GdK dK CG 3') and 1492R (5' dKGdP TAC CTT GTT ACG ACT T 3') with identical conditions as previously reported (Rassa et al., 2009). Three replicate PCR reactions were pooled, desalted using Amicon® Pro Purification System (EMD Millipore, Darmstadt, Germany), and split between eight restriction enzyme treatments using *AluI*, *BstUI*, *HaeIII*, *HhaI*, *HinfI*, *MboI*, *MspI*, and *RsaI* (New England BioLabs, Ipswich, MA). Reactions were visualized with an internal LIZ-500 (Applied Biosystems, Foster City, CA) size standard by capillary electrophoresis on an ABI 3130xl genetic analyzer (50-cm capillary array, POP-6; Life Technologies, Grand Island, NY).

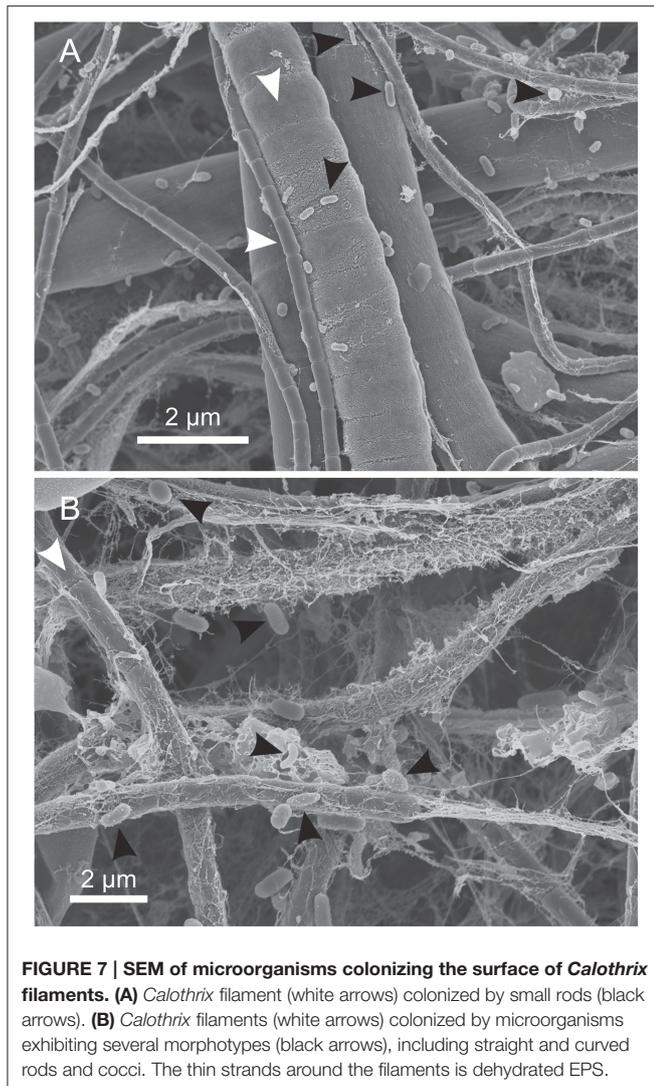


FIGURE 7 | SEM of microorganisms colonizing the surface of *Calothrix* filaments. (A) *Calothrix* filament (white arrows) colonized by small rods (black arrows). **(B)** *Calothrix* filaments (white arrows) colonized by microorganisms exhibiting several morphotypes (black arrows), including straight and curved rods and cocci. The thin strands around the filaments is dehydrated EPS.

Clone Library

Five replicate SSU rRNA gene PCR reactions were pooled and cleaned as above, with the modification that the forward primer did not contain a 5' fluorescent label. Desalted amplicons were cloned with a CloneJET PCR Cloning Kit following manufacturer's instructions (Thermo Fisher Scientific, Waltham, MA). All clones were streaked to isolation and assayed for the correct size fragment using PCR with pJET1.2 forward and reverse primers. Clones were then grown up in Terrific Broth with 100 $\mu\text{g mL}^{-1}$ ampicillin and were sequenced at Beckman Coulter Genomics (Danvers, MA). Operational taxonomic unit (OTU) analysis was initially determined with 5' reads of the SSU rRNA gene (Tartof and Hobbs, 1987). Sequences were aligned using the ARB-SILVA database with SINA Webaligner (Pruesse et al., 2007), masked, and binned into OTUs based on 97% minimum similarity. At least one clone from each OTU was chosen for full-length sequencing and checked for chimeras using Pintail (Ashelford et al., 2005) and Mallard (Ashelford et al., 2006). Using unambiguously aligned full-length

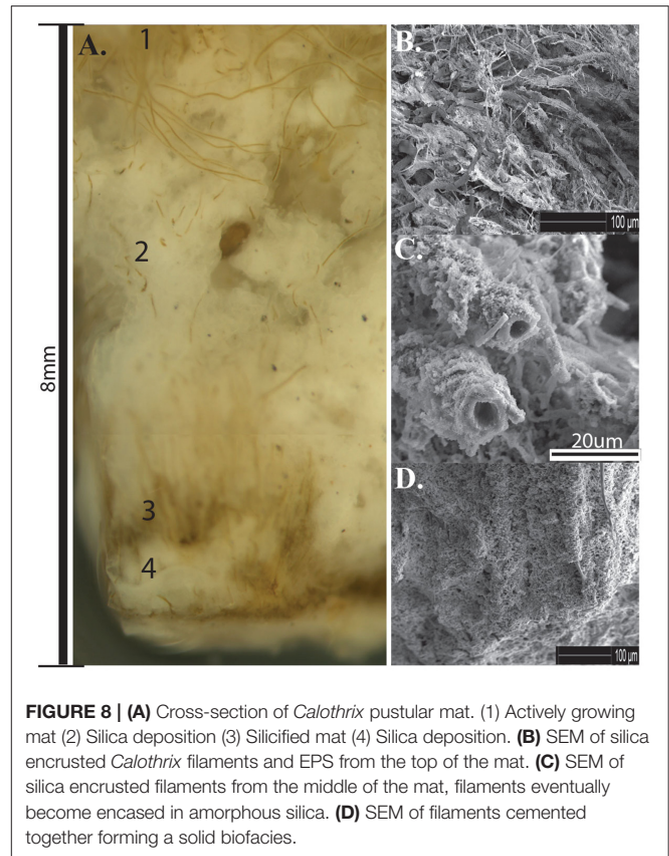


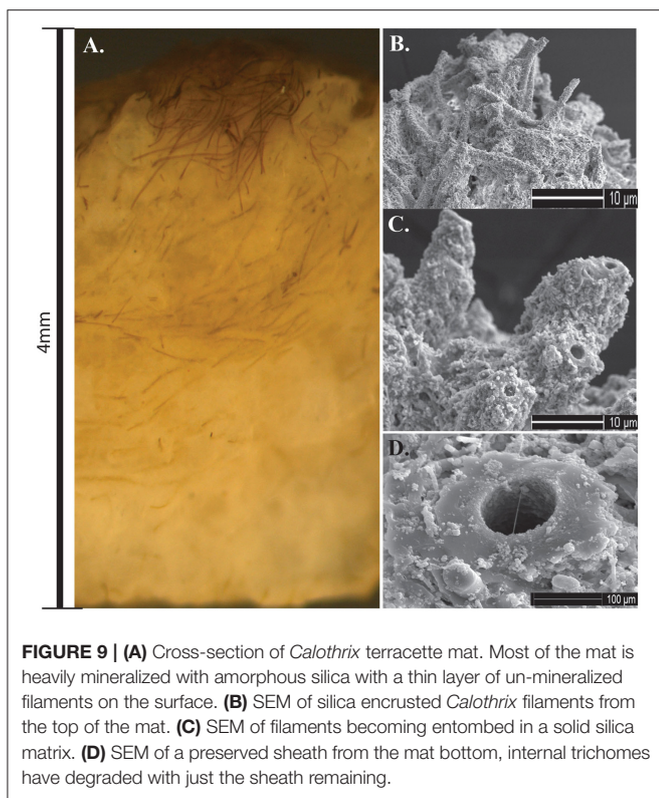
FIGURE 8 | (A) Cross-section of *Calothrix* pustular mat. (1) Actively growing mat (2) Silica deposition (3) Silicified mat (4) Silica deposition. **(B)** SEM of silica encrusted *Calothrix* filaments and EPS from the top of the mat. **(C)** SEM of silica encrusted filaments from the middle of the mat, filaments eventually become encased in amorphous silica. **(D)** SEM of filaments cemented together forming a solid biofacies.

sequences, phylogenetic placements according to maximum likelihood methods were calculated using RAxML version 7.2.6 (Stamatakis, 2006) with the General Time Reversible (GTR) model of nucleotide substitution, optimized substitution rates, and GAMMA model of rate heterogeneity; bootstrap values were calculated using a 1000 replicates. Full-length sequences were submitted to the National Center for Biotechnology Information (NCBI), accession numbers KU382114 through KU382143.

Estimating Relative Abundance of *Calothrix* using T-RFs

Relative proportions of *Calothrix* from each of the three biofacies were calculated using restriction maps from the T-RFLP restriction enzymes (Engelbreton and Moyer, 2003; Rarnette, 2009). Estimates of community composition were generated *in silico* from *Calothrix* and cyanobacterial sequences found from clone library analysis. Upon examining the eight restriction enzymes we found that restriction enzyme *Bst*UI had the most specific identifier signature for *Calothrix* by comparison of the terminal-restriction fragments (T-RFs). Comparison of T-RFs showed that peaks at 57 bp were shared with the least amount of other cyanobacteria from the *Calothrix* biofacies.

To estimate the relative abundance of *Calothrix* within the community all bands between 50 and 500 bp were counted for the *Bst*UI digests. The height of the band representing the 57 bp *Calothrix* T-RF was divided by the sum of the heights from all the bands in the electropherogram above a 3% background



relative fluorescence maximum. This method gave us an estimate of *Calothrix* relative abundance. Abundance estimated from T-RFs is only semi-quantitative. For example, while this method estimated a 76% abundance of *Calothrix* sp., clone library analysis found only 47% *Calothrix* sp.

RESULTS

We identified three distinct stages of silicification using microscopy: (1) encrustation, when the sheath of individual filaments is overlain with fragile silica minerals that easily fracture and dissociate when manipulated (Figures 4A,B); (2) encasement, when cells have a rind of silica minerals forming around individual filaments and EPS developing a thick, tough semisolid outer mineral rind (Figure 4C); and (3) entombment of individual filaments occurs when cells are cemented together in a solid silica matrix (Figures 4D,E). EDS analysis of *Calothrix* biofacies revealed that silicon was the most abundant element in the mineral matrices (Figure 5).

Nodular *Calothrix* Biofacies

Nodular *Calothrix* mats formed loose, spherical nodules on the bottom of shallow pools. Nodular mats were pigmented dark green to brown. The dark coloration is due to the presence of the pigment scytonemin in the sheath and is typically found to be synthesized in environments where cyanobacteria are exposed to intense light and UV irradiation (Ehling-Schulz et al., 1997; Dillon and Castenholz, 2003). Filaments in the nodular mats were oriented vertically from the base of the mat fanning out toward

the surface of the mat which was encased in a semi-solid EPS-mineral matrix of amorphous silica (Figure 6A). Nodular mats were found growing in shallow pools covering a wide distribution of temperatures between 75.9 and 14.3°C. *Calothrix* filaments at the surface of the mat serve as a substrate for colonization by other microorganisms (Figure 7), filaments in the center of the mat had accumulated more silica in their EPS with the outer sheath becoming semi-solid, and filaments at the base of the mat were entombed in amorphous silica with the extensive outer sheath becoming completely fossilized (Figures 6B–D respectively).

Pustular *Calothrix* Biofacies

Pustular *Calothrix* mats formed attached to the bottom of shallow terracette pools along the length of the outflow apron. Mats had an irregular morphology sitting atop a semi-solid core of compacted silica grains, in which filaments grow radially from the core. Pustular mats were dark brown-green pigmented, forming an extensive mat along the bottom of pools (Figure 8A). *Calothrix* filaments were cemented together in a semi-solid EPS-silica matrix, with the tapered ends of the filament anchored in the underlying sinter and the filament body and terminal heterocyst directed outwards. The surface of the mat had a hirsute appearance from protruding filaments (Figure 8B). The mat was comprised of alternating layers of *Calothrix* filaments and amorphous silica, with layers alternating between the tapered and broad ends of the filament (Figure 8C). Sinter at the base of the mat contained completely entombed filaments that were oriented vertically throughout the mat perpendicular to the attachment site at the base of the structure (Figure 8D). Pustular mats formed in spring fluids from 61.2 to 7°C.

Stratiform *Calothrix* Biofacies

Stratiform terracette mats formed slightly raised ridges that formed a continuous meandering narrow mat across the width of the ridge. Mats formed along the length of the outflow apron over a broad temperature range, from 62.2 to 8.5°C. Microscopic observations of cross-sections through terracette mats exposed multiple laminae composed of alternating layers of filaments that have an upward growth and dense silica layers (Figure 9A). *Calothrix* filaments in the actively growing upper portion of the mat exhibited tan to dark brown pigmentation, filaments were heavily encrusted in a semi-solid silica matrix (Figures 9B,C). Sheaths at the base of the mat were completely entombed and fossilized (Figure 9D).

Community Analyses

T-RFLP analyses confirmed the presence of *Calothrix* as the dominant bacterial species (or operational taxonomic unit, OTU) in these biofacies samples, making up on average 43% of the community by T-RFLP. From clone library analysis, the dominant *Calothrix* sp. (at 47% of the community) from that single sample were related to other *Calothrix* spp. (AY147029-30) found across YNP (Dillon and Castenholz, 2003). The next most abundant species was an unclassified cyanobacterium at 12% of the community. Further, the unclassified cyanobacterium

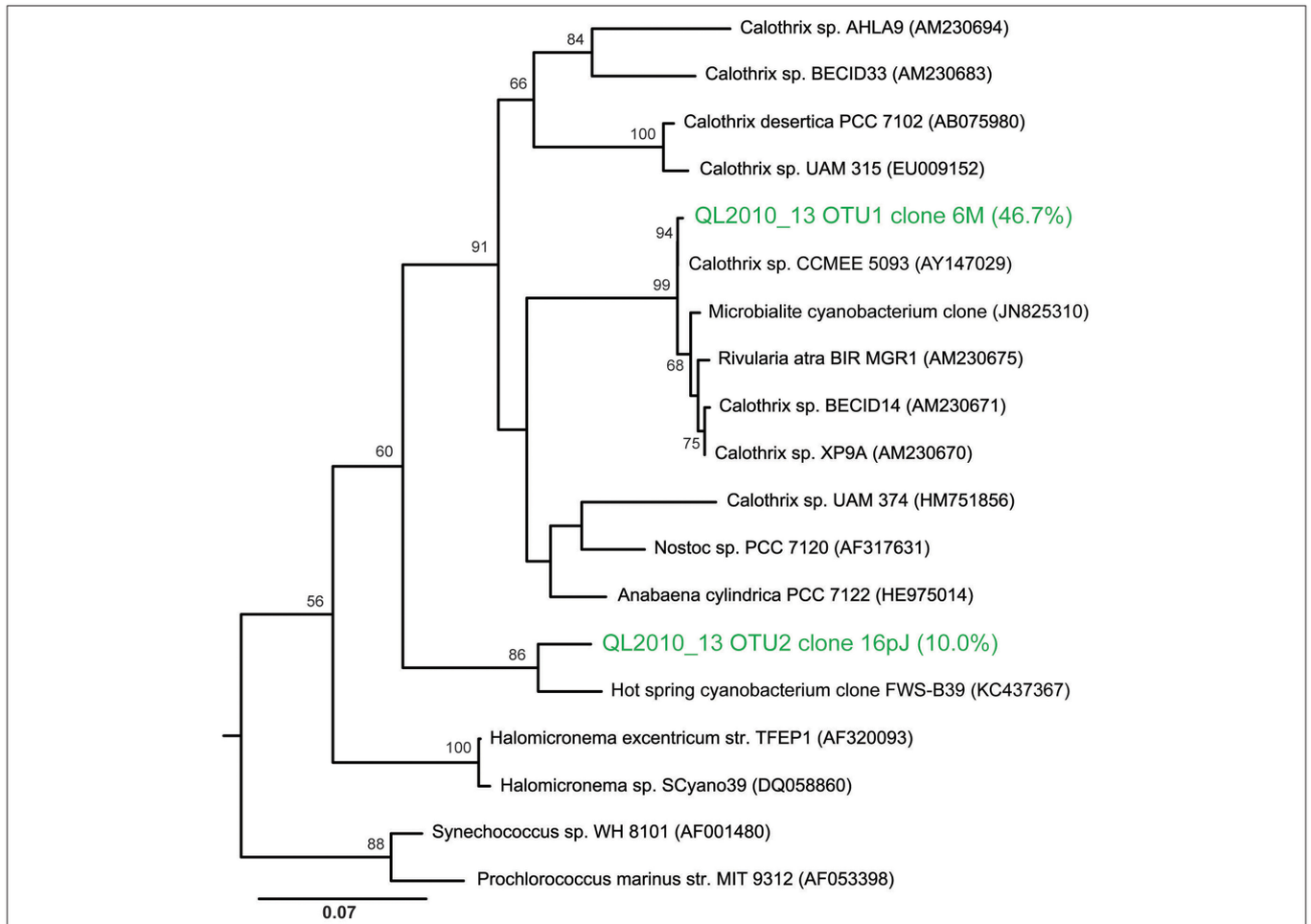


FIGURE 10 | Maximum likelihood phylogenetic tree showing the placement of the two most abundant OTUs from a nodular biofacies (QL2010_13). *Calothrix* clone 6M13 made up 47% of the bacterial community, with the next most abundant clone, 16pJET (12%), being an unclassified cyanobacteria. The tree is rooted using *Aquifex pyrophilus*.

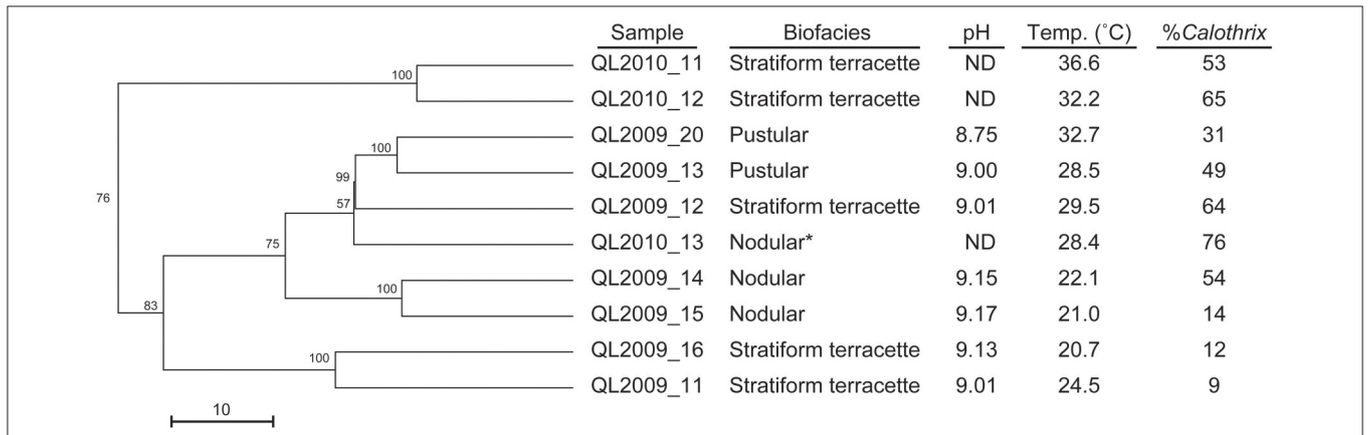


FIGURE 11 | Dendrogram illustrating the cluster analysis of bacterial communities from T-RFLP fingerprints, showing the clustering of our three distinct biofacies at different temperature optima. Starred (*) sample was selected for clone library analysis.

was most closely related (**Figure 10**) to an uncultivated clone from a hot-spring in China, with the closest cultivated relative within the GpIV genus of the Cyanobacteria (92% similar to *Halomicronema* sp.). The majority of the remaining members of the clone library bacterial community were heterotrophic lineages within the *Bacteroidetes* at 27% of the community.

Grouping of biofacies samples by T-RFLP fingerprints showed clustering based primarily by temperature regime and then by biofacies type and the temperature regime and biofacies type (**Figure 11**). Clustering of samples was also largely dependent on changes in the relative abundance of secondary microbial communities which were identified by their T-RF fingerprint, in agreement with observations made via microscopy (**Figure 7**). The changes in heterotroph abundance is seen as a decline in *Calothrix* relative abundance (%) with temperature in **Figure 11**. Biofacies samples ranged from 76 to 9% *Calothrix* (**Figure 10**) (Dicksved et al., 2007). Terracette biofacies clustered into three communities based on different sampling temperature niches (high 34°C, middle 30°C, and low 22°C). With three clusters of stratiform terracette communities, the diversity of this biofacies type was much greater than the other two biofacies types (55 vs. 83% minimum similarity, respectively).

DISCUSSION

In YNP *Calothrix* biofacies appear to be constrained to shallow (few cm) regions of the spring across a broad temperature range (7–75.9°C). The specific environmental conditions (i.e., flow rate and temperature) in which these biofacies form have implications for the interpretation of *Calothrix* biosignatures preserved in ancient cherts, not unlike those identified by House et al. (2000) in the Gunflint Formation. Previous studies have shown that the extensive fibular sheath around *Calothrix* filaments plays a significant role in mineral templating as dissolved silica ions nucleate on the sheath and associated EPS and LPS, thereby promoting microfossil formation (Gilbert et al., 2005; Hugo et al., 2011). The addition of reactive side chains to the EPS and LPS challenges the idea of passive mineral nucleation, favoring instead active, directed templating of biogenic minerals (Bhaskar and Bhosle, 2005; Hugo et al., 2011). Directed nucleation of minerals may serve several purposes for the survival of *Calothrix* in extreme environments. For example the silica rind may offer UV protection, deter predation, and prevent dehydration (Phoenix et al., 2006). In contrast to directed mineral nucleation, spontaneous nucleation occurs as silica-rich hydrothermal fluids erupt to the surface and undergo rapid changes in temperature and pH (Yee et al., 2003). Our findings suggest that soluble silica ions, similar in molecular structure to water, have the ability to cross through the fibular sheath resulting in the accumulation of silica ions in the extracellular space. The accumulation of silica ions results in spontaneous nucleation, forming silica colloids that become trapped in the extracellular space, further entombing the internal trichomes in silica, and promoting their preservation as microfossils.

In this study we identified and characterized three distinct biofacies along the outflow apron of Queen's Laundry Hot-Spring, each of which formed distinct textures specific to the flow regime in which they formed. Nodular biofacies formed on the bottom of pools; these biofacies experience little to no fluid motion as they typically form in meanders when fluid discharge from the hot-spring is high. Pustular biofacies form in extremely shallow pools along the outer edge of the outflow apron; mats forming here are attached to the bottom of the pool and experience continual but slow fluid motion. Stratiform terracette biofacies form slightly raised ridges along the outer edge of the outflow apron and experience rapid sheet flow fluid motion. Due to the rapid flow rate, stratiform biofacies are comprised of thin compact layers of *Calothrix* filaments overlain by relatively massive silica deposits.

We identified and characterized three distinct stages of silicification: encrustation, encasement, and entombment. Close examination of these stages indicate that early silicification and encrustation of *Calothrix* occurs on the surface of the extensive outer sheath, and with continued silica deposition filaments become encased and eventually cemented together. With continued silica deposition filaments become entombed and advanced mineralization occurs with the infiltration of silica ions in the extracellular space.

Community fingerprinting coupled with clone library construction confirmed that biofacies were *Calothrix*-dominated. In addition, we found that within the Queen's Laundry Hot-Spring there was variation in community composition between each distinct biofacies as a result of environmental conditions (i.e., temperature and pH). A prior study by Boyd et al. (2012) found that the patchy distribution of phototrophs in diverse ecological niches was a result of adaptations enabling habitation in diverse environments as a function of temperature, pH, and sulfide concentrations. In addition we would suggest that flow regime in which the biofacies form is equally important as it has a profound impact on the delivery of dissolved chemical constituents (e.g., nutrients, carbon, sulfide, etc.), and influences temperature, pH, and morphology of the biofacies.

AUTHOR CONTRIBUTIONS

WS conceived of the study, with research developed and directed by PIs BT and CM. Samples were collected by WS, SM, KRH, and CM. Morphological analysis was conducted by WS. Molecular analysis was conducted by WS, SM, KRH, KWH, and CM. WS wrote the manuscript, with major contributions from SM and CM, and edits from all coauthors.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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