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**Sun exposure drives Antarctic cryptoendolithic community structure and composition**

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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

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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  
17 conditions become incompatible with active life on rock surfaces. Due to the thermal inertia of  
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 scarcely 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  
25 shaping microbial abundance and diversity, while sun exposure was hypothesized as significant  
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  
30 qPCR techniques, of 48 rocks with varying sun exposure, collected in Victoria Land along an  
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.

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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  
44 regions (Bekryaev et al., 2010). The impact of climate change is, therefore, particularly intense at  
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  
52 Antarctica, is also documented; researchers from the British Antarctic Survey report a warming  
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  
55 reported an apparent contrast between strong warming of the Antarctic Peninsula and slight cooling  
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).

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

65  
66 Endolithism is a specialized colonization by microbes to enable dwelling inside airspaces of rocks.  
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  
75 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 lichen-  
79 dominated communities (Nienow and Friedmann, 1993). These self-supporting microbial  
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 (Zuconni 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 parameters 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  
101 Weed, 1987). Rocks with varying sun exposure (north and south faced surfaces) were collected in  
102 all the localities surveyed. Based on the prior experiences (Selbmann et al., 2017; Coleine et al.,  
103 2018a), it is suggested that sampling strategy and environmental parameters considered may have  
104 important consequences on a proper and exhaustive biodiversity description; thus, before  
105 proceeding with metabarcoding and metagenomics experiments to develop a detailed picture of  
106 microbial diversity and functionality, we have selected a suite of more rapid and cheaper  
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  
114 collected along a latitudinal transect from 74°10'44.0''S 162°30'53.0''E (Mt. New Zealand,  
115 Northern Victoria Land) to 77°54'43.6''S 161°34'39.3''E (Knobhead, Southern Victoria Land),  
116 ranging from 834 (Battleship Promontory, Southern Victoria Land) to 3100 m a.s.l. (Mt. New  
117 Zealand) (Table 1). All rock samples were excised aseptically and collected in triplicate, transported  
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.

120

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  
125 and quantified by Quant-iT dsDNA HS assay kit (Invitrogen molecular probes- Eugene, Oregon,  
126 USA). Microbial diversity was screened by Denaturing Gel Gradient Electrophoresis (DGGE). A  
127 semi-nested PCR was performed using primers with a GC-clamp (Muyzer et al., 1993); fungal and  
128 algal ITS rRNA was amplified from environmental DNA with primers ITS1F-GC/ITS2 (Gardes  
129 and Bruns, 1993; White et al., 1990), following the protocol reported in Selbmann et al. (2017),  
130 while 341F-GC/518R primers (Muyzer et al., 1993) were utilized for 16S rRNA amplification  
131 (Valášková and Baldrian, 2009) (Table 2).

132 Amplicons were purified using NucleoSpin® Gel and PCR Clean-up (Macherey-Nagel, GmbH &  
133 Co. KG, Duren, Germany). One-hundred ng of final DNA concentration were loaded into each well  
134 for DGGE and runs were performed on DGGE Electrophoresis System (C.B.S. Scientific, Del Mar,  
135 California, USA). Gels at 7.5% polyacrylamide (37.5:1 acrylamide:bisacrylamide) were mixed in a  
136 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× TAE Buffer at a constant temperature of 60 °C and 200 V.

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

147 Triplicate samples were processed and analyzed independently.

148

149

### 150 *2.3. NMDS ordination plots*

151 Multivariate statistical analyses were performed to determine the effects of sun exposure on  
152 microbial diversity composition using PAST software (PAleontological STatistics, ver. 2.17) The  
153 effect of this abiotic parameter was tested displaying changes in communities' composition with

154 Non-Metric Multidimensional Scaling (NMDS) based both on abundance data (bands intensity),  
155 calculating Bray-Curtis distance index and presence–absence data, using Jaccard index (Clarke,  
156 1993). Means of abundance data were square-root transformed and analyses were carried out with  
157 999 permutations as described in Coleine et al. (2018b). NMDS were plotted using the combined  
158 occurrence and abundance data of the three replicates from each site. Permutational multivariate  
159 analyses of variance (PERMANOVA,  $p < 0.05$ ) based on the Euclidean distance were utilized to  
160 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)  
165 using NS91F (5'-GTCCCTGCCCTTTGTACACAC-3') and ITS51R (5'-  
166 ACCTTGTTACGACTTTTACTTCCTC-3') for total fungi and Eub338 (5'-  
167 ACTCCTACGGGAGGCAGCAG-3') and Eub518 (5'-ATTACCGCGGCTGCTGG-3') for total  
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*  
170 ITS rRNA gene for fungi and *Escherichia coli* 16S rRNA gene for bacteria. The amplicons were  
171 generated in 100  $\mu$ l reactions, containing 50  $\mu$ l of 2x PCR BioMix™ (Bioline, London, UK), 5  
172 pmol of both forward and reverse primers, 43  $\mu$ l of DEPC water and 5  $\mu$ l template of genomic  
173 DNA. For fungi, amplification of ITS region was carried out as follows: 95 °C for 3 min and the 35  
174 cycles of 95 °C 40 s, 55 °C 30 s, and final extension at 72 °C 40 s; while bacterial 16S region was  
175 amplified as follows: 95 °C for 3 min and the 35 cycles of 95 °C 40 s, 53 °C 30 s, and final  
176 extension at 72 °C 40 s. PCR products were then purified using the NucleoSpin® PCR Clean-up kit  
177 (MACHEREY-NAGEL, GmbH & Co. KG), quantified with Qubit dsDNA HS Assay kit and  
178 cloned using the pGEM®-T Easy Vector Systems (Promega, Madison, Wisconsin, US). Plasmids  
179 were isolated using the NucleoSpin Plasmid kit (Macherey-Nagel, GmbH & Co. KG).

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

181 The 25  $\mu$ l qPCR reactions contained 12.5  $\mu$ l iQ™ SYBR® Green Supermix (Bio-Rad, Hercules,  
182 California, US), 1  $\mu$ l of each forward and reverse primers, 0.3 ng of environmental DNA or standard  
183 and 9.5  $\mu$ l nuclease-free water. The reactions were carried out on Quantitative real-time BioRad  
184 CFX96™ 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  
186 products were of the appropriate size. Fungal and bacterial gene copy numbers were generated  
187 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  
205 also supported by the dendrograms, generating DGGE patterns that grouped together as most  
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  
208 scattered exceptions. In particular, in the clustering generated on fungal profiles, samples were split  
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  
211 bacterial profiles (Fig. 2); as for fungi, the splitting was obtained in the communities from Pudding  
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*

217 To investigate the similarity of the fungal and bacterial communities' composition amongst  
218 different sun exposures, a Non-metric Multi-Dimensional Scaling (NMDS) analysis was computed.  
219 NMDS ordination plots were generated both with the only presence-absence matrix using the  
220 Jaccard index and with the combined frequency of occurrence using the Bray-Curtis index. Because  
221 both approaches produced similar results, we showed results based on abundance only.

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

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

232

233 *3.3. Abundance of fungal and bacterial communities*

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

239 and Table 1S).  
240 Overall, both fungal and bacterial abundances varied between the two sun-exposures, except in the  
241 case of Finger Mt. and University Valley in Dry Valleys, where abundance of the two major  
242 biological compartments was similar. Furthermore, microbial abundance was generally higher in  
243 north-exposed rocks, with only few exceptions: fungi were most abundant in south sun-exposed  
244 samples at Battleship Promontory, while bacteria were larger in number at Mt. New Zealand south  
245 (Fig.4).  
246 The fungal-to bacterial ratio (F/B) (based on log-copy numbers) showed a slightly higher  
247 dominance of fungi in these ecosystems in both sun-exposed sampled localities and were  
248 significantly different in all samples ( $p < 0.05$ ). In north-exposed rocks F/B was  $1.24 \pm 0.2$  (mean  $\pm$   
249 SD), while in south-exposed surface it varied between  $1.37 \pm 0.31$  (data not shown).  
250

251 **4. Discussions**

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

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

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

267  
268 We found that DGGE-based similarity assessment visualized as dendrograms indicated a clear  
269 grouping of samples according mostly by sun exposure, highlighting an evident relationship  
270 between microbial composition and this environmental parameter. This trend is consistent in all  
271 biological compartments examined. Particularly, a clear clustering was observed for DGGE profiles  
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.

276 Considering all the data, higher similarities in fingerprinting profiles were observed within samples  
277 collected in the same site, with very few scattered exceptions, likely due to variability among  
278 samples.

279 These findings highlighted the effectiveness of sampling strategy applied for the Antarctic  
280 Expedition 2015-16 and did not reflect what was observed by Selbmann et al. (2017). In that study,  
281 based on a previous sampling (PNRA, Antarctic Expedition 2010-11), the largest Antarctic  
282 sampling to date of rocks hosting lithic communities, including different rock typologies (i.e.  
283 sandstone, granite, dolerite) from 46 different localities in Victoria Land, was investigated. Results  
284 indicated a remarkable local variability found even in rocks from the same site, maybe due to the  
285 size of sampled area (about 100 mq<sup>2</sup>) 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.

295

296 It was previously hypothesized that the distribution of endoliths reflects the degree of insulation on  
297 the rock surfaces; in northern exposed rocks, environmental conditions are more favourable than  
298 southern exposed faces, and cryptoendolithic colonization is more often observed and favoured  
299 (Friedmann, 1977; Friedmann and Weed, 1987). The capability to maintain biological activity may  
300 depend on sufficient insulation of the rock to allow an efficient photosynthetic process; moreover,  
301 warmer temperatures will allow metabolic activity and more water due to snow melt (McKay and  
302 Friedmann, 1985; Deegenaaers 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

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  $6.1 \times$   
308  $10^4$  to  $9.8 \times 10^6$  gene copies) and bacterial ( $7.7 \times 10^5$  to  $1.3 \times 10^3$  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 significant 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  
315 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  
334 considered the extremophile par excellence and gold-medalist of radiation resistance (Battista et al.,  
335 1997; Venkateswaran et al., 2000), survives up to 20 kGy of gamma radiation.

336

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

339 effective method for determining shapes in microbial community composition (Zheng et al., 2013;  
340 Kovalski Mitter et al., 2018). This technique gave us the advantage to rapidly and inexpensively  
341 analyze a great number of rock samples and three biological domains (fungi, algae and bacteria),  
342 providing a clear trend of how sun exposure influences cryptoendolithic community biodiversity  
343 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.  
347 With the advent of -omics approaches such as metabarcoding, metagenomics and metabolomics, we  
348 may explore the composition deeply focusing on a targeted selection of rock samples. These  
349 samplings will characterize taxa identity and abundance in rocks hosting endolithic communities,  
350 examine the complex community dynamics, their stress-adaptation strategies, and potential  
351 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.

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359 acknowledged for financial support to the Mycological Section on the MNA for preserving rock Antarctic  
360 samples analysed in this study and stored in the Culture Collection of Fungi from Extreme Environments  
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362 **Figure captions**

363 **Figure 1.** Dendrogram obtained by Unweighted Pair Group Method using Arithmetic average (UPGMA)  
364 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  
377 assays. Error bars are the standard errors. Significant differences are calculated by Tukey test with p  
378 value<0.05 and indicated by different letters.

379

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**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

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

Amplified group	Primer	Sequence 5 <sup>1</sup> -3 <sup>1</sup>
Fungi	ITS1F-GC <sup>a</sup> / ITS2	GCACGGGGGGCTTGGTCATTTAG/ GCTGCGTTCTTCATCGATGC
Bacteria	341F-GC <sup>b</sup> /518R	GCACGGGGGGCCTACGGGAGGC/ ATTACCGCGGCTGCTGG

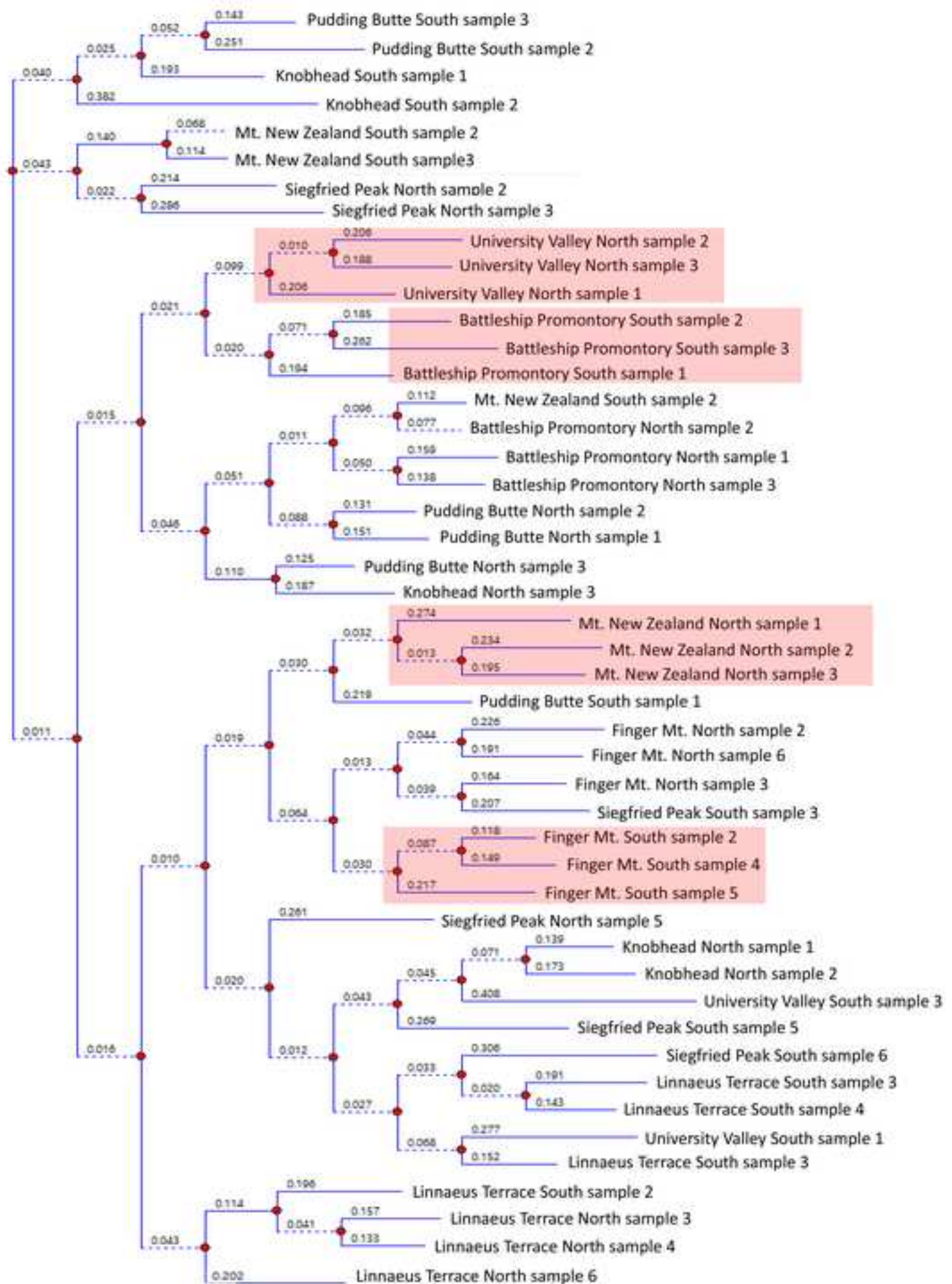
<sup>a</sup>5'-CGCCCGCCGCGCCCCGCGCCCGTCCCGCCGCCCCGCCCG-3';

<sup>b</sup>5'-CGCCCGCCGCGCGCGGCGGGCGGGGCGGGGGCACGGG-3'.



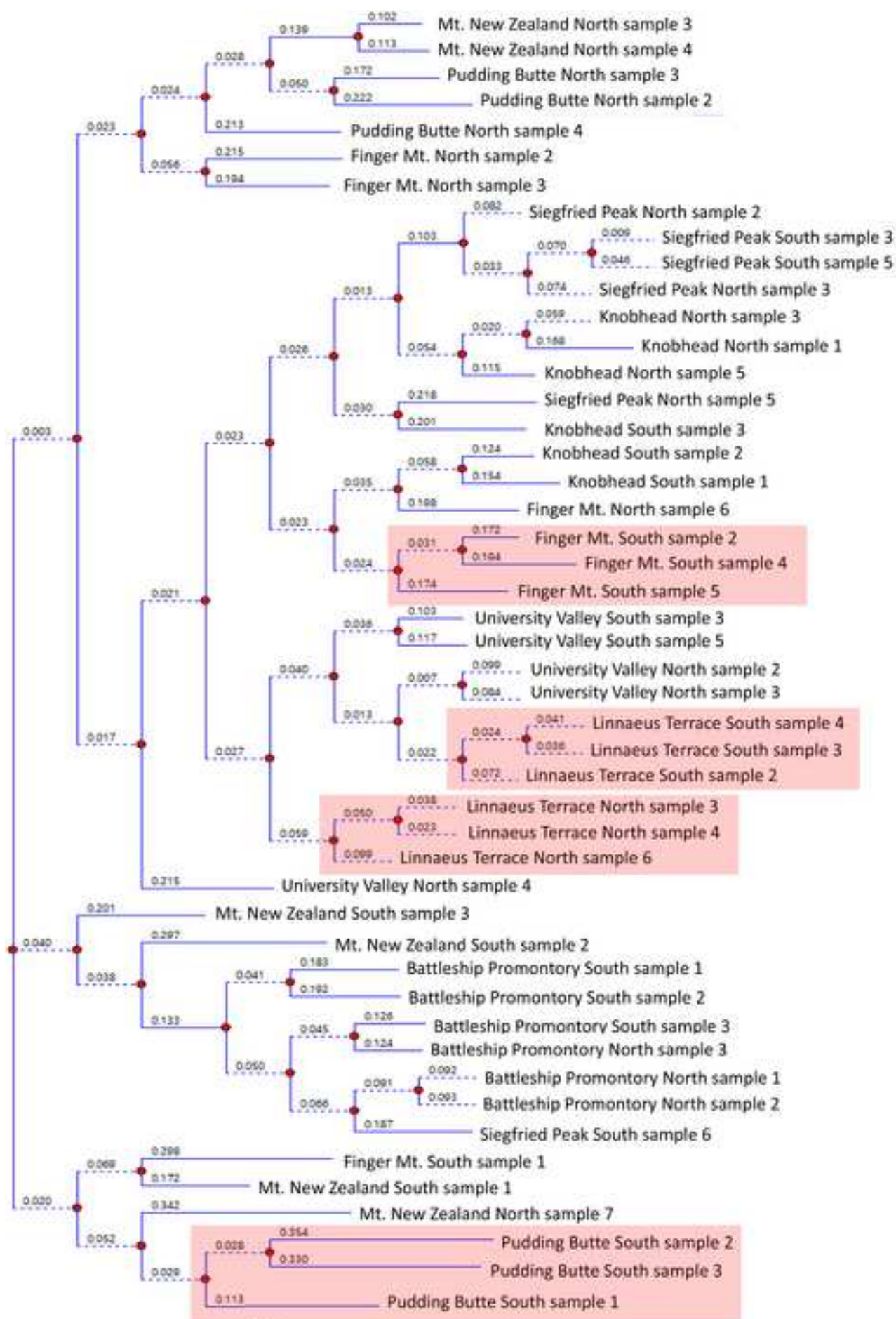
Figure

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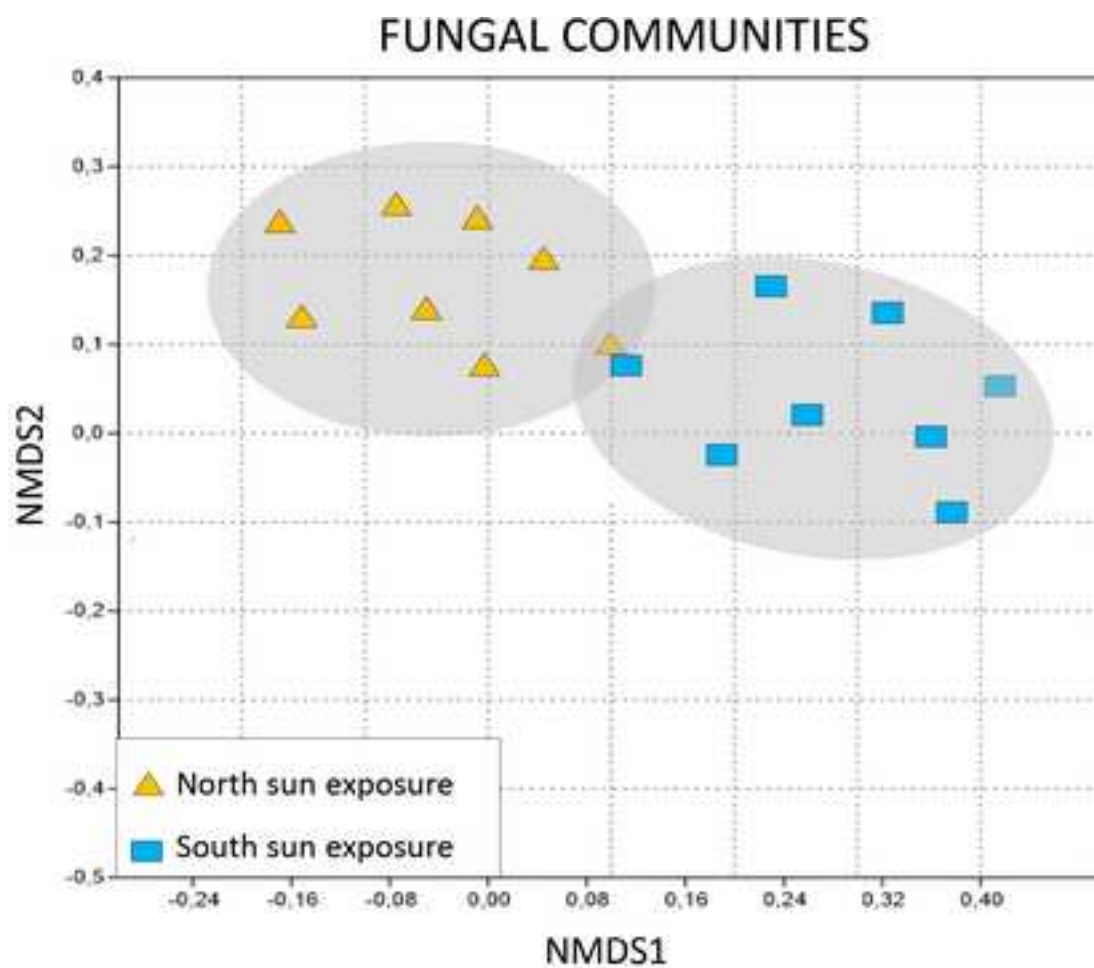


Figure

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**A**



**B**

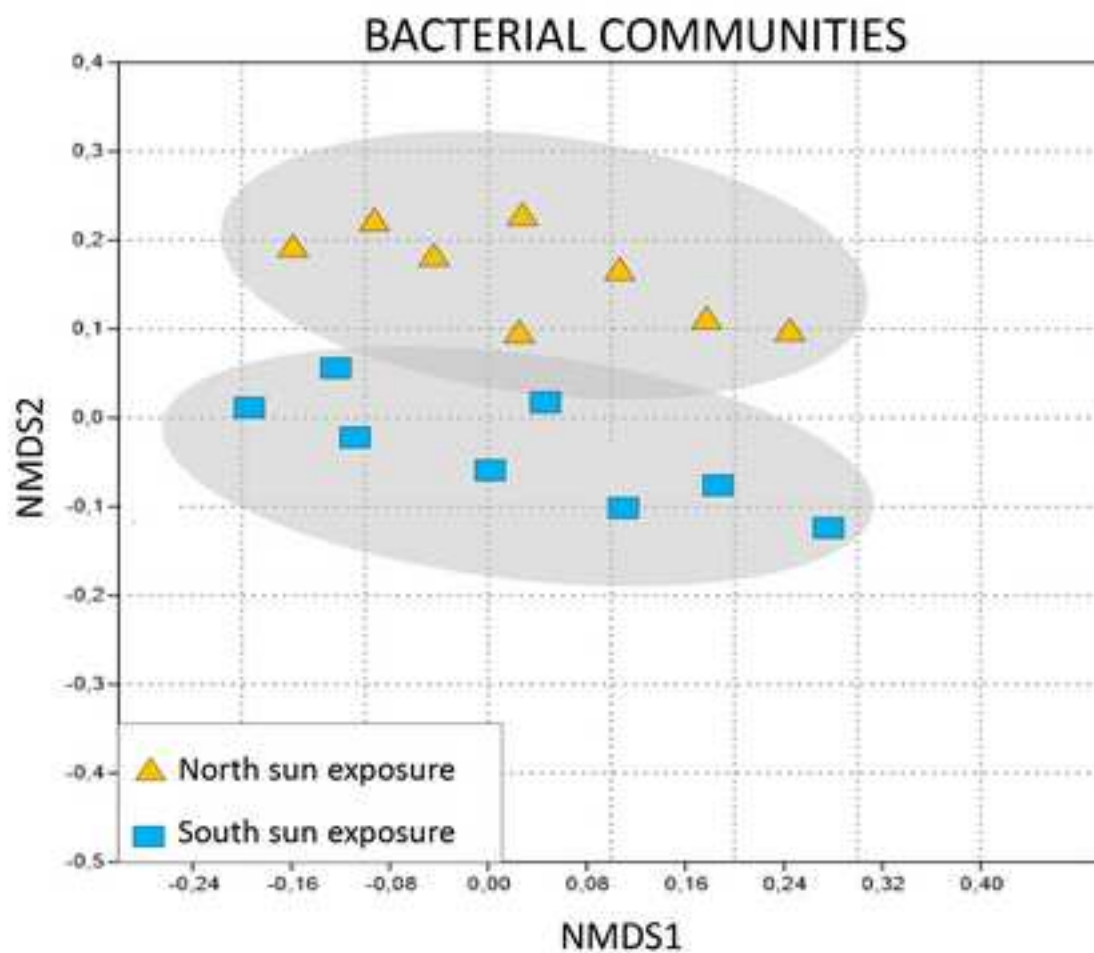
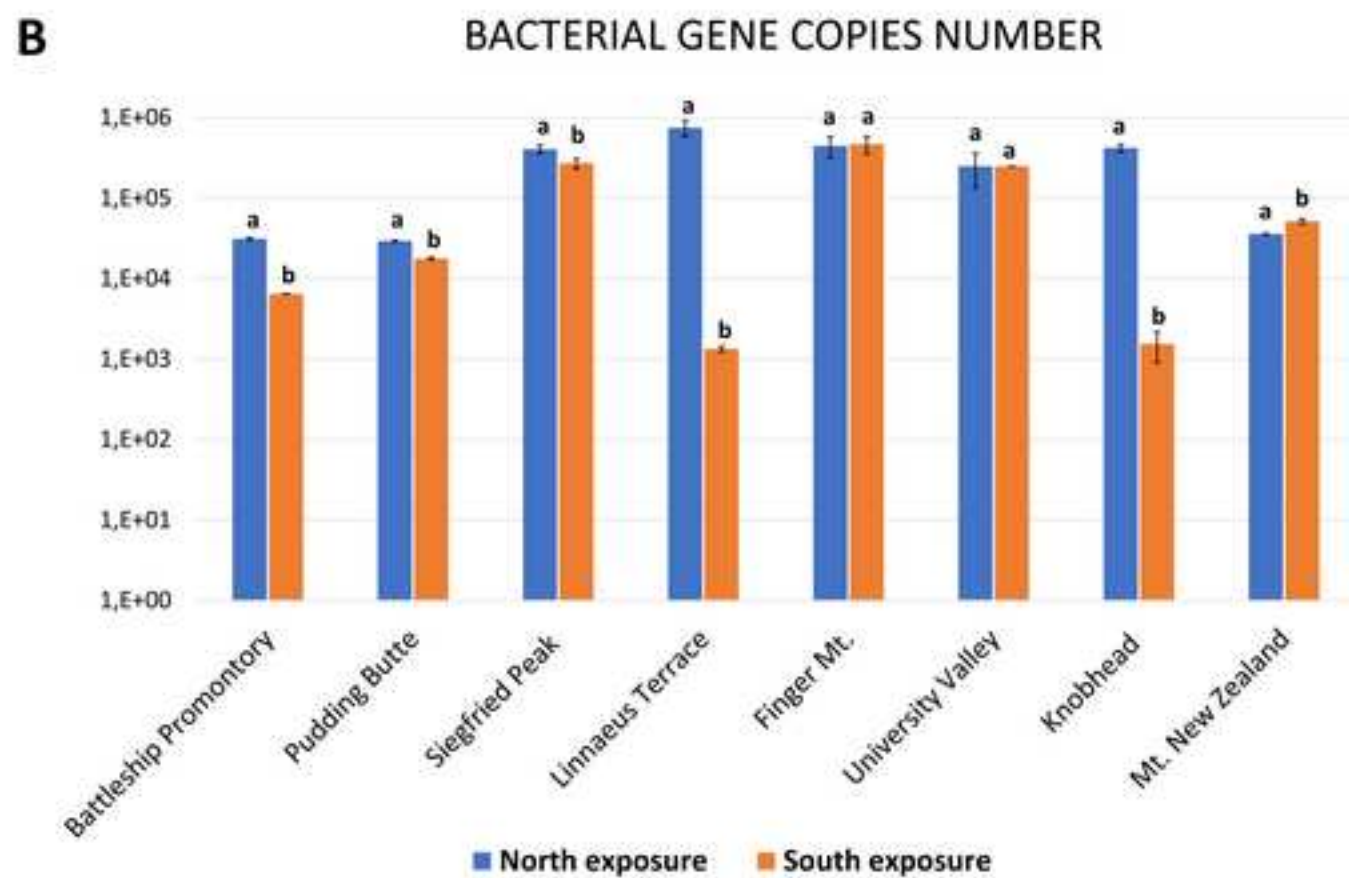
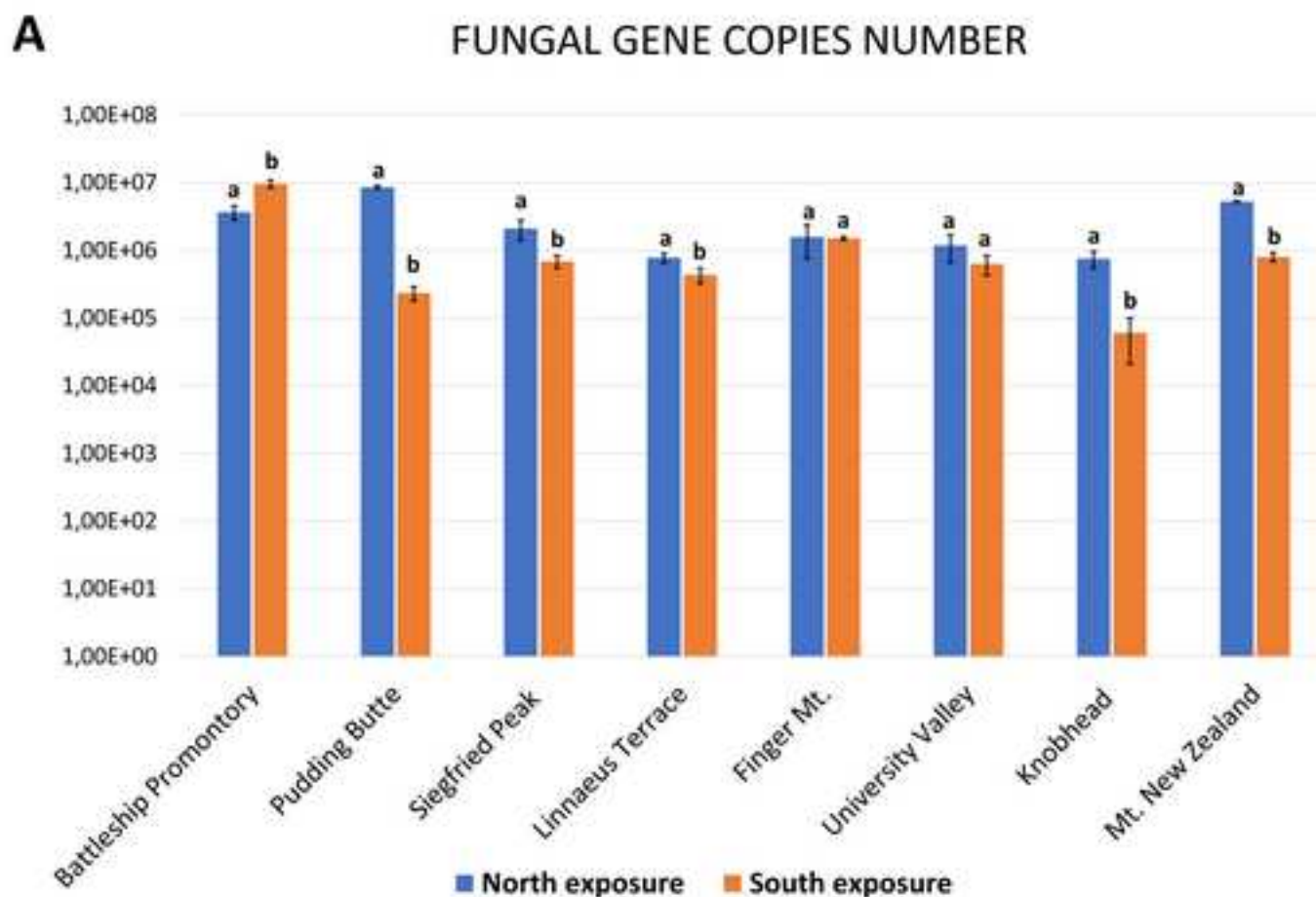




Figure  
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