- 1 Lichen photobiont diversity and selectivity at the southern limit of the maritime Antarctic
- 2 region (Coal Nunatak, Alexander Island)
- 3
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25 Abstract

26 Antarctic ice-free inland sites provide an unique perspective on the strategies coevolving 27 organisms have developed for survival at the limits of life. Here, we provide the first combined 28 description of the ecological and genetic diversity of lichen photobionts colonising an isolated 29 Antarctic inland site, Coal Nunatak, on south-east Alexander Island (Antarctic Peninsula). 30 Photobionts of 14 lichen species (42 samples) representing the entire lichen community of Coal 31 Nunatak were investigated using the internal transcribed spacer region (ITS) of the nuclear 32 ribosomal DNA. The study attempted to address the hypothesis that mycobiont selectivity for 33 the photobiont partner is lower in more extreme environments. This hypothesis did not appear 34 to hold true for the entire lichen community except one species. Another aspect focuses on the 35 relevance of the reproduction modus concerning the distribution of photobiont haplotypes in 36 the lichen community. Dispersal of generative mycobiont diaspores depends on lichenisation 37 processes while by dispersal of vegetative diaspores both symbiotic bionts get dispersed.

38

39 Keywords: inland site, extreme environment, symbiotic association, community, genetic 40 diversity, photobiont haplotypes

42 Introduction

43 Antarctica is the windiest, coldest and highest continent on Earth. Less than 0.5% of the 44 Antarctic continent is permanently or seasonally free of ice cover (British Antarctic Survey, 45 2004). Many lichens are successful colonisers of extreme environments, and can be found 46 worldwide in deserts, high mountain ranges, tropical and polar regions. Lichens colonising 47 Antarctic habitats are exposed to some of the most extreme environmental conditions faced in 48 terrestrial environments on Earth (Peck *et al*., 2006), including high levels of UV radiation, 49 both extremely low and very variable temperatures, lack of liquid water and desiccation stress, 50 and high wind speeds. With two flowering plants and approximately 50 liverworts and 104 51 bryophytes known, the roughly 427 recorded lichen species form the dominant element in the 52 diversity of the Antarctic flora (Ochyra, 1998; Bednarek-Ochyra *et al.*, 2000; Øvstedal & 53 Smith, 2001; Convey, 2013).

54

55 The success of lichens under extreme environmental conditions is based on a remarkable 56 symbiotic relation between, at least, two bionts. Approximately 21% of all fungi are known to 57 form lichens (Hawksworth, 1988). In this obligate symbiosis the fungus (mycobiont) is 58 associated with one or more photosynthesizing symbionts, the photobionts, which either can be 59 eukaryotic green algae or prokaryotic cyanobacteria (Hawksworth, 1988). The green algae 60 most commonly found as photobionts in lichens belong to the genus *Trebouxia* (Peveling, 61 1988). Although between 14 000 and 20 000 lichen-forming mycobionts, mainly ascomycota, 62 are estimated to exist (Feuerer & Hawksworth, 2007) they are associated with only a few 63 different photobiont species (Tschermak-Woess, 1988).

64

65 To date, studies of the diversity of lichen photobionts in Antarctica have concentrated on 66 coastal regions along the Antarctic Peninsula (Romeike *et al*., 2002; Brinkmann, 2002;

67 Langohr, 2004; Siegesmund, 2005). Inland sites, where environmental conditions are generally 68 more extreme and terrestrial diversity and levels of community development much lower 69 (Convey & Smith, 1997) have received little attention (Neuburg, 2007; Pérez-Ortega *et al.*, 70 2012). Until recently most studies have been carried out based on traditional taxonomic 71 approaches. However, few studies have addressed the degree of selectivity between bionts and 72 any correlation this may have with ecological factors in more extreme environments as typified 73 by Antarctic inland sites (Pérez-Ortega *et al.*, 2012).

74

75 In this study diversity data are reported from Coal Nunatak (south-eastern Alexander Island, 76 70°03´S 068°31´W), an inland nunatak ecosystem at the extreme southern limit of the maritime 77 Antarctic. A very limited lichen and bryophyte flora is present with only 14 lichen species 78 recorded. These 14 species were collected on Coal Nunatak, in order to investigate the 79 hypothesis that mycobiont selectivity for the photobiont partner will be lower in more extreme 80 environments (Romeike *et al*. 2002). The slow rates of community development at locations 81 such as Coal Nunatak, which is characterized by very harsh environmental conditions, provide 82 an opportunity to study lichen photobiont diversity. A second hypothesis relates to the 83 reproductive tactics of the mycobionts, where it is postulated that the distribution of photobiont 84 haplotypes is dependent on either the asexual or sexual reproduction of the mycobiont.

85

86 Coal Nunatak is located on south-eastern Alexander Island off the west coast of the Antarctic 87 Peninsula. It is protected from the direct influence of the open sea, over 200 km due west in 88 summer, by the high landmass of Alexander Island and the ice shelves that fringe its west coast, 89 and by the permanent ice shelf that occupies George VI Sound to the east (6 km from the study 90 site) and south (20 km from the study site). Located at the extreme southern limit of the 91 maritime Antarctic, this region's climate is considered to be intermediate between that of the

92 more moist maritime region and the colder and drier continental zone (Smith, 1988; Convey & 93 Smith, 1997). Coal Nunatak is snow-free during the Antarctic summer for approximately three 94 months. The ecosystem at this site is characterised by its low developmental level (Brinkmann 95 *et al.*, 2007; Engelen *et al.*, 2008). Small and often barely visible populations of different lichen 96 species, occasionally associated with the few recorded bryophyte species, can be found in 97 microniches restricted to rock surfaces and crevices and to the margins of soil polygons.

98

99 Reproductive tactics:

100 Lichens disperse over long distances by utilising two fundamentally different mechanisms. 101 There may be joint asexual dispersal of both symbionts in specific structures, either by means 102 of a vegetative thallus fragment or by specialized dispersal organs as soredia, which are small 103 (100 - 150 µm in diameter) dispersal units that are produced in specialized cup-like structures 104 called soralia composed of both myco- and photobiont cells. These diaspores can easily be 105 distributed over long distances by wind at high altidtudes. The sexual mechanism of lichen 106 dispersal involves the independent dispersal of the mycobiont (as ascospores) and the 107 photobiont (as vegetative cells). Both can grow individually in a new habitat, before coming 108 into contact through a recognition process and forming a new lichen thallus at that location *de* 109 *novo* by lichenisation (Ott, 1987). Dispersal of the bionts separately clearly requires the process 110 of relichenisation. Environmental conditions and physiological factors influence the success of 111 the recognition process (Meeßen & Ott, 2013; Meeßen *et al.*, 2013).

112

113 Selectivity and specificity:

114 The species diversity of lichen-forming fungi is much greater than that of the photobionts,

115 especially if only green-algal partners, that constitute the photobionts in the majority of lichens,

116 are considered. Algal lineages are widely shared among taxonomic mycobiont groups.

118 Previous studies have demonstrated that mycobionts and photobionts cannot simply be 119 combined randomly (Ahmadjian & Jacobs, 1981, 1982, 1983), indicating a degree of selectivity 120 between the two bionts. Successful and complete lichenisation can only take place when both 121 symbionts possess the appropriate adaptations (Schaper & Ott, 2003). The degree of specificity 122 and selectivity of the mycobiont partner for particular photobionts varies between species. 123 Rambold *et al.* (1998) defined specificity as the taxonomic range of photobionts associated 124 with a mycobiont and selectivity as the exclusiveness with which specific photobionts are 125 selected as partners. Galun & Bubrick (1984) defined 'selectivity' as the preferred interaction 126 between two bionts, and 'specificity' as the exclusive interaction between photo- and 127 mycobiont. Some fungi are only able to lichenise if a specific algal species is available (Galun, 128 1988) while, in contrast, other species of fungi are able to form a lichen thallus with several 129 members of the same genus of photobiont, and sometimes with partners related at an even 130 higher systematic level (Piercey-Normore & DePriest, 2001; Helms *et al*., 2001; Beck *et al.*, 131 2002; Brinkmann, 2002; Romeike *et al*., 2002). Symbiont selectivity and specificity are not 132 only species-specific, but also can vary during the life-cycle of the partners and due to partner 133 availability and environmental conditions. Graduated selectivity is expressed in the form of 134 symbiotic contact achieved between myco- and photobiont. All stages, ranging from the 135 intimate mutualistic contact of both symbionts in a well-developed lichen thallus to a loose-136 fitting parasitic contact, where the fungus penetrates the algal cells using haustoria and 137 subsequently even kills the algae, are possible (Schaper & Ott, 2003). 138 139 High levels of selectivity shown by a mycobiont are linked with a low diversity of suitable

140 photobionts being present in a lichen genus as, for example, found in the family Cladoniaceae

141 (Piercey-Normore & DePriest, 2001), the genus *Physcia* (Helms *et al*., 2001) and the genus

164 Coal Nunatak belongs to the Le May Group and is composed mainly of greywacke, a coarse

165 grained sedimentary rock type (Burn, 1983). Surface geomorphology is characterised by

166 extensive development of patterned ground and other typical periglacial features (e.g. frost-

167 sorted soil polygons, stone stripes) and bare rocks (Brinkmann *et al.*, 2007; Engelen *et al.*, 168 2008).

169

170 Coal Nunatak experiences a continental rather than a maritime climate. From March until mid-171 December the study site is covered by snow, becoming mostly snow-free during the short 172 summer period from mid-December to early March. Terrestrial ecosystems at this site are at a 173 very low or early stage of development. Much of the ground is barren to the naked eye, with 174 colonisation by macroscopic vegetation restricted to small and generally sheltered micro-niches 175 on rocks, in crevices, and on soil sheltered by rocks or associated with longer-lying snow 176 patches. Investigations of the vegetation on the nunatak revealed a total of 14 lichen species 177 and a small number of mosses are known from the nunatak (Brinkmann *et al.*, 2007; Engelen *et* 178 *al.*, 2008).

179

180 Lichen material:

181 Most lichen species were collected from the north-eastern part of the study area on the north-182 east of Coal Nunatak. *Xanthoria elegans* was obtained from the west exposed part of the study 183 area. Three independent samples were obtained for each lichen species, with the quantities 184 sampled being limited by the requirement not to damage the lichen community. After short-185 term storage at ambient conditions at the field site, lichen samples were transported to the 186 British Antarctic Survey's Rothera Research Station (Adelaide Island). There the samples were 187 stored at -20°C and returned frozen to the laboratory in Düsseldorf. Determination of the lichen 188 species was carried out by taxonomic experts (H. Hertel, Munich; D. Øvstedal, Bergen; N. 189 Wirtz, Frankfurt).

190

191 Mycobionts included in the study:

192 The lichen species examined in this study are listed in Table 1. Seven species of the 14

193 obtained from Coal Nunatak were epilithic (crustose lichens: *Tephromela disciformis*,

194 *Tephromela atra*, *Caloplaca johnstonii*, *Lecidella pataviana*; macro lichens: *Usnea lambii,*

195 *Pseudephebe minuscula*, *Xanthoria elegans*) and seven colonised soil-surface habitats (crustose

196 lichens: *Buellia papillata*, *Candelariella flava, Caloplaca lewis-smithii, Lepraria cacuminum,*

197 *Lepraria borealis, Ochrolechia frigida* and *Psoroma cf. tenue*)*. Psoroma cf. tenue* was the only

198 lichen colonising soil sites that has a well differentiated thallus.

199

200 Laboratory procedures:

201 Identification of the unicellular green algal photobionts using morphological and anatomical

202 characters is known to be challenging. Therefore, we used a molecular approach to assess

203 photobiont diversity. The nuclear internal transcribed spacer (ITS) region of the rDNA was

204 analysed, including ITS1, ITS2 and the gene coding for the 5.8S ribosomal subunit. The region

205 is located between the genes coding for the 18S and 26S ribosomal units in the ribosomal DNA

206 tandem repeats and has been used routinely in molecular studies of green algal photobionts

207 (Friedl & Rokitta, 1997; Rambold *et al*., 1998; Beck, 1999; Helms *et al.*, 2001; Kroken &

208 Taylor, 2000; Piercey-Normore & DePriest, 2001; Romeike *et al*., 2002; Schaper & Ott, 2003;

209 Yahr *et al*., 2004; Yahr *et al*., 2006).

210 To obtain photobiont DNA, conglomerates of photobiont cells were first carefully removed

211 from the lichen thalli. This avoided the disruption of molecular procedures by secondary lichen

212 metabolites such as phenolic substances. The clusters of photobiont cells were fragmented

213 using liquid nitrogen and quartz sand. For DNA extraction the DNeasy Plant Mini Kit (Qiagen,

214 Hilden, Germany) was used. After extraction the isolated DNA was stored at -20°C.

216 For a 25 µl PCR reaction, 2.5 µl template, 9 µl sterilized water, 12.5 µl HotStartTaq TM Master 217 Mix (Qiagen) and 0.5 µl of each primer were used. The green alga specific primer with 5^{\sim} 218 3´orientation is Al 1700f (Helms *et al*., 2001). The primer used with 3´-5´orientation (LR3, 219 http://www.biology.duke.edu/fungi/mycolab/primers.htm) is not specific for green algae (Freidl 220 & Rokitta, 1997). For the amplification of the photobiont ITS-region a thermocycler (Biometra, 221 Goettingen, Germany) was used as follows. The taq-polymerase was activated for one minute 222 95 $^{\circ}$ C. The DNA was denatured for one minute at 94 $^{\circ}$ C. The annealing temperature of the 223 primers was set to 53^oC for one minute. The elongation of the annealed primers by taq-224 polymerase took place for 1.5 minutes at 72°C. The denaturation, annealing and elongation 225 steps were repeated 35 times, after which the final extension of partially elongated products 226 took 10 minutes at a temperature of 72°C. After final extension the PCR product was cooled at 227 4°C. The amplified PCR products were purified using the OIAquick PCR Purification Kit 228 (Qiagen, Hilden, Germany).

229

230 DNA sequencing was carried out by GATC-Biotech (Konstanz, Germany) using an ABI 3730

231 XL Sequencer. Non algal specific primers used for sequencing were 1800f (5´-3´orientation)

232 (Friedl, 1996) and ITS4 (3´-5´orientation) (White *et al*., 1990). The resulting ITS rDNA

233 sequences were edited using the application 'Bioedit for Windows'

234 (http://www.mbio.ncsu.edu/bioedit/bioedit.html). NCBI-BLAST searches of GenBank records

235 were performed to confirm that the amplified and sequenced DNA fragments originated from

236 the photobiont and to identify the taxonomic classification of the closest hit.

237 The alignment of all sequences was carried out using the online application MAFFT version 7

238 (http://mafft.cbrc.jp/alignment/server/) based on the HKY substitution model (Hasegawa *et al.*,

239 1985). The calculation of a phylogenetic maximum likelihood tree using PhyML 3.0 (Guindon

240 *et al.*, 2010) was supported by 1000 bootstrap steps. All sequences obtained from samples of

241 lichen photobionts from Coal Nunatak have been added to the database of the National Center

242 for Biotechnology Information (acession numbers: FJ426284 - FJ426299).

243

244 Results

245 In 14 lichen species of 11 genera from Coal Nunatak we found seven different haplotypes of 246 the genus *Trebouxia* and one haplotype of the genus *Asterochloris*. Sexual reproduction was 247 noted only in crustose species, with six lichens producing fruiting bodies (Table 1). The 248 reproduction of these lichen species using ascospores in Antarctic habitats has previously been 249 noted by Øvstedal & Smith (2001). In most of the lichens included in this study the mycobiont 250 was associated with green algal photobionts, with the exception of *Psoroma cf. tenue*. This 251 lichen species also forms thallus structures (cephalodia) with cyanobacteria of the genus 252 *Nostoc*. The mycobiont of this lichen is, therefore, associated with both a green algal species 253 and a cyanobacterial species.

254

255 In the lichen species examined, haplotype 8 was found as photobiont of three lichen species: 256 *Lepraria borealis*, *Usnea lambii* and *Pseudephebe minuscula. Trebouxia* haplotype 7 was the 257 dominant haplotype in the samples investigated at this locality. Six species (*Lecidella* 258 *pataviana*, *Lepraria borealis*, *Lepraria cacuminum*, *Tephromela atra*, *Tephromela disciformis* 259 and *Xanthoria elegans*) were associated with this photobiont. Several haplotypes were 260 restricted to a single lichen species, including haplotypes 2 (*Buellia papillata*), 3 (*Candelariella* 261 *flava*), 4 (*Psoroma cf. tenue*), 5 (*Caloplaca lewis-smithii*) and 6 (*Caloplaca johnstonii*). The 262 algal genus *Asterochloris* (haplotype 1) was found as photobiont in the two lichen species 263 *Lepraria borealis* and *Ochrolechia frigida* (Table 1). 264

275 in a BLAST search were to the genus *Asterochloris*. The sequences of the second clade 276 belonging to haplotype 8 showed highest BLAST hits with taxa identified as *Trebouxia* 277 *jamesii*. The third clade consisted of two subclades. One included haplotypes 6 and 7, and had 278 highest BLAST similarities with taxa also identified as *T. jamesii*, and the other consisted of 279 haplotypes 2 to 5, which were most similar to *Trebouxia impressa*. Most substitutions were 280 found in the subclade consisting of haplotypes 2 to 5 (*T. impressa)*. The other clades, involving 281 the two different groups of *Trebouxia jamesii* and one group of *Asterochloris* sp*.*, were 282 composed of almost identical sequences within each clade.

283

284 With the exception of *Lepraria borealis*, the photobiont clades were correlated to groups of 285 lichens characterised by sharing particular morphological and ecological features. The clade 286 consisting of the photobiont haplotype 8 (*T. jamesii*) was associated with fruticose lichens 287 growing on rocks, whereas photobionts of the clade consisting of haplotypes 6 and 7 (*T.* 288 *jamesii*) were found in crustose lichens on rocks. Crustose lichens growing on soil and/or 289 mosses were either associated with photobionts of the clade consisting of haplotypes 2-5 (*T.*

290 *impressa*) or with the photobiont clade of haplotype 1 (*Asterochloris*). Photobionts of *L.*

291 *borealis* were present in all clades with the exception of the *T. impressa* group.

292

293 Discussion

294 This study is the first to document lichen photobiont diversity in the southern region of the

295 Antarctic Peninsula. Molecular studies on photobionts have shown that identical haplotypes are

296 widespread and can be found across geographic regions (Kroken & Taylor, 2000; Yahr *et al.*,

297 2004; Yahr *et al.*, 2006), continents (Piercey-Normore & DePriest, 2001) and even

298 hemispheres. *Trebouxia jamesii* identified here at Coal Nunatak, has been described from a

299 range of other localities in the maritime and continental Antarctic (Romeike *et al.*, 2002; Pérez-

300 Ortega *et al.*, 2012) as well as from Europe (Beck, 1999), supporting effective dispersal across

301 distances up to intercontinental and global scales.

302

303 Patterns of haplotype distribution being unique to specific habitats suggest a process of local 304 adaptation or might be pre-adapted to local environmental conditions and communities (Pérez-305 Ortega *et al.*, 2012). Such an interpretation is in line with the conclusions drawn by Yahr *et al.* 306 (2004) in an extensive community study, who found a homogeneous photobiont pool across 307 geographic distances and proposed that local adaptation of the photobiont was of importance at 308 some sites.

309

310 All localities analyzed by Yahr *et al.* (2004) shared the same habitat and vegetation type, 311 reflected by the occurrence of similar *Cladonia sp.* communities. Such a sampling design 312 enhances the detection of environmental selection acting in photobiont lineages that might 313 cause ecological specialization.

315 The ITS sequence of photobiont haplotype 8 found on Coal Nunatak in this study was also 316 found in lichens from the maritime Antarctic sites Lagoon Island, Rothera Point and Charcot 317 Island by Romeike *et al*. (2002). It is also identical to the sequence of *T. jamesii* cultured from 318 *Lecidea silacea*, collected from siliceous and heavy-metal containing rocks at localities in 319 Austria (Beck, 1999). It seems that haplotype 8 (*T. jamesii)* is a generalist with a bi-320 hemispherical distribution that occurs ranging from extreme habitats at the limits of vegetation 321 to more moderate maritime and polar habitats to comparably benign temperate localities. 322

323 Romeike *et al*. (2002) noted that this haplotype has only been described to date from iron-rich 324 sites (Beck, 1999). However, the study on Coal Nunatak does not have atypically high iron 325 concentration. Haplotype 8 was the second most abundant photobiont at Coal Nunatak.

326

327 At Coal Nunatak photobionts are associated with a relatively low number of mycobionts. 328 Haplotype 7 was the most abundant photobiont amongst 14 lichen species recorded on Coal 329 Nunatak, being present in 6 different species while haplotypes 1-6 and 8 were distributed 330 amongst 9 lichen species (Table 1). This is suggestive of selectivity and specificity of the 331 symbionts of the respective lichen species. Eight different photobiont haplotypes were found in 332 the 14 lichen species (Table 1). With one exception *Lepraria borealis* (Engelen *et al.*, 2010) all 333 mycobionts were associated with a single photobiont haplotype.

334 This might be due to the harsh environmental conditions and the overall short growing season 335 at the inland site that limits photosynthetic activity and thus photobiont productivity. In such a 336 life-averse habitat which effects a limited primary production (Sadowsky & Ott, 2012) 337 symbiont interactions can be expected to be fine-tuned for the holobiont to survive and to 338 successfully colonise and keep the habitat.

340 To form an own thallus *L. borealis* takes over the photobiont haplotypes 1, 7 and 8. Our data 341 suggest that the species may be able to obtain these photobionts from physically adjacent thalli 342 of other lichen species, such as *Ochrolechia frigida* (Pertusiales), *Tephromela disciformis* and 343 *Usnea lambii* (Lecanorales s. str.) (Table 1). When growing close to *T. disciformis* or *U. lambii*, 344 *L. borealis* incorporated the identical *Trebouxia jamesii* haplotypes 7 and 8 as the photobionts 345 of these immediately adjacent lichens. Similarly, when growing in association with 346 *Ochrolechia frigida* the same *Asterochloris* haplotype 1 was present in both lichens (Engelen *et* 347 *al.*, 2010). Thus, only the *L. borealis* mycobiont shows low selectivity towards potential 348 photobionts consistent with the prediction of Romeike *et al*. (2002). 349 350 The diversity of the photobiont haplotypes within the lichen community may be influenced by 351 the mechanism of reproduction. Sexual reproduction requires a lichenisation process during 352 which the mycobiont must encounter a suitable algal partner amongst those available in its

353 immediate vicinity. The data obtained in the current study give no indication of any difference 354 in diversity of photobiont haplotypes between the mycobionts reproducing asexually or

355 sexually (Table. 1).

356

357 For the degree of selectivity of the mycobiont to the photobiont partner characteristic features 358 of the respective lichen symbiosis may be primarily responsible at extreme habitats. 359 Environmental conditions might also effect the degree of selectivity (Romeike *et al.* 2002). 360 However, based on the results presented the potential of the symbiotic state of lichens seems to 361 be the more relevant factor considering the success of colonisation processes particularly at 362 extreme environments.

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- 527 Tab. 1: Reproductive mode and distribution of photobiont haplotypes in lichen species found
- 528 on Coal Nunatak.
- 529 * haplotype 1: genus *Asterochloris*; haplotypes 2-8: genus *Trebouxia*
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- 533

545 Figure captions

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547 Fig. 1: a: arrow=location of Coal Nunatak on Alexander Island (70°03'S 68°31'W). b:

548 circle=location of the research area on Coal Nunatak.

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- 550 Fig. 2: ML-tree of the *Asterochloris* haplotype and the 7 *Trebouxia* haplotypes as shown in 551 Tab. 1.
- 552 The photobiont sequences are named as follows: abbreviation of the photobiont_abbreviation
- 553 of the mycobiont_number of the haplotype as in Tab.1

- 555 Abbreviations of the mycobiont: Bupa: *Buellia papillata*, Cafl: *Candelariella flava*, Cajo:
- 556 *Caloplaca johnstonii*, Cale: *Caloplaca lewis-smithii*, Lebo: *Lepraria borealis*, Leca: *Lepraria*
- 557 *cacuminum*, Lepa: *Lecidella pataviana*, Ocfr: *Ochrolechia frigida*, Psmi: *Pseudephebe*
- 558 *minuscula*, Pste: *Psoroma cf. tenue*, Teat: *Tephromela atra*, Tedi: *Tephromela disciformis*,
- 559 Usla: *Usnea lambii*, Xael: *Xanthoria elegans*
- 560 Abbreviations of the photobionts: Ast: *Asterochloris spec.*, Tja: *Trebouxia jamesii*, Tim:
- 561 *Trebouxia impressa*

