

Revisiting Australian Ectocarpus subulatus (Phaeophyceae) from the Hopkins River: distribution, abiotic environment, and associated microbiota

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Simon M. Dittami, Akira F Peters, Bezhin Rosko, John West, Thierry Cariou, et al.. Revisiting Australian Ectocarpus subulatus (Phaeophyceae) from the Hopkins River: distribution, abiotic environment, and associated microbiota. Journal of Phycology, 2020, 56 (3), pp.719-729. hal-02340942v2

HAL Id: hal-02340942 https://hal.sorbonne-universite.fr/hal-02340942v2

Submitted on 9 Dec 2020

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REVISITING AUSTRALIAN *ECTOCARPUS SUBULATUS* 1 (PHAEOPHYCEAE) FROM THE HOPKINS RIVER: DISTRIBUTION, 2 ABIOTIC ENVIRONMENT, AND ASSOCIATED MICROBIOTA¹ 3

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23 Abstract

In 1995 a strain of *Ectocarpus* was isolated from Hopkins River Falls, Victoria, Australia, 24 25 constituting one of few available freshwater or nearly freshwater brown algae, and the only one 26 belonging to the genus *Ectocarpus*. It has since been used as a model to study acclimation and 27 adaptation to low salinities and the role of its microbiota in these processes. To provide more 28 background information on this model, we assessed if *Ectocarpus* was still present in the Hopkins river twenty-two years after the original finding, estimated its present distribution, 29 described its abiotic environment, and determined its in situ microbial composition. 30 We sampled for *Ectocarpus* at 15 sites along the Hopkins River as well as 10 neighboring sites 31 32 and found individuals with ITS and *cox*1 sequences identical to the original isolate at three sites 33 upstream of Hopkins River Falls. The salinity of the water at these sites ranged from 3.1-6.9 psu, 34 and it was rich in sulfate (1-5 mM). The diversity of bacteria associated with the algae *in situ* 35 (1312 operational taxonomic units) was one order of magnitude higher than in previous studies of the original laboratory culture, and 95 alga-associated bacterial strains were isolated from algal 36 37 filaments on site. In particular, species of *Planctomycetes* were abundant in situ but rare in 38 laboratory-cultures.

Our results confirmed that *Ectocarpus* was still present in the Hopkins River, and the newly
isolated algal and bacterial strains offer new possibilities to study the adaptation of *Ectocarpus* to
low salinity and its interactions with its microbiome.

42 Keywords: *Ectocarpus subulatus*; distribution; low salinity adaptation; freshwater colonization;
43 microbiota.

44 Introduction

Brown algae (Phaeophyceae) are widespread in the tidal and sub-tidal zone of rocky shores in 45 46 temperate marine environments, but they are rarely found in fresh water (Dittami *et al.* 2017). 47 While there are ca. 2,000 known species of marine brown algae, covering a large range of 48 morphologies from small filamentous algae to large and morphologically complex kelp species. 49 there is only a handful of known freshwater brown algae, all of them small and with simple morphology (crust-forming or filamentous). Among these freshwater brown algae the genus 50 51 *Ectocarpus* has a unique position because it corresponds to a predominantly marine genus. 52 which, on two occasions, has been recorded also in rivers: One occurrence of Ectocarpus 53 *crouaniorum* Thuret in a highly salt-contaminated section of the Werra river in Germany 54 (Geissler 1983), and one occurrence of *Ectocarpus subulatus* Kützing in a nearly freshwater 55 habitat (salinity 1ppt) in the Hopkins River, Victoria, Australia (West and Kraft 1996; Peters et al. 2015). 56

The isolate from the latter site (Culture Collection of Algae and Protozoa accession 1310/196), 57 58 constitutes a potential model system to study marine-freshwater transitions in brown algae. The 59 species E. subulatus (Peters et al. 2015) is related to the genomic model species Ectocarpus siliculosus (Dillwyn) Lyngbye (Cock et al. 2010) and has previously been found in highly 60 61 variable environments, including environments with high levels of abiotic stressors. Its occurrence was reported, for instance, at Port Aransas, Texas, USA, where monthly average 62 63 water temperatures reach 30°C in July (Bolton 1983). More recently, the nuclear genome of E. 64 subulatus has been sequenced, revealing that E. subulatus, in comparison to Ectocarpus siliculosus, has lost members of gene families down-regulated in low salinities, and conserved 65 66 those that were up-regulated (Dittami et al. 2018, preprint). The E. subulatus strain from Hopkins

67 River Falls has further been used for physiological experiments: it can grow in both seawater and fresh water and its transcriptomic and metabolic acclimation to these conditions has been 68 69 examined (Dittami et al. 2012) along with the composition of its cell wall with regard to sulfated 70 polysaccharides (Torode et al. 2015). Moreover, the capacity of the freshwater strain to grow in 71 low salinities has been shown to depend on its associated microbial community, although the 72 nature of this dependence is still unknown (Dittami et al. 2016). Extensive efforts have been 73 made to develop a collection of cultivable bacteria to study this phenomenon (KleinJan et al. 74 2017).

Despite this increasing quantity of data on the physiology of the Hopkins River Falls strain of E. 75 76 subulatus, we currently know little about its abiotic environment in situ. The original paper 77 describing its isolation (West and Kraft 1996) states that it was isolated on March 24th, 1995 from 78 cracks between the basalt rock of the Hopkins River, just above the Hopkins River Falls. Water temperature was 16°C, salinity was approximately 1 ppt, and conductivity approximately 3 mS·s⁻ 79 80 ¹. However, it is unknown if *E. subulatus* is still present at Hopkins River Falls, and if so what its current distribution is. Furthermore, the culture has undergone > 20 years of cultivation in 81 82 different laboratories, potentially having a strong impact on its associated microbiota.

In this study, we address both of these knowledge gaps by returning to the Hopkins River and searching for this alga for the first time since its discovery 20 years ago. We found *E. subulatus* individuals at three locations along the Hopkins River, examined its associated microbiome *in situ*, and isolated several novel alga-associated bacterial strains from these samples. These data provide important background information for the use of *E. subulatus* as a model to study low salinity acclimation/adaptation and the role of microbes in these processes.

89 Materials and methods

90 **Biological samples**

The sampling campaign was carried out from March 21st to March 27th, 2017 and covered several 91 92 locations along the Hopkins River between Warrnambool and Ararat (sites 1-15; selected due to 93 their accessibility and to cover the entire length of the river), as well as ten sites selected arbitrarily along the Southern Australian Coastline between Port Fairy and Avalon (Figure 1, 94 95 Table 1). At each sampling site, we manually searched for filamentous algae resembling a member of the Ectocarpales within a range of ca. 50m and for at least 30 min. If filaments were 96 97 found, small amounts of live samples were taken and rinsed three times in sterile 50 mL Falcon 98 tubes with 0.2µm-filtered local water (3 replicates). A small piece of each sample was stored at max. 20°C in sterile 2mL Eppendorf tubes filled with the surrounding water for live algal 99 100 cultures. The second part of the samples was ground on-site according to Tapia et al. (2016), with 101 50 µL of 0.2 µm-filtered local water in a sterile mortar and the proximity of a Bunsen burner. 102 One, seven, and 35 μ L of the ground alga were diluted with 0.2 μ m-filtered local water to a final 103 volume of 50 μ L and spread immediately onto pre-prepared R2A agar plates (Sigma-Aldrich, St. 104 Louis, MO, USA) for isolation of culturable bacteria. These plates were kept at ambient 105 temperature (max. 25°C) and were monitored for two weeks. Newly emerging colonies were 106 purified once more on fresh R2A plates and then put into culture in liquid Zobell medium (Zobell 107 1941) with 8-fold reduced salt concentration, identified by 16S rRNA gene sequencing (see 108 below), and put into stock at -80°C in 40% Glycerol. The remaining sample was dried using 109 silica gel for downstream analysis of the microbial community composition, and frozen at -20°C 110 after the sampling campaign.

111 For all sites, we also collected samples for germling emergence experiments to detect the 112 presence of *Ectocarpus* spores. Three to seven sediment samples including small pieces of solid 113 substrate (shells, pebbles, branches) if present. Approximately 0.1 mL of sediment were kept as 114 live samples in sterile 2 mL Eppendorf tubes. After two weeks these samples were transferred to 115 fresh Provasoli-enriched (Starr and Zeikus 1993) medium based on 5%, 25%, or 100% seawater, 116 depending on the salinity of the water at the sampling site. Seawater for culture media was 117 collected in Roscoff (48°46'40" N, 3°56'15" W), 0.45 µm filtered, and autoclaved at 120°C for 20 118 min) prior to use. The sediment samples were then kept at 13°C in a 14/10 light-dark cycle at an irradiance level of 25 μ mol PAR·m⁻²·s⁻¹, and the emergence of *Ectocarpus*-like germlings was 119 120 monitored over four months.

Both live algae collected *in situ* and those recovered from germling emergence experiments were
cleaned by rigorous pipetting with a Pasteur pipette and several transfers to fresh, sterile,
medium. Any diatoms that remained attached to the algal filaments were removed via treatment
with 3mg·L⁻¹ GeO₂ for 3 weeks.

125 Water samples

126 Approximately 100 mL of water were taken from each site, immediately filtered with 0.45 127 μ M syringe filters to remove particulate matter, and then pasteurized for 1h at 95-100°C to 128 remove any remaining bacterial activity. Filtered samples were stored at ambient temperature until the end of the sampling campaign (max. 2 weeks) and then frozen at -20° C until analysis. 129 130 The conductivity of water samples was determined using a Type CD78 conductivity meter 131 (Tacussel Electronique, Villeurbanne, France) and converted to practical salinity units (psu) 132 according to Fofonoff and Millard (1983). Phosphate, nitrite, and nitrate concentrations were 133 determined using an AA3 auto-analyser (SEAL Analytical, Southampton, UK) following the

method of Aminot and Kérouel (2007) with an accuracy of 0.02 µmol·L⁻¹, 0.01 µmol·L⁻¹, and 134 0.01 μ mol L⁻¹ for NO₃⁻, NO₂⁻, PO₄³⁻, respectively. Sulfate concentrations were determined by 135 136 high-performance anion-exchange chromatography (HPAEC), according to a protocol adapted 137 from Préchoux et al. (2016). After suitable dilution, water samples were injected onto an 138 IonPacTM AS11-HC column (4 x 250 mm) equipped with an AG11-HC guard column (4 x 50 mm), using an ICS-5000 Dionex system (SP-5 & Analytical CD Detector, Thermo Fisher 139 140 Scientific, Waltham, MA, USA). Elution was performed with isocratic 12mM NaOH at a flow rate of 1 mL·min⁻¹, and sulfate ions were detected in conductimetry mode (ASRS 500, 4 mm) and 141 142 quantified using a standard calibration curve. To test if variations in sulfate concentration merely 143 mirrored variations in the overall salinity, a Pearson correlation coefficient was calculated 144 between both variables.

145 Barcoding of algal and bacterial isolates

146 Algal isolates were identified to a species level using the mitochondrial *cox*1 and the nuclear 147 ITS1+2 markers. Algal DNA was extracted from the cleaned cultures using the Macherey Nagel 148 (Düren, Germany) NucleoSpin Plant II kit according to the manufacturer's instructions (PL1 149 protocol with two 25 μ L elutions), and 1 μ L of DNA (10-30 ng) was used in subsequent PCRs. 150 For the ITS region, we used the AFP4LF (3'- CAATTATTGATCTTGAACGAGG-5') and 151 LSU38R (5'-CGCTTATTGATATGCTTA-3') primers (Lundholm et al. 2003; Peters et al. 152 2004), and for the 5'cox1gene the GAZF2 (3'-CCAACCAYAAAGATATWGGTAC-5') and 153 GAZR2 (3'-GGATGACCAAARAACCAAAA-5') primers (Lane et al. 2007), each at a final 154 concentration of 0.5 µM. PCRs were carried out using a GoTag polymerase and the following program: 2 min. 95 °C followed by 30 cycles [1 min 95 °C; 30 sec. 50°C for ITS or 55°C for 155 156 cox1; 3 min 72 °C] and a final extension of 5 min 72 °C.

157 Bacterial cultures were identified by partial sequencing of their 16S rRNA gene. Fifty µL of

dense bacterial culture were heated to 95° C for 15 min, spun down for 1 min, and 1 μ L of

supernatant was used as a template in a PCR reaction with the 8F (5'-

160 AGAGTTTGATCCTGGCTCAG-3') and 1492R 5'-GGTTACCTTGTTACGACTT-3')

161 (Weisburg *et al.* 1991) at a final concentration of 0.5 µM. Except for the annealing temperature

162 (53°C here), the same PCR protocol as above was employed.

163 All PCR products were purified using ExoStar (Thermo Fisher Scientific) and the purified 16S 164 rRNA gene amplicons were sequenced with Sanger technology at the GENOMER platform 165 (FR2424, Roscoff Biological Station), using the BigDye Xterminator v3.1 cycle sequencing kit 166 (Applied Biosystems, Waltham, MA, USA). For bacterial strains, sequencing was carried out 167 only in one direction using the 8F primer, and for algal sequences both the forward and the 168 reverse strand were sequenced and manually assembled. Sequence identification was carried out 169 using RDP classifier (Wang et al. 2007) for bacterial 16S rRNA gene sequences, and BLAST 170 searches against the NCBI nt database (July 2017) for algal sequences. They were further aligned 171 together with reference sequences from the NCBI nt database using the MAFFT server (Katoh et 172 al. 2002) and the G-INS-i algorithm. All positions with less than 95% site coverage were 173 eliminated. Phylogenetic analyses were carried out with MEGA 7 (Kumar et al. 2016) using the 174 Maximum Likelihood method based on the GTR+G+I model and 1,000 bootstrap replicates.

175 Amplicon sequencing of in situ bacterial communities

Amplicon sequencing of bacterial communities was carried out to assess the *in situ* composition of the *E. subulatus* microbiome. Sufficient material for these analyses was obtained at two of the three sites with *Ectocarpus* individuals: sites 8 and 9 (Figure 1). Approximately 20mg dry weight for each of the three replicate samples for each site were ground twice for 45 sec. at 30 Hz in a

180	TissueLyser II (Qiagen, Hilden, Germany). DNA was then extracted using the Qiagen DNeasy
181	Plant mini kit according to the manufacturer's instructions. Approximately 50 ng of DNA (as
182	estimated using a NanodropONE, Thermo Fisher Scientific), were then used to amplify the V3-
183	V4 region of the 16S rRNA gene. Furthermore, a mock community comprising a mix of DNA
184	from 26 bacterial genera cultivated in our laboratory (see Thomas et al. 2019 for details) as well
185	as a negative control, were added alongside the samples. PCR amplification, indexing, and
186	library construction were carried out following the standard "16S Metagenomic Sequencing
187	Library Preparation" protocol (Part # 15044223 Rev. B). Final library concentrations were
188	measured using a BioAnalyzer (Agilent, Santa Clara, CA, USA) before pooling. Libraries for
189	each sample were then pooled in an equimolar way, diluted to 5nM final concentration and
190	supplemented with 20%PhiX to add sufficient diversity for sequencing on an Illumina MiSeq
191	using a 2x300bp cartridge. Raw data were deposited at the European Nucleotide Archive (ENA)
192	under project accession number PRJEB34906
193	(https://www.ebi.ac.uk/ena/data/search?query=PRJEB34906).
194	Raw reads were first trimmed and filtered using the fastx_quality_trimmer script
195	(http://hannonlab.cshl.edu/fastx_toolkit/), assembled using Pandaseq 2.11 (Masella et al. 2012)
196	and further processed with mothur according to the Miseq SOP (version April 4th, 2018; Kozich
197	et al. 2013). Sequences were aligned to the non-redundant SSU ref database version 132,
198	chimeric sequences removed using Vsearch (Rognes et al. 2016), and operational taxonomic
199	units (OTUs) defined based on a 97% identity threshold (Stackebrandt and Goebel 1994). Rare
200	sequences (<5 reads across all samples) were removed from the final analyses. Taxonomic
201	assignments were generated for both the raw reads and the final OTUs using the RDP classifier
202	method (Wang et al. 2007). Non-metric multidimensional scaling (NMDS) of the OTU matrix

was carried out in R 3.5.1 using the isoMDS function of the Vegan package and Bray-Curtis 203 dissimilarity as a distance measure. An Analysis of Similarity (ANOSIM) was used to test for 204 205 differences in the overall community composition between the two sites (3 replicates each, 719 206 permutations). Statistical differences between the two sites at the level of individual OTUs were 207 assessed by multiple two-sided t-tests (one test per OTU) on log-transformed abundance data 208 with subsequent correction for multiple testing according to Benjamini and Hochberg (1995). 209 Alpha diversity was estimated using the Shannon index with e as a base and the diversity() 210 function of the VEGAN package. A two-sided t-test was used to compare these indexes obtained 211 for the replicate samples of both sites. Differences between sites were considered significant if 212 the Type I error rate was below 0.05.

213 **Results**

214 Distribution of Ectocarpus subulatus along the Hopkins River

We found live *E. subulatus* at three of the 15 sampled sites along the Hopkins River, and 215 216 germlings of other Ectocarpales emerged from four additional marine sites along the Victorian 217 coastline, including at the mouth of Hopkins River (Table 1). Despite extensive searches, no 218 traces of *Ectocarpus* were found at the original isolation site of *E. subulatus* at Hopkins River 219 Falls (site 7, Figure 1). Ectocarpus was, however, abundant at two sites (Framlingham Forest 220 reserve and Kent's Ford, sites 8 and 9, Figure 1), which were approximately 12km and 37 km upstream of Hopkins River Falls. The third finding of *Ectocarpus* was registered 83 km upstream 221 222 (site 10), although only a few filaments were found at this site. The *cox*1 and ITS sequences 223 obtained from *Ectocarpus* cultures from all three sites were identical to those of the strain

224	isolated from Hopkins River Falls in 1995 (Figure 2). We found no <i>E. subulatus</i> individuals in
225	other sampled rivers, along the coastline, or in germling emergence experiments.

226 Water chemistry

227 The salinity of the Hopkins River was highest close to the source (8.4 psu; approximately $\frac{1}{4}$ that

of seawater), and then gradually decreased towards the mouth of the river, where it dropped to ca.

- 1 psu, before re-spiking due to the influence of seawater (Figure 3). This decrease corresponded
- to an increase in the flow of water masses towards the mouth river. Sulfate concentrations

followed the same pattern as salinity (Pearson correlation r=0.995, p<0.001) and decreased from

nearly 7 mM to approximately 0.4 mM close to the river mouth. Finally, phosphate and

233 nitrite/nitrate concentrations were variable along the river. They were highest at the Chatsworth

site (site 11), reaching 3.3 and 22.8 μ M, respectively, and then strongly decreased at sites where

235 *Ectocarpus* was found (PO_4^{3-} 0.5-0.8 µM, NO_2^{-}/NO_3^{-} 0.8-1.7 µM; Figure 3).

236 Bacterial communities associated with algae

237 In situ bacterial community composition was determined by 16S rRNA gene amplicon

sequencing for field samples taken at Framlingham Forest reserve (site 8) and Kent's Ford (site 9)

239 (Figure 4A). We detected 1312 OTUs across the three sampled individuals from both sites

240 (Supporting Information File S1). The bacterial communities of both sites were dominated by

241 *Alphaproteobacteria* (25% of reads), *Bacteriodetes* (20%), *Gammaproteobacteria* (8%),

- 242 Planctomycetes (8%), and Actinobacteria (8%) (Figure 4a,b), and there was a significant
- difference in the community structure between the two sites (ANOSIM p=0.001, Figure 4B).
- Examining the OTUs individually, we identified 86 OTUs that were specific to Framlingham
- Forest reserve (including 31 Proteobacteria and 22 Planctomycetes), and 60 more had a higher
- relative abundance there. At Kent's Ford, 27 OTUs were site-specific (including 13

247 *Proteobacteria* and 6 *Bacteroidetes*), and 13 more exhibited higher relative abundance. In three cases, site-specific OTUs from both sites were found to belong to the same genera: Rickettsiales 248 249 of the SM2D12 group, Flavobacterium, and Luteolibacter. A detailed list of these OTUs is 250 provided in Supporting Information File S1. Alpha-diversity (Shannon index) was also slightly 251 higher at the Framlingham Forest reserve (t-test p=0.03; Figure 4C). Amplicon sequencing 252 analyses of bacterial communities were complemented by *in situ* isolation of bacterial strains 253 from the algae after thorough rinsing with sterile river water (Figure 5). They comprise 254 Gammaproteobacteria (48 isolates, including 28 Pseudomonas), Firmicutes (27 isolates), 255 Actinobacteria (8 isolates), Alphaproteobacteria (7 isolates), and Bacteriodetes (5 isolates). No 256 members of the *Planctomycetes* were isolated.

257 **Discussion**

258 The data presented in this paper confirm that the original finding of *E. subulatus* by West and 259 Kraft was not the result of a transient "contamination", but that algae with identical ITS sequences are present in the river at three sites upstream of the original location. At the time of 260 261 sampling, the water at these sites was saline and contained high levels of sulfate for a river, but 262 low levels of nitrite/nitrate and phosphate compared to other upstream and downstream sites. 263 Furthermore, the bacterial community associated with the algae in situ comprised over 1300 264 OTUs, which is highly diverse compared to laboratory cultures. It also, it included a high 265 diversity of *Planctomycetes*. Each of these findings, discussed in more detail below, provides 266 valuable background information when using *E. subulatus* cultures as a model system to study 267 acclimation, adaptation, or interactions with their associated microbiome.

268 Based on these observations, it seems likely that *E. subulatus* has persisted in the Hopkins River 269 for over 20 years, maintaining a population despite the water currents. *Ectocarpus* spores and gametes are motile, but swimming speeds reported are only in the range of 150-270 µm s⁻¹ 270 271 (Müller 1978). This implies that *E. subulatus* in Hopkins River either (1) does not rely on gamete 272 releases for reproduction, (2) that its gametes are able to remain close to the substratum as has 273 been suggested for male gametes (Müller 1978) and direct their movement upstream, or (3) that 274 gametes rely on zoochory, as has been proposed in the case of red algae (Žuljević *et al.* 2016). 275 Our findings thus open interesting perspectives for population genetics studies as well as more 276 detailed studies of the reproductive biology of *Ectocarpus* in this area. Furthermore, the fact that 277 no traces of E. subulatus were found in nearby rivers or along the coastline suggests that it may 278 be restricted to the Hopkins River, although the range of colonization within the river may have 279 been subject to variation, notably because individuals of *E. subulatus* were no longer found at the 280 original isolation site.

281 Although limited to a single point in time, our sampling campaign also provides novel 282 information on the chemical parameters in the Hopkins River at the time of sampling. Notably, 283 the observed salinity at sites with E. subulatus between 3.1 and 6.9 psu leads us to classify the 284 water in the Hopkins River at the sites with E. subulatus at the time of sampling as low salinity 285 brackish water rather than fresh water (usually defined by a salinity <0.5 psu. International 286 Symposium for the Classification of Brackish Waters 1958). This may be one of the factors enabling E. subulatus to be competitive in this environment, a hypothesis which is supported by 287 288 the fact that no individuals were found in the lower portions of the river with lower salinity. High 289 salinity in our samples also positively correlated with high sulfate concentrations between 1 and 5 mM - average sulfate concentrations in fresh water are 0.12 mM (vs. 28 on average in the ocean; 290

291 Wetzel 2001). Sulfated polysaccharides are typical components of the cell walls of marine plants and algae (Popper et al. 2011) and require sulfate for their synthesis, but their importance for 292 293 *Ectocarpus* remains to be explored. In the same vein the question remains open to what extent the 294 low nitrate concentrations at sites with *E. subulatus* compared to upstream and downstream sites. 295 are related to the presence of the algae, either as a cause or as an effect. It should be noted, though, that a direct metabolomic comparison of E. subulatus and the marine E. siliculosus 296 297 revealed markers for high nitrogen status (total amino acids, ratio of glutamine to glutamate) in E. 298 subulatus (Dittami et al. 2012). Regardless of the physiological implications of the composition 299 of the Hopkins River water, we argue that it may be more appropriate to refer to the *E. subulatus* 300 strains isolated from the Hopkins River as "fluviatile", *i.e.* "river" strains rather than freshwater 301 strains, despite their capacity to grow in fresh water in laboratory conditions (Dittami et al. 2012). 302

303 In addition to these facts about the distribution and environment of *E. subulatus*, the present 304 study provides insights into its associated microbiome – a component likely connected to the 305 capacity of this species to grow in low salinity (Dittami et al. 2016). The number of OTUs 306 associated with E. subulatus in our in vivo study was one order of magnitude higher than in a 307 previous study of the laboratory strain after 20 years of cultivation (1312 OTUs for six samples 308 from two sites vs 84 OTUs for six samples in two conditions) (Dittami et al. 2016). Moreover, a 309 direct taxonomic comparison of these two studies at the genus level revealed only 5 genera 310 (Acinetobacter, Phycisphaera, Maribacter, Marinoscillum, and Gaiella) that were found in both 311 studies. All of them were rare *i.e.* supported by < 0.01% of reads in our study; Supporting 312 Information File S1). Both studies were based on sequencing runs with similar depth and employed similar analysis pipelines, yet many technical factors could contribute to such 313

314 differences: the sampling protocol, the primers used, library preparation, the sequencing platform 315 and chemistry (Illumina Miseq V2 vs V3), etc.. Nevertheless, the profoundness of the observed 316 differences suggests that either the microbiome of E. subulatus in the Hopkins River has evolved 317 and diversified over time or that the cultivation of algae in the laboratory has impacted its 318 microbiome, leading to a reduction of diversity and a change in composition. In a context of the 319 development of new laboratory models for the study of marine holobionts (Dittami et al. 2019, 320 preprint), a targeted examination of these potential changes, *e.g.* by following the evolution of 321 alga-associated microbiomes in the field as well as over several cultivation cycles may yield 322 important insights on possible limitations of laboratory model systems. If confirmed, such biases 323 would underline the necessity of devising targeted experiments to test the validity of laboratory 324 findings in the field.

325 The availability of parallel amplicon sequencing data of bacterial communities and untargeted cultivation efforts further allows us to identify under-sampled lineages in cultivation experiments. 326 327 In this study particularly *Planctomycetes* stand out, as they constituted 176 OTUs and 8% relative 328 abundance of all algae-associated reads but did not have a single associated culture. 329 *Planctomycetes* are notoriously difficult to cultivate, partially due to their long doubling time of 330 up to one month. They require low organic content in media, physical separation from fast-331 growing competitors e.g. via dilution to extinction experiments, and they may benefit from the 332 use of fungicides (Lage and Bondoso 2012). In contrast to the present study, previous barcoding 333 data (Dittami et al. 2016) on cultivated E. subulatus revealed the presence of very few 334 *Planctomycetes* (0.1% of reads), implying that these culturing techniques would need to be put into place using freshly collected material. In contrast, the high abundance of *Firmicutes* in the 335

isolation experiments although they account for only 1% of the reads in the amplicon sequencingdata may be because these bacteria were particularly amenable to the culture condition.

The present study enhances our knowledge on *E. subulatus* from the Hopkins River and its associated microbiome. It furthermore provides a new set of microbes for coculture experiments and thus strengthens the use of *E. subulatus* both as a model for the study of acclimation and adaptation to low salinity and of algal-bacterial interactions.

342 Acknowledgments

343 This work was funded partially by ANR project IDEALG (ANR-10-BTBR-04) "Investissements d'Avenir, Biotechnologies-Bioressources", the European Union's Horizon 2020 research and 344 345 innovation Programme under the Marie Sklodowska-Curie grant agreement number 624575 346 (ALFF), and an internal call for proposals from the UMR8227 (CNRS, Sorbonne University). We 347 thank Cécile Hervé, Amandine Simeon, and Agnieszka P. Lipinska for helpful discussions; 348 Gwenn Tanguy and Erwan Legeay from the GENOMER platform, Roscoff for support during the 349 library construction and sequencing; and the ABIMS platform for providing the computational 350 facilities for the amplicon sequencing analyses.

351 **Conflict of interest statement**

352 The authors declare no conflict of interest.

353 Author's contributions

- 354 SD, AFP, HK, SE, JW, CB planned the study; TC performed nutrient analyses; AP measured
- 355 sulfate concentrations; BBD performed amplicon sequencing analyses of bacterial communities ;

SD performed sampling, culturing, *in silico* analyses, and wrote the manuscript; All authorscorrected the manuscript and approved the final draft.

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447

Tables

Table 1: Overview of samples taken and species identified. The numbers in the location column

451	correspond to	site numbers	in Figure	1.
451	correspond to	site numbers	in Figure	1

Data	Location	Material	<i>Ectocarpus</i> tissue	GPS coordinates	Germling	Sequence
2017	1 Morri Island	nabblag/muggalg	Tounu	28 402116	emergence	
2017-		in tide neel	110	-36.402110,	-	-
2017	2 Daint Ditahia	in the pool		142.4/1340	Estasamour	I D725221 (aav1)
2017-	2. Point Ritchie	sand/ sandstone	по	-38.401275,	Eciocarpus	LK/35221(COX1)
2017	2 Mahamar			142.309318	siliculosus	LK/35414 (115)
2017-	5. Manoney	sand,	по	-38.392103,	-	-
03-22	Road	wood/plastic		142.531644		
2017-	4. Smith Lane	mud, wood	no	-38.39/984,	-	-
03-22		1 ())		142.578286		
2017-	5. Allan's Ford	rock (granite)	no	-38.385325,	-	-
03-22				142.587398		
2017-	6. Donovans	granit, clay	no	-38.355110,	-	-
03-22	Lodge			142.599250		
2017-	Hopkins	volcanic rock	no	-38.333509,	-	-
03-22	River Falls			142.621352		
2017-	8. Framlingham	rock, pebbles	E. subulatus Au8	-38.297064,	-	LR735222 (cox1)
03-23	Forest Reserve		(abundant)	142.668291		LR735415 (ITS)
2017-	9. Kent's Ford	rock, pebbles	E. subulatus Au9	-38.191574,	-	LR735223 (cox1)
03-23		, 1	(abundant)	142.698058		LR735416 (ITS)
2017-	10. Hexham	mud. wood	E. subulatus	-37,995732	-	LR735224 (cox1)
03-23			Au10 (rare)	142,689141		LR735417 (ITS)
2017-	11 Chatsworth	mud sand	no	-37 856357	-	-
03-23	11. Chuisworth	detrirus	110	142 650644		
2017	12 Wickliffe	cond detritus	n 0	37 60/3/8		
02 22	12. WICKIIIIC	sanu, ueunus	110	-37.094340, 142.726074	-	-
2017	12 Decembridge	roals nabblas		27 490217		
2017-	15. Rossbildge	Tock, peobles	110	-57.460217,	-	-
03-24	14 4	1		142.849405		
2017-	14. Ararat	sand	no	-37.300211,	-	-
03-23	1.5.0			142.9/39/9		
2017-	15. Green Hill	pebbles	no	-37.295320,	-	-
03-24	Lake			142.979170		
2017-	16. Merri River,	concrete	no	-38.362077,	-	-
03-24	Warnambool			142.484414		
2017-	17. Killarney	sand, lava	no	-38.357966,	<i>Kuckuckia</i> sp.	LR735225 (cox1)
03-24	beach	pebbles, shells		142.306880		LR735418 (ITS)
2017-	Belfast	sand, wood	no	-38.361889,	-	-
03-25	Lough airport	pollar		142.262045		
2017-	19. Curdies	rock	no	-38.519965,	-	-
03-25	River			142.833558		
2017-	20. Port	tide pool. sand	no	-38.620681,	<i>Feldmannia</i> sp.	LR735226 (cox1)
03-25	Campbell	1		142.992981	1	()
2017-	21. Gellibrand	mud. detritus	no	-38.727482	-	-
03-25	River	indu, uttitus		143 250932		
2017-	22 Aire River	mud nebbles	no	-38 763797	_	
$03_{-}25$	22. / me nover	inda, peooles	110	1/13 /7/727		
2017	23 Wild Dog	cand	n 0	28 725011		
03 25	25. White Dog Creek	sanu	110	-30.733711,	-	-
2017	21 Smithala	nabblas biafilm	n 0	143.003343		
201/-	24. Smythe's	peoples, blomm	110	-38./04048,	-	-
03-25	Creek	(1.1		145./62856	4	L D 72 5 2 2 7 (1)
2017-	25. Lorne	tide pool, sand	no	-38.531281,	Acinetospora	LR/35227(cox1)
03-26	A (11	shells		143.980994	sp.	
2017-	26. Hovell's	clay	no	-38.018825,	-	-
03-26	Creek			144.402156		

Figure legends



454 Figure 1: Sites sampled. Panels A-C: Map of sampling sites at an increasing scale. Each
455 sampling site is numbered (1-26, see Table 1). *Ectocarpus* individuals were found at (sites 8, 9,
456 10), and Panel D contains photos taken at these sites. Hopkins River Falls, the original site of
457 isolation, corresponds to site 7.



0.020

459 Figure 2: Maximum-likelihood tree of *Ectocarpus* isolates from Hopkins River and related

E. subulatus CCAP 1310/109 FR668795 E. subulatus CCAP 1310/123 FR668804 99 E. subulatus CCAP 1310/282 FR668909 1.E. subulatus CCAP 1310/196 FR668844

E. subulatus Au10 66 E. subulatus Au8 E. subulatus Au9

- 460 strains. Panel A displays a tree based on the COXI gene (alignment of 677 bp after curation),
- 461 and Panel B on the ITS region (860bp after curation). Blue color indicates isolates from this
- 462 study. Support values correspond to the percentage of support using 1000 bootstrap replicates.



Figure 3: Water chemistry at the different sampling sites along the Hopkins River (see
Figure 1). Panel A displays salinity (gray bars) and nutrient concentrations (blue lines), and Panel
B shows sulfate concentrations. Yellow background indicates sites with the occurrence of *Ectocarpus*. Each measurement corresponds to a single sample collected between 2017-03-22
and 2017-03-24.

A) Taxonomic compostion



470 Figure 4: Bacterial community at isolation sites determined by 16S rDNA amplicon

471 sequencing of bacterial communities. Panel A shows the taxonomic distribution of the bacterial

472 communities associated with each replicate at the two sampling sites with sufficient material, uc

473 = unclassified. Panel B shows the distances between the communities (NMDS plot based on

- 474 Bray-Curtis dissimilarity matrix); communities at both sites differed significantly (ANOSIM
- 475 p=0.001). Panel C shows the alpha-diversity of bacterial communities in the two sites (mean of
- 476 three replicates \pm SD; * indicates a significant difference, two-sided t-test, p=0.03).







483 Supporting Information

484 Supporting Information File S1 – Amplicon sequencing results of bacterial communities.
 485