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Amplicon Sequencing of Rock-Inhabiting Microbial Communities from Joshua Tree National Park, USA

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ABSTRACT

Endolithic microorganisms have been reported to date in hot and cold drylands worldwide, where they represent the prevailing life forms ensuring ecosystem functionality, playing a paramount role in global biogeochemical processes. We report here an amplicon sequencing characterization of rocks collected from Joshua Tree National Park (JTNP), USA.

Understanding the microbial community structure and diversity in dryland regions is of paramount importance in both microbial ecology and evolution. Under these conditions, where most life-forms cannot survive, rocks represent the main refuge for life. Indeed, the endolithic habitat provides thermal buffering, physical stability, and protection against UV and solar radiation, ensuring the ecosystem functionality and creating positive feedback for further colonization (1). Although endoliths are ubiquitous in drylands worldwide, including the hyperarid drylands, from Atacama Desert to the Antarctic Dry Valleys (for examples, see references 2 to 8), a comprehensive assessment of worldwide rock diversity remains insufficiently studied, and comparisons between geographic locations and different climatic conditions are still missing (9).

The present study, to our knowledge, represents one of the few available amplicon sequencing reports within the hot desert of Joshua Tree National Park (JTNP) in California (for an example, see reference 10).

Ten rock samples were collected from three different sites in the JTNP (GPS, 34.10, −115.45; the elevation for all sites is between 1,040 and 1,340 m); the presence of endolithic colonization was assessed by direct observation in situ. Rocks were collected under sterile conditions using a geologic hammer and a chisel, placed in sterile bags, transported in a cooler with ice, and stored at −80°C at the University of California, Riverside (CA), until downstream analysis. DNA extraction was performed using 0.5 g of powdered rock with a PowerSoil DNA extraction kit (MoBio Laboratories, Carlsbad, CA, USA).

The fungal internal transcribed sequence 1 (ITS1) region was amplified using the primers ITS1F and ITS2 (11), while the bacterial 16S rRNA V4 region was amplified using 515F and 806R (12), according to the Earth Microbiome Project protocols (https://earthmicrobiome.org/protocols-and-standards/). Two replicates were amplified for each sample. PCRs were carried out with a total volume of 25 µL, containing 1 µL of each primer, 12.5 µL of Taq DNA polymerase (Thermo Fisher Scientific, Waltham, MA, USA), 9.5 µL of nuclease-free water (Sigma-Aldrich, St. Louis, MO, USA), and 5 ng of DNA template using an automated thermal cycler (Bio-Rad, Hercules, CA, USA). The ITS1 locus was amplified following an initial denaturation at 94°C for 1 min and 35 cycles at 94°C for 30 s, annealing at 52°C for 30 s, and extension at 68°C for 90 s, followed by a final extension at 68°C for 7 min. The PCR for the V4 region followed a protocol of an initial denaturation at 94°C for 3 min, 35 cycles at 94°C for 45 s, annealing at 50°C for 1 min, and extension
at 72°C for 90 s, followed by a final extension at 72°C for 10 min. The amplicons were quantified using the Qubit double-stranded DNA (dsDNA) high-sensitivity (HS) assay kit (Life Technologies, USA), purified using the Qiagen PCR cleanup kit (Macherey-Nagel, Hoerd, France), and then pooled to produce an equimolar mixture. Sequencing was performed at the Institute for Integrative Genome Biology (IIGB; https://iigb.ucr.edu/), University of California, Riverside (USA), with the MiSeq reagent kit v3 on the Illumina MiSeq platform (2 × 300-bp paired-end format).

The ITS1 and 16S rRNA data sets were processed separately using the AMPtk v1.4.3 tool (https://github.com/nextgenusfs/amptk) (13), which removed barcodes and primers, removed chimeras, denoised sequences, and eliminated sequences of <250 bp using default parameters. Amplicon sequence variants (ASVs) were generated using DADA2 v1.16.0 (14); singletons and rare taxa (ASVs represented by ≤5 reads) were discarded as recommended by Lindahl et al. (15). Finally, AMPtk was used for taxonomic identification by a “hybrid” taxonomy assignment, which calculates a consensus last common ancestor (LCA) taxonomy based on the results of a global alignment using the USEARCH v9.2.64 (16), UTAX, and SINTAX databases.

The fungal ITS resulted in 960,527 quality-filtered reads clustered in 869 valid ASVs, while a total of 1,059,591 valid output reads were obtained for 16S rRNA, resulting in 3,221 ASVs. The majority of the identified ITS sequences recovered from all samples belonged to the phylum Ascomycota, followed by Basidiomycota, while Actinobacteria and Proteobacteria predominated in the 16S rRNA data set.

Data availability. The bacterial 16S rRNA and fungal ITS1 data sets generated and analyzed in the current study are available in the NCBI Sequence Read Archive (SRA) under BioProject accession number PRJNA706555 and in the Zenodo repository (https://doi.org/10.5281/zenodo.3960774).

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