UC Riverside UC Riverside Previously Published Works

Title

Sun exposure drives Antarctic cryptoendolithic community structure and composition

Permalink

https://escholarship.org/uc/item/7qm8k72d

Journal

Polar Biology, 43(5)

ISSN 0722-4060

Authors

Coleine, Claudia Stajich, Jason E Zucconi, Laura <u>et al.</u>

Publication Date

2020-05-01

DOI

10.1007/s00300-020-02650-1

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at https://creativecommons.org/licenses/by/4.0/

Peer reviewed

Sun exposure drives Antarctic cryptoendolithic community structure and composition

Claudia Coleine¹, Jason E. Stajich^{2*}, Laura Zucconi¹, Silvano Onofri¹, Laura Selbmann^{1,3}

¹Department of Ecological and Biological Sciences (DEB), University of Tuscia, Viterbo, Italy

²Department of Microbiology and Plant Pathology and Institute of Integrative Genome Biology, University of California, Riverside, CA, USA

³Italian National Antarctic Museum (MNA), Mycological Section, Genoa, Italy

*Corresponding author: Jason Stajich: jason.stajich@ucr.edu; University of California-Riverside, 900 University Ave, Riverside, CA 92521



Cryptoendolithic lichen dominated community colonizing a sandstone rock sample at Linnaeus Terrace (McMurdo Dry Valleys, Southern Victoria Land), Continental Antarctica.

- 1. DGGE profiles indicated a clear grouping of samples according by sun exposure.
- 2. Higher similarities were observed within samples collected in the same site.
- 3. Separation in abundance and community composition in response to sun exposure.
- 4. The F/B dominance showed a significative dominance of fungi.

1	Sun exposure drives Antarctic cryptoendolithic community structure and composition
2	
3	Claudia Coleine ¹ , Jason E. Stajich ^{2*} , Laura Zucconi ¹ , Silvano Onofri ¹ , Laura Selbmann ^{1,3}
4 5 6 7 8	¹ Department of Ecological and Biological Sciences (DEB), University of Tuscia, Viterbo, Italy ² Department of Microbiology and Plant Pathology and Institute of Integrative Genome Biology, University of California, Riverside, CA, USA ³ Italian National Antarctic Museum (MNA), Mycological Section, Genoa, Italy
9	
10	

- 11 *Corresponding author: Jason Stajich: jason.stajich@ucr.edu; University of California-Riverside,
- 12 900 University Ave, Riverside, CA 92521

13 Abstract

14 The harsh environmental conditions of the ice-free regions of Continental Antarctica are considered 15 one of the closest Martian analogues on Earth. There, rocks play a pivotal role as substratum for life 16 and endolithism represents a primary habitat for microorganisms when external environmental conditions become incompatible with active life on rock surfaces. Due to the thermal inertia of 17 18 rock, the internal airspace of lithic substratum is where microbiota find a protected and buffered 19 microenvironment, allowing life to spread throughout these regions with extreme temperatures and 20 low water availability. The high degree of adaptation and specialization of the endolithic 21 communities makes them highly resistant but scarsely resilient to any external perturbation and 22 thus, any shifts in microbial community composition may serve as early-alarm systems of 23 environmental perturbation, including climate change.

24 Previous research concluded that altitude and distance from sea do not play as driving factors in shaping microbial abundance and diversity, while sun exposure was hypothesized as significant 25 26 parameter influencing endolithic settlement and development. This study aims to explore our 27 hypothesis that changes in sun exposure translate to shifts in community composition and 28 abundances of main biological compartments (fungi, algae and bacteria) in the Antarctic 29 cryptoendolithic communities. We performed a preliminary molecular survey, based on DGGE and qPCR tecniques, of 48 rocks with varying sun exposure, collected in Victoria Land along an 30 31 altitudinal transect from 834 to 3100 m a.s.l.

32 Our findings demonstrate that differences in sun radiation between north and south exposure 33 influence temperature of rocks surface, availability of water and metabolic activity and also have 34 significant impact on community composition and microbial abundance.

- 35
- 36
- 37

38 Key words: Antarctica, Cryptoendolithic communities, Sun exposure, DGGE, qPCR, Sampling

39

40 **1. Introduction**

41 The rate of warming due to increased levels of greenhouse gases in the atmosphere is amplified 42 with elevation and at high latitudes due to the polar amplification phenomenon. Polar amplification 43 predicts that as global mean temperature climbs, the greatest warming will occur at the Polar regions (Bekryaev et al., 2010). The impact of climate change is, therefore, particularly intense at 44 45 the Poles and in mountain environments, nowadays known as the Third Pole (Yao et al., 2012; 46 Yang et al., 2014). As consequence of warming, range-restricted species, particularly polar and 47 mountain top species, have already shown severe contractions and have been the first groups in 48 which entire species have gone extinct due to recent climate change (Parmesan, 2006; Descamps et 49 al., 2017; Bhatta et al., 2018).

50 The Arctic regions are melting faster than the Antarctic and, if the heating trend continues, studies 51 forecast an ice-free North Pole in summer by mid-century. Strong evidence of warming in Antarctica, is also documented; researchers from the British Antarctic Survey report a warming 52 53 trend up to 2.5 °C since the 1940s in the Antarctic Peninsula and Maritime Antarctica, the most 54 rapid changes in mean air temperatures on Earth (e.g. Turner et al. 2005, 2007). Previous research reported an apparent contrast between strong warming of the Antarctic Peninsula and slight cooling 55 56 of the Antarctic continental interior; there are now evidences that significant warming extends well 57 beyond the Antarctic Peninsula and covers most of West Antarctica with a warming exceeding 0.1 58 °C per decade over the past 50 years (Steig et al., 2009).

Progressions of this warming trend will influence Antarctica's biodiversity by the introduction of allochthonous, competitive species and the consequent extinction of highly specialized and less competitive autochthonous ones (Farrell et al., 2011; Olech and Chwedorzewska, 2011; Selbmann et al., 2012), which will have impacts on the ecosystem functions of glaciers, freshwater systems and atmosphere. Thus, it is urgent to develop a strong base knowledge for Antarctic terrestrial ecosystems and use this to identify ecosystem changes (NAS, 2011).

65

Endolithism is a specialized colonization by microbes to enable dwelling inside airspaces of rocks. 66 67 This lifestyle represents adaptation at the edge inhabitable conditions. Airspaces within rocks offer 68 to microbiota a protected and buffered microenvironment, allowing life to expand into different 69 extreme conditions, i.e., hot and cold deserts or geothermal environments (Friedmann and Ocampo, 70 1976; Friedmann, 1982; Bell, 1993; Walker et al., 2005). Rocks are the prevailing substratum for 71 life in the ice-free areas of Antarctica, supporting the highest standing biomass in the Antarctic ice-72 free desert and mountain tops emerging from the Polar Plateau (Cowan and Tow, 2004; Cary et al., 73 2010; Cowan et al., 2014; Selbmann et al., 2017). Endolithic microbial life represents the 74 predominant recorded life-form in these areas (Nienow and Friedmann, 1993). The harsh conditions

are considered one of the closest analogues to Mars on Earth (Quintal et al., 2018).

76 Different from soil microbial communities of these areas, endolithic microbes develop as very tiny 77 and stable communities thanks to the stable and concrete nature of rocks. Various typologies have 78 been observed for the microbial composition and the most complex and widespread are the lichendominated communities (Nienow and Friedmann, 1993). These self-supporting microbial 79 80 ecosystems are composed of algae, mainly lichenized fungi, bacteria and cyanobacteria, many of 81 which are endemic species to the regions (Nienow and Friedmann, 1993; Selbmann et al., 2005, 82 2008; Egidi et al., 2014). The high degree of adaptation and specialization in exploiting such 83 ultimate niches makes these communities very susceptible to physical and climatic alteration

84 (Selbmann et al., 2017) and any shift in microbial communities composition may serve as early-85 alarm system of environmental perturbation. Based on a substantial sampling of different typologies 86 (volcanic and sedimentary) of colonised rocks in the Victoria Land, Antarctica, sandstone was 87 determined to be the most suitable substratum for microbial endoliths, allowing them to spread and 88 persist under stronger environmental pressure (Zucconi et al., 2016; Selbmann et al., 2017). To get 89 clues to the future effects of climate change on these unique ecosystems, the response of the 90 communities to increasing environmental pressure, due to altitude (from sea level to 3600 m a.s.l.) 91 and sea distance (up to 100 km) was recently investigated. The results suggested that these two 92 paramenters alone do not play as driving factors in shaping the community diversity and 93 composition, and highlighted the needs to consider additional environmental parameters to elucidate 94 how, in the long run, future environmental changes will impact these unique communities (Coleine 95 et al., 2018a).

96

97 With this in mind, a new sampling campaign (Dec.2015- Jan. 2016) of sandstones was performed in 98 the Victoria Land (Antarctica) in the frame of the Italian National Program for Antarctic Researches 99 (PNRA), along an altitudinal gradient from 834 to 3100 m a.s.l., adding sun-exposure as new 100 parameter to investigate its influence on endolithic settlement and development (Friedmann and Weed, 1987). Rocks with varying sun exposure (north and south faced surfaces) were collected in 101 all the localities surveyed. Based on the prior experiences (Selbmann et al., 2017; Coleine et al., 102 2018a), it is suggested that sampling strategy and environmental parameters considered may have 103 important consequences on a proper and exhaustive biodiversity description; thus, before 104 105 proceeding with metabarcoding and metagenomics experiments to develop a detailed picture of microbial diversity and functionality, we have selected a suite of more rapid and cheaper 106 107 complementary approaches. We have used Denaturing Gradient Gel Electrophoresis (DGGE) and 108 quantitative PCR, to test hypothesis that changes in sun exposure impacts community composition 109 and abundances of primary biological assemblages.

110 **2. Materials and methods**

111 *2.1. Study area*

112 Eight localities were visited in Victoria Land (Continental Antarctica) during the XXXI Italian 113 Antarctic Campaign (Dec. 2015 - Jan. 2016). North and south exposed sandstone samples were collected along a latitudinal transect from 74°10'44.0''S 162°30'53.0''E (Mt. New Zealand, 114 115 Northern Victoria Land) to 77°54'43.6''S 161°34'39.3''E (Knobhead, Southern Victoria Land), ranging from 834 (Battleship Promontory, Southern Victoria Land) to 3100 m a.s.l. (Mt. New 116 Zealand) (Table 1). All rock samples were excised aseptically and collected in triplicate, transported 117 118 at -20 °C at the Tuscia University (Viterbo, Italy) and stored at Mycological Section of the Italian 119 Antarctic National Museum (MNA) until downstream analysis.

119

121

122 2.2. Environmental DNA extraction and Denaturing Gel Gradient Electrophoresis

123 Environmental DNA was extracted from 0.3 g of crushed rocks using NucleoSpin® Plant II Kit 124 (Macherey-Nagel, Gmbh & Co. KG, Duren, Germany) according to the manufacturer's instructions and quantified by Quant-iT dsDNA HS assay kit (Invitrogen molecular probes- Eugene, Oregon, 125 126 USA). Microbial diversity was screened by Denaturing Gel Gradient Electrophoresis (DGGE). A semi-nested PCR was performed using primers with a GC-clamp (Muyzer et al., 1993); fungal and 127 128 algal ITS rRNA was amplified from environmental DNA with primers ITS1F-GC/ITS2 (Gardes and Bruns, 1993; White et al., 1990), following the protocol reported in Selbmann et al. (2017), 129 while 341F-GC/518R primers (Muyzer et al., 1993) were utilized for 16S rRNA amplification 130 (Valášková and Baldrian, 2009) (Table 2). 131

132 Amplicons were purified using NucleoSpin® Gel and PCR Clean-up (Macherey-Nagel, Gmbh &

133 Co. KG, Düren, Germany). One-hundred ng of final DNA concentration were loaded into each well

for DGGE and runs were performed on DGGE Electrophoresis System (C.B.S. Scientific, Del Mar,
 California, USA). Gels at 7.5% polyacrylamide (37.5:1 acrylamide:bisacrylamide) were mixed in a

gradient maker (Hoefer, USA), using two different concentrations: for fungi and algae (0% and 137 70%) and for bacteria (0% and 60%). The electrophoresis run was performed for 5 h for fungi and 138 algae and 3.5 h for bacteria in $1 \times TAE$ Buffer at a constant temperature of 60 °C and 200 V.

algae and 5.5 If for bacteria in 1^{1} TAE Burlet at a constant temperature of 60° C and 200 V.

139 Bands were visualized by staining for 40 min with GelRed solution (Biotiuminc, CA, USA) (1.34 g NaCl, 66.7 pi GelRed and 200 mL dw) and then visualized with an UV transilluminator (Chemidoc, 140 141 Bio-Rad). Scanned gels were analyzed with TotalLab Ouant Software (Cleaver Scientific Ltd: 142 United Kingdom): bands were assigned and matched automatically and then checked manually. Profile similarity was calculated by determining Dice's similarity coefficient, considering 143 presence/absence of bands, for the total number of lane patterns from the DGGE gels. The 144 145 similarity coefficients calculated were then used to generate the dendrograms utilizing the clustering method Unweighted Pair Group Method using Arithmetic mean (UPGMA). 146

147 Triplicate samples were processed and analyzed independently.

148 149

150 2.3. NMDS ordination plots

Multivariate statistical analyses were performed to determine the effects of sun exposure on microbial diversity composition using PAST software (PAleontological STatistics, ver. 2.17) The effect of this abiotic parameter was tested displaying changes in communities' composition with Non-Metric Multidimensional Scaling (NMDS) based both on abundance data (bands intensity), calculating Bray-Curtis distance index and presence–absence data, using Jaccard index (Clarke, 1993). Means of abundance data were square-root transformed and analyses were carried out with 999 permutations as described in Coleine et al. (2018b). NMDS were plotted using the combined occurrence and abundance data of the three replicates from each site. Permutational multivariate analyses of variance (PERMANOVA, p<0.05) based on the Euclidean distance were utilized to establish differences in the two differently sun-exposed rock-inhabiting communities.

161 162

163 2.4. Quantitative PCR

164 Total fungal and bacterial abundances were measured for all samples by quantitative PCR (qPCR) NS91F (5'-GTCCCTGCCCTTTGTACACAC-3') and 165 using ITS51R (5'-ACCTTGTTACGACTTTTACTTCCTC-3') fungi Eub338 (5'-166 for total and ACTCCTACGGGAGGCAGCAG-3') and Eub518 (5'-ATTACCGCGGCTGCTGG-3') for total 167 168 bacteria (Fierer et al., 2005). To determine relative gene-copies abundances, standard curves were 169 generated using a 10-fold serial dilution of a plasmid containing a copy of Cryomyces antarcticus ITS rRNA gene for fungi and Escherichia coli 16S rRNA gene for bacteria. The amplicons were 170 generated in 100 µl reactions, containing 50 µl of 2x PCR BioMix[™] (Bioline, London, UK), 5 171 pmol of both forward and reverse primers, 43 µl of DEPC water and 5 µl template of genomic 172 173 DNA. For fungi, amplification of ITS region was carried out as follows: 95 °C for 3 min and the 35 cycles of 95 °C 40 s, 55 °C 30 s, and final extension at 72 °C 40 s; while bacterial 16S region was 174 175 amplified as follows: 95 °C for 3 min and the 35 cycles of 95 °C 40 s, 53 °C 30 s, and final extension at 72 °C 40 s. PCR products were then purified using the NucleoSpin® PCR Clean-up kit 176 177 (MACHEREY-NAGEL, GmbH & Co. KG), guantified with Qubit dsDNA HS Assay kit and cloned using the pGEM®-T Easy Vector Systems (Promega, Madison, Wisconsin, US). Plasmids 178 were isolated using the NucleoSpin Plasmid kit (Macherey-Nagel, GmbH & Co. KG). 179

180 Five standards were utilized for qPCR in series from 10^7 to 10^3 copies.

181 The 25 μl qPCR reactions contained 12.5 μl iQ[™] SYBR® Green Supermix (Bio-Rad, Hercules,

182 California, US), 1 µl of each forward and reverse primers, 0.3 ng of environmental DNA or standard

and $9.5 \,\mu$ l nuclease-free water. The reactions were carried out on Quantitative real-time BioRad

184 CFX96TM PCR detection system (Bio-Rad, Hercules, California, US). Primers and amplification

185 protocols are mentioned before. Melting curves were generated to confirm that the amplified

products were of the appropriate size. Fungal and bacterial gene copy numbers were generated using a regression equation for each assay relating the cycle threshold (C_t) value to the known

188 number of copies in the standards.

189 Each assay included no-template controls (NTC). All qPCR reactions were run in triplicate.

190 Means and standard deviations were calculated and statistical analysis were performed using one-

191 way analysis of variance (Anova) and pairwise multiple comparison procedures, carried out using

192 the statistical software SigmaStat 2.0 (Jandel, USA). Significant differences were calculated by

193 Tukey test (p<0.05).

194 The fungal-to bacterial ratio was calculated from log-transformed abundance values.

195

196 **3. Results**

197 3.1. DGGE profiles

198 DNA was efficiently extracted from almost all rock samples, with few exceptions (i.e. Linnaeus 199 Terrace south) where DNA extraction was not quantifiable, but PCR-DGGE worked out for all 200 samples.

201 Because results were similar when analysing fungi and algae, only data based on fungi are reported.

202 Profile similarity based only on band presence/absence, was calculated by the Dice coefficient and 203 UPGMA was used to create dendrograms describing pattern similarities (Figs. 1, 2). Overall, the 204 banding patterns of the replicates showed a high degree of similarity (data not shown), which was also supported by the dendrograms, generating DGGE patterns that grouped together as most 205 206 similar to each other. A coherent grouping according to locations and to sun exposure was 207 generated in the clustering based on fungal, and, even more clearly, on bacterial profiles, with few scattered exceptions. In particular, in the clustering generated on fungal profiles, samples were split 208 209 according to the localities for University Valley, Pudding Butte, Battleship Promontory, Mt. New 210 Zealand and Finger Mt. (Fig. 1). The grouping was also evident in the clustering based on DGGE bacterial profiles (Fig. 2); as for fungi, the splitting was obtained in the communities from Pudding 211 212 Butte (south), Battleship Promontory (south) and Finger Mt. (south), but also for Linnaeus Terrace 213 and Knobhead for both sun expositions.

- 214
- 215

216 *3.2. NMDS ordination plots*

To investigate the similarity of the fungal and bacterial communities' composition amongst
 different sun exposures, a Non-metric Multi-Dimensional Scaling (NMDS) analysis was computed.

NMDS ordination plots were generated both with the only presence-absence matrix using the Jaccard index and with the combined frequency of occurrence using the Bray-Curtis index. Because both approaches produced similar results, we showed results based on abundance only.

When stress values are <0.1, the NMDS plot is considered to be an acceptable representation of the original data; in fact, a stress value below 0.1 indicates a reliable ordination of data, without a real probability of misinterpretation (Clarke, 1993). In this analysis, the stress value was 0.09 for fungi and 0.07 for bacteria, fitting with the ideal ordination.

NMDS plots generated from the DGGE profiles of amplified fungal ITS and bacterial 16S rDNA revealed that only small changes occurred among samples collected in the same sun-exposed rock surface (p>0.05) and did not exhibit any changes endolithic communities by sampled localities (data not shown). On the contrary, a major change (1-way NPMANOVA, p<0.05) occurred between north and south sun-exposed communities, showing a strong structuring of fungal and bacterial communities according to the sun exposure (Fig. 3).

232

233 3.3. Abundance of fungal and bacterial communities

Both ITS and 16S rRNA gene copy numbers varied mostly significantly (p <0.05) along the sampled sites, ranging from 6.1 x 10^4 (Knobhead south sun exposure, Dry Valleys, Southern Victoria Land) to 9.8 x 10^6 fungal copies (Battleship Promontory south sun exposure, Dry Valleys); conversely, the differently north and south exposed rock surfaces of Linnaeus Terrace (Dry Valleys) showed the highest (7.7 x 10^5) and the lowest (1.3 x 10^3) bacterial gene-copies, respectively (Fig. 4 and Table 1S).

Overall, both fungal and bacterial abundances varied between the two sun-exposures, except in the case of Finger Mt. and University Valley in Dry Valleys, where abundance of the two major biological compartments was similar. Furthermore, microbial abundance was generally higher in north-exposed rocks, with only few exceptions: fungi were most abundant in south sun-exposed samples at Battleship Promontory, while bacteria were larger in number at Mt. New Zealand south (Fig.4).
The fungal-to bacterial ratio (F/B) (based on log-copy numbers) showed a slightly higher dominance of fungi in these ecosystems in both sun-exposed sampled localities and were

dominance of fungi in these ecosystems in both sun-exposed sampled localities and were significantly different in all samples (p<0.05). In north-exposed rocks F/B was 1.24 ± 0.2 (mean \pm

SD), while in south-exposed surface it varied between 1.37 ± 0.31 (data not shown).

250

4. Discussions

The Antarctic cryptoendolithic communities host among the most resistant microorganisms on Earth, being able to survive under environmental conditions once accounted as incompatible with life (Horowitz et al., 1972).

Recent studies have improved knowledge of the microbial diversity and composition of these ecosystems (Wei et al., 2016; Archer et al., 2017) and provided important insights to clarify the complex relationship between environmental parameters (i.e. altitude and distance from sea) and biodiversity (Selbmann et al., 2017; Coleine et al., 2018a, b). The lack of consistent microbial diversity patterns along altitudinal and sea distance gradients has suggested that new hypotheses about which can act as driving factor shaping biodiversity must put forward (Selbmann et al., 2017; Coleine et al, 2018a).

- Towards this end, the aim of this study was to test the potential effects on the structure and composition of the bacterial, fungal and algal assemblages of a new parameter, namely sun exposure, as significant variable influencing endolithic settlement and development, using a preliminary molecular screening approach on a considerable amount of rock samples collected along the Victoria Land (Continental Antarctica).
- 267

268 We found that DGGE-based similarity assessment visualized as dendrograms indicated a clear grouping of samples according mostly by sun exposure, highlighting an evident relationship 269 270 between microbial composition and this environmental parameter. This trend is consistent in all biological compartments examined. Particularly, a clear clustering was observed for DGGE profiles 271 272 obtained from samples collected in Pudding Butte, Battleship Promontory, Mt. New Zealand and 273 Finger Mt. in fungi and algae (Fig. 2, data not shown); as for eukaryotes, in bacteria, clear groups 274 were observed in samples belonging to Battleship Promontory, Finger Mt., Pudding Butte, 275 Knobhead and Linnaeus Terrace.

- Considering all the data, higher similarities in fingerprinting profiles were observed within samples
 collected in the same site, with very few scattered exceptions, likely due to variability among
 samples.
- These findings highlighted the effectiveness of sampling strategy applied for the Antarctic Expedition 2015-16 and did not reflect what was observed by Selbmann et al. (2017). In that study, based on a previous sampling (PNRA, Antarctic Expedition 2010-11), the largest Antarctic sampling to date of rocks hosting lithic communities, including different rock typologies (i.e. sandstone, granite, dolerite) from 46 different localities in Victoria Land, was investigated. Results indicated a remarkable local variability found even in rocks from the same site, maybe due to the size of sampled area (about 100 mq²) and to variability of rock typologies collected.
- 286 DGGE band profiles were analysed by statistical analysis and, consistently to clustering analysis, 287 samples were mainly grouped by sun exposure along the three analysed biological groups. The 288 observed shift in community composition correlated to sun exposure was also confirmed by Non-289 Metric Multidimensional scaling (NMDS) analysis, which organizes data into 2-D spatial graphs by 290 reducing dimensionality. An apparent gradient in abundance and community composition in 291 response to sun exposure was previously observed in Coleine et al. (2018b), where authors 292 investigated biodiversity and composition of functional groups of fungi in Antarctic endolithic 293 communities and reported the absence of any correlation with altitude and sea distance, while a 294 remarkable variability was observed considering the sun exposure parameter.

- It was previously hypothesized that the distribution of endoliths reflects the degree of insulation on the rock surfaces; in northern exposed rocks, environmental conditions are more favourable than southern exposed faces, and cryptoendolithic colonization is more often observed and favoured (Friedmann, 1977; Friedmann and Weed, 1987). The capability to maintain biological activity may depend on sufficient insulation of the rock to allow an efficient photosynthetic process; moreover, warmer temperatures will allow metabolic activity and more water due to snow melt (McKay and
- 302 Friedmann, 1985; Deegenaars and Watson, 1998).
- 303 Our findings support the hypothesis that sun radiation not only affects temperature of rocks surface, 304 availability of water and metabolic activity, but even the biodiversity.
- 305

295

- 306 Community structure was also shaped by north and south exposure when fungal and bacterial 307 abundances were estimated with a qPCR assay. In the most localities, fungal (ranging from 6.1 x 308 10^4 to 9.8 x 10^6 gene copies) and bacterial (7.7 x 10^5 to 1.3 x 10^3 gene copies) abundances changed 309 significantly according to sun exposure.
- 310 We also calculated the fungal:bacterial (F/B) ratio, a metric to assess environmental impacts and the
- 311 functional implications of microbial communities (Raeymaekers, 2000; Fierer et al., 2005). The F/B
- 312 dominance was significantly affected by sun exposure and showed a significative dominance of
- 313 fungi in these microbiomes. Fungi predominate in the south-exposed rocks (with the only exception
- 314 of Finger Mt. in McMurdo Dry Valleys) where conditions are much more extreme; indeed, we
- found a greater fungal dominance (F/B, 1.37) respect than north-exposed sites (F/B, 1.24).
- 316

317 We propose that further studies are needed to elucidate the relationship between fungi and bacteria, 318 and to reveal their functions in these ecosystems. Nevertheless, this result is not surprisingly: 319 extreme environments (i.e. the ice-free desert of Victoria Land in Continental Antarctica) are not an 320 only prerogative of archaea and bacteria. Among eukaryotes, fungi (alone or in symbiosis with 321 cyanobacteria or algae forming lichens) are the most versatile and ecologically successful 322 phylogenetic lineage; they evolved to survive and proliferate in the extremes (Magan, 2007), even 323 in habitats normally precluded to the most (Selbmann et al., 2013). Fungi are, for instance, 324 generally more resistant to desiccation than bacteria with hyphae that may cross air-filled soil pores 325 to access nutrients and water (Gordon et al., 2008; de Vries et al., 2012), showing a remarkable 326 ability to survive in stress conditions (e.g. low water availability) of Antarctic ice-free areas (Onofri 327 et al., 2004).

328 The most remarkable example for stress resistance is, in fact, given by the fungus Cryomyces 329 antarcticus, a cryptoendolithic black fungus isolated from the McMurdo Dry Valleys in Antarctica, 330 chosen as the best eukaryotic test organisms for astrobiological investigations for its stunning 331 resistance to temperature cycles (-20/+20 °C), high temperature (+90 °C) and saline concentration 332 (up to 25% NaCl) (Onofri et al., 2008; 2012; 2015). It was found to resist to ionizing radiation up to 333 55.81 kGy (Selbmann et al., 2018), while the bacterium Deinococcus radiodurans, widely considered the extremophile par excellence and gold-medalist of radiation resistance (Battista et al., 334 335 1997; Venkateswaran et al., 2000), survives up to 20 kGy of gamma radiation.

336

In conclusion, even though DGGE-based approach does not show individual taxa responses and gives limited insight into taxonomy, it is still currently considered a powerful, rapid and costly-

- effective method for determining shapes in microbial community composition (Zheng et al., 2013;
 Kovalski Mitter et al., 2018). This technique gave us the advantage to rapidly and inexpensively
 analyze a great number of rock samples and three biological domains (fungi, algae and bacteria),
 providing a clear trend of how sun exposure influences cryptoendolithic community biodiversity
 and composition.
- 344 This study expands existing knowledge on the relationship between environmental parameters and
- 345 Antarctic endolithic biodiversity and examines endolithic community structure. The results inform
- 346 planning of future Antarctic campaigns to maximize identification of active endolithic communities.
- With the advent of -omics approaches such as metabarcoding, metagenomics and metabolomics, we may explore the composition deeply focusing on a targeted selection of rock samples. These samplings will characterize taxa identity and abundance in rocks hosting endolithic communities, examine the complex community dynamics, their stress-adaptation strategies, and potential functions across an environmental variation gradient that is influenced by sun exposure.
- 352 A detailed picture on how these communities respond to increasing environmental pressures will
- 353 also give clues for predicting and monitoring the effects of global change on these unique border
- 354 ecosystems and their vulnerable biodiversity.

355 ACKNOWLEDGMENTS

L.S., C.C. and L.Z. wish to thank the Italian National Program for Antarctic Researches (PNRA) for funding sampling campaigns and researches activities in Italy in the frame of Projects 2009/A1.11, 2013/AZ-17, 2015/AZ1.02 and AMunDsEN PNRA_00006. The Italian Antarctic National Museum (MNA) is acknowledged for financial support to the Mycological Section on the MNA for preserving rock Antarctic samples analysed in this study and stored in the Culture Collection of Fungi from Extreme Environments

361 (CCFEE), University of Tuscia, Italy.

362 Figure captions

- Figure 1. Dendrogram obtained by Unweighted Pair Group Method using Arithmetic average (UPGMA)
 based on the DGGE profiles of the fungal component of the endolithic communities.
- 365 The relationships among samples are based on similarity, evaluated by the Dice coefficient.
- 366 Triplicate samples were analysed.
- 367 Figure 2. Dendrogram obtained by Unweighted Pair Group Method using Arithmetic average (UPGMA)
- 368 based on the DGGE profiles of the bacterial component of the endolithic communities.
- 369 The relationships among samples are based on similarity, evaluated by the Dice coefficient.
- 370 Triplicate samples were analysed.
- 371 Figure 3. Non-Metric Multidimensional Scaling (N-MDS) ordination plots for fungal (A) and bacterial
- 372 endolithic communities (B), calculating the Bray–Curtis index, based on square-root transformed abundance
- 373 data.
- 374 Stress value (A): 0.09.
- 375 Stress value (B): 0.07
- 376 Figure 4. Abundances of total fungi and bacteria in endolithic communities, as estimated using the qPCR
- assays. Error bars are the standard errors. Significant differences are calculated by Tukey test with p
 value<0.05 and indicated by different letters.
- 379

380 References

- Archer, S.D., de los Ríos, A., Lee, K.C., *et al.* 2017. Endolithic microbial diversity in sandstone and granite
 from the McMurdo Dry Valleys, Antarctica. Polar Biol. 40(5), 997–1006.
- Battista, J. R. 1997. Against all odds: the survival strategies of *Deinococcus radiodurans*. Ann. Rev.
 Microbiol. 51(1), 203–224.
- Bekryaev, R.V., Polyakov, I.V., Alexeev, V.A. 2010. Role of polar amplification in long-term surface air temperature variations and modern Arctic warming. J. Climate 23(14), 3888–3906.
- 387 Bell, R.A. 1993. Cryptoendolithic algae of hot semiarid lands and deserts. J. Phycol. 29, 133–139.
- Bhatta, K.P., Grytnes, J.A., Vetaas, O.R. 2018. Downhill shift of alpine plant assemblages under contemporary climate and land- use changes. Ecosphere 9(1).
- Cary, S.C., McDonald, I.R., Barrett J.E. *et al.* 2010. On the rocks: the microbiology of Antarctic dry valley
 soils. Nat. Rev. Microbiol. 8, 129–138.
- Clarke, K.R., 1993. Non-parametric multivariate analyses of changes in community structure. Aust. J. Ecol.
 18, 117–143.
- Coleine, C., Stajich, J.E., Zucconi, L., *et al.* 2018a. Antarctic cryptoendolithic fungal communities are highly
 adapted and dominated by Lecanoromycetes and Dothideomycetes. Front. Microbiol. 9.
- 396 Coleine, C., Zucconi, L., Onofri, S., *et al.* 2018b. Sun exposure shapes functional grouping of fungi 397 cryptoendolithic Antarctic communities. Life 8(2).
- 398 Cowan, D., Tow, L. 2004. Endangered Antarctic environments. Annu. Rev. Microbiol. 58:649–690.
- Cowan, D., Makhalanyane, T.P., Dennis, P.G., *et al.* 2014. Microbial ecology and biogeochemistry of
 continental Antarctic soils. Front. Microbiol. 5, 154.
- 401 Deegenaars, M. L., Watson, K. 1998. Heat shock response in psychrophilic and psychrotrophic yeast from
 402 Antarctica. Extremophiles 2(1), 41–50.
- 403 Descamps, S., Aars, J., Fuglei, E., *et al.* 2017. Climate change impacts on wildlife in a High Arctic 404 Archipelago–Svalbard, Norway. Global Change Biol. 23(2), 490–502.
- De Vries, F.T., Liiri, M.E., Bjørnlund, L., *et al.* 2012. Land use alters the resistance and resilience of soil
 food webs to drought. Nat. Clim. Change 2(4), 276.
- 407 Egidi, E., De Hoog, G.S., Isola, D., *et al.* 2014. Phylogeny and taxonomy of meristematic rock-inhabiting
 408 black fungi in the dothidemycetes based on multi-locus phylogenies. Fungal Divers. 65, 127–165.
- Farrell, R.L., Arenz, B.E., Duncan, S.M., *et al.* 2011. Introduced and indigenous fungi of the Ross Island
 historic huts and pristine areas of Antarctica. Polar Biol. 34, 1669–1677.
- Fierer, N., Jackson, J.A., Vilgalys, R., *et al.* 2005. Assessment of soil microbial community structure by use
 of taxon-specific quantitative PCR assays. Appl Environ Microbiol. 71(7), 4117–4120.
- 413 Friedmann, E.I., Ocampo, R. 1976. Endolithic blue-green algae in dry valleys-primary producers in Antarctic

- 414 desert ecosystem. Science 193, 1247–1249
- 415 Friedmann, E.I. 1977. Microorganisms in Antarctic desert rocks from dry valleys and Dufek Massif.
 416 Antarctic J. US 12(4), 26–29.
- 417 Friedmann, E.I. 1982. Endolithic microorganisms in the Antarctic cold desert. Science 215, 1045–1053
- 418 Friedmann, E.I., Weed, R. 1987. Microbial trace-fossil formation, biogenous, and abiotic weathering in the 419 Antarctic cold desert. Science 236(4802), 703-705.
- 420 Gardes, M., Bruns, T.D. 1993. ITS primers with enhanced specificity for basidiomycetes- application to the 421 identification of mycorrhizae and rusts. Mol. Ecol. 2(2), 113–118.
- 422 Gordon, H., Haygarth, P.M., Bardgett, R.D. 2008. Drying and rewetting effects on soil microbial community 423 composition and nutrient leaching. Soil Biol. Biochem. 40(2), 302–311.
- Horowitz, N.H., Cameron, R.E., Hubbard, J.S. 1972. Microbiology of the dry valleys of Antarctica. Science
 176(4032), 242–245.
- 426 Kovalski-Mitter, E., de Freitas, R., Germida, J.J. 2018. Microbial communities associated with barley 427 growing in an oil sands reclamation area in Alberta, Canada. Can. J. Microbiol. 64(12), 1004–1019.
- 428 Magan, N. 2007. Fungi in extreme environments. The Mycota 4, 85–103.
- 429 McKay, C.P., Friedmann, E.I. 1985. The cryptoendolithic microbial environment in the Antarctic cold 430 desert: temperature variations in nature. Polar Biol. 4(1), 19–25.
- Muyzer, G., de Waal, E.C., Uitterlinden, A.G. 1993. Profiling of complex microbial populations by
 denaturing gradient gel electrophoresis analysis of polymerase chain reaction amplified genes coding for 16S
 rRNA. Appl. Environ. Microbiol. 59, 695–700.
- 434 NAS 2011. Future science opportunities in Antarctica and the Southern Ocean (report in brief). National
 435 Academy of Sciences <u>http://wwwdels.nas.edu/prb</u>.
- 436 Nienow, J.A., Friedmann, E.I. 1993. Terrestrial lithophytic (rock) communities. In: Friedmann EI (ed)
 437 Antarctic microbiology. Wiley-Liss, New York, 343–412.
- Olech, M., Chwedorzewska, K.J. 2011. Short note. The first appearance and establishment of an alien
 vascular plant in natural habitats on the forefield of a retreating glacier in Antarctica. Antarct. Sci. 23, 153–
 154
- 441 Onofri, S., Selbmann, L., Zucconi, L., Pagano, S. 2004. Antarctic microfungi as models for exobiology.
 442 Planet. Space Sci. 52(1-3), 229–237.
- Onofri, S., Barreca, D., Selbmann, L., *et al.* 2008. Resistance of Antarctic black fungi and cryptoendolithic
 communities to simulated space and Mars conditions. Stud Mycol 61, 99–109.
- Onofri, S., de la Torre, R., de Vera, J.P., *et al.* 2012. Survival of rock-colonizing organisms after 1.5
 years in outer space. Astrobiology 12, 508-516.
- 447 Onofri S., de Vera J.P., Zucconi L., et al. 2015. Survival of Antarctic Cryptoendolithic Fungi in Simulated

- 448 Martian Conditions nn Board the International Space Station. Astrobiology 15, 1052–1059.
- Parmesan, C. 2006. Ecological and evolutionary responses to recent climate change. Annu. Rev. Ecol. Evol.
 Syst. 37, 637–669.
- Quintal, H., Head, J., Palumbo, A., Dickson, J. 2018. McMurdo Dry Valleys: Exploring Antarctica as a Mars
 Analogue. In Lunar and Planetary Science Conference, 49.
- 453 Raeymaekers, L. 2000. Basic principles of quantitative PCR. Molecular Biotechnol. 15(2), 115–122.
- 454 Selbmann, L., de Hoog, G.S., Mazzaglia, A. *et al* 2005. Fungi at the edge of life: cryptoendolithic black 455 fungi from Antarctic deserts. Stud. Mycol. 51, 1–32.
- Selbmann, L., de Hoog, G.S., Zucconi, L. *et al* 2008. Drought meets acid: three new genera in a dothidealean
 clade of extremotolerant fungi. Stud. Mycol. 61, 1–20.
- Selbmann, L., Isola, D., Fenice, F., 2012. Potential extinction of Antarctic endemic fungal species as a
 consequence of Global Warming. Science Total Environ. 438, 127–134.
- 460 Selbmann, L., Egidi, E., Isola, D., 2013. Biodiversity, evolution and adaptation of fungi in extreme 461 environments. Plant. Biosyst 147(1), 237–246.
- Selbmann, L., Onofri, S., Coleine, C., *et al.* 2017. Effect of environmental parameters on biodiversity of the
 fungal component in the lithic Antarctic communities. Extremophiles 21, 1069–1080.
- Selbmann, L., Pacelli, C., Zucconi, L., *et al.* 2018. Resistance of an Antarctic cryptoendolithic black fungus
 to radiation gives new insights of astrobiological relevance. Fungal Biol. 122(6), 546–554.
- 466 Steig, E.J., Schneider, D.P., Rutherford, S.D., *et al.* 2009. Warming of the Antarctic ice-sheet surface since 467 the 1957 International Geophysical Year. Nature 457(7228), 459.
- 468 Turner, J., S.R. Colwell, G.J. Marshall, *et al.* 2005. Antarctic climate change during the last 50 years. Int. J.
 469 Climatol. 25, 279–294.
- 470 Turner, J., Overland J.E., Walsh, J.E. 2007. An Arctic and Antarctic perspective on recent climate change.
 471 Int. J. Climatol. 27, 277–293.
- 472 Valášková, V., Baldrian, P. 2009. Denaturing gradient gel electrophoresis as a fingerprinting method for the
 473 analysis of soil microbial communities. Plant. Soil Environ. 55(10), 413–423.
- Venkateswaran, A., McFarlan, S.C., Ghosal, D., *et al.* 2000. Physiologic determinants of radiation resistance
 inDeinococcus radiodurans. Applied Environ. Microbiol. 66(6), 2620–2626.
- Walker, J.J., Spear, J.R., Pace, N.R. 2005. Geobiology of a microbial endolithic community in the
 Yellowstone geothermal environment. Nature 434, 1011–1014.
- Wei, S.T., Lacap-Bugler, D.C., Lau, M.C., *et al.* 2016. Taxonomic and functional diversity of soil and
 hypolithic microbial communities in Miers Valley, McMurdo Dry Valleys, Antarctica. Front. Microbiol. 7,
 1642.
- 481 White, T.J., Bruns, T., Lee, S., Taylor, J. 1990. Amplification and direct sequencing of fungal ribosomal

- 482 RNA genes for phylogenetics. In: Innis MA, Gelfand DH, Sninsky JJ, White TJ (eds) PCR Protocols: a
 483 Guide to Methods and Applications, 315-322. Academic Press, New York.
- 484 Yao, T., Thompson, L.G., Mosbrugger, V., *et al.* 2012. Third pole environment (TPE). Environmental
 485 Development 3, 52–64.
- Yang, K., Wu, H., Qin, J., *et al.* 2014. Recent climate changes over the Tibetan Plateau and their impacts on
 energy and water cycle: A review. Glob. Planet. Change 112, 79–91.
- Zheng, J., Liang, R., Zhang, L., *et al.* 2013. Characterization of microbial communities in strong aromatic
 liquor fermentation pit muds of different ages assessed by combined DGGE and PLFA analyses. *Food Res. Int.* 54(1), 660–666.
- Zucconi, L., Onofri, S., Cecchini, C., *et al.* 2016. Mapping the lithic colonization at the boundaries of life in
 Northern Victoria Land, Antarctica. Polar Biol 39(1), 91–102.

Table 1. Table lists sampling details of eight visited localities in Victoria Land, Antarctica: sun exposure, altitude, air temperature (measured when sampling), relative humidity and geographic coordinates.

Locality	Sample	Sun	Altitude	Temperature	Humidity	Coordinates
Battleship Promontory	1	North	834	-4.4	22.9	76°54'04.0''S 160°54'36.6''E
Battleship Promontory	2	North	834	-4.4	22.9	76°54'04.0''S 160°54'36.6''E
Battleship Promontory	3	North	834	-4.4	22.9	76°54'04.0''S 160°54'36.6''E
Battleship Promontory	1	South	834	-4.4	22.9	76°54'04.0''S 160°54'36.6''E
Battleship Promontory	5	South	834	-4.4	22.9	76°54'04.0''S 160°54'36.6''E
Battleship Promontory	6	South	834	-4.4	22.9	76°54'04.0''S 160°54'36.6''E
Pudding Butte	2	North	1573	8.5	32.4	75°51'30.2''S 159°58'25.7''E
Pudding Butte	3	North	1573	8.5	32.4	75°51'30.2''S 159°58'25.7''E
Pudding Butte	4	North	1573	8.5	32.4	75°51'30.2''S 159°58'25.7''E
Pudding Butte	1	South	1588	8.5	32.4	75°51'33.0''S 159°58'26.6''E
Pudding Butte	2	South	1588	8.5	32.4	75°51'33.0''S 159°58'26.6''E
Pudding Butte	3	South	1588	8.5	32.4	75°51'33.0''S 159°58'26.6''E
Siegfried Peak	2	North	1620	-9.3	52.8	77°34'43.3''S 161°47'11.7''E
Siegfried Peak	3	North	1620	-9.3	52.8	77°34'43.3''S 161°47'11.7''E
Siegfried Peak	5	North	1620	-9.3	52.8	77°34'43.3''S 161°47'11.7''E
Siegfried Peak	3	South	1620	-6.8	54.9	77°34'39.9''S 161°47'17.4''E
Siegfried Peak	5	South	1620	-6.8	54.9	77°34'39.9''S 161°47'17.4''E
Siegfried Peak	6	South	1620	-6.8	54.9	77°34'39.9''S 161°47'17.4''E
Linnaeus Terrace	3	North	1649	-9.6	58.6	77°36'01.3''S 161°05'00.5''E
Linnaeus Terrace	4	North	1649	-9.6	58.6	77°36'01.3''S 161°05'00.5''E
Linnaeus Terrace	6	North	1649	-9.6	58.6	77°36'01.3''S 161°05'00.5''E
Linnaeus Terrace	2	South	1761	-12.6	68.2	77°37'09.9''S 161°11'50.8''E
Linnaeus Terrace	3	South	1761	-12.6	68.2	77°37'09.9''S 161°11'50.8''E
Linnaeus Terrace	4	South	1761	-12.6	68.2	77°37'09.9''S 161°11'50.8''E
Finger Mt.	2	North	1720	-6.4	35.1	77°45'0.9"S 160°44'44.5" E
Finger Mt.	3	North	1720	-6.4	35.1	77°45'0.9"S 160°44'44.5" E
Finger Mt.	6	North	1720	-6.4	35.1	77°45'0.91"S 160°44'42.9"
Finger Mt.	2	South	1720	-6.4	35.1	77°45'10''S 160°44'40''E
Finger Mt.	4	South	1720	-6.4	35.1	77°45'10"S 160°44'44.39.7" E
Finger Mt.	5	South	1720	-6.4	35.1	77°45'10.1"S 160°44'45.1" E
University Valley	2	North	2090	-14.3	18	77°52'28.6''S 160°44'22.6''E
University Valley	3	North	2090	-14.3	18	77°52'28.6"S 160°44'22.6" E
University Valley	4	North	2090	-14.3	18	77°52'29"S 160°44'22.3" E
University Valley	3	South	2200	-11.2	39.1	77°52'21.5''S 160°45'19.2''E
University Valley	5	South	2200	-11.2	39.1	77°52'21.5''S 160°45'19.2''E
University Valley	6	South	2200	-11.2	39.1	77°52'21.5''S 160°45'19.2''E
Knobhead	1	North	2150	-12.5	50	77°54'37.8''S 161°34'48.8''E
Knobhead	2	North	2150	-12.5	50	77°54'37.8''S 161°34'48.8''E

Knobhead	3	North	2150	-12.5	50	77°54'37.8''S 161°34'48.8''E
Knobhead	1	South	2150	-8.9	38.9	77°54'43.6''S 161°34'39.3''E
Knobhead	2	South	2150	-8.9	38.9	77°54'43.6''S 161°34'39.3''E
Knobhead	3	South	2150	-8.9	38.9	77°54'43.6''S 161°34'39.3''E
Mt. New Zealand	1	North	3100	-17.2	47.6	74°10'44.0''S 162°30'53.0''E
Mt. New Zealand	2	North	3100	-17.2	47.6	74°10'44.0''S 162°30'53.0''E
Mt. New Zealand	3	North	3100	-17.2	47.6	74°10'44.0''S 162°30'53.0''E
Mt. New Zealand	1	South	3100	-17.2	47.6	74°10'44.0''S 162°30'53.0''E
Mt. New Zealand	2	South	3100	-17.2	47.6	74°10'44.0''S 162°30'53.0''E
Mt. New Zealand	3	South	3100	-17.2	47.6	74°10'44.0''S 162°30'53.0''E

Amplified group	Primer	Sequence 5 ¹ -3 ¹			
Fungi	ITS1F-GC ^a / ITS2	GCACGGGGGGGCTTGGTCATTTAG/			
		GCTGCGTTCTTCATCGATGC			
Bacteria	341F-GC ^b /518R	GCACGGGGGGGCCTACGGGAGGC/			
		ATTACCGCGGCTGCTGG			

Table 2. PCR primers used for DGGE in the present study.

Figure Click here to download high resolution image



Figure Click here to download high resolution image



Figure Click here to download high resolution image



NMDS1





Supplementary material for on-line publication only Click here to download Supplementary material for on-line publication only: Table 1S.docx