

Algal Species and Light Microenvironment in a Low-pH, Geothermal Microbial Mat Community

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Received 27 October 2004/Accepted 1 July 2005

Unicellular algae are the predominant microbial mat-forming phototrophs in the extreme environments of acidic geothermal springs. The ecology of these algae is not well known because concepts of species composition are inferred from cultivated isolates and microscopic observations, methods known to provide incomplete and inaccurate assessments of species in situ. We used sequence analysis of 18S rRNA genes PCR amplified from mat samples from different seasons and different temperatures along a thermal gradient to identify algae in an often-studied acidic (pH 2.7) geothermal creek in Yellowstone National Park. Fiber-optic microprobes were used to show that light for algal photosynthesis is attenuated to <1% over the 1-mm surface interval of the mat. Three algal sequences were detected, and each was present year-round. A *Cyanidioschyzon merolae* sequence was predominant at temperatures of $\geq 49^\circ\text{C}$. A *Chlorella protothecoides* var. *acidicola* sequence and a *Paradoxia multisita*-like sequence were predominant at temperatures of $\leq 39^\circ\text{C}$.

Nondescript spherical unicellular red algae (Rhodophyta) predominate in microbial mats in high-temperature (45 to 55°C) acidic (pH of <3) geothermal habitats in Yellowstone National Park and similar habitats worldwide (10, 49). Unicellular *Chlorella*-like algae (Chlorophyta) whose cells closely resemble those of thermophilic red algae are also present in acidic geothermal mats; however, *Chlorella* is not known to thrive at high temperatures (>45°C) (17). Knowledge of the ecology of these algae is based largely on traditional methods, i.e., cultivated species and microscopic descriptions of algal cells in environmental samples (27). However, microscopic identification of simple unicellular algal species is difficult (53). Indeed early studies refer to thermophilic red algae as *Chlorella* (3, 4). Moreover, cultivation-independent (molecular) analyses of microbial communities have repeatedly demonstrated that cultivated isolates are inadequate to describe the ecology of microbes in natural habitats (47, 48, 51, 52).

Recognizing the limitations of traditional algal classification schemes, researchers have used sequence analysis of 18S rRNA and ribulose biphosphate carboxylase genes to establish phylogenetic relationships among cultivated strains of unicellular thermoacidophilic and *Chlorella*-like algae (1, 13, 24, 25, 30–33, 53). In many instances traditional generic designations based on cytological and physiological characteristics are incongruent with genetic phylogeny, and generic nomenclature of some strains requires revision. In the case of the *Chlorella*-like algae, sequence analyses indicate that the ability to thrive in low-pH environments has evolved independently in

diverse lineages of green algae (30) and that the genus “*Chlorella*” has been applied to phylogenetically divergent but morphologically similar unicellular algae (30–33, 53). In the case of thermoacidophilic red algae, three genera and at least four species have been described based on traditional characteristics, *Galdieria maxima*, *Galdieria sulphuraria*, *Cyanidium caldarium*, and *Cyanidioschyzon merolae*. In general, *G. sulphuraria* cells are spherical and indistinguishable from those of *C. caldarium*; however, *G. sulphuraria* strains are distinguished by their ability to grow readily under dark heterotrophic conditions. *G. maxima* strains grow slowly under dark heterotrophic conditions and are distinguished by large cell size. *C. merolae* is a strict autotroph distinguished by crescent- or elliptically shaped cells. Genetic studies indicate that all *G. sulphuraria* strains form a well-supported clade (nomenclature of some strains within this clade likely requires revision [25]) while species of the three remaining morphologically and physiologically diverse genera, *C. caldarium*, *C. merolae*, and *G. maxima*, are closely related to each other but genetically distinct from *G. sulphuraria* (13, 25). Genetic sequence analysis has become a generally acceptable method of identifying and classifying microorganisms. Much additional effort will be required to resolve ambiguities in generic concepts and nomenclature of algae based on traditional criteria and sequence data. However, the taxonomic ambiguities themselves and the discovery of new uncultivated thermoacidophilic algae in situ using molecular sequencing (13) indicate that sequence identification of “*Chlorella*” and thermoacidophilic algae inhabiting acidic geothermal environments is warrantable.

The Norris Geyser basin in Yellowstone National Park contains some of the most extensively studied acidic algal mat communities (10, 15). One acidic stream in particular, Nymph Creek, has been the focus of numerous investigations (7, 8, 10, 14, 16, 17,

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TABLE 1. Cloned 18S rRNA gene sequences of algae detected from a microbial mat in an acidic geothermal creek in samples collected from a warm upstream (~45 to 50°C) and a cool downstream (~25 to 35°C) site at different times of the year and in samples collected at different temperatures along the length of the mat at increasing distances from the thermal source on a single day in August

Type of algal 18S rRNA gene	No. of sequences detected ^a												
	Warm upstream site				Cool downstream site				Temp (°C)				
	Apr	Jun	Aug	Dec	Apr	Jun	Aug	Dec	51	47	38	35	30
I	9	7	8	7	0	0	0	0	12	7	0	0	0
II	0	0	3	3	2	6	26	5	0	1	5	13	3
III	0	0	1	2	15	6	5	6	0	0	1	2	7

^a Number of cloned algal sequences detected from seasonal samples collected over an annual cycle from upstream and downstream regions of the mat in April (Apr), June (Jun), August (Aug), and December (Dec). Transect samples were collected on the same day in August at different temperatures along the length of the mat.

19, 22, 38–40, 42, 43); however, microbial species in the Nymph Creek algal mat have been characterized using only traditional microbiological methods. We are using molecular analyses and microsensors to examine microbial populations and processes in the Nymph Creek mat (19, 42–44). Here, we use PCR amplification and 18S rRNA gene sequencing to identify thermoacidophilic red algae and *Chlorella*-like algae that predominate in the mat. We examine algal species composition over an annual cycle and across temperature gradients and use fiber-optic microprobes to describe the microenvironment for algal photosynthesis through the vertical aspect of the mat.

MATERIALS AND METHODS

Site description. Nymph Creek is an acidic (pH, ~2.7) stream located near the Norris Geyser basin of Yellowstone National Park, Wyoming. The creek originates from several permanent and ephemeral 56°C to 60°C hot springs. The shallow (1- to 2-cm-deep), warm spring effluents combine to form a creek that is about 1 to 2 m wide and flows for approximately 130 m before it empties into Nymph Lake. The upstream region of the creek is quite dynamic and changes considerably over time. Shallow springs emerge and recede due to seasonal changes in groundwater. In the shallow, warm effluent springs, temperature gradients from ~52°C to 37°C can occur over short distances (~10 cm) as measured from the center of the effluent channel to the stream edge. Sections of the algal mat that develop in the shallow, warm effluents are impacted by wide temperature changes caused by stochastic factors such as silt deposition, fallen arboreal debris, hail, or trampling by bison, all of which can alter warm water flow over sections of the mat. A temperature gradient also forms along the length of the mat as water flows downstream away from the thermal sources. Nondescript spherical green algal cells ~2 μm to 4 μm in diameter are the predominant cells in mat samples. Near the high-temperature limit of algal growth (~52°C) the "mat" is a thin green biofilm (<1 mm thick). At lower temperatures (~49°C to 35°C) the mat is typically 3 to 5 mm thick.

Temperature data loggers. Temperature data for an upstream and a downstream site in the mat were collected using HOBO XT data loggers (Onset Computer Corporation, Bourne, MA) recording at 2-hour intervals from small cylindrical probes (~3-mm diameter) placed horizontally in the upper layer of the mat. The temperature at the upstream site when the probe was installed was ~50°C. Another probe was installed ~100 m downstream in a low-temperature region of the mat (~30°C). The depth of the water over the probes at both sites was ~1 cm. The probes were periodically repositioned due to shifts in the streambed or other disturbances.

Sample collection for molecular analyses and cultivation. Samples for PCR analysis were collected in 2-ml plastic microcentrifuge tubes by scraping portions of the mat from the streambed using clean weighing spatulas. Samples were frozen on dry ice and stored at -20°C. To evaluate seasonal variations in algal species, samples were collected in December of 1999 and in April, June, and August of 2000 from upstream and downstream regions of the mat near areas containing the temperature probes. To examine the influence of temperature on algal species composition, a set of samples was collected on the same day in August 2001, from sites along the mat at increasing distances downstream from the warm thermal source springs where temperatures were 51°C, 49°C, 38°C, 35°C, and 30°C. Nucleic acids were obtained from mat samples and from culti-

vated algal isolates, using a bead beating cell lysis, nucleic acid extraction protocol as previously described (43).

Samples were collected from a ~50°C region of the mat for enrichments of thermoacidophilic algae. These enrichments were performed in medium 17, pH 2.9, described on the website of the Culture Collection of Algae (SAG), Göttingen, Germany (<http://www.epsag.uni-goettingen.de/html/culturemedia.html>). Cultures were incubated at 50°C under an irradiance of 100 μmol photons m⁻² s⁻¹ provided by cool, white fluorescent lamps in 5% CO₂ in air. Single algal colonies were repeatedly restreaked on SAG agar plates to obtain pure cultures. Pure culture isolates and algal mat samples from an ~50°C site were incubated at 50°C in SAG medium supplemented with 0.2% D-glucose and 0.2% yeast extract without light for 2 weeks to test for the ability of thermoacidophilic algae to grow heterotrophically.

Chlorella-like isolates were obtained by spreading 10-fold serially diluted mat samples collected at ~35°C onto CA agar (37) and successively picking portions of colonies and replating them to obtain pure cultures. The plates were incubated at room temperature under fluorescent lamps. A Nymph Creek *Chlorella*-like isolate, ROF/NC15, and a thermoacidophilic red algal isolate, PC15, were deposited in the University of Oregon Culture Collection of Microorganisms from Extreme Environments (Ecology and Evolution Program, Department of Biology, University of Oregon, Eugene).

18S rRNA gene PCR primers. Unless otherwise indicated, sequences were amplified using the universal eukaryotic primers 5'-GTCAGAGGTGAAATTCTTGATTA-3' and 5'-AGGGCAGGGACGTAATCAACG-3' (25), which amplify an ~700-bp region of the 18S rRNA gene from nucleotides 912 to 1593 of *C. caldarium*, GenBank reference AB090833.

PCR, cloning, sequencing, and taxonomic analysis. Cloning of PCR-amplified rRNA gene sequences was performed using a TOPO-TA PCR cloning kit (Invitrogen, Carlsbad, Calif.), plasmid isolation was performed using a QIAprep Spin Miniprep kit (Qiagen, Valencia, Calif.), and sequencing was performed using a BigDye Ready Reaction Termination Mix (PE ABI, Foster City, Calif.) as previously described (18, 42). Raw sequence data were analyzed using Sequencher software (Gene Codes Corporation, Ann Arbor, Mich.). Preliminary identification of sequences was done using the BLASTn algorithm at the NCBI web site (5). Sequences were aligned using CLUSTAL X (11, 29). Dendrograms were generated using PAUP* 4.0b8 software (46) as described previously (28) using neighbor-joining analyses with uncorrected distance. Bootstrap values were from 1,000 replicates.

Microscale light measurements. The penetration of photosynthetically active radiation (PAR) for oxygenic photosynthesis (λ = 400 to 700 nm) in the algal mat was measured with a fiber-optic scalar irradiance meter (34). Small, undisturbed core samples of the mat were transferred within 1 h to a nearby laboratory, where the light measurements were conducted. The sample was immersed in Nymph Creek water and illuminated with collimated light from a fiber-optic halogen lamp (Schott KL1500). Using a manually operated micromanipulator (Märzthäuser MM33), a scalar irradiance microprobe (35) was inserted into the mat under a zenith angle of 135 and the intensity of PAR was measured in steps of 100-μm vertical distance (34).

Nucleotide sequence accession numbers. GenBank reference numbers of 18S rRNA gene sequences detected in this study are AY422216 through AY422223.

RESULTS

Algal species detected. Three algal 18S rRNA gene phylogenies designated types I, II, and III were repeatedly de-

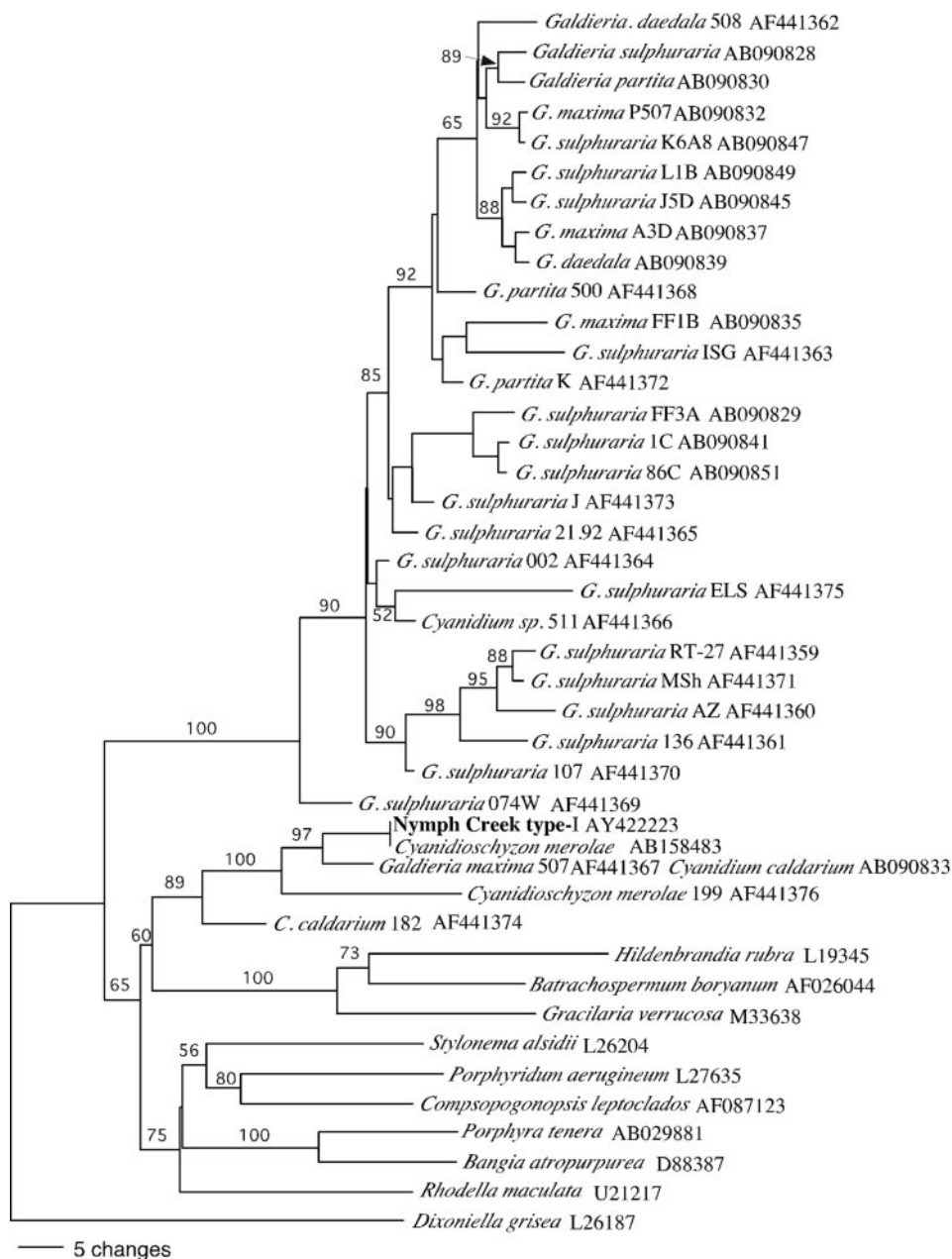


FIG. 1. Dendrogram based on ~680 nucleotides of 18S rRNA genes relating the predominant Nymph Creek thermoacidophilic alga (type I) to other red algal sequences. Bootstrap values (>50%) are indicated at nodes. GenBank reference numbers are listed next to species.

tected among clones from PCR-generated libraries of mat samples (Table 1). Sequence variation within each phylo-type was <1%. Some of this variation may be attributed to unique microbial populations (20, 21); to artifacts associated with the PCR amplification, cloning, and sequencing approach to microbial community analysis (45); or to intragenomic sequence variation (9, 12). Since we have no evidence that suggests that the variation within each group represents distinct populations, we tentatively infer the presence of three algal species in Nymph Creek.

The type I sequence is most similar (>99% to 100% sequence similarity) to three (nonidentical) 18S rRNA gene

sequences within the genome of *Cyanidioschyzon merolae* 10D (36) (GenBank accession no. AB158483, AB158484, and AB158485). The sequences of all algal isolates (six were sequenced) obtained from a 50°C mat sample from the upstream region of the mat were all type I. The spherical, 2- to 4- μ m-diameter cells of each isolate were indistinguishable, by light microscopy, from cells in the mat. The dendrogram (Fig. 1) illustrates the type I sequence groups in a clade containing the sequences of *C. merolae*, *C. caldarium*, and *Galdieria maxima* (Fig. 1).

Type II sequences are most similar (>99% to 100% sequence similarity) to the 18S rRNA gene sequence of the

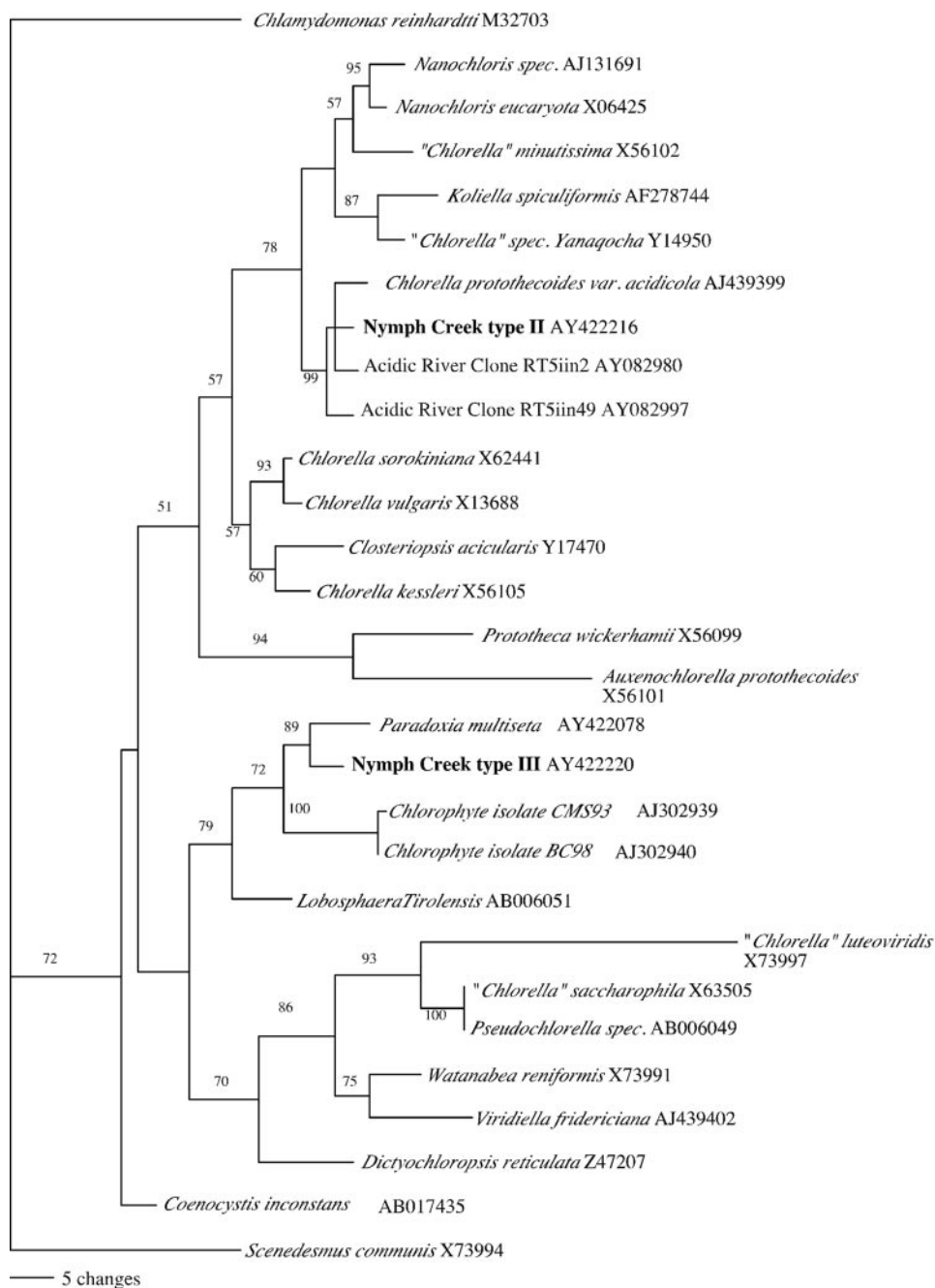


FIG. 2. Dendrogram tree based on ~680 nucleotides of 18S rRNA genes relating predominant Nymph Creek *Chlorella*-like algae (types II and III) to other *Chlorella*-like algal sequences. Bootstrap values (>50%) are indicated. GenBank reference numbers are listed next to species.

acid-tolerant species "*Chlorella*" *protothecoides* var. *acidicola* (2, 30) (Fig. 2). Type II sequences are also nearly identical (>99% sequence similarity) to algal 18S rRNA gene sequences detected by PCR in an acidic river in Spain (6) (Fig. 2). Type III sequences are most similar (98% sequence similarity) to the 18S rRNA gene of *Paradoxia multiseta* and to the 18S rRNA gene sequences of unidentified *Chlorellaceae* strains CMS93 and BC98 (Fig. 2). Cultivated algal isolates whose 18S rRNA gene sequences matched type II clones were obtained from a 35°C mat sample. Type II cells are spherical and indistinguish-

able by light microscopy from cells in the mat. Cultivated isolates representing type III 18S rRNA gene sequences were not obtained.

Absence of *G. sulphuraria*. *G. sulphuraria* strains form a highly supported clade of thermoacidophilic algae that is distantly related to other thermoacidophilic algal species (13, 25) (Fig. 1). Traditional studies suggest that facultative heterotrophic *G. sulphuraria*-like thermoacidophilic algae are abundant in Nymph Creek and similar habitats in the Norris Geysers basin (10, 16, 17). However, none of the 18S rRNA gene sequences de-

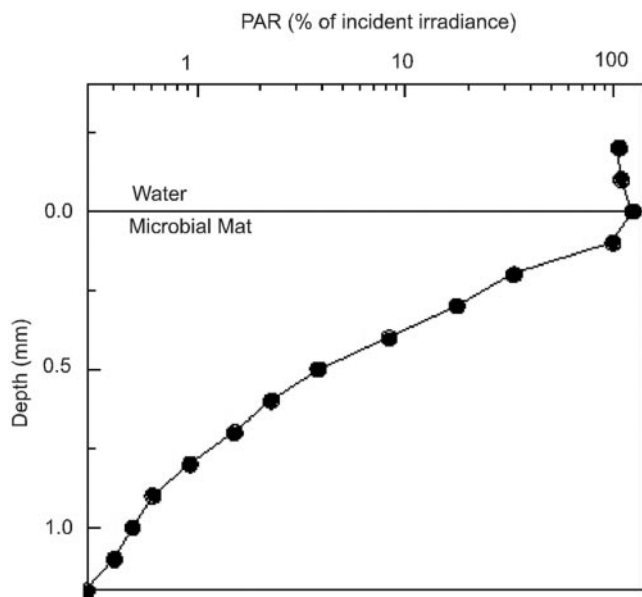


FIG. 3. Light penetration depth profile of photon scalar irradiance (PAR, $\lambda = 400$ to 700 nm) available for oxygenic photosynthesis in the Nymph Creek mat. Data were normalized to the down-welling photon irradiance at the mat surface and are shown on a logarithmic intensity scale.

tected in our analyses grouped within the *G. sulphuraria* clade and type I algal isolates did not grow heterotrophically under conditions where a control *G. sulphuraria* strain (University of Texas at Austin Culture Collection of Algae strain UTEX 2393) grew readily.

It seemed unlikely that PCR primer bias explained the lack of *G. sulphuraria*-like algal sequences in the mat, since complementary priming sites exist on *G. sulphuraria* 18S rRNA gene sequences in public databases and the primer set amplified *G. sulphuraria* 18S rRNA gene sequences from control *G. sulphuraria* genomic DNA. Nonetheless, a second set of universal eukaryotic primers, NS1 and NS2 (50, 53), targeting an alternate region of the 18S rRNA gene was used to amplify DNA from a 48°C sample collected from the upstream region of the mat and DNA from a type I isolate. The sequences of all algal clones from the NS1/NS2 library (10 were sequenced) were $>99\%$ to 100% similar to the sequence of the type I isolate. No *G. sulphuraria* or other algal sequences were detected.

Template competition for universal eukaryotic primers likely favors detection of abundant algal species. Since *G. sulphuraria* sequences were not detected in our clone libraries, we hypothesized that *G. sulphuraria* cells are rare in the mat. *G. sulphuraria* is known to survive in dark endolithic environments via cryptic growth (26); we speculated that they might occupy a dark subsurface niche in the Nymph Creek mat. However, the extent of light (PAR) penetration in the Nymph Creek mat was unknown. We used a light microsensor to examine light penetration. The results show that light available for algal photosynthesis was strongly attenuated over the first 1-mm vertical interval of the mat (Fig. 3), indicating that an extremely low-light subsurface niche exists below this interval. This result is consistent with previous oxygen microsensor

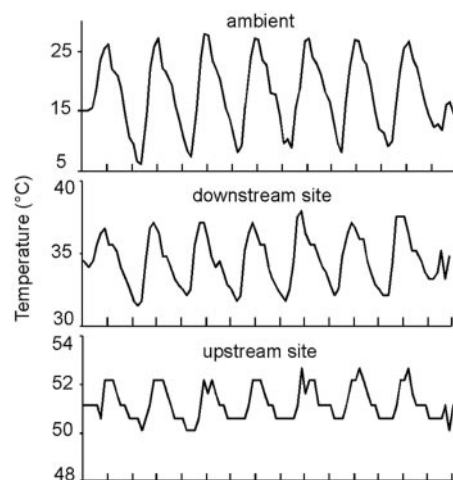


FIG. 4. Example of typical August temperatures recorded every 2 hours by data loggers connected to sensors placed in the surface of the algal mat in an upstream high-temperature region of Nymph Creek, a downstream lower-temperature region, and a shaded site along the creek. Tick marks represent noon and midnight; peaks correspond to midday readings.

analyses of the Nymph Creek mat showing that oxygenic photosynthesis is restricted to the first 1-mm surface interval (38). We attempted to selectively enrich *G. sulphuraria*-like algae by inoculating heterotrophic enrichment broths with upstream ($\sim 48^{\circ}\text{C}$) mat samples and incubating them under dark conditions. No algal growth was observed after 2 weeks. Control broths inoculated with *G. sulphuraria* grew well under the same conditions.

Temperature and algal species composition. To examine the influence of temperature on algal species composition, we analyzed sequences in samples collected at different temperatures from sites at increasing distances downstream from the thermal source springs (Table 1). All 51 sequences obtained from the temperature distribution study were either type I, II, or III. Nineteen of 20 sequences from $\geq 47^{\circ}\text{C}$ samples were type I. All 31 sequences from $\leq 38^{\circ}\text{C}$ samples were type II or type III. This suggests that the distribution of the thermoacidophilic red algae is largely confined to the first ~ 20 m of mat downstream from the thermal source springs, where temperatures are $\geq 39^{\circ}\text{C}$.

Geothermal streams can experience brief surges in temperature due to, for example, geyser eruptions or decreases in temperature caused by runoff from snowmelt (10). Sampling during or following such events can skew interpretations of temperature influence on algal population distributions (10). Data logger readings suggest that Nymph Creek temperature patterns were stable prior to collection of samples for algal temperature distribution analyses. Graphs of temperatures illustrate typical daily temperature dynamics for a week in August prior to sampling (Fig. 4). Diurnal peaks and troughs in stream temperatures correlate with those of ambient temperatures as previously reported (39). The temperature time series confirms that air temperature impacts mat temperatures even in the upstream region despite the continuous input of warm water into the creek channel. Changes in upstream and downstream temperature that occurred over extended time periods

were also noted. These were mainly decreasing temperatures due to shifts in the flow of warm water over the probes caused by sediment deposition or due to cold winter conditions (10).

Seasonal algal species composition. To explore the possibility that algal species composition varied over an annual cycle, we collected samples from both the warm upstream and the cooler downstream regions of the mat four times during an annual cycle (Table 1). Type I sequences were the most abundant clones in each of the four upstream seasonal samples and represented 31 of 40 total algal clones (Table 1). Nine *Chlorella*-like (six type II and three type III) sequences were also detected in upstream seasonal samples. Since *Chlorella*-like algae are not known to thrive at high temperatures, we suspect that the high proximity of cooler niches and shifts in warm water flows account for their sequences in upstream samples. Only *Chlorella*-like, type II and III sequences were detected in samples from the downstream site, 39 of type II and 32 of type III (Table 1). No other algal sequences were detected, and algal species in the Nymph Creek mat did not vary over an annual cycle despite variations in photoperiod that range from ~8 daylight hours in December to ~16 daylight hours in June.

DISCUSSION

Our analyses of 18S rRNA gene sequences from mat samples and cultivated isolates indicate that a single species of strictly photoautotrophic red algae with a spherical morphology, 2- to 4- μm cell diameter, and 18S rRNA gene sequence matching that of a *Cyanidioschyzon merolae* strain (sequence type I) is predominant in the upstream, high-temperature (47°C to 51°C) region of the Nymph Creek mat (Table 1). Unlike *Cyanidioschyzon merolae*, algal cells in the Nymph Creek mat, and isolates with type I sequences cultivated from the mat, are not oval, crescent, or club shaped (1, 41). This incongruity may suggest that not all *C. merolae* strains are aspherical; however, ambiguities between molecular taxonomic analyses and thermoacidophilic algal nomenclature based on determinative characteristics have been noted (1, 13, 25). For example, *Galdieria maxima* is a facultative heterotroph, and its cells are spherical and notably large (10- to 16- μm diameter) compared to those of other thermoacidophilic algal species (1); however, 18S rRNA (25) and *rbcL* (13) gene sequence analyses indicate that *Galdieria maxima* is much more closely related to *Cyanidioschyzon merolae* than to other *Galdieria* species (13). The nomenclature of thermoacidophilic algae is further confounded by instances of identical *rbcL* and 18S rRNA gene sequences that are associated with different genera/species in public databases, for example, *Galdieria maxima* and *Cyanidium caldarium rbcL* genes AY391370 and D63676 and 18S rRNA genes AB090833 and AF441367. Such ambiguities underscore the need for continued clarification of nomenclature and taxonomy of these algae. We tentatively refer to the Nymph Creek (type I) alga as *Cyanidioschyzon merolae* based on 18S rRNA gene sequence.

Nomenclature aside, a predominance of strictly autotrophic spherical thermoacidophilic algae in Nymph Creek is consistent with a traditional microbiological study describing predominantly strictly autotrophic, spherical thermoacidophilic algae in acidic springs in the Norris Geyser basin (15) and with a recent molecular analysis of a low-pH, endolithic algal com-

munity in the Norris Geyser basin describing abundant spherical algae with 18S rRNA gene sequences >99% similar to those of the closest type I relative, the *G. maxima/C. caldarium* sequence (Fig. 1) (46a). A predominance of strict autotrophic thermoacidophilic algae contrasts with other traditional studies suggesting that a single species of facultative heterotrophic (*G. sulphuraria*-like) thermoacidophilic algae predominates in Nymph Creek and springs throughout the Norris Geyser basin (10, 16, 17). We suspect that a bias toward the isolation of *G. sulphuraria* on heterotrophic plating medium (10) and the microscopic resemblance between *G. sulphuraria* and *C. caldarium* cells likely account for inferences of *G. sulphuraria* being widespread in the Norris Geyser basin.

Nymph Creek is routinely described in terms of a *C. caldarium* mat. It is notable that *Chlorella*-like algal sequences were the predominant type detected over much of the length of the Nymph Creek mat, ~100 of 130 m downstream from the 38°C site (Table 1). The reason for the scarcity of type I sequences below the 38°C site is unclear. We speculate that a low washout rate of type I algal cells from upstream and competition from *Chlorella*-like algae at lower temperatures are significant factors. A shift from red algae to *Chlorella*-like algae at ~39°C is consistent with a traditional study of a similar Yellowstone stream showing a sharp transition from thermoacidophilic red algae to "*Chlorella*" species at ~38°C due to seasonal shifts in temperature (17). Abrupt transitions from red algae (green-pigmented mat) to diatoms (brown-pigmented mat) at 39°C have also been noted in geothermal springs in Japan (10).

Although *Chlorella*-like algae have been noted in Nymph Creek and other acidic areas of the Norris Geyser basin, species have not been identified (10, 39). The 18S rRNA gene sequence of the type II strain suggests that it is *Chlorella protothecoides* var. *acidicola*, an acid-tolerant alga that was originally isolated from low-pH soil (2, 30). Sequences nearly identical to type II have also been detected in an acidic river in Spain (6). This suggests that *C. protothecoides*-like algae may be widespread in low-pH aquatic habitats (23). A second *Chlorella*-like alga (type III) also appears to be abundant in the Nymph Creek mat. The 18S rRNA gene sequence of this alga suggests that it is a relative of *Paradoxia multisita* AY422078 (98% sequence similarity). However, the alga harboring the type III sequence has not been isolated. Sequence information suggests that neither Nymph Creek *Chlorella*-like alga is a "*Chlorella*" species *sensu stricto* (30, 32, 33).

Molecular analyses are beginning to provide the first few cultivation-independent assessments of algal species that thrive in acidic geothermal habitats (13, 25). Accurate information about the identity of algae and other microbial species *in situ*, and an understanding of their distribution, provides a base on which realistic concepts of the ecology of these extreme ecosystems can be constructed. Knowledge of species composition will likely enhance interpretations of past and future studies of bulk properties in the often-studied Nymph Creek mat.

ACKNOWLEDGMENTS

We thank the National Science Foundation (Microbial Observatory Grant 9977922), the Research Institute for Children at New Orleans, the Danish Natural Science Research Council, and the Thermal Biology Institute at Montana State University for financial support.

This project was conducted under the direction of the Yellowstone Center for Resources following the guidelines for research in Yellowstone National Park.

We thank John Varley and Christie Hendrix for assistance.

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