Cyanobacteria and chloroflexi-dominated hypolithic colonization of quartz at the hyper-arid core of the Atacama Desert, Chile

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Abstract Quartz stones are ubiquitous in deserts and are a substrate for hypoliths, microbial colonists of the underside of such stones. These hypoliths thrive where extreme temperature and moisture stress limit the occurrence of higher plant and animal life. Several studies have reported the occurrence of green hypolithic colonization dominated by cyanobacteria. Here, we describe a novel red hypolithic colonization from Yungay, at the hyper-arid core of the Atacama Desert in Chile. Comparative analysis of green and red hypoliths from this site revealed markedly different microbial community structure as revealed by 16S rRNA gene clone libraries. Green hypoliths were dominated by cyanobacteria (*Chroococcidiopsis* and Nostocales phylotypes), whilst the red hypolith was dominated by a taxonomically diverse group of chloroflexi. Heterotrophic phylotypes common to all hypoliths were affiliated largely to desiccation-tolerant taxa within the Actinobacteria and Deinococci. Alphaproteobacterial phylotypes that affiliated with nitrogen-fixing taxa were unique to green hypoliths, whilst Gemmatimonadetes phylotypes occurred only on red hypolithon. Other heterotrophic phyla recovered with very low frequency were assumed to represent functionally relatively unimportant taxa.

Keywords Atacama · Chloroflexi · *Chroococcidiopsis* · Desert · Hyper-arid · Hypolith

Introduction

The Atacama Desert is an ancient coastal arid zone that extends across a large area of coastal Chile in South America (Thomas 1997). The desert is, like most arid regions on Earth, typified by a stony desert pavement terrain. This extends approximately 1,000 km along the Pacific coast of South America. At the hyper-arid core of the Atacama at Yungay (24°06′ S 70°01′ W with 1,051-m elevation), the mean annual rainfall (MAR) is around 2.4 mm making it the driest zone in the desert. Long-term climate data indicate that the hyperarid core of the Atacama Desert at Yungay is the driest non-polar desert location on Earth (Warren-Rhodes et al. 2006; Kuhlman et al. 2008).

Hyper-arid desert pavements are characterized by a lack of higher plants or animals (Thomas 1997), and microorganisms are often the only discernible life. This lends significance to microbial colonization as the predominant form of primary productivity and nitrogen input in such extreme desert regions. A ubiquitous feature of desert pavement microbial ecology has been the occurrence of hypolithic cyanobacterial colonization on quartz rocks both in hot (Schlesinger et al. 2003; Pointing et al. 2007; Warren-Rhodes et al. 2006; Warren-Rhodes et al. 2007) and in cold deserts (Broady 2005; Wood et al. 2008; Pointing et al. 2009; Cowan et al. 2010; Wong et al. 2010a, b). Whilst early studies identified cyanobacteria as the predominant colonist (Cameron and Black 1996; Friedmann and Galun 1974; Berner and Evenari 1978), several recent studies have used DNA fingerprinting...
techniques to inform community structure at a molecular level (Warren-Rhodes et al. 2006; Wood et al. 2008; Pointing et al. 2007). These have generally lacked a quantitative approach and so beta diversity remains uncharacterized, although such molecular studies indicated that hypolithic communities comprise significantly more complex diversity than previously appreciated. More recently, hypolith community structure has been elucidated using sequence data derived from clone libraries and this has yielded valuable data on beta diversity for cold desert hypoliths (Pointing et al. 2009; Wong et al. 2010a). Whilst clone libraries have well-documented limitations, they remain a widespread tool for inferring community structure but have, as yet, not been applied to Atacama hypoliths. Besides the value in more fully resolving microbial community assembly of ‘typical’ green hypoliths at the arid limit for life, the observation that red pigmentation of a quartz rock supported filamentous microbial colonization created a comparative motive for the current study between green and red hypoliths.

Materials and methods

Micro-environmental data

Environmental data that included long-term mean annual temperature and precipitation were collected. Historical rainfall data were sourced from Dirección General de Aguas (http://www.dga.cl), Dirección Meteorológica de Chile (http://www.meteochile.com or http://docs.lib.noaa.gov/rescue/data_rescue_chile.html) and field measurements (McKay et al. 2003). Nanoclimate data for the hypolithic niche were collected in situ from October 2001–2005, including: (1) air and soil temperature and relative humidity at 1 m above the ground surface, at the ground surface and at ~2–5 mm, 2–5 cm, 10 cm, and 20 cm below the soil surface, using Onset Computer HOBO Pro® dataloggers; (2) presence of liquid water condensed on rock and soil surfaces from dew or fog, using a Campbell 237® leaf wetness moisture sensing grid and/or a Spectrum® moisture sensing grid; and (3) rainfall using an Onset Computer tipping bucket rain gauge. An Onset weather station was also installed, with additional sensors to measure solar flux, photosynthetically active radiation (PAR), barometric pressure, and wind speed and direction. The occurrence of liquid water in the soil sufficient to support photosynthesis was defined as a soil relative humidity of ≥95% (Schlesinger et al. 2003) and on soil or stone surfaces by the Campbell moisture sensing grid as a rise in voltage above the 0.005 V ‘dry’ baseline (or a drop in voltage from the 2.095 V ‘dry’ baseline for the Spectrum sensor). Measurements of photosynthetically active radiation (PAR), UV-A, and UV-B radiation were carried out at solar noon using a Li-Cor LI-1400 datalogger (Li-Cor Inc, Nebraska, USA) for PAR, and a UV UVX radiometer (UVP Inc, California, USA) for UV irradiance. For transmittance studies through rock substrates, the sensor was placed in situ under quartz rocks and edges sealed using plasticine.

Molecular community characterization

Colonized rocks were collected from the hyperarid zone, Yungay area (24°06′ S 70°01′ W with 1,051-m elevation), of the Atacama Desert. The three rocks within the sampling zone at Yungay supporting hypolithic biomass (from the same sampling regime conducted by Warren-Rhodes et al. 2006) were used in the present study. Community DNA recovery was achieved following the protocol for environmental DNA recovery from lithic biofilms described in Warren-Rhodes et al. 2006. The entire colonized surface area of each rock was sampled. 16S rRNA genes were PCR amplified using bacteria-specific primers 8F (AGAGTT TGATCCTGTCAG) and 1391R (TGYACWCCGCG CCGTTC) (Lane 1991). Archaeal-specific primers 8Fa (TCYSGTGTACCTCAGCS) and 1492R (GTTACCC GTTACGACTT) (Costello and Schmidt 2006) and eukaryal-specific primers NS1 (GTAGTCATATGCCTGTTC) and NS4 (CTTCCGCTAATCTCCTTAAAG) (White et al. 1990) were also used, but did not give positive amplification. Purified PCR product was used to construct clone libraries for each rock sample (CloningPlus, Qiagen, Valencia, CA, USA). A minimum of 100 clones per library were screened by RFLP (MSP1 and CFO1, Amersham, Bucks, UK) and a total of 104 unique RFLP-defined phylotypes were sequenced (ABI 3730 Genetic Analyzer, Applied Biosystems, Foster City, CA, USA) to generate 70 unique 16S rRNA gene sequences (Operational Taxonomic Unit, OTU) of approximately 1,446 bp each. Phylogenetic OTU’s were determined using the computer program DOTUR described in Schloss and Handelsman (2005), with a 97% sequence similarity cutoff. The sufficiency of sampling from each library was estimated using non-parametric rarefaction and OTU Richness determined by Chao1 using Estimate S software (http://viceroy.eeb.uconn.edu/estimateS). Coverage estimates, Shannon’s Diversity Index (Llyod et al. 1968), and Pielou’s Evenness Index (Pielou 1993) were calculated using Primer v6 (Clarke 1993).

Chimera_Check software (Ribosome Database Project, http://rdp.cme.msu.edu.html) was used to check possible chimeric structures among the sequence data. Approximate phylogenetic affiliations were then determined using BLAST searches of NCBI GenBank database. Multiple alignments were created using ClustalX v.1.81 (Thompson...
et al. 1997) with reference to sequences selected from the GenBank database. All phylogenetic analysis was performed using both GARLI (Genetic Algorithm for Rapid Likelihood Inference) 0.96 beta (Zwickl 2006) and PAUP* 4.0b10 (Swofford 2001). Maximum likelihood analysis was used to illustrate relationships of the 16S rRNA gene sequences to representative taxa. Bayesian posterior probabilities and bootstrap values of 1,000 replications were calculated and are shown for branch nodes supported by more than 50% of the trees. The phylogenetic differences among communities were quantified using the $F_{st}$ statistic calculated using Arlequin v3.0 (Excoffier and Scheider 2005). Sequence data have been submitted to NCBI GenBank database with accession numbers FJ890991 to FJ891059.

**Results and discussion**

Long-term micro-environmental monitoring at the study site carried out by our group indicated that the hypolithic niche beneath quartz rocks received an average of 395±18 h/year liquid water (Table 1) although of this only 75 h were under conditions that were estimated to support photosynthetic activity (sufficient PAR and temperature) (Warren-Rhodes et al. 2006). This moisture input was derived from fog/dew (68%) and rainfall (32%). We have also conducted radiocarbon dating studies which indicated that Atacama hypoliths may be very long lived and possibly persisted for thousands of years (Warren-Rhodes et al. 2006). Hypolithic colonization can thus be envisaged as a slow, but persistent colonization, and although they clearly tolerate significant environmental stochasticity in terms of water availability and temperature, we have observed that disturbed hypoliths seldom survive, thus suggesting that substrate stability is important.

The extremely low colonization frequency of quartz rocks at the hyper-arid core of the Atacama Desert mean that locating hypolithic colonized rocks that can be analyzed is highly challenging (i.e. only 3 out of 3,723 stones were colonized at our site, across an area >2,000 m$^2$) (Warren-Rhodes et al. 2006). The study reported here includes all three colonized rocks located within the survey area and therefore represents the total recoverable hypolithic diversity within this area. Despite equal effort in DNA recovery and cloning, the three rocks yielded significantly different rRNA gene-defined communities, and this heterogeneity was supported by microscopy that indicated major phylotypes recovered in each library were also the most commonly encountered morphotypes. The data revealed two major findings; that the two ‘green’ *Chroococcidiopsis*-dominated hypoliths supported a highly similar diversity although abundance levels varied; and that the ‘red’ hypolith supported chloroflexi rather than cyanobacteria as the dominant taxon plus a markedly different and more speciose heterotrophic component.

The two green hypoliths were the same samples that have been previously analyzed using a DGGE DNA-fingerprinting approach (Warren-Rhodes et al. 2006). Unsurprisingly, our clone libraries revealed more than twice the number of phylotypes than the fingerprinting approach, and also allowed a phylogenetic assignment to near-full length rRNA gene phylotypes. The limitations of the DGGE technique are now well-documented (Jackson and Churchill 1999; Sekiguchi et al. 2001; Kirk et al. 2004; Nocker et al. 2007), and the rationale for the present study was to more comprehensively understand diversity of hypoliths at the driest non-polar desert location on Earth. Our libraries allowed us to more fully define the community composition and generate relatively quantitative beta diversity data (Table 2). For both green hypoliths, the community was dominated by diverse phylotypes within the cyanobacterial genus *Chroococcidiopsis* and the *Nostocales* (Fig. 1). The former is a near-ubiquitous colonist of arid deserts worldwide (Bahl et al. 2010), also known to fix nitrogen under certain conditions (Boison et al. 2004). The latter a nitrogen-fixing family that contains several UV-screening pigment-secreting taxa recovered from temperate (Warren-Rhodes et al. 2006) and cold deserts (Pointing et al. 2009; Wong et al. 2010a, b) regions. Production of UV-protective pigments by cyanobacteria, such

**Table 1 Summary of beta diversity for green and red hypoliths**

| Phylum                        | Green 1 (AY1) | Green 2 (AY5) | Red 1 (AY6) |
|------------------------------|--------------|--------------|-------------|
| Cyanobacteria                | 77           | 85           | 39          |
| *Acaryochloris*              | 1            |              |             |
| *Chroococcidiopsis*          | 27           | 47           | 29          |
| *Nostocales*                 | 48           | 37           | 10          |
| Oscillatoriales              | 2            |              |             |
| Actinobacteria               | 5            | 4            | 11          |
| Alphaproteobacteria          | 13           | 5            |             |
| Bacteroidetes                | 2            | 1            |             |
| Betaproteobacteria           | 2            |              |             |
| Chloroflexi                  | 32           |              |             |
| Deinococci                   | 3            |              |             |
| Gammaproteobacteria          | 4            | 4            |             |
| Gemmatimonadetes             | 1            |              |             |
| Planctomycetes               | 1            |              |             |
| Unknown bacteria             | 11           |              |             |
| *Thermobaculum*              | 6            |              |             |

Phylotypes were assigned to a given taxonomic group based upon phylogenetic analysis of 16S rRNA gene sequences.
as scytonemin (Proteau et al. 1993) can be envisaged as an
adaptive advantage in high UV desert environments. Both
of these cyanobacterial groups are noted for their radiation
tolerance (Billi et al. 2000), a proxy for desiccation
tolerance.

The second most abundant phylum was the Alphaproteobacteria (Fig. 2). These phylotypes were specific to
green hypoliths and whilst phylogenies for this phylum are
skewed toward human associates as a result of availability
of identified environmental taxa among published sequen-
ces, they did also affiliate with the nitrogen fixing genus
Agrobacterium. Other studies have revealed that N₂-fixing
organisms are often the abundant and early colonizers of
arid soil environments that give way to other species over
time (Yeager et al. 2004; Garcia-Pichel et al. 2001; Walker
1993), presumably as they create nitrogen sufficiency
within these otherwise nitrogen-depleted desert soils. Other
phylotypes were recovered with very low abundance
(Table 1; Fig. 2) and so can be postulated to represent
functionally relatively unimportant taxa, although their
latency might indicate the capacity of the community to
respond to environmental change.

The red hypolith represented an opportunity to describe
a novel hypolithic colonization, often previously assumed
to represent an abiotic red/brown patina on rocks. Our
study revealed surprisingly different community structure
between red and green hypoliths (Table 1). The most
striking feature was the dominance by a collection of
phylogenetically relatively diverse chloroflexi (Fig. 2). The
chloroflexi have traditionally been associated with hot
spring environments (Jing et al. 2005; Lau et al. 2006,
2008, 2009; Lacap et al. 2007), but recently have emerged
colonists of hot (Pointing et al. 2007) and cold (Wong
et al. 2010a) desert hypoliths, in addition to cold endoliths
(Wong et al. 2010b). As a phylum they may therefore display a
general adaptation to arid environments. The phylotypes
affiliated with the genera Roseiflexus, Anaerolinea and also
formed a deep-branching cluster of uncharacterized Chlo-
roflexi (Fig. 2), possibly indicating a novel group of green
non-sulfur bacteria. The cyanobacterial genus Chroococ-
cidiopsis was the second most abundant group, whilst the
Nostocales comprised only 10% of the community.

Interestingly, no other putative nitrogen fixing taxa were
indicated from the phylogenies, and this may reflect a
higher degree of nitrogen sufficiency within this micro-
niche, or generally more abundant nutrients since the
chloroflexi are heterotrophically capable and may have out-
competed the cyanobacteria under such conditions. The
difference is unlikely a function of moisture availability
since the sampled area experiences very similar water input
and residence times (Warren-Rhodes et al. 2006).

We measured light transmittance (PAR, UVA, and
UVB) through all rocks from the point of soil surface to the
depth of colonization, with no marked differences between
green and red colonized rocks. We therefore discount a
selective effect from light transmittance. Whilst low
abundances of commonly encountered arid soil heterotro-
phic bacteria (Yeager et al. 2004; Garcia-Pichel et al. 2001;
Walker 1993) from the phyla Actinobacteria and Bacte-
riodetes were shared between green and red hypoliths, the
red hypolith also supported additional unique heterotrophic
phylotypes (Table 2; Fig. 2). The actinobacterial phylo-
types affiliated with the genus Rubrobacter, a noted radia-
tion tolerant desert bacterium (Ferreira et al. 1999). This
included the Deinococci, which affiliated with those
recovered from arid soils in other hot deserts (Rainey et al.
2005; Chanal et al. 2006; Pointing et al. 2007). The Dei-
nococci are noted for their desiccation and radiation toler-
ance (De Groot et al. 2005) although they appear to be a
feature of hot rather than cold desert hypoliths. The
remaining red hypolith phylotypes affiliated with the
Gemmatimonadetes, which are emerging as a common
taxon in hypoliths from hot (Pointing et al. 2007) and cold
(Pointing et al. 2009) deserts, plus unidentified beta pro-
teobacteria and other unidentified phyla. The greater
diversity and abundance of heterotrophic phylotypes may

### Table 2 Summary of diversity data for clone libraries generated from green and red hypoliths

| Sample     | No. of positive RFLP-defined phylotypes | No. of OTU (<97% sequence similarity) | Coverage (C) (%) | Chao1 richness ± SD | Shannon’s diversity index (H) | Pielou’s evenness (J) | Fstab statistic |
|------------|----------------------------------------|---------------------------------------|------------------|---------------------|-------------------------------|----------------------|-----------------|
| Green 1 (AY1) | 108                                    | 32                                    | 22               | 90                  | 20.8 ± 5.2                    | 3.15                 | 0.71            | 0.1401         |
| Green 2 (AY5) | 101                                    | 29                                    | 19               | 88                  | 17.3 ± 5.2                    | 2.73                 | 0.64            | 0.14164        |
| Red 1 (AY6)  | 102                                    | 43                                    | 29               | 83                  | 32.4 ± 9.7                    | 4.04                 | 0.83            | 0.13528        |

**Fig. 1** Phylogenetic relationships among cyanobacterial 16S rRNA phylotypes recovered from Atacama hypoliths. Phylotypes recovered during this study are shown in **bold type**. Phylotypes indicated by an *asterisk* share >97% sequence similarity with their most closely affiliated phylotype indicated by a NCBI GenBank accession number (in brackets) on the tree. NCBI GenBank accession tree topologies are supported by Bayesian posterior probabilities (first number) and bootstrap values for 1,000 replications (second number). Scale bar 0.1 nucleotide changes per position.
indicate that the chloroflexi-dominated hypolith represents a community structured under more nutrient sufficient conditions.

Statistical analyses of the phylogenetic data (Table 2) indicated that whilst the red hypolith community was most biodiverse, all three hypoliths displayed significantly different diversity and evenness indices (Table 2). The phylogenetically informed community structure was demonstrated as significantly different between all three communities (Table 2). This is interesting, since given the high coverage of our libraries (due to the relatively low diversity of hypoliths); this indicates that hypolith communities can be highly heterogenous within a given climatic biome. This is likely due to unmeasured differences in the microclimate beneath each quartz rock.

The Atacama Desert supports several distinct microbial oases, including quartz hypoliths, halite endoliths, and microbial mats in hypersaline waters (Weirzchos et al. 2009). It is noteworthy that similar cyanobacterial taxa occur in all of these niches and so they may act as local reservoirs for ‘extreme’ colonists in a variety of challenging niches. It will be interesting to establish in future work the extent to which common cyanobacterial taxa are involved in community assembly within each niche, and whether non-cyanobacterial taxa, such as the chloroflexi are also widespread in these niches.

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