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1 ***Schizoxylon* as an experimental model for studying interkingdom symbiosis**

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13

14 **Abstract**

15 Experiments to resynthesize lichens so far focused on co-cultures of fungal and algal partners.

16 However, recent studies have revealed that bacterial communities colonize lichens in a stable and

17 host-specific manner. We were therefore interested in testing how lichenized fungi and algae

18 interact with selected bacteria in an experimental setup. We selected the symbiotic system of

19 *Schizoxylon albescens* and the algal genera *Coccomyxa* and *Trebouxia* as a suitable model. We

20 isolated bacterial strains from the naturally occurring bacterial fraction of freshly collected

21 specimens and established tripartite associations under mixed culture experiments. The bacteria

22 belong to Actinobacteria, Firmicutes and Proteobacteria and corresponded to groups already found

23 associated with fungi including lichens. In mixed cultures with *Coccomyxa* the fungus formed a

24 characteristic filamentous matrix and tightly contacted the algae; the bacteria distributed in small

25 patches between the algal cells and attached to the cell walls. In mixed cultures with *Trebouxia*, the

26 fungus did not develop the filamentous matrix, but bacterial cells were observed to be tightly

27 adhering to the fungal hyphae. Our experiments show that this tripartite fungal-algal-bacterial

28 model system can be maintained in culture and can offer multiple opportunities for functional

29 studies based on experiments under controlled conditions in the laboratory.

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32 **Keywords:** algae, bacteria, culture, fungi, scanning electron microscopy, system.

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36 **Introduction**

37

38 Most textbooks describe lichens as a partnership of two organisms, a fungus and an alga. This two-
39 partner concept of the lichen symbiosis was introduced by Schwendener (1869) but was challenged
40 very early, not least by contemporary researchers who could not at first accept the idea that lichens
41 were not a separate group of organisms. Cengia-Sambo (1924) suggested the term “polysymbiosis”
42 for cases where cyanobacteria are present in addition to the green algal symbionts, now known as
43 tripartite lichen symbioses (Kaasalainen *et al.*, 2009; Magain and Sérusiaux, 2014). Recently,
44 culture-independent analyses and microscopic approaches revealed that in lichens bacterial
45 communities occur in a stable and host-specific manner (e.g., Cardinale *et al.*, 2008; Grube *et al.*,
46 2009; Hodkinson & Lutzoni, 2009; Hodkinson *et al.*, 2011; Bates *et al.*, 2011; Mushegian *et al.*,
47 2011; Wedin *et al.*, 2016). While algal and cyanobacterial partners have clear functional
48 assignments concerning their provision of fixed carbon or nitrogen to the symbiosis, the role of
49 other bacteria is still unknown. Grube *et al.* (2015) used metagenomic and metaproteomic
50 approaches to highlight potential functions of the bacterial fraction in lichen symbioses. To what
51 extent bacterial functions actually influence the symbiotic community of lichens remains a
52 challenge. Such functional studies require controlled experimental conditions, which can potentially
53 be achieved if the interacting partners can be co-cultured.

54 Lichen symbionts have been cultured since the 19th century, but due to their slow growth
55 they never became popular as experimental systems. The early attempts to resynthesize lichens by
56 co-culturing their partners had primarily the intent to test the dual hypothesis of the lichen
57 symbiosis (e.g., Bonnier, 1886), which was still very controversial at that time. Re-synthesis
58 experiments of lichens were accomplished repeatedly with success, especially in the second half of
59 the 20th century (e.g., Ahmadjian, 1973; Honegger, 1990; Yoshimura *et al.*, 1993; Stocker-
60 Wörgötter, 2001 and references therein), and these experiments also demonstrated the specificity of
61 the algal-fungal relationship to form the characteristic thallus morphology (e.g., Ahmadjian *et al.*,
62 1980; Etges *et al.*, 2003). Bacterial species were never included in any of these experiments, and if
63 they emerged, they were generally regarded as contaminations. In nature, however, lichen fungi are
64 in contact with bacteria almost continuously. So far, the only reported co-culture of bacteria with
65 lichen mycobionts was provided by Seneviratne and Indrasena (2006). Their experiments showed
66 increased solubilisation of mineral substrate in co-cultures of an isolated lichen fungus with N-
67 fixing *Bradyrhizobium elkanii*. Their images also showed tight attachment of the bacteria onto the
68 fungal hyphae in culture. Alternatively, Lőrincz *et al.* (2010) studied an artificial symbiosis by co-

69 culturing the green algae *Chlamydomonas*, the bacterium *Azotobacter* and the fungus *Alternaria*
70 and demonstrating that the basis of the interdependence is the complementation of photosynthetic
71 CO₂ assimilation and atmospheric nitrogen fixation. We are not aware, though, of any other study
72 which attempted cultivation of lichen fungi, algae and bacteria together. We were therefore
73 interested in testing whether lichenized fungi and algae interact with selected bacteria under culture
74 conditions, to assess the influence of the latter on the development of the resynthesized thallus.

75 Here, we present our results from experiments done to investigate if a tripartite system
76 (fungus-algae-bacteria) indeed was possible to establish for experimental symbiosis purposes. We
77 used the lichenized fungus *Schizoxylon albescens*, the associated algae *Coccomyxa*, other lichenized
78 algae of the genus *Trebouxia* and bacterial strains isolated in cultures from the naturally occurring
79 bacterial fraction of freshly collected *Schizoxylon* specimens. *Schizoxylon* belongs to the
80 Stictidaceae, a group of ascomycetes where lichenized species have evolved within clades of
81 closely related, saprotrophic and parasitic species (Baloch *et al.*, 2009, 2010, 2013) and where
82 several species either can live as lichens or as saprotrophs (“optional lichenization”, Wedin *et al.*,
83 2004). In contrast to other lichens, which require algal partners to develop symbiotic thalli as a
84 prerequisite for fungal sexual reproduction, *Schizoxylon* also grows as saprotroph on dead wood
85 (Wedin *et al.*, 2006). Phylogenetic and haplotype analyses have shown that these two morphs of
86 *Schizoxylon* neither are genetically differentiated nor correlate with geographical origin in any of
87 the markers analysed (Muggia *et al.*, 2011). In nature *Schizoxylon albescens* forms simple
88 lichenized stages with a consortium of algae of the genus *Coccomyxa*. Previous co-culture
89 experiments also indicated that the fungus shows certain selectivity for some of the isolated algal
90 strains and form lichenization structures in mixed cultures (Muggia *et al.*, 2011). Ongoing analyses
91 on environmental samples revealed a diversity of bacteria in both the lichenized and saprotrophic
92 morphs of *Schizoxylon*.

93 The aims of our study were: i) to establish and maintain “three-partner associations” in
94 culture by growing the *Schizoxylon* fungus together with different algal and bacterial strains, and if
95 successful, using such three-partner mixed cultures for small pilot studies; ii) to study the
96 morphological organization of these associations; iii) to study whether the presence of bacteria
97 induce *Schizoxylon* to develop further symbiotic stages with *Coccomyxa*; iv) to study whether the
98 presence of bacteria triggers *Schizoxylon* to interact also with *Trebouxia*, a common lichen
99 photobiont.

100

101

102 **Material and Methods**

103

104 *Sampling* – A total of 20 fresh specimens of the lichenized (15, L1-L10, MW9634-MW9638) and
105 saprotrophic (5, S1-S5) forms of *Schizoxylon albescens* were collected in Sweden (Hälsingland
106 province, Färila parish, South of lake Skålvallssjön, 61°56'24.6" N/ 15°36'32.6" E, 225 m a.s.l., in
107 deciduous-tree dominated succession forest on former agricultural land, 15/09/2013, leg. S.
108 Fernández-Brime and M. Wedin) and were used for bacterial isolation.

109 The fungus and algae used in the experiments are all stored as fresh cultures or cryostocks in
110 the culture collection of the first author at the University of Graz (numbered as "LMCCxxxx").
111 *Schizoxylon* was isolated from the lichenized specimen MW7645 and has been maintained in living
112 cultures since 2008 (Muggia *et al.*, 2011; cryostock n. LMCC0016). The two *Coccomyxa* algal
113 strains PL2-1 and MW8233 were also isolated in 2008; they were available from the cryostocks
114 (LMCC0023 and LMCC0038) and were freshly plated two months before starting the experiments.
115 Two additional strains of *Trebouxia* photobionts, isolated from the lichen *Tephromela atra* and kept
116 in living cultures, *Trebouxia* 'sp. 1' (LMCC0107) and *Trebouxia* 'TR1' (LMCC0121) - according
117 to the nomenclature followed by Muggia *et al.* (2014) - were used for the mixed culture
118 experiments.

119

120 *Culture isolations and mixed culture experiments* – The fungal and the algal strains were isolated in
121 2008 as described in Muggia *et al.* (2011). Of the 20 freshly collected samples of *Schizoxylon*
122 *albescens*, ten lichenized (L1-L10) and the five saprotrophic (S1-S5) were selected for the isolation
123 of bacteria. The samples were analysed under the stereo microscope and for five lichenized and the
124 five saprotroph two types of pieces were distinguished and dissected for culture isolations: (A) the
125 pieces containing *Schizoxylon* ascomata together with the algae clumps and a portion of the
126 surrounding substrate in the lichenized one, or containing only the ascomata in the saprotrophic
127 ones, (B) the pieces of the bark (for the lichenized samples) and of the wood (for the saprotrophic
128 samples), without ascomata and algae. In doing this, we aimed at maximizing the isolation of the
129 whole spectrum of culturable bacteria which are associated with the fungus, the fungus and the
130 algae, and the growth substrates (Fig. 1a). The selected pieces were put in 1.5 ml tubes and washed
131 with a 0.9% of NaCl solution by vortexing the tube for 2 minutes. The solution was plated both
132 undiluted and 1/2 diluted on R2a agar (Reasoner *et al.*, 1979) plates, as this medium is the minimal
133 one to isolate universal bacteria (Zachow *et al.*, 2013). The plates were incubated at room
134 temperature for 2-4 days for allowing bacteria to grow. Once bacterial colonies grew, they were
135 picked and individually sub-cultured to obtain single-colony isolates. We picked for each plate, a

136 maximum of 12 different bacterial colonies, distinguishing them according to their different colour
137 (Fig. 1b, c), in order to include a broad diversity of the strains.

138 The remaining five specimens of lichenized *Schizoxylon* (MW9634-MW9638) were used to
139 isolate bacteria directly from the algal clumps surrounding the ascomata of the fungus (Fig. 1a).
140 Algal clumps were picked with a sterile needle and inoculated on BBM medium. All the different
141 bacteria that grew out of the inocula were further individually picked and sub-cultured to obtain
142 single-colony isolates.

143 All the successfully grown single-colony isolates were prepared for cryostock storage and
144 molecular analyses for their identification. Cryostock preparation consisted in inoculating a single-
145 cell colony in 500 µl liquid R2a medium, adding the same volume of 40% glycerol and storing
146 them in tubes at -80 °C.

147 Mixed cultures of multiple combinations of *Schizoxylon*, algae and bacteria were set on five
148 different growth media: malt yeast (MY; Ahmadjian, 1967), *Trebouxia* medium (TM; Ahmadjian,
149 1967), Sabouraud-Agar (SAB; Sigma), Lilly and Barnett's Medium (LBM; Lilly and Barnett, 1951)
150 and water-agar medium (a medium without nutrients solidified only with agar). The fungus was
151 ground in water in a sterile mortar with a pestle and fragments were pipetted on the medium. Algae
152 and bacteria were suspended in 1 ml water and for each mixed culture 200 µl were pipetted on the
153 medium on the same spot of the fungus. The mixed cultures were set as follow: the same fungus
154 was combined with each of the four algae strains (two *Coccomyxa*, PL2-1 and MW8233, and two
155 *Trebouxia*, 'sp.1' and 'TR1'). Each fungal-algal combination (four in total) was subsequently
156 combined with a total of 14 different bacterial strains. These were randomly taken from those both
157 isolated from the fragments of the lichenized (four strains: LIC-1B/1, LIC-4A/10, LIC-4A/11, LIC-
158 4B/12) and the saprotrophic (four strains: SAP-4A/2, SAP-5B/2, SAP-5B/1r, SAP-5B/1w)
159 specimens, either from the type A or type B pieces, and those isolated directly from the algal
160 clumps (six strains: MW9636.1, MW9636.2, MW9636.5-.8). A total of 280 plates were therefore
161 prepared. No replicate on the same medium was performed for any of the fungus-algae-bacteria
162 combination.

163
164 *Scanning electron microscopy (SEM) analyses* – The mixed cultures that successfully developed the
165 intertwined growth of the three organisms were selected for electron microscopy analyses. A piece
166 of about 1 cm² was taken from each culture and fixed with a glutaraldehyde (2%) cacodylate 0.2 M
167 pH 7.4 buffer for 2 hours at room temperature. The fragments were washed three times with the
168 cacodylate 0.1 M pH 7.4 buffer and dehydrated with an ethanol series consisting of three fast
169 washes with EtOH 30%, three fast washes with EtOH 50%, two washes of 5 min with EtOH 70%,

170 two washes of 12 min with EtOH 90% and two final washes of 15 min with EtOH 100%. The
171 samples were then dried at the CO₂ critical point and gold sputtered. For each samples both vertical
172 sections and upper surfaces were arranged on the stubs.

173

174 *Molecular analyses of the bacterial strains* – Bacterial colonies were first transferred to
175 microcentrifuge tubes containing 200 µl of ddH₂O and then samples were spun for 10 minutes at
176 7500 rpm in order to precipitate the bacterial cells. Once the supernatant was discarded, the DNA
177 was extracted using the KingFisher™ Cell and Tissue DNA Kit (Thermo Fisher Scientific) or the
178 DNeasy Plant Mini Kit (Qiagen) following the manufacturers' instructions. The primers F515 and
179 R806 were used to amplify part of the III and the IV 16S variable regions (Chakravorty *et al.*, 2007;
180 Caporaso *et al.*, 2011). We sequenced this fragment in order to incorporate the data obtained from
181 these strains into a broader study using MiSeq Illumina data targeting the same fragment
182 (Fernández-Brime *et al.*, in prep). The PCR was performed using Illustra™ Ready-To-Go PCR
183 Beads (GE Healthcare), according to the manufacturer's instructions and the following PCR cycle
184 was applied: 5 min at 95 °C followed by 37 cycles, each of 45 s at 95 °C, 45 s at 50 °C, 90 s at 72 °C
185 and a final extension of 10 min at 72 °C. PCR products were then purified with the enzymatic
186 method Exo-sap-IT© (USB Corporation) and the sequencing of both strands was performed with
187 the Big Dye Terminator technology kit v3.1 (ABI PRISM, U.S.A.) using the same PCR primers.
188 Sequences fragments were edited and assembled using Sequencer v.4.9 (Gene Codes Corp., Ann
189 Arbor, MI). Primers were truncated at this point. The dataset was processed in QIIME v.1.9.1
190 (Caporaso *et al.*, 2010) with the script for closed-reference OTU picking in order to pick operational
191 taxonomic units with a 0.97 similarity threshold, using as reference the Greengenes core set.
192 Taxonomy was assigned with the uclust classifier accepting a maximum of three hits and a
193 minimum similarity of 0.90.

194

195

196 **Results**

197

198 *Culture isolations and mixed culture experiments* – A total of 121 bacterial colonies isolated from
199 the five lichenized and the five saprotrophic samples and a total of 18 bacterial colonies isolated
200 from the algal clumps, were successfully sub-cultured and prepared for cryostock storage and
201 molecular sequence identification. Of the 139 colonies, 14 (eight out of the 121 and six out of the
202 18) were arbitrarily taken for the mixed culture experiments and were identified by sequencing (see
203 below). After two months, two-thirds of the mixed cultures were discarded due to overgrowth of the

204 bacteria or failed growth. Mixed cultures failed to grow on 95% of the water-agar plates; on the
205 remaining water-agar plates, algae and bacteria grew inconspicuously and very slow, whereas the
206 fungus did not grow at all. These cultures were excluded from further analyses. About 20% of the
207 mixed culture experiments developed successfully and eight of them were selected for SEM
208 analyses after 6 months of co-growth (Table 1).

209 In mixed culture with *Coccomyxa*, the fungus developed within a homogeneous mass
210 formed by the intermixed growth of algal and bacterial cells (Fig. 1d, g). In mixed culture with
211 *Trebouxia*, algae and bacteria formed well delimited clumps of tightly connected cells. The fungal
212 hyphae either grew among the algal and the bacterial colonies (Fig. 1f) or they built a more compact
213 mycelial mass with filamentous hyphae in contact with bacterial and algal cells (Fig. 1e). In squash
214 preparations mounted in water of *Schizoxylon* grown with *Trebouxia* sp.1 and the bacterial strain
215 LIC-1B/1 on MY medium, it was possible to observe bacteria attached to the fungal hyphae (Fig.
216 1h, i).

217 In axenic culture, a developing mycelium can first be observed c. 2 months after plating the
218 *Schizoxylon* inoculum, growing at a rate of 0.3 mm/month on MY medium. When the fungus was
219 co-cultured with algae and bacteria, mycelium growth was initiated considerably earlier, c. 1 month
220 after inoculation, but grew at a similar rate. No difference in growth morphology, growth rate, or
221 fitness of the mixed cultures was observed amongst the bacterial strains.

222

223 *SEM analyses* – The mixed cultures which succeeded in growth and were analysed by SEM, were
224 set with bacteria (Actinobacteria, Firmicutes and Gammaproteobacteria, as it follows) isolated from
225 the lichenized samples, both from the pieces of type (A) and directly from the algal clumps. In
226 mixed cultures with the two *Coccomyxa* strains, the hyphae were in tight contact with the algae
227 (Fig. 2a, b) and the fungus formed a characteristic filamentous matrix (Fig. 2c) concordant with the
228 results in Muggia *et al.* (2011). This filamentous matrix enclosed the *Coccomyxa* cells and appeared
229 locally well developed covering the algal layer entirely (Fig. 2e). Bacteria were detected in small,
230 irregular patches between the algal cells and attached to the cell walls (Fig. 2d). In some parts of the
231 mixed cultures the fungus overgrew and algal cells were scattered in low amount between the
232 hyphae (Fig. 2f). In mixed cultures with *Trebouxia*, the fungus did not develop any filamentous
233 matrix. The hyphae spread among the algal colonies, entangling single cells or algal clumps. The
234 observed clumped growth of *Trebouxia* is well known and it is due to the clonal growth by
235 autospores inside the parental gelatinous sheet (Fig. 3b, d). In these mixed cultures, bacterial cells
236 were observed to be tightly adhering to the fungal hyphae (Fig. 3a-e), and distributed according to

237 the branching of the hyphae, also where the three organisms grew in close contact. We seldom
238 observed bacteria adhering to *Trebouxia* cells.

239 In the culture set with the saprotrophic strain SAP-5B/2 (which was not identified due to the
240 bad quality of the sequence) we did not observed mixed growth of the three organisms (not shown).

241

242 *DNA analyses of bacterial identity* – From the 14 bacterial strains used for the mixed culture
243 experiments, PCR products were not obtained for the strain MW9636.6 and sequences of two
244 strains (MW9636.5 and SAP-5B/2) were excluded from further analyses due to low quality. The
245 remaining 11 sequenced strains were processed in QIIME and their putative identities are
246 summarized in Table 2. A total of eight OTUs were recognised within the phyla Actinobacteria
247 (Geodermatophilaceae and Microbacteriaceae), Firmicutes, Alphaproteobacteria
248 (Sphingomonadaceae) and Gammaproteobacteria (Xanthomonadaceae). The identity of the strains
249 was assessed according to the hits agreeing with the lowest taxonomic level: one strain, SAP-4A/2,
250 was identified at the species level as *Luteibacter rhizovicinus*. Six strains were classified to the
251 genus level, MW9636.1 as *Acinetobacter*; LIC-1B/1 as *Bacillus*; LIC-4A/11 and MW9636.7 as
252 *Frigoribacterium*; MW9636.2 and MW9636.8 as *Sphingomonas*, and four to the family level, LIC-
253 4B/12, SAP-5B/1r and SAP-5B/1w in Microbacteriaceae, and LIC-4A/10 in Geodermatophilaceae.

254

255

256 **Discussion**

257

258 *A model for interkingdom microbial symbioses* – Lichen symbioses represent suitable subjects for
259 studying interaction among distantly related organisms (Berg, 2015). Axenic cultures of lichen
260 mycobionts have earlier been established to study patterns of secondary metabolite production
261 under different controlled conditions (e.g. Brunauer *et al.*, 2007; Stocker-Wörgötter and Elix, 2009;
262 Fazio *et al.*, 2009, 2014). Yet, only few co-cultivation experiments with photobionts (green algae)
263 and attempts of re-synthesis of lichen thalli have been reported in recent times (e.g., Stocker-
264 Wörgötter, 1995; Joneson and Lutzoni, 2009; Guzow-Kreminska and Stocker-Wörgötter, 2013),
265 probably due to the slow growth of lichen symbionts. Thus, *in vitro* re-synthesis experiments of
266 lichen thalli have focused on the first stages of thallus formation, which are highly dependent on the
267 compatibility of the fungal and the algal symbionts (Ahmadjian *et al.*, 1978; Schaper and Ott, 2003,
268 Joneson and Lutzoni, 2009; Guzow-Kreminska and Stocker-Wörgötter, 2013). Compatible partners
269 are commonly first characterized by molecular analyses and later co-growth experiments are set up
270 (Guzow-Kreminska and Stocker-Wörgötter, 2013; Muggia *et al.*, 2011). In most cases, however,

271 the development of more differentiated thallus structures, such as lobes or fungal fruiting bodies,
272 has hardly ever been reported on solid agar media. For this purpose, the use of selected grow
273 substrates, such as sterilized soil, and controlled growth conditions proved to be triggering (Bubrick
274 and Galun, 1986; Stocker-Wörgötter and Türk, 1991, 1993; Stocker-Wörgötter, 2001). Re-synthesis
275 experiments of lichens starting from their aposymbiotic partners is in general challenging, as the
276 experimental system has to be optimized for each case, usually by simulating some ecological
277 factors as close as possible by culture conditions. Modified media (Guzow-Kreminska and Stocker-
278 Wörgötter, 2013), adjusted light-dark regimes and temperature can be applied to balance the joint
279 growth of the different organisms, which indeed favour different conditions (Muggia, pers. obs.).

280 Co-culture experiments of lichens have hitherto only considered two partners. Because non-
281 photosynthetic bacteria have been increasingly recognized to be regularly present in lichen
282 symbioses, it is tempting to include these in symbiotic resynthesis experiments as well (Grube *et*
283 *al.*, 2009, 2015; Aschenbrenner *et al.*, 2014). Bacteria are localized in lichen thalli by *in situ*
284 hybridization, which also show different spatial arrangements at different ages of the thalli
285 (Cardinale *et al.*, 2008, 2012). Generally, only a minor fraction of the lichen-associated bacteria can
286 successfully be isolated by axenic cultures and further characterized for their biological potential
287 without the hosting lichen (Cernava *et al.*, 2015; Erlacher *et al.*, 2015). Previous studies of lichen-
288 associated bacteria have focused on well-structured lichen thalli of model species (e.g. *Lobaria*
289 *pulmonaria*) which develop three-dimensional structures. However, the mycobionts of these model
290 lichens grow slowly in axenic culture and do not offer suitable systems for long-term and repeatable
291 culture experiments. Further, the use of these species for re-synthesis experiments would first
292 require axenic cultures of all individual symbionts, which is not always straightforward.

293 *Schizoxylon albescens*, which is facultatively lichenized and able to either associate with
294 *Coccomyxa* or to live as a saprotroph, was thoroughly investigated some years ago. We earlier
295 proposed this fungus as a suitable model species for studies of the lichen symbiosis (Muggia *et al.*,
296 2011). The availability of axenic cultures of both the fungus and the associated algae, and the
297 successful isolation of a bacterial fraction associated with the apothecia and the algal clumps enable
298 us to set up pilot experiments with mixed cultures including three partners. *Schizoxylon* grows faster
299 than other lichen mycobionts in culture, and this reduces the risk of being overgrown by faster-
300 growing algae. As the system *Schizoxylon-Coccomyxa* was available, we used it as a reference to
301 test whether the addition of bacteria as third partner group would induce changes during the joint
302 growth. We clearly find faster initiation of growth of fungal inocula during co-cultivation with
303 symbionts (algae and bacteria in this case). This phenomenon indicates that a symbiotic effect in the
304 co-cultures could be expressed by more rapid initiation of growth, rather than by elevated growth

305 rates. However, we still need to find out whether the effect is due to co-cultivation with algae
306 together with bacteria, or due to one of these alone. Here we focused on the morphological
307 development of the mixed cultures in the presence of different bacterial strains and whether the
308 tested bacteria preferentially grow in certain parts of the cultures to give rise to more complex
309 symbiotic architectures. We were also interested to test whether the presence of bacteria would
310 induce or facilitate fungal association with another photobiont than *Coccomyxa*. Nevertheless, we
311 are aware that the isolated bacteria were arbitrarily selected for the mixed culture experiments and
312 represent only a subset of the whole bacterial community associated with *Schizoxylon*. Strains that
313 by chance were not selected so far or that are not possible to isolate and grow in axenic culture may
314 of course also play a significant role during the lichenization phases.

315

316 *Plasticity of the fungal-algal-bacterial association in mixed cultures* – In accordance to previous
317 observations (Muggia *et al.*, 2011), our analyses showed that *Schizoxylon albescens*, when cultured
318 with *Coccomyxa*, forms a characteristic filamentous matrix and tightly adheres to algal cells. We
319 reproduced this phenomenon and the formation of initial fungal-algal symbiotic stages (Ahmadjian
320 *et al.*, 1978; Galun, 1988; Joneson and Lutzoni, 2009). In contrast to the initial experiments by
321 Muggia *et al.* (2011), this time *Schizoxylon* was also able to grow with and contact *Trebouxia* algae.
322 However, no filamentous matrix was observed in the *Schizoxylon-Trebouxia* cultures and further
323 analyses are needed to demonstrate if the fungus forms haustoria with this photobiont, as commonly
324 observed in lichen fungi associated with *Trebouxia* (Honegger, 1984). Bacterial cells are distributed
325 irregularly in the *Schizoxylon-Coccomyxa* mixed cultures, but only in the *Schizoxylon-Trebouxia*
326 cultures are they pronouncedly attached to the fungal hyphae. The successfully grown mixed
327 cultures that were analysed by SEM, were set with bacteria isolated from the lichenized samples,
328 both from the pieces type (A) and directly from the algal clumps. In the culture set with the
329 saprotrophic strain SAP-5B/2 (so far unidentified due to bad quality of the sequence) we did not
330 observe mixed growth of the three organisms (not shown). Further mixed culture experiments
331 should therefore be set with additional bacterial strains isolated from the lichenized pieces (A) to
332 confirm these first results. While we have not observed differences in the *Schizoxylon-Coccomyxa*
333 cultures which could have been induced by the addition of bacteria, future analyses will address
334 whether the tight fungal-bacteria interaction triggers the capacity of *Schizoxylon* to associate with
335 *Trebouxia*. Further, the tri-partite associations were grown on different media, of which only MY,
336 SAB and TM worked properly. These should be utilized in future experiments to test different
337 *Schizoxylon-Coccomyxa/Trebouxia*-bacteria associations and to standardize the mixed cultures.

338 Among the bacteria we isolated from *Schizoxylon*, we have identified genera known to be
339 associated with fungi and which also belong to groups already found in lichen microbiota
340 (Cardinale *et al.*, 2006, 2011; Parrot *et al.*, 2015; Grube *et al.*, 2016; Aschenbrenner *et al.*, 2016).
341 Several strains were identified as Actinobacteria and assigned to the family Microbacteriaceae, a
342 large group that is widespread in various terrestrial and aquatic environments. Within the
343 Gammaproteobacteria, which include N₂-fixers, one strain was assigned to *Acinetobacter* and
344 another was identified as *Luteibacter rhizovicinus*, which has earlier been isolated from *Cladonia*
345 *coniocraea* and *Peltigera membranacea* (Cardinale *et al.*, 2006; Sigurbjörnsdóttir *et al.*, 2015). In
346 the mixed cultures, bacterial colonies grow only in small patches on the filamentous matrix or are
347 attached to the hyphae, sometimes covering the hyphae completely, and rarely to the algal cell
348 walls. Even though organisms may develop different phenotypes when inoculated on different
349 media, we cannot correlate the various detected growth patterns to a certain medium, or to the age
350 of the culture. Medium variations, which may affect the interaction patterns, need to be assessed by
351 extended culture experiments. This may also include liquid cultures, which could represent a
352 feasible approach either for easier propagation of the symbiotic cultures or for maintenance of long-
353 term cultures for time series analyses.

354
355 *Outlooks of multipartite experimental model systems* – Co-culture and mixed culture approaches are
356 increasingly applied to test working hypotheses for transcriptomics and metabolomics analyses, to
357 explore biosynthetic potentials, and to test the latent capacity to form symbiosis (Goers *et al.*, 2014;
358 Hom and Murray, 2014; Hays *et al.*, 2015). Because co-culture approaches require extensive
359 optimization to subsequent analyses, the application of an established model system for further
360 exploration of symbiotic interactions, e.g. by omics approaches, is essential. The main result from
361 our study is that we have now successfully established a *Schizoxylon*-algal-bacterial system which
362 offers further opportunities to study lichenization *in vitro*. As *Schizoxylon* facultatively forms
363 lichenized associations in nature, it represents a useful subject for transcriptomic studies comparing
364 the symbiotic and saprotrophic life styles. Furthermore, a culture system, which allows the re-
365 combination of multiple symbiotic partners, is useful for exploring specificity patterns, symbiotic
366 robustness and, eventually, the establishment of artificial symbioses (Hays *et al.*, 2015). In the case
367 of the *Schizoxylon*-algal-bacterial system, further stabilization of the mixed cultures could indeed be
368 performed by modified media or by introducing periodic disturbance or variation in hydratic
369 conditions, which would promote tighter interactions between the symbionts. While bacteria did not
370 influence the growth rate directly according to the present results, they might play a role for the
371 integrity of the symbiosis under the varying, natural conditions, or for establishing the symbiotic

372 phenotype. With our approach we were able to keep mixed cultures of members from different
373 kingdoms in balance, and we are now able to explore the lichen symbiosis and its relation with
374 associated bacteria using controlled experiments.

375

376

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381

382

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532

533 Captions to figures

534

535 **Figure 1.** Habit of *Schizoxylon albescens* in nature and co-cultured with bacteria and algae (a=
536 algae, b= bacteria, f= fungus): A) *Schizoxylon* on *Populus* bark, green algal colonies are visible
537 around the fungal fruiting body (arrow); B, C) bacterial colonies on agar plates; D-I) mixed cultures

538 of *Schizoxylon* with different algae and bacteria strains: D) *Schizoxylon*, *Coccomyxa* PL2-1 and
539 SAP-5B/1w on MY medium; E) *Schizoxylon*, *Trebouxia* sp.1 and LIC-4A/10 on TM medium; F)
540 *Schizoxylon*, *Trebouxia* sp.1 and LIC-4A/10 on MY medium; G) *Schizoxylon*, *Coccomyxa*
541 MW8233 and MW9636.7 on MY medium; H, I) squash preparations mounted in water of
542 *Schizoxylon*, *Trebouxia* sp.1 and LIC-1B/1 on MY medium, bacteria are visibly attached to the
543 fungal hyphae (arrows). Scale bars: A, E = 0,5 mm; B, C = 2,5 cm; D = 4 mm; F, G = 1 mm; H, I =
544 20 μm .

545

546 **Figure 2.** Scanning electron microscopy (SEM) microphotograph of *Schizoxylon albescens* co-
547 cultured with *Coccomyxa* algae and bacterial strains (a= algae, b= bacteria, f= fungus). A, B) Algal
548 cells intertwined by fungal hyphae. B, D) Bacterial cells are spread among fungal hyphae and algal
549 cells, but are locally distributed only in some parts of the mixed cultures (arrow). C, E) The
550 filamentous matrix formed by the fungus when co-grown with *Coccomyxa* algae is visible (arrow).
551 A, B) *Schizoxylon*, *Coccomyxa* PL2-1 and MW9636.7 on MY medium; C, D) *Schizoxylon*,
552 *Coccomyxa* MW8233 and MW9636.7 on MY medium; E, F) *Schizoxylon*, *Coccomyxa* PL2-1 and
553 MW9636.1 on TM medium. Scale bars: A-C, F = 10 μm ; D = 6 μm ; E= 15 μm .

554

555 **Figure 3.** Scanning electron microscopy (SEM) microphotograph of *Schizoxylon albescens* co-
556 cultured with *Trebouxia* algae and bacterial strains (a= algae, b= bacteria, f= fungus): A-C)
557 *Schizoxylon*, *Trebouxia* sp. 1 and LIC-4A/10 on MY medium; D, E) *Schizoxylon*, *Trebouxia* sp. 1
558 and LIC-4A/12 on MY medium. Bacteria are tightly adhering to the fungal hyphae (A, D) or spread
559 among algal cells and fungal hyphae (B, C, E). Scale bars: A-E = 6 μm .

560