Unexpected diversity and complexity from the Guerrero Negro hypersaline microbial mat

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ABSTRACT

We applied nucleic acids-based molecular methods, combined with estimates of biomass (ATP), pigments and microelectrode measurements of chemical gradients, to map microbial diversity vertically on the mm-scale in a hypersaline microbial mat from Guerrero Negro, Baja California Sur, Mexico. To identify the constituents of the mat, small-subunit ribosomal RNA genes were amplified by PCR from community genomic DNA extracted from layers, cloned and sequenced. Bacteria dominated the mat and displayed unexpected and unprecedented diversity. The majority (1336) of 1586 bacterial 16S rRNA sequences generated were unique, representing 752 species (≥97% rRNA sequence identity) in 42 of the main bacterial phyla, including 15 novel candidate phyla. The diversity of the mat samples differentiated according to the chemical milieu defined by concentrations of O$_2$ and H$_2$S. Chloroflexi formed the majority of the biomass by percentage of bulk rRNA and of clones in rRNA gene libraries. This result contradicts the general belief that Cyanobacteria dominate these communities. Although Cyanobacteria constituted a large fraction of the biomass in the upper few mm (>80% of total rRNA and photosynthetic pigments), Chloroflexi sequences were conspicuous throughout the mat. Filamentous Chloroflexi were identified by fluorescent in-situ hybridization within the polysaccharide sheaths of the prominent cyanobacterium Microcoleus chthonoplastes in addition to free-living in the mat. The biological complexity of the mat far exceeds that observed in other polysaccharide-rich microbial ecosystems, such as human and mouse distal guts, and suggests that positive feedbacks exist between chemical complexity and biological diversity.
INTRODUCTION

Microbial mats are benthic aquatic ecosystems fueled by light energy and composed of microbial cells attached to extracellular polymeric material and mineralized scaffolds in visible mm-scale layers (12). Unlike stromatolites, which have a biotic mechanism for calcification (49), microbial mats become layered due to occasional sedimentation and re-growth. Microbial mats and stromatolites are found in the fossil record dating back 3.4 billion years (60), and are thought to have significantly influenced the composition of the atmosphere with production of O$_2$, H$_2$ and CH$_4$ (21). Ancient and modern mats share properties inherent to their structure. For instance, different wavelengths of light penetrate differentially, gas exchange with the atmosphere occurs at the surface, an organic carbon-based matrix provides a scaffold for growth, and sedimentation occasionally buries the surface, which is then overgrown, leading to layering. Comparisons of biosignatures in modern and fossilized mats seek to describe ancient biogeochemical cycles and the microbial activities of ancient communities (34, 57, 58).

The microbial mats within the hypersaline lagoons of the Exportadora de Sal SA saltern in Guerrero Negro, Baja California Sur, Mexico, cover an extensive area of artificial shallow lagoons protected from tidal disturbance by levees. Biogeochemical studies of these mats have shown that oxygen and light, as well as photosynthetic capacity, are rapidly depleted with depth. Degradation of organic matter occurs largely with two processes, fermentation and substrate oxidation through sulfate reduction (12). Both processes take on unusual characteristics in the mats. Fermentation contributes to molecular hydrogen release into the overlying water column even in the presence of
oxygen, such that bubbles of mixed oxygen and hydrogen gases form on the surface of the mat (21). In addition, the highest rates of sulfate reduction occur in the upper, oxygen-rich layers of the mat (5). Known sulfate-reducing bacteria (SRBs) of the δ-Proteobacteria occupy the anoxic zone, however, which suggests that novel groups of SRBs reduce sulfate in the mats aerobically (51).

Despite the intriguing biogeochemistry of hypersaline microbial mats, and their importance as model systems for studies of the early Earth (21), the composition of the microbiota has not been surveyed comprehensively with culture-independent molecular methods. Classic microbiological studies and limited molecular studies have shown that Cyanobacteria dominate the surface layers and revealed 5 other of the main bacterial phyla (phylogenetic divisions: Chloroflexi, Spirochaetes, Proteobacteria, Bacteroidetes, Firmicutes), and thus indicated a relatively simple community with little deep evolutionary diversity (9, 39-42, 51, 57). The biological “simplicity” of the mats was the basis for their recommended use as model systems for metagenomic analyses (4). The dominance of Cyanobacteria and the biological simplicity of the community have not been verified using culture-independent methods, however.

The aim of this study was a more comprehensive description of the microbial diversity within the mats and how that diversity is distributed in relation to depth and chemical gradients characteristic of the depth profile. We studied a limited area of one mat intensively. In situ gradients of O₂ and H₂S concentrations and pH, measured on a μm-scale with microelectrodes, provided a backdrop of vertical chemical gradients onto which we mapped the biological data. ATP concentrations were measured to provide an estimate of living biomass distribution throughout the mat. Pigment concentrations
measured by high-pressure liquid chromatography (HPLC) offered a view of the distribution of oxygenic chlorophyll $a$ (Chl $a$) containing Cyanobacteria in relation to bacteriochlorophyll (BChl)-bearing anoxygenic photosynthetic bacteria. A survey of rRNA genes provided a culture-independent assessment of dominant organisms. For each of 10 layers that divided the mat into a mm-cm scale depth profile, the composition and diversity of communities was determined by sequence analysis of 16S rRNA genes generated by PCR with universal and bacteria-specific primers from community genomic DNA. We used RNA extraction and quantitative hybridization with group-specific probes as a PCR-independent verification of the abundance of the dominant group identified by sequence analysis, and of the Cyanobacteria. In addition, we visualized the morphologies and associations of these bacteria using fluorescent in-situ hybridization (FISH) with tyramide signal amplification to overcome the intrinsic fluorescence of the mat. These studies collectively revealed unexpected diversity, complexity and structure within the mat.

MATERIALS AND METHODS

Sample collection. We studied the microbial mat underlying Pond 4 (near 5) of the Exportadora de Sal SA, a solar salt works located at Guerrero Negro, Baja California Sur, Mexico (see ref. (38) for site details). The mat is covered by ~1 m of brine with a salinity of ~80‰. Collections were made in June and October of 2001, at 4 am (night) and 1 pm (day). We collected replicate cores (1 cm x 6 cm) from a 0.4 m$^2$ area of mat harvested approximately 10 m from the levee. Upon collection, cores were treated randomly in one of three different ways. Cores destined for DNA extraction, ATP and pigment analyses were sliced horizontally in 1 mm increments (top to a depth of 6 mm)
and 1.5-cm depth increments (remainder of core, to a depth of approximately 60 mm), then the corresponding layers from 5 different cores were pooled, homogenized manually with a polypropylene pellet pestle (Thomas Scientific, Swedesboro, NJ) and frozen in liquid N\textsubscript{2}. Cores intended for RNA extraction and analysis were sliced length-wise and frozen in liquid N\textsubscript{2}. Those intended for microscopy assays were fixed in 4\% paraformaldehyde in pond brine for 4 hours and preserved in 50:50 10X PBS:ethanol at 4°C.

**In situ microelectrode measurements.** O\textsubscript{2} concentration, H\textsubscript{2}S concentration and pH within the microbial mat were measured at 200-μm intervals on the sample collection days in October 2001 using a diver-operated microprofiler (Unisense, Århus, Denmark). A total of 31 profiles were taken over the course of the diel; the data shown are representative of eight qualitatively similar mid-afternoon profiles taken at three different mat locations. All profiles were made within the same 0.25 m\textsuperscript{2} area from which the mat cores were taken. The Clark-type oxygen microelectrode and amperometric sulfide microelectrode were calibrated as previously described (3, 38). A three-point calibration (pH 4, 7, 10) was performed for the pH microelectrode.

**ATP analysis.** We measured ATP concentrations, a biomarker for biomass (28), in each section of October and June daytime mat by using the luciferase enzyme assay (Molecular Probes, Carlsbad, CA). ATP was extracted from homogenized mat samples by treatment with 0.5 M H\textsubscript{2}PO\textsubscript{4} for 20 min on ice followed by centrifugation at 420 rcf to remove suspended material and neutralization with addition of 1 M NaOH to a final pH of 7.4. Mat samples were washed twice with 1.7 M glucose solution, by centrifugation at
420 rcf followed by replacement of the supernatant with the glucose solution, prior to extraction, to remove salts that were found to inhibit the luciferase reaction. Light emitted from the ATP-activated luciferase reaction was quantified with a scintillation counter (Beckman Coulter, Inc., Fullerton, CA).

**Pigment analysis.** Pigments were extracted from June and October daytime samples by sonication in acetone:methanol (7:2 v/v) in the dark, followed by centrifugation, filtration (0.45μm, Whatman) and injection into an HPLC column (25 cm-by-4.6 mm 5-μm Discovery C18 (Supelco, Bellefonte, PA)) as described in (37) with these modifications: the time at 100% solution B was extended to 16 min to ensure that all long-tailed quinones were completely eluted. The HPLC-DAD (diode array detection) system consisted of an Agilent 1100 series binary pump (model G1312A), vacuum degasser (model G1379A), manual injector (model G1328A), and diode array detector (model G1315B). The data were analyzed with ChemStation for LC 3D software (version A.08.03, Agilent technologies, Waldbronn, Germany). Pigments were identified by absorption spectra and retention times. The concentrations of undegraded (bacterio)chlorophylls were determined from the areas of the elution peaks using the equation \[ m = F \cdot A \left( e_m d \right)^{-1}, \] where \( m \) = the mass of (B)Chl in mg, \( F \) = rate of solvent flow through the column (1ml min\(^{-1}\)), \( A \) = the area of the elution peak, \( e_m \) = the extinction coefficient in units of L mg\(^{-1}\) cm\(^{-1}\), and \( d \) = the detection length of the diode array detector (1 cm). The following extinction coefficients (L mg\(^{-1}\) cm\(^{-1}\)) were used: Chl \( a \), 79.2(33), BChl \( a \), 60 (45); BChl \( c \), 86 (55); and BChl \( d \), 82(55).

**DNA extraction, polymerase chain reaction (PCR) and cloning.** DNA was extracted from frozen mat samples using a bead-beating protocol (14). For each sample,
six replicate 25 μL PCR operations were performed, each containing 100-200 ng purified genomic DNA, 100 mM Tris-HCl (pH 8.3), 500 mM KCl, 20 mM MgSO₄, 200 μM dNTPs, 200 μM each forward and reverse primer, 1M betaine, 800 μg/ml bovine serum albumin, and 1U Taq DNA polymerase (Invitrogen). We used the bacteria-specific forward primer 8F 5’-AGA GTT TGA TCC TGG CTC AG-3’, universal primer 515F 5’-GTG CCA GCM GCC GCG GTA A-3’, archaeal-specific primer 333Fa 5’-TCC AGG CCC TAC GGG-3’, all coupled with universal reverse primer 1391R 5’- GAC GGG CGG TGW GTR CA-3’ (31). Cycling conditions were 94°C for 2 min; followed by 20 cycles of 94°C for 1 min, 45 seconds at 55°C, and 2 min at 72°C, with a final extension period of 20 min at 72°C. Replicate PCR products were pooled, and amplicons were gel-purified (Qiagen, Valencia, CA), cloned into TOPO TA pCR4.0, and transformed into E. coli TOP10 cells (Invitrogen, Carlsbad, CA). The majority of the sequences were generated from October daytime samples, after a pilot study of June day and night samples did not show statistically significant diurnal differences due to the unexpectedly high diversity (these sequences are included in the final dataset). A subset of sequences was generated with the forward primer 333Fa that is usually employed to amplify Archaea. Extraction controls (no material added) did not produce visible PCR products or colonies. For each sample/primer pair combination, 96 colonies were selected; strands of plasmids were sequenced using vector-specific primers with an ABI 377 DNA sequencer (BigDye terminator ready reaction mixture, PE Applied Biosystems, Inc.).

**Sequence and phylogenetic analysis.** SSU rRNA sequences were edited and assembled into consensus sequences using PHRED and PHRAP aided by XplorSeq (Dr. 9
Daniel Frank, unpublished), and bases with a PHRAP quality score of < 20 were trimmed. Chimeras were detected using Bellerophon (23). Non-chimeric consensus sequences were named according to the layer they originated from (01-10), the time of collection (D for Day, N for night), the month of collection (1=June, 2=October), the primer pair (X, Z or ZZ= 8F-1391R, Y or YY= 515F-1391R, B= 333Fa-1391R; different symbols for the same primer pair indicate different authors generated the sequence), and the clone number (01-96). The layers correspond to specific depths (1: 0-1 mm, 2: 1-2 mm, 3: 2-3 mm, 4: 3-4 mm, 5: 4-5 mm, 6: 5-6 mm, 7: 6-13 mm, 8: 13-26 mm, 9: 26-39 mm, 10: 39-60 mm). For example, clone 08D2Z44 was collected during the day in October from 13-26 mm depth and generated with 8F-1391R. Sequences have been deposited under GenBank accession numbers DQ329539-DQ331020 and DQ397339-397511.

Non-chimeric sequences were aligned using the Arb software package (36), based on an initial alignment described in (26). The alignment is available as Supplementary Online Information. Distance matrices generated in Arb (with Olsen correction) were used to cluster sequences into operational taxonomic units (OTU's) by pair-wise identity (\%ID) with a furthest-neighbor algorithm and a precision of 0.01 implemented in DOTUR (52). We used DOTUR to determine OTU frequencies in mouse cecal and human colonic 16S rRNA gene sequence datasets using Arb alignments provided by the authors (15, 32). Simpson’s diversity index and collector’s curves were calculated using EstimateS (8).

Assignment of the majority of sequences to their respective phyla was based on their position after parsimony insertion to the Arb dendrogram (omitting hypervariable
portions of the rRNA gene with the lanemaskPH provided with the database; see Supplementary Online Material). Sequences that did not fall within described phyla were further characterized. Phylogenetic trees including the novel sequences and reference taxa (20) were constructed by evolutionary distance (test version 4.0b2 of PAUP*, a neighbor-joining algorithm with either Kimura two-parameter correction or maximum-likelihood correction with an empirically determined gamma distribution model of site-to-site rate variation and empirically determined base frequencies), parsimony (test version 4.0b2 of PAUP*; heuristic search), and maximum likelihood (fastDNAml) analyses. Bootstrap resampling was used to test the robustness of inferred topologies. Novel candidate phyla, designated “GN” 01-15, were defined by generally accepted criteria that 1) there must be three or more sequences from independent PCR products; 2) sequences must be a minimum of 1000 bp; and 3) there must be high levels of support in phylogenetic analyses (20, 24, 46). However, four of the novel candidate phyla (GN6, GN12, GN13, GN14) met only two of these criteria (the sequences were <1000bp) and await confirmation with longer sequence reads.

To cluster the communities from each layer, we used the UniFrac computational tool (35). The Arb alignment (excluding hypervariable regions) containing all 1586 sequences was used to construct a maximum likelihood (ML) tree using RAxML (54). The ML tree was annotated according to the layer from which each sequence was derived, and the fraction of tree branch length unique to any one layer in pair-wise comparisons (the UniFrac metric) was calculated. Microbial communities from individual layers were clustered by application of the unweighted pair-group method with arithmetic mean method (UPGMA) to the UniFrac metric matrix.
**Fluorescent in situ hybridization of Chloroflexi.** Mat samples were disrupted gently with a pestle, dehydrated in an ethanol series and adhered to silane-coated glass slides using Cell-Tak™ (BD Biosciences, Inc.). Cells were permeabilized with lysozyme (10 mg/mL, 30 min, 37°C), followed by achronapeptidase (60 U/mL, 1 Hr, 37°C) and mutalysin (10X, 30 min, 37°C) (Sigma Aldrich, Inc.). Hybridizations were performed with a buffer containing 30% formamide, 0.9 M NaCl, 20 mM Tris-HCl pH 8.0, 0.01% SDS, and 50 ng biotin-labeled probe for 2 h at 46°C. Signal intensity was boosted using the “TSA™ Kit #22 with HRP-streptavidin and Alexa Fluor® 488 tyramide” following the manufacturer’s instructions (Molecular Probes, Carlsbad, CA). The probes used in this study were Chloroflexi-specific ChloroflexiB941 5’-AAA CCA CAC GCT CCG CT-3’ (18) and bacteria-specific EUB338 5’- GCT GCC TCC CGT AGG AGT-3’ (1). Samples were counter-stained with 10 μg 4’6’-diamidino-2-phenylindole (DAPI)/ml, and mounted with antifadent (CitiFluor Ltd., Leicester, England). Images were generated by laser confocal microscopy (Leica Microsystems, Bannockburn, IL) using a 488 nm excitation laser and a 350 nm excitation laser (DAPI). Probes and protocols were tested against reference strains (Chloroflexi: *Thermomicrobium roseum* ATCC27502, *Chloroflexus aurantiacus* ATCC23779, *Roseiflexus castenholzii* obtained from S. Hanada (19), *Herpetosiphon aurantiacus* ATCC23781; Chlorobi: *Chlorobium tepidum* ATCC49652; Proteobacteria: *Pseudomonas aeruginosa* ATCC14205, *Escherichia coli* ATCC29181; Firmicutes: *Clostridium sporogenes* ATCC13663, *Bacillus subtilis* ATCC12432; Archaea: *Sulfolobus acidocaldarius* ATCC33909).

**Quantitative dot-blot hybridization of rRNA from Chloroflexi and Cyanobacteria.** To extract bulk RNA from the mat, triplicate mat samples (100 mg)
from three depths (0-4mm, 5-9mm, 20-30mm) were washed with 1.7M glucose amended with Rnase OUT (Sigma Aldrich, St. Louis, MO) to remove excess salts, and resuspended in 750 µL of pH 4.7- buffer containing 100mM EDTA, 20mM Na-Acetate and 0.5 uL/mL Rnase OUT. Aliquots (100 µL) were further homogenized by bead-beating (2 min on high with 0.1 mm zirconium beads, Mini-BeadBeater-8, BioSpec Products, Inc., Bartlesville, OK) in Tri-Reagent (Sigma Aldrich, St. Louis, MO). RNA was extracted similarly but without the glucose wash, from the pure cultures previously mentioned under the FISH protocol. In addition, RNA was obtained from the following cyanobacteria: *Synechococcus* sp. UTCC477, *Lyngbya* sp. UTCC592, and *Oscillatoria* sp. UTCC487. To obtain SSU rRNA from reference organisms without cultured representatives, RNA was synthesized by transcription from rRNA gene clones. We transcribed RNA from 2 cyanobacterial clones (01D2Z20, 05D2Z68), and five Chloroflexi clones (10D2Z49, 08D2Z44, 05D2Z83, 05D2Z90, 05D2Z56) chosen to represent the diversity of those groups. Plasmids with cloned rRNA gene inserts were prepared from *E. coli* cultures using a Qiagen mini-prep kit. Plasmids were linearized by restriction enzyme digest (*SpeI*, 37°C, 2 h). Transcription reactions at 37°C overnight contained per 100 µL reaction: 6-8 µg of linearized plasmid, 52.5 U yeast pyrophosphatase, 7 µL Rnase OUT, T7 RNA polymerase, 28 mM each ribonucleotide, 0.175 M MgCl₂ buffer. Newly synthesized RNA was ethanol-precipitated and isolated by electrophoresis in denaturing polyacrylamide gels, followed by nondenaturing ion exchange chromatography on HiTrap Q HP Sepharose (Amersham Pharmacia) (29).

RNA from mat samples and reference strains was blotted in triplicate onto nylon membranes (Magna Charge, Micron Separation Inc., Westboro, MA), and bound by
baking at 80°C. Probes were labeled with γ-ATP with T4 polynucleotide kinase following the manufacturer’s instructions (Invitrogen, Carlsbad, CA). Washing temperatures for each probe were determined empirically according to ref. (11), with modifications: we used 200 µL assay volumes in 96-well plates and controlled temperature with a thermocycler. Membranes were hybridized as described in (11). Signal intensity of the blots was determined with a Phosphorimager (Molecular Dynamics, Carlsbad, CA) and the program ImageQuant (Amersham Biosciences, Piscataway, NJ). Signal intensities were analyzed as described in (47, 48).

RESULTS

Depth gradients of H$_2$S, O$_2$ and pH. The *in situ* chemical environment of the mat was profiled with microelectrode measurements of O$_2$ and H$_2$S concentrations and of pH from the surface of the mat to a depth of 6 mm in 200 µm increments. The high spatial resolution of these measurements provides a view of the variability of the local chemistry. Based on our microelectrode measurement profiles, the mat can be divided into three distinct habitats that serve as a backdrop for the spatial organization of the microbial community (Fig 1A). These zones are: the “oxic zone”, ranging from the top of the mat to a depth of 2 mm and characterized by diurnally fluctuating concentrations of O$_2$; the “low-H$_2$S zone”, ranging in depth from 2 mm to 6 mm, where H$_2$S levels are drawn down diurnally; and the “H$_2$S-rich zone”, the largest zone ranging from 6 mm to the bottom (~60 mm), where concentrations of H$_2$S are permanently high (Fig 1A). The coefficient of variation for the O$_2$ measurements was greatest at an average depth of 2.5 mm, where the average O$_2$ concentration was low (28 µM, Fig 1A). This is an indication
that the deepest penetration of oxygen into the mat is also the depth at which oxygen concentrations are the most variable (measured range, 0 to 522 μM) during the day. At night, the mat is completely anoxic.

**Biomass and pigment distributions by depth.** To determine the depth distribution of biomass in the mat, we measured the concentration of ATP per gram, from the top of the mat to the bottom, in June and October. ATP is the energy currency of all cells, and it therefore serves as a proxy for biomass (28). ATP concentrations were highest in the oxic zone (1343 ± 400 ng ATP g⁻¹; Fig 1B), and tapered off rapidly with depth, to 322 ± 50 ng ATP g⁻¹ in the lower, H₂S-rich, zone. Integrating by depth, the substantially greater volume of the lower H₂S-rich zone results in 5-fold greater overall biomass per unit surface area: 27.8 μmole ATP cm⁻² in the lower zone versus 5.0 μmole ATP cm⁻² for the combined oxic and low-H₂S zones. The ATP concentrations and depth profiles of the October and June sampling dates were similar.

We determined concentrations and distributions of photosynthetic pigments using HPLC. Pigment concentrations were highest in the oxic zone, as expected, yet we detected pigments at all depths (Fig 1C). The cyanobacterial pigment Chl a was an order of magnitude more abundant than BCHls (BChl a, d and c) in the oxic zone.

**Estimated richness and coverage.** Two rRNA gene clone libraries were constructed from each of the 10 layers, one with bacteria-specific primers (8F, 1391R) and one with universal primers (515F-1391R). RFLP screening revealed unexpectedly high levels of diversity for each 96-clone library, therefore all clones were sequenced bi-directionally. When the universal libraries from all layers were combined, the Bacteria:Archaea:Eukarya ratio was 57:7:1. In this report we focus on the bacterial
sequences. Archaeal and eucaryal sequence data are described elsewhere (J. Spear, R. Ley and N.R. Pace, unpublished). Bacterial sequences encountered in the libraries constructed with archaeal primers were included in the final dataset discussed here. The two forward primers (515F, 8F) used to generate the majority of the sequences yielded an equivalent proportion of bacterial phyla ($\chi^2$, p<0.05), an indication that our coverage was not significantly biased by the primers pairs that we used. For subsequent analyses, all sequences were combined on the basis of the layer from which they originated.

To assess the coverage and the richness of clone libraries combined by layer, we employed collector’s curves for coverage, the non-parametric estimators Chao1 and Ace1 for richness, and computed Simpson’s Index to estimate the evenness of community composition (see (52)). The total bacterial 16S rRNA gene sequences (n= 1586) consisted of 1336 unique sequences and 752 phylotypes defined by a minimum threshold of 99% ID. In all subsequent analyses, we report masked sequence pairwise identities (%ID). The hypervariable regions of sequences were masked for alignment purposes because these regions cannot be aligned with certainty, particularly across the large phylogenetic distances encountered in this dataset. Generally, 99% ID is equivalent to ~97% ID when the entire sequence length is used (e.g., see dataset of 11,831 bacterial 16S rRNA genes in (15)), often taken to indicate species-level variation. Figure 2A shows the frequency of observed taxa with %ID thresholds ranging from 90% to 100%. The Chao1 and the Ace1 richness estimates for phylotypes ranging from 90% ID to 100% ID yielded equivalent richness curves (Fig 2A). Both richness estimators indicated >10,000 unique sequences based on the distribution of observed sequences. Collector’s curves for taxa with %ID’s greater than 90% indicate that our coverage of the diversity
was not comprehensive, since the curves did not begin to asymptote (Fig 2B). Together, the richness estimators and the collector’s curves indicate a high degree of diversity, most of which likely remains undescribed. Bacterial diversity was uneven in the upper layers sampled, which had a subset of dominant bacteria, and became more even with depth (Simpson’s Index vs. depth, $R^2=0.63$, $p<0.005$).

**Bacterial diversity and distribution.** The phylum Chloroflexi dominated clone libraries numerically and included the highest proportion of sequences on average (Fig 3A) from each chemically defined zone (Fig 3B,C,D). The Proteobacteria and the Bacteroidetes were the second most represented phyla in each zone. Cyanobacterial sequences comprised at most 10% of the total sequences in a chemical zone (Fig 3B) and were obtained from the oxic zone only. Figure 4 shows the phylogenetic relationships of the Chloroflexi and cyanobacterial sequences obtained from the mat in relation to cultured representatives and clones obtained by culture-independent methods from other environments. 24% of the Chloroflexi sequences were obtained from the surface layers and were close relatives of known photosynthetic organisms, such as *Chloroflexus* and *Chlorothrix* spp. previously described for these mats (30, 41) (Fig 4A). Another group of Chloroflexi sequences (4.5% of Chloroflexi sequences) with no cultured close relative was only obtained from the surface layers (Fig 4A). This result indicates that the Chloroflexi may include additional unrecognized photosynthetic members. The other Chloroflexi groups recovered from the mat had average depth distributions extending below the oxic zone. These groups include oxygen-tolerant members, however, since members of all groups were obtained from the oxic zone. The majority of sequences have no closely related cultured representatives from which properties can be inferred.
61% of the cyanobacterial sequences obtained from the mat were members of a novel group of Cyanobacteria whose closest relative was obtained from a hot spring mat (Fig 4B). 16% formed a group affiliated with *Leptolyngya* spp.

In addition to the Chloroflexi and Cyanobacteria, the 16S rRNA gene sequence analysis revealed 28 other previously described phyla (Fig 3). Half of these were candidate phyla, so termed because they are known only by 16S rRNA gene sequences and do not contain representatives, which have been cultured in the laboratory and from which physiological and metabolic properties can be inferred. This study expanded the known habitat space and diversity of several candidate phyla. For example, candidate phylum KSB1, previously represented by <10 sequences encountered in surveys of brackish water sediments (59), cave sediments (22) and a bioreactor treating 4-methylbenzoate (62), was expanded significantly with the addition of 35 sequences from the mat.

The majority of sequences could be assigned unambiguously to known phyla, however, 119 sequences remained (7.5% of total) that did not have affiliate with any known phylum. Phylogenetic analysis of these unaffiliated sequences revealed 15 novel candidate phyla termed GN01 – GN15. Several of the candidate phyla include sequences that were previously deposited in GenBank and for which there is no described affiliation (Table 1; a dendogram is available in the Arb database in the Supplementary Online Material). Most of the novel GN candidate phyla include sequences derived from several different layers, and therefore several separate PCR products. Novel GN candidate phyla were detected at all layers in the mat, and over half were detected in the oxic zone (Table 1).
Figure 5 shows the depth distribution of members of the 14 most commonly observed bacterial phyla. The abundance of each is shown as the percentage of sequences at each depth, calculated as a fraction of the total number of sequences obtained for any given phylum. The Chloroflexi are distributed fairly evenly with depth, except for a relative reduction in abundance in the transitional, low-$H_2S$ zone. Phyla that were most abundant in the oxic zone included the Cyanobacteria, Proteobacteria, Bacteroidetes, Spirochaetes, Verrucomicrobia and candidate phylum GN01. Several known candidate phyla (KSB1, OP10, OP5, OD1) had highest abundances in the low-$H_2S$ zone, just below the oxic layer. The Firmicutes exhibited a marked bimodal distribution, abundant in the oxic zone and in the lower portion of the mat.

**Community similarities.** In order to compare the communities within the chemical zones characterized by the microelectrode measurements of $O_2$ and $H_2S$, we used the recently-developed UniFrac metric analysis (35). UniFrac measures the phylogenetic distance between pairs of communities represented by sequences in a phylogenetic tree as the fraction of branch length of the tree that leads to descendants from either one community or the other but not both (35). A phylogenetic tree containing all of the microbial mat sequences ($n=1585$) was generated, each sequence was annotated according to the analytical layer from which it was derived, and the UniFrac for all combinations of pairs of communities ($10 \times 10$) was computed. The layers were then clustered according to their pair-wise UniFrac metrics using UPGMA. This analysis revealed three main clusters of related communities that match the zones delimited by the chemistry of the mat. The communities in layers 0-1 mm and 1-2 mm clustered together, as did the 4 communities obtained from layers 2-6 mm, and the 4 communities from the
lower layers clustered together (6-60 mm, Fig 3E). The robustness of the inferred UniFrac tree topology to the presence of specific communities represented was confirmed by jackknife analysis (p<0.001).

**Quantitative RNA hybridizations with probes specific for Chloroflexi and Cyanobacteria.** In order to test the abundances of Chloroflexi and Cyanobacteria indicated by the clone library results that may be subject to PCR and other biases (24), we conducted quantitative rRNA dot-blot hybridizations with oligonucleotide probes based on the sequences. We used pure cultures of members of the Chloroflexi and Cyanobacteria to generate reference rRNA. To represent novel Chloroflexi clades detected by rRNA gene analysis for which no cultures are available, we synthesized rRNA by *in vitro* transcription from selected clones. Overall, the majority of the community rRNA extracted from the mat hybridized with the Chloroflexi-specific probe, confirming the dominance of the Chloroflexi observed in the rRNA clone libraries. However, the majority of the total rRNA in the oxic zone was cyanobacterial (87%±2.7), while the Chloroflexi likely comprised the majority of the remainder (22%±1.7). Together these percentages exceed 100%, which indicates that some non-specific binding of the probes may have occurred. In the low-H$_2$S zone, the proportion of cyanobacterial rRNA dropped to 28%±3.6 while the proportion of Chloroflexi RNA increased to 41% (0.8). Deep in the H$_2$S-rich zone of the mat, cyanobacterial abundance declined further to 10%±2.2, while Chloroflexi remained a dominant proportion of the overall biomass (32%±4.5).

**Visualization of Chloroflexi by fluorescent *in-situ* hybridization.** Because of their abundance, the Chloroflexi bacteria must have a major influence on the nature of
this mat. In order to gain insight into the ecology and spatial organization of these organisms, we visualized the Chloroflexi bacteria in upper and lower layers of the mat using Chloroflexi-specific fluorescent oligonucleotide probes with in situ hybridization and confocal microscopy. Tyramide signal amplification combined with the use of Alexa fluors boosted the signal above the high levels of background autofluorescence. Chloroflexi bacteria were seen within the exopolysaccharide sheaths of the conspicuous filamentous cyanobacterium *Microcoleus chthonoplastes* in the upper layers of the mat (Figs. 6 A, B, C). The invasion of the trichomes seemed specific for Chloroflexi bacteria. No other bacteria were seen within the sheaths, although many were visualized by DAPI staining on the outside of the sheaths (not shown). The filaments of cyanobacterial cells were often observed disrupted in the presence of the Chloroflexi bacteria (Fig. 6B). In the lower, anoxic zones of the mat, the Chloroflexi were visualized as filamentous bacteria pervading the polymeric matrix of the mat, but they were not observed inside of sheaths (Fig. 6D).

**DISCUSSION**

Microbial mats occur worldwide in shallow aquatic environments where high salinity or temperature precludes the establishment of algae or aquatic plants that may overgrow the mats, or of grazers that may otherwise consume them (17). Prior to the evolution of such organisms, microbial mat-like structures were widespread around shallow seas and lakes and are thought to have contributed significantly to the evolution of Earth’s atmosphere (21). The biogeochemical cycling of carbon, oxygen and hydrogen have been studied extensively in the hypersaline mats of Guerrero Negro, Baja
California Sur, Mexico (5, 12, 13, 61), yet the identities of the organisms that drive these cycles are known primarily from culture and microscopy studies (9) and only limited molecular analyses have been performed (39-41, 51). This study is the most extensive rRNA-based survey of a microbial mat so far conducted. Furthermore, we integrate a variety of molecular and chemical analytical approaches to characterize the depth profile of the mat. In contrast to the long-held view that these mats are dominated by Cyanobacteria and are biologically simple, our results indicate an overall unexpected dominance by Chloroflexi, and a remarkably high level of diversity. We found that Cyanobacteria dominate the biomass in the upper 2 millimeters, however, the dominance of Chloroflexi below 2 mm, where the majority of the biomass resides makes them the most abundant type of bacterium overall. In the upper two millimeters, the Chloroflexi can be seen intertwined within the exopolysaccharide sheaths of the dominant cyanobacterium. Together, our results indicate that the photosynthetic activity of the Cyanobacteria sustains a highly diverse and structured community.

Chemically-defined niches and biomass distribution. Our microelectrode measurements of H$_2$, O$_2$ and pH provided a detailed view of the chemical environment in the upper 6 millimeters of the mat. These chemical profiles defined three distinct chemical niche spaces onto which to map the biological diversity. These three zones, the oxic zone, low-H$_2$S zone, and the H$_2$S-rich zone, have been described previously with coarser spatial resolution (5).

In previous studies, the density of the cells was observed qualitatively to be highest in the upper few mm and to taper off with depth (9). Bulk RNA concentrations have also been shown to be higher by an order of magnitude in the upper 1 mm than in
the underlying layers (51). We measured ATP concentrations in the mat as a proxy for biomass. ATP concentrations approached those of pure cell paste in the uppermost millimeters of the mat. The upper few millimeters are considered to be the highly active zone of the mat, where carbon flux driven by photosynthesis is highest. However, even though ATP concentrations were lowest below the oxic zone, the total amount of ATP present in the lower portion of the mat is highest per unit area below the oxic zone. Therefore, more overall biomass resides in the dark sulfidic part of the mat than in the cell-dense upper few millimeters where most photosynthesis occurs.

Pigment depth profiles showed that the cyanobacterial pigment Chl \( a \) was by far the most abundant photosynthetic pigment. It is therefore likely that Cyanobacteria fix more inorganic C than the non-oxygenic phototrophic bacteria (e.g., Chloroflexi and Proteobacteria) and thus the Cyanobacteria presumably provide the main sustenance of the mat, as previously thought. We measured peak concentrations of the bacteriochlorophylls directly under the peak concentration of Chl \( a \). This segregation with depth is consistent with the specialization of bacteriochlorophylls for longer, more deeply penetrating wavelengths, and the previously reported vertical stratification of different phototrophic organisms (9, 56). The photosynthetic Chloroflexi, (e.g., Chlorothrix halophila and relatives) which are similar in properties to Green Sulfur bacteria by having chlorosomes, may also contribute in a substantial way to the primary productivity in the mat. Taken together, the biomass and pigment depth distributions indicate that photosynthetic activity within the oxic zone fuels the underlying, larger biomass. Such an inverted pyramid model of trophic levels, where consumer biomass apparently outweighs producer biomass, has recently been described for the stromatolites.
(calcified mats) of Sharks’s Bay, Australia (44), and may be common in microbial ecosystems where primary producers exude polysaccharides.

**Diversity mapped onto chemical gradients.** We generated 1586 bacterial 16S rRNA gene sequences from the entire depth profile of the mat although this number appears to under-sample woefully the diversity present in the mat. Together, the diversity estimates based on our sequence coverage and the collectors’ curves all indicate a highly diverse community, most of which still awaits description. Despite the low sampling coverage, the high proportion of known and new bacterial phyla in this hypersaline mat makes it the most biologically diverse environment yet characterized. The study has encountered the highest number of confirmed and potential candidate phyla so far observed in a single environment (25). This is remarkable, particularly at a time when the discovery rate of new phyla was thought to be tapering off (46), and suggests that diversity and complexity are a feature of the microbial mat ecosystem.

Previous microscopy-based studies have indicated a vertical stratification of the microbial community with depth, down to about 6 mm, the maximum depth studied (9). Cyanobacteria, particularly the filamentous, exopolysaccharide sheath-forming *Microcoleus chthonoplastes*, were observed by microscopy in the upper 2 millimeters. Other, anoxygenic photosynthetic bacteria such as * Chloroflexus*-like Chloroflexi were observed below the cyanobacteria along with other undefined microbiota with various morphologies (9). We found cyanobacterial rRNA genes in the uppermost layers only, and the majority of the RNA in the upper few millimeters was cyanobacterial. Furthermore, cyanobacteria Chl *a* was far more abundant in the upper few millimeters than BChl *c* and *d*, ascribed to the Chloroflexi. The discrepancy between Chloroflexi and
cyanobacterial representation in 16S rRNA gene libraries vs. bulk rRNA and pigment profiles in the upper millimeters could be due to the comparatively large size of cyanobacteria. For instance, *Microcoleus chthonoplastes* cells are 2-3 times wider than the Chloroflexi cells, which may result in a higher ratio of rRNA expressed to rRNA gene copy number. Nonetheless, overall, the Chloroflexi dominated the mat biomass, with the exception the top 2 mm where Cyanobacteria were a larger proportion of the biomass. Popular reference to the mats as “cyanobacterial” reflects the focus of most previous work on the top few mm.

The majority of the novel candidate phyla that we describe from the mat were detected at a variety of depths. Candidate phyla are known from their constituent 16S rRNA gene sequences alone, and therefore the physiological attributes of the bacteria within them can be gleaned only from the context in which they were discovered. The depth distributions of the novel GN candidate phyla provide testable hypotheses about the physiologies of the organisms. More importantly, the spatial distribution of the candidate phyla can help direct efforts to bring representatives into culture for physiologic studies. The middle, low-H$_2$S zone harbored 9 of the 15 GN candidate phyla, indicating that such organisms are anaerobic and H$_2$S-tolerant, yet may rely on the diurnal variation in H$_2$S levels and the H$_2$S-O$_2$ interface. In contrast, candidate phyla GN4 and GN12 were detected only well below the oxic layer, and are likely to be composed of strict anaerobes. GN14 was observed only in the top layer (day and night libraries), which suggests that it could be a novel photosynthetic group.

The three distinct zones delineated by concentration gradients of O$_2$ and H$_2$S harbored distinct bacterial communities. The UniFrac analysis of the phylogenetic tree
representing the entire dataset, resulted in clusters of the 10 separate samples according to the depth distributions of the three chemical habitats delineated by our in-situ microelectrode measurements. This observation implies that related bacteria occupy similar chemical niches. Therefore, despite the potential for horizontal gene transfer to confer many physiological traits to distantly related bacteria, phylogenetic groups appear to share physiological properties that are manifested as chemical niche preferences at particular depths in the mat. For many of the novel phylotypes observed, the depth at which they were found is consistent with what is known about the physiologies of their close relatives. For instance, close relatives of known photosynthetic organisms, such as Chloroflexi relatives of the photosynthetic Chloroflexus and Chlorothrix spp. previously described for these mats (30, 41), were abundant in the upper zone where light penetrates. Similarly, close relatives of known sulfate-reducing members of the delta-Proteobacteria were most abundant in the oxic zone, which also is where rates of sulfate reduction are highest (39). Members of the Bacteroidetes phylum, known to degrade polysaccharide anaerobically (63), were abundant throughout the dark anoxic zone.

**Chloroflexi and Cyanobacteria: Symbiosis or Antagonism?** A previously described feature of hypersaline microbial mats is the close physical interaction between the cyanobacterium Microcoleus chthonoplastes and a thinner filamentous bacterium also inside the polysaccharide sheath. The thinner partner was thought to be a member of either the Proteobacteria or the Chloroflexi based on physiological properties and TEM microscopy (10). We used fluorescent in situ hybridization with Chloroflexi-specific probes and laser confocal microscopy to confirm that the filamentous bacteria inside the M. chthonoplastes sheaths were Chloroflexi (Fig. 6 A-C). The association is most often
observed at 0.3-1.2 mm (5, 10), corresponding to the zone below maximal oxygenic photosynthesis. Indeed, lower in the mat, Chloroflexi are free-living (Fig 6D). The depth location of the association together with physiological experiments (10) have suggested a co-metabolism of sulfur: the Chloroflexi may draw down levels of H$_2$S stressful for the cyanobacterium (38), which excretes organic carbon used by the Chloroflexi. However, we observed that *M. chthonoplastes* filaments were often disrupted when Chloroflexi were present (Fig 6B), suggesting an alternative view: the Chloroflexi may parasitize the cyanobacterium under H$_2$S stress. The tight physical association of these bacteria from deeply divergent lineages is an example of the physical and chemical microbial interactions that build complexity in the mat.

**Comparisons with Other Systems: Chemical Diversity drives Microbial Diversity.** Comparisons of diversity levels between environments with contrasting characteristics can yield valuable insight and testable hypotheses regarding the rules that govern diversity distributions. For instance, as shown in Figure 7, the very high level of phylum diversity that we found in this study is a sharp contrast to the relatively low level encountered in the human and mouse distal gut. Furthermore, whereas the diversity in the distal gut is characterized by a few deep lineages that diversified at phylogenetically “shallow” levels (more closely related, e.g., at the species/strain level) (2), the microbiota of the microbial mat is far more diverse at all phylogenetic levels (Fig 7). The difference in phylogenetic structure between these two microbial systems is most likely due to the far greater diversity of chemical niches in the mat, which allow more opportunities for specialization within the microbiota. The mammalian distal gut, like microbial mats, is an energy-rich microbial ecosystem where polysaccharides form scaffolds that provide
attachment sites and nutrients for bacteria (53). Fermentation dominates organic matter
diogenesis in both systems, although sulfate reduction and methanogenesis also occur in
both systems (12, 43). In microbial mats where sulfide production levels are high,
however, a larger proportion of electrons produced by diogenesis is likely shunted to
sulfate. The prominence of the sulfur cycle in mats allows the exploitation by the
microbiota of many intermediate chemical moieties (61). Another niche available in the
mats but not in the distal gut is phototrophy: the differential depth penetration of light
wavelengths creates a stratification of phototrophs adapted to using particular
wavelengths of light (6, 7, 27, 50, 56). Additionally, the frequent washout inherent to the
gut undoubtedly is a powerful selection force against slow-growing microbes and the
accretion of community structures important in mats. Thus, the relative complexity of
the microbiota in microbial mats probably correlates with broad niche space.
Furthermore, biological diversity itself can drive diversity through niche creation (16),
therefore chemical and biological diversity are expected to form positive feedback loops.

Microbial mats have been called simple systems based on microscopy and culture
studies. However, our molecular analysis revealed that the hypersaline microbial mats of
Guerrero Negro harbor the most complex bacterial assemblage documented to date in
any environment, with 42 phyla including 15 novel candidate phyla. Microbial mats are
hot-spots of bacterial diversity, and constitute a rich reservoir of gene diversity for future
studies of bacterial evolution and genomic diversity.

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56. **Stolz, J. F.** 1990. Distribution of phototrophic microbes in the flat laminated microbial mat at Laguna Figueroa, Baja California, Mexico. Biosystems **23**:345-57.


Table 1. Mat layers from which sequences forming candidate phyla GN01-15 were obtained. Numbers indicate the depth in the mat from which candidate phylum clones were obtained, the number of sequences from each depth is in parentheses. The GenBank accession numbers are those of clones from other studies without prior phylum affiliations that are included in GN candidate phyla. See Supplementary Online Material for the Arb dendogram showing these phyla in the context of previously described phyla.

<table>
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<th>Phylum</th>
<th>Depth in mm (# sequences)</th>
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FIG. 1. Chemical and biochemical characteristics of the mat as a function of depth. (A) Microelectrode measurements of O$_2$ and H$_2$S concentrations and pH. (B) ATP concentrations. Means of 3 independent measurements are plotted; the bars show standard errors. (C) Pigment concentrations. Abbreviations: Chl $a$, chlorophyll $a$; BChl, bacteriochlorophylls. For all measurements, October values are plotted; the June values were equivalent (data not shown).
FIG. 2: Bacterial diversity within the hypersaline microbial mat. (A) Observed and predicted (Chao1, Ace1) number of taxa with minimum thresholds ranging from 90 to 100% %ID for masked sequences. (B) Collector’s curves for taxa (OTU's) with minimum thresholds of 90, 95, 97, 98, 99 and 100% %ID.
Fig. 3. Bacterial diversity in the mat. Proportion of bacterial phyla: (A) in the total dataset, (B) in the oxic zone (0-2 mm), (C) in the low- H₂S zone (2-6 mm), (D) in the H₂S-rich zone (6-60 mm). Others: Cyanobacteria, KSB1, OP10, GN03, OP5, GN1, Firmicutes, OP11, GN04, GN05, GN09, GN10, WS1, WS2, GN2, Deinococcus-Thermus, GN07, Haloanaerobiales, GN06, GN11, BRC1, OP8, OS-K, GN12, GN13, GN14, Actinobacteria, GN15, WS3, GN8, OP9, TM6, VadinBE97. Abbreviations: Chloro: Chloroflexi, Cyano: Cyanobacteria, Verr: Verrucimicrobia, Planct: Planctomycetales, Spiro: Spirochaetes, Firm: Firmicutes, Bact: Bacteroidetes, Proteo: Proteobacteria. (E) Bacterial community clustering by layer studied (UPGMA tree of UniFrac metric based on 1585 16S rRNA gene sequences). Shaded areas refer to the different chemical milieu’s identified by the microelectrode measurements in Fig 1A.
sequences of cultured representatives are shown in italics. Wedges represent groups of microbial mat sequences, and single sequences are indicated by their clone names. The length of the top and bottom edges represents the range of sequence divergence. The average depth from which sequences were obtained is indicated next to the wedge, with the total depth range in parentheses. (A) Chloroflexi sequences. Percentages indicate the fraction of Chloroflexi sequences within a given sequence cluster. “Oxic zone” indicates clusters of sequences obtained from surface layers exclusively. (B) Cyanobacteria. Percentages indicate the fraction of Cyanobacteria sequences within a given sequence cluster.
FIG 5. Depth distributions of the ten most abundant phyla in the mat. Points indicate the percentage of sequences within each phylum (not the percentage of total sequences) obtained at each depth. The bar indicates 20% of sequences within each group. Shaded areas: see legend in Fig 2E.
FIG 6. Chloroflexi and the cyanobacterium *Microcoleus chthonoplastes* in the mat visualized by laser confocal microscopy. A. Chloroflexi (red, fluorescent *in situ* hybridization (FISH) Chloroflexi-probe) entwined with *M. chthonoplastes* (green, DAPI)) at 1 mm depth. The arrow indicates the edge of the polysaccharide sheath. B. Chloroflexi (green, Chloroflexi-probe) and *M. chthonoplastes* (green, autofluorescence of Chl a). C. Chloroflexi (thin filaments) and *M. chthonoplastes* (thick filaments), DAPI-stained. D. Chloroflexi filaments (red, Chloroflexi -probe) polysaccharide material (dull green) at 50 mm depth. Non- Chloroflexi are seen as bright green spots (Arrow 1). Arrow 2 indicates a buried *M. chthonoplastes* filament. Scale = 10 µm.
FIG. 7. Phylogenetic structure of the microbial mat (n=1585, this study), human colonic (n=11,831, (15)) and mouse cecal (n=5088, (32)) 16S rRNA gene sequence datasets. Sequences were clustered into phylotypes based on % sequence ID (OTU’s with similarity thresholds ranging from 65% Id to 100%ID). The ratio of phylotypes at each threshold to total sequence in each dataset is plotted.