## 1 Microbial 'gardening' by a seaweed holobiont: surface metabolites attract

- 2 protective and deter pathogenic epibacterial settlement
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- 15 Running Head: Chemical 'gardening' of beneficial epibacteria by an invasive seaweed
- 16 Summary
- Epimicrobial communities on seaweed surfaces usually contain not only potentially
   pathogenic, but also potentially beneficial microorganisms. Capacity of terrestrial
   plants for chemically mediated recruitment i.e. 'gardening' of bacterial communities
   in the rhizosphere was recently demonstrated. Empirical evidence directly linking
   such chemical 'gardening' with the beneficial role of gardened microbes in terrestrial
   plants is rare and largely missing for aquatic macrophytes.
- Here we demonstrate that our model invasive seaweed holobiont *Agarophyton vermiculophyllum* possesses beneficial microbiota on its surface that provide

protection from bacterial pathogens. Metabolites from the algal holobiont's surface
 reduced settlement of opportunistic pathogens but attracted protective epibacterial
 settlement.

3. We tested 58 different bacterial species (isolated from the surface of A. 28 29 vermiculophyllum) individually in tip bleaching assays. Kordia algicida was identified as a 'significant pathogen' inducing a bleaching disease. In addition, 9 other species 30 significantly reduced the risk of algal bleaching and were thus 'significantly 31 32 protective'. Additionally, 2 'potential pathogens' and 10 'potential protectors' were identified. When 19 significant and potential protectors and 3 significant and 33 potential pathogens were tested together, the protective strains fully prevented 34 bleaching, suggesting that a component of A. vermiculophyllum's epimicrobiome 35 provides an associational defence against pathogens. Chemically mediated selective 36 37 recruitment of microbes was demonstrated in bioassays, where A. vermiculophyllum 38 surface metabolites attracted the settlement of protective strains, but reduced settlement of pathogens. 39

40 4. Synthesis: The capacity of an aquatic macrophyte to chemically 'garden' protective
41 microorganisms to the benefit of strengthened disease resistance is demonstrated
42 for the first time. Such a role of surface chemistry in 'gardening' of microbes as
43 found in the current study could also be applicable to other host plant – microbe
44 interactions. Our results may open new avenues towards manipulation of the
45 surface microbiome of seaweeds via chemical 'gardening', enhancing sustainable
46 production of healthy seaweeds.

Key words: Agarophyton vermiculophyllum, Macrophyte, Chemical Defence, Plant-microbe
interactions, *Gracilaria vermiculophylla*, Bleaching, Gardening, Invasive, Seaweed,
Holobiont.

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#### 51 Introduction

All eukaryotes including terrestrial plants and aquatic macrophytes are influenced by complex interactions with microbial communities. The animal gut microbiome is very well known to influence the health and nutritional status of its host (Hooper *et al., 2002;* Flint *et al., 2012*), ultimately forming a metaorganism or holobiont that consists of the host and associated microbiomes (Bordenstein & Theis, 2015). These microbes form an integral part of a plant or animal phenotype, influencing the fitness and ecological traits of their hosts.

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The outer body surface is the primary physiological and ecological interface of multicellular 59 aquatic organisms like water plants or seaweeds with the environment (Wahl, 2008). Apart 60 61 from exchange and uptake of nutrients, this interface is involved in the exchange of chemical cues and signals that mediate the recognition of an organism by a partner, a 62 63 parasite, an epibiont or a predator. This surface is often colonized by complex microbial 64 communities, a biofilm-like epimicrobiome that has also been denoted as 'second skin' (Wahl *et al.*, 2012). Marine macroalgae i.e. seaweeds have an additional diffusive boundary 65 66 layer (Hurd, 2000) along with their 'second skin' that serves as the micro-niche of chemically mediated ecological interactions. This micro-niche is analogous to the rhizosphere of plant 67 roots (Hartmann *et al.*, 2009) or the phycosphere of phytoplankton (Bell & Mitchell, 1972). 68

This niche is an ecological interface of seaweed-microbe relationships, modulates most of the interactions between the seaweed host and the environment and is typically characterized by a specific chemical fingerprint.

Seaweeds are omnipresent organisms in photic coastal zones, play key roles in carbon 72 fixation, biogeochemical cycling and food web formation. They can drive the biogeochemical 73 74 pump and release climate cooling gases like dimethyl sulphide (Van Alstyne & Houser, 2003). They act as nursery ground and protective shelters for many animals (Schiel & Foster, 75 76 2006; Pereira et al., 2017). Seaweeds also provide substrate for numerous sessile organisms, 77 ranging from bacteria to macro-invertebrates (Wahl, 1989). Epibacteria that colonize the surfaces of seaweeds vary taxonomically with host, space and time (Cundell et al., 1977; 78 79 Lachnit *et al.*, 2011) and can affect the well-being of their host in multiple ways.

The epimicrobial communities on seaweeds consist not only of pathogenic species but also 80 of potentially beneficial ones. Interactions with the surface epimicrobiome have the 81 82 potential to influence seaweed health and development in two different ways: they can be 83 detrimental, as seaweeds can be plagued by bacterial and eukaryotic pathogens (see Gachon et al., 2010; Egan et al., 2013 and references therein). The epimicrobiome also often 84 provides inductive settlement cues to algal spores and invertebrate larvae, causing heavy 85 detrimental fouling (see Wahl et al., 2012 and references therein). Alternatively, seaweed 86 87 epimicrobiomes can also be beneficial, supplying essential nutrients (see Hollants et al., 88 2013 and references therein) and chemical cues for morphogenesis (see Wichard et al., 89 2015 and references therein).

90 A suspected yet relatively undemonstrated beneficial role of the epimicrobiome is the 91 protection from pathogens and other detrimental microorganisms (but see Longford *et al.*,

2019). A certain component of these epimicrobial communities on seaweed surfaces is quite 92 host-specific (Lachnit et al., 2009; Bengtsson & Ovreas, 2010) and the same is true for the 93 rhizosphere of terrestrial plant roots (Raaijmakers et al., 2009). However, the principles 94 95 governing the assemblages of microbes on surfaces of seaweeds or any other aquatic 96 macrophytes are unclear. Based upon recent independent studies with terrestrial plants and 97 aquatic macrophytes the following models for the association of microbial communities 98 have been proposed: 1. The 'neutral' hypothesis assumes that species are ecologically 99 equivalent, and the community structure is determined randomly (Hubbel 2001, 2006; Woodcock, 2007). 2. The 'niche' model stresses that only microorganisms which are 100 adapted to the specific conditions on a host surface will be able to settle on it (Dumbrell et 101 al., 2010). 3. The 'lottery' hypothesis combines both neutral and functional aspects and 102 predicts that multiple microorganisms could make use of the same niche, but those that 103 104 reach it first have a larger chance of settlement success (Burke et al., 2011). 4. Untargeted 105 recruitment of microorganisms by the host via the release of exuded nutrients has also been 106 proposed, as well as targeted deterrence by processes like induced defence (Weinberger, 107 2007).

108 By comparing bacterial root microbiomes between wildtype Arabidopsis thaliana and 109 mutants that could not produce the defence phytohormone salicyclic acid, Lebeis et al., 2015 recently demonstrated that salicylic acid signalling can modulate root microbial 110 111 communities. While such studies on the role of chemical manipulation of root microbiota 112 have started to appear for land plants, no parallel study exists for aquatic macrophytes that demonstrates an active 'deliberate' recruitment or 'gardening' of beneficial microbes. 113 114 Surface associated metabolites may shape the microbial communities on seaweed surfaces. 115 For example, halogenated furanones excreted by the host Delisea pulchra were

demonstrated to shape the microbiome of the seaweed (Longford et al., 2019). Also, in the 116 brown alga Fucus vesiculosus surface metabolites were found to have an effect on the 117 biofilm composition both under field and lab conditions (Lachnit et al., 2010). The authors 118 used an experimental system that simulated the delivery of Fucus surface associated 119 120 metabolites on artificial substrates and tested the effect of algal surface chemistry on 121 bacterial community composition. Bacterial communities that developed on test surfaces 122 loaded with Fucus surface metabolites were found to be quite similar to communities on the 123 surfaces of Fucus, but different from communities on solvent controls, which hinted at the strong selective force of these surface metabolites of *Fucus*. However, for the investigation 124 with Fucus vesiculosus no evidence could be demonstrated for the beneficial role of such 125 microbes and thus the purpose of such chemically mediated recruitment of microbes. Also, 126 studies of the rhizosphere of terrestrial plants already reported selective 'gardening' of 127 128 microbes (Currier & Strobel, 1976; Bacilio-Jim'enez et al., 2003). For example, root exudates 129 of different developmental stages of Arabidopsis promoted the formation of microbial communities with different compositions when the influence of environmental and soil 130 131 edaphic factors was experimentally excluded (Yuan et al., 2015). Although there have been demonstrations of the possible beneficial roles of such active microbial gardening for plant 132 growth and development in terrestrial environments (Lebeis et al., 2015) and marine 133 134 environments (Kessler et al., 2018), none of the studies in the aquatic realm have yet been able to empirically link chemically mediated microbial 'gardening' with resistance to disease. 135

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Thallus whitening, bleaching or *'ice-ice* disease' is a common problem in certain farmed and
wild red seaweeds, such as *Gracilaria 'conferta'* (Weinberger *et al.,* 1994; Weinberger *et al.,*

139 1997), Kappaphycus and Eucheuma or Delisea pulchra (Case et al., 2011; Campbell et al., 2011). It was repeatedly shown that this depigmentation symptom can be induced by 140 multiple opportunistic bacterial pathogens and in the case of G. 'conferta' a component of 141 142 the microbiome was shown to prevent the disease. Also, in Delisea pulchra early successional epibacterial strains protected the host from a later successional strain that was 143 pathogenic when the host microbiome was experimentally disturbed (Longford *et al.*, 2019). 144 145 In the context of the 'gardening' hypothesis the present study investigated whether (a) 146 epibacteria originating from healthy specimens of the invasive red seaweed Agarophyton vermiculophyllum can also induce thallus bleaching in A. vermiculophyllum, whether (b) a 147 148 subset of epibacterial strains of the algal microbiome offers protection towards pathogenic strains and whether (c) A. vermiculophyllum has a capacity for chemically mediated 149 recruitment of such protective microbes while deterring the settlement by pathogens. 150

#### 151 Materials and Methods

#### 152 Isolation and identification of epibacterial strains

Five invasive and five native populations of Agarophyton vermiculophyllum (Gurgel et al., 153 2018) (Synonym: Gracilaria vermiculophyllum (Ohmi) Papenfuss, hereafter: Agarophyton) 154 155 were sampled along the Danish-German Peninsula of Jutland and Schleswig-Holstein and the South Korean peninsula, respectively (see Table S1 in Supporting Information). Using 156 standard protocols, bacterial strains were isolated from the surface of Agarophyton. Thus, 157 158 the tested bacterial strains were ecologically relevant. 5 g of pooled algal individuals arising 159 from each population were rinsed three times in 35 ml of Bacto Marine Broth (MB; Difco 2216, Becton Dickinson and Company, Heidelberg, Germany) to remove loosely attached 160 161 bacteria. Then, the samples were immediately transferred to 10 ml of MB and vortexed vigorously for 20 s to detach the epibacteria. The suspension was subsequently diluted in 162

MB using the log dilution method and plated out directly on MB agar (37.3 g<sup>-1</sup> MB, 15.0 g<sup>-1</sup> 163 agar; pH 7.6) in standard Petri dishes. Incubation was performed in the dark at 28°C for 7 164 days. Pure cultures were obtained through several subsequent picking and culturing steps 165 for individual colonies on MB agar plates. The isolates were cryopreserved at -80°C using the 166 Cryobank System (Mast Diagnostica GmbH, Reinfeld, Germany) according to the 167 manufacturer's instructions, until processed further. Strains were identified by 16S rRNA 168 169 sequencing as described in Saha et al., (2016) and tested in the bioassays described below. 170 For methodological details see Appendix S1 in Supporting Information.

#### 171 Agarophyton tip bleaching assay with epibacterial strains

#### 172 (A) Bleaching assay with single isolates

To test the potential capacity of epibacterial strains for induction of thallus bleaching in *Agarophyton*, 58 of the cryopreserved bacterial strains were reanimatedin November 2015. They were then maintained on MB agar medium in darkness. All cultures were incubated at 25°C, except *Psychroserpens mesophilus* and *Pseudoalteromonas lipolytica* as they exhibited no growth at this temperature and were incubated at 15°C.

In November 2015, *Agarophyton* individuals were sampled from Nordstrand (53°29'10.25"N, 8°38'35.33"E) and brought to the laboratory in a cooler box. They were maintained in 20 L aquaria at a salinity of 33 psu (approximate salinity value at the collection site) and a temperature of  $16^{\circ}$  C under constant aeration and a photon flux density of 75 µmol m<sup>2</sup> s<sup>1</sup> (12 h of light per d).

For the experiment, *Agarophyton* thallus tips (n = 6 in total for each bacterial strain, each tip was ca. 2-3 cm long) were individually placed into separate wells of 24 well plates (Sarstedt, GmbH) containing 2 ml of sterile sea water (SSW, 33 psu). To eliminate epibacteria from the algal surface, two antibiotics, Vancomycin and Cefotaxim (each at concentration of 0.1 mg  $ml^{-1}$ ) were added to each well (Weinberger et al., 1997). The wells were then incubated for 2 days at 16 °C and a photon flux density of 75 µmol m<sup>-2</sup> s<sup>-1</sup>. Following this pre-treatment, the wells were carefully emptied of SSW and antibiotics. Remaining antibiotics were removed from *Agarophyton* tips and the wells by washing with 1 ml of SSW. Finally, 2 ml of SSW were again added into each well and bacteria cultures were immediately inoculated.

Prior to inoculation all bacterial cultures were grown in sterile MB medium for 3-7 d at the 192 same temperature that was used for their maintenance (25°C or 15°C, see above) in 193 194 darkness until they had reached an  $OD_{610}$  of 0.2 to 0.3. A volume of 20 µl bacterial cells along with the medium was then added into the wells containing Agarophyton tips (n = 6). 195 Controls consisted of the same volume of sterile bacterial culture medium added into the 196 wells containing Agarophyton tips (n = 6) and treated in a similar manner as above. 197 Following five days of incubation (16 °C and a photon flux density of 75  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>) all wells 198 199 were checked under the binocular microscope (magnification factor: 20, see Supporting 200 Information Fig. S1) and numbers of bleached and non-bleached tips in each well were counted, using a black background (Weinberger et al., 1997). 201

202 Relative risk of thallus tip bleaching in treatments with addition of bacteria relative to 203 control treatments were calculated as odds ratios of numbers of bleached and non-204 bleached tips.

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## 206 Relative risk of bleaching = (Bleached tips in treatments + healthy tips in treatments)

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(Bleached tips in controls + healthy tips in controls)

208 95% Confidence intervals of these ratios were constructed following Fisher and Van Belle
209 (1993). The divergence of these ratios from 1 (= no effect of the treatment on the risk) was
210 tested for significance, using Fisher's exact test (Fisher & Van Belle, 1993). Isolates that

significantly induced thallus tip bleaching were retested in one or two independent 211 repetitions of the experiment to confirm the result. The Mantel-Haenßel-extention of 212 Fisher's exact test for replicated test designs was used for the statistical analysis in these 213 cases (Fisher & Van Belle, 1993; Weinberger et al., 1997). To reduce the risk of type I error a 214 Bonferroni correction was applied if multiple tests had to be done (Fisher & Van Belle, 215 216 1993). Isolates that turned out to be significantly pathogenic after applying Bonferroni 217 correction (i.e. p<0.00086) were called 'significant pathogens' while the ones which were 218 non-significant after Bonferroni correction (i.e. p<0.05) were called 'potential pathogens'. Isolates that reduced the risk of thallus tip bleaching were all designated as 'protectors'. 219 However, isolates that significantly reduced the risk of thallus tip bleaching after applying 220 221 Bonferroni correction (i.e. p<0.00086) were called 'significant protectors', while the other isolates that also reduced tip bleaching (p<0.05) but were not significant after applying 222 223 Bonferroni correction were designated as 'potential protectors'.

#### (B) Bleaching assay with combined 'protectors' and 'pathogens'

The combined effect of all 'protectors' ('significant' and 'potential' protectors) on the 225 226 virulence of confirmed pathogens ('significant' and 'potential' pathogens) identified in the above experiment was tested in an additional experiment. In order to observe any 227 community effect of these epibacteria, we included all the significant and potential strains 228 229 because bonferroni correction is known to be relatively conservative (Moran 2003). The 230 general design was as described above, but the method of inoculation differed: bacterial cultures were incubated until their OD<sub>610</sub> was between 0.1 and 0.5. Different aliquots of all 231 identified 'protectors' were then pooled so that each culture contributed the same OD<sub>610</sub> to 232 the mixture, which had a final OD<sub>610</sub> of 0.25. Cells in the mixture were separated from the 233 234 medium by centrifugation (10 000 g, 20 min) and resuspended in SSW. A mixture of three

235 pathogens was prepared in an analogous manner. Agarophyton was then inoculated with 10 µl of either (i) all of the 'protectors' (19 bacterial strains, thereof 10 'significant protectors' 236 and 9 'potential protectors') or (ii) all of the pathogens (3 bacterial strains, thereof 1 237 238 'significant pathogen' and 2 'potential pathogens') or (iii) pathogens and protectors together 239 (22 bacterial strains). Final volumes of either protectors or pathogens were brought up to 20  $\mu$ l with SSW, while controls received just 20  $\mu$ l of SSW. This experiment was repeated in one 240 241 fully independent repetition ( $n = 2 \times 6$ ). Numbers of bleached thallus tips relative to all tips 242 were counted and significant differences were identified using Kruskal-Wallis-ANOVA and Dunn's post hoc test. 243

#### 244 Extraction of surface associated metabolites of Agarophyton

To generate surface associated metabolites, Agarophyton individuals (n=5) were collected 245 from the same location as above, i.e. Nordstrand. Surface-associated metabolites 246 247 originating from single Agarophyton specimens were extracted immediately upon collection 248 according to Saha et al., (2016). Briefly, Agarophyton branches were dipped into a solvent mixture of dichloromethane and hexane 1:4 (v/v) for 5 s. This process is benign and does not 249 250 harvest intracellular metabolites (see Saha et al., (2016) for details). The prepared extract (n=5) containing the surface associated metabolites was filtered through GF/A filter paper 251 (Whatman  $\emptyset$  = 15 mm) to remove particles, and the solvent was evaporated under a 252 253 vacuum at 20°C, using a rotary evaporator (Laborota 3000, Heidolph, Germany). The extract 254 was then taken up in acetonitrile in such a way that 1.5  $\mu$ l contained metabolites extracted from an algal surface of 99.64 mm<sup>2</sup>. The extract was used to coat each replicate well with a 255 surface area 99.64 mm<sup>2</sup>. Acetonitrile was then evaporated and metabolites originating from 256 the surface of the alga remained on the surface of the well, allowing us to test at an 257 258 ecologically realistic 1-fold concentration.

# 260 Defence capacity test of *Agarophyton* surface metabolites against pathogens and 261 protectors

Inhibition or reduction of bacterial settlement and attachment represents the first line of 262 defence against microbial challenge (Lane & Kubaneck, 2009). Thus, an antisettlement assay 263 was employed as the most relevant criterion for determining antimicrofouling defence 264 265 because it quantifies both repellent and toxic effects (Wahl, Jensen & Fenical, 1993). The 266 assay was performed according to Saha et al. (2016). Briefly, the assay was conducted in 96well plates that were surface-impregnated with Agarophyton surface extract metabolites at 267 268 a 1-fold natural concentration (Saha et al., 2016) and with solvent residue as a control. In 269 total five extracts – originating from five Agarophyton individuals - were tested and regarded as replicates. Each Agarophyton extract was then subdivided and tested in three 270 271 pseudo replicates against each bacterial isolate to account for the variability in the bacterial 272 settlement rates. Results obtained for pseudo replicates were averaged before statistical 273 analyses were conducted. The tested target strains were chosen based on results from the 274 tip bleaching assay described above. All three pathogens (both the 'significant pathogen' Kordia algicida and the 'potential pathogens' Croceitalea eckloniae, Pseudoalteromonas 275 arctica) were tested in the anti-settlement assays, but due to shortage of surface extracts 276 277 was it not possible to test all nineteen 'protective' strains. Thus, only five 'significant i.e. Ralstonia sp., Shewanella aquimarina, Tenacibaculum skagerrakense, 278 protectors' Alteromonas stellipolaris, Tenacibaculum aestuarii and two 'potential protectors' i.e. 279 Cobetia marina and Nonlabens dokdonensis were tested. 106 µL suspensions of these 280 281 bacterial strains (O.D. 0.6-0.8) precultured in MB liquid medium (as described above) were 282 pipetted into the wells. The bacteria were allowed to settle for 3 h, and the settled cells that could not be removed by rinsing two times with 110μL sterile seawater were stained with
the fluorescent DNA-binding dye Syto 9 (Invitrogen, GmbH). Fluorescence was subsequently
measured (excitation, 477–491 nm; emission, 540 nm) with a plate reader as a proxy for
bacterial settlement in terms of the attached cell density. All tested strains were allowed to
settle on all extracts.

288 The defence strength of Agarophyton surface metabolites is expressed as the 'log effect ratio,' i.e., the logarithm of the fluorescence attributable to the settled bacteria of strain Y in 289 290 the presence of surface metabolites, divided by the fluorescence attributable to the settled bacteria of strain Y in the absence of surface metabolites. A log effect ratio value of 0 (i.e., 291 an equal number of settled bacteria in wells with and without surface metabolites) 292 indicated that the tested surface metabolites had no effect on settlement, whereas a 293 294 negative log effect ratio value indicated a deterrent effect, and a positive log effect ratio value indicated an attractant effect. Thus, a log effect ratio of -1 represents a 10-fold 295 296 reduction, whereas a value of +1 represents a 10-fold enhancement of bacterial settlement 297 caused by surface metabolites.

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299	Defence strength = log (b	acterial settlement in presence of	Agarophyton surface metabolites)
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(bacterial settlement in absence of Agarophyton surface metabolites)

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302 Results

303 Agarophyton tip bleaching assay with epibacterial strains

304 (A) Bleaching assay with single isolates

Out of 58 tested bacterial isolates *Kordia algicida* was found to significantly increase the risk 305 of tip bleaching (Table 1; Fig. 1, p<0.00086), compared to control treatments without 306 inoculation of bacteria and was a 'significant pathogen' after Bonferroni correction. Two 307 additional isolates (Pseudoalteromonas arctica and Croceitalea eckloniae) had the same 308 309 effect but were not significantly pathogenic after Bonferroni correction (Table 1; Fig. 1, p<0.05) and were thus 'potential pathogens'. Out of the remaining 55 isolates, 9 were found 310 311 to significantly reduce the risk of tip bleaching (Table 1; Fig. 1, p<0.00086) and were grouped 312 under 'significant protectors'. 10 others had the same effect, although they were not significantly protective after Bonferroni correction (Table 1; Fig. 1, p<0.05) and were called 313 314 'potential protectors'. The remaining 36 isolates were found to be neutral, neither inducing 315 nor preventing bleaching (see Table S2; Fig. S2 in Supporting Information).

Similar numbers of microbiota that originated from native and non-native populations (30 and 28, respectively, see Table 1 and Table S2) of *Agarophyton* were tested in our bleaching assay and double numbers of protective microbiota were detected from the non-native range (3 from the native range and 6 from the non-native range, respectively, Table 1).

### 320 (B) Bleaching assay with combined 'protectors' and 'pathogens'

When all three isolates (one 'significant pathogen' and two 'potential pathogen') that induced bleaching individually at least with p < 0.05 were combined, a significant increase in bleaching relative to the control was again observed (Fig. 2, p<0.05). However, combined application of these three 'pathogens' and the nineteen 'protective' isolates that prevented bleaching individually at least with p < 0.05 resulted in no such increase (Fig. 2, p<0.05). No bleaching was observed when all 19 'protectors' and no 'pathogens' were inoculated.

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#### 328 Defence capacity test of Agarophyton against pathogens and protectors

The effect of *Agarophyton* surface associated metabolites on bacterial settlement differed significantly between the two groups of bacteria, i.e. 'protectors' and 'inducers' (Fig. 3, Welch-corrected t-test, p < 0.0001). While the surface associated metabolites significantly increased the settlement of 'protectors', the settlement of the bleaching 'inducers' was significantly reduced by the surface associated metabolites.

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#### 335 Discussion

336 The data presented here demonstrate for the first time that aquatic macrophytes can use surface associated chemicals not only to directly reduce settlement of pathogenic bacteria, 337 but also to recruit bacterial strains that provide protection from such pathogens. The 338 epimicrobiome of Agarophyton contains a component that protects the alga from 339 340 pathogens in a similar way as earlier demonstrated for two other seaweeds (G. conferta (Weinberger et al., 1997) and D. pulchra (Longford et al., 2019)), and the settlement of such 341 342 protective bacteria on the surface of Agarophyton is not random. Surface associated metabolites from the Agarophyton holobiont significantly deterred three strains that were 343 significantly and potentially pathogenic, while the metabolites had a probiotic effect 344 345 towards seven significantly and potentially protective strains that were tested. This confirms the surface chemistry of Agarophyton has a similarly strong selective effect on bacterial 346 colonization as in Fucus vesiculosus (Lachnit et al., 2010) or Delisea pulchra (Longford et al., 347 348 2019). Moreover, it demonstrates for the first time that this selection is not only targeted to 349 exclude pathogens, but also targeted to attract protectors. Together with Lachnit et al., (2010) and Kessler et al., (2018) our data strongly support the concept of chemically 350 mediated recruitment of microbes and not the 'neutral' hypothesis, according to which the 351 microbial community structure is determined randomly. Our data clearly support the 352

targeted deterrence hypothesis, as settlement of detrimental bacteria was chemically suppressed. On the other hand, we cannot reject the 'niche' model, as multiple microbiota were attracted by *Agarophyton* and possibly able to make use of resources provided by it. Also, the 'lottery' hypothesis cannot be currently rejected, since the capacity of attracted microbiota to coexist and share host resources is unknown.

358 Only 5% of the marine bacterial strains are cultivable (Haglund et al., 2002) and to date no alternative technique has been developed to separate selected microbial components from 359 360 natural microbial communities and to test them in infection assays. Thus, only a small fraction of all bacteria that are associated with the surface of Agarophyton could be isolated 361 and tested in our study. One representative out of 58 tested bacterial species, Kordia 362 algicida, was significantly capable to induce the tip bleaching symptom in Agarophyton. K. 363 364 algicida is already known to be detrimental to other organisms. It can kill diatom blooms in a protease mediated molecular interaction (Paul & Pohnert, 2011) and a similar mechanism 365 366 cannot be excluded in the present case. Bleaching is often correlated with microbial cell wall matrix degrading activity (Weinberger et al., 1994, 1997), but this was not the case in the 367 present study, as *Kordia* is incapable of agar degradation. 368

Two other isolates – which were also not agar degraders - also exhibited the potential to induce bleaching symptoms in *Agarophton*, which strongly suggests that this capacity is not unique. Interestingly, all three detrimental isolates originated from virtually healthy host specimens. Thus, a relevant fraction of *Agarophyton*'s surface microbiome is obviously composed of opportunistic pathogens that can induce bleaching symptoms under certain conditions, similar as in several red seaweeds belonging to other species (Case *et al.*, 2011; Weinberger *et al.*, 1994; Weinberger *et al.*, 1997). Given that three out of 58 culturable

strains were significant or potential pathogens this fraction can be estimated to include
approximately 5 % of the microbiome. However, this percentage calculation is based on the
culturable proportion which is just a representative sample of the whole microbiome.

Of the remaining strains, 19 (9 'significant protectors' and 10 'potential protectors') could 379 380 reduce the risk of 'spontaneous' bleaching in thalli that were not intentionally inoculated 381 with pathogenic bacteria (Fig. 1). All the specimens of Agarophyton tested in our bleaching induction assays were subjected to a pretreatment with two antibiotics that inhibited 382 383 bacterial cell wall synthesis, with the dual goal to remove opportunistic pathogens and to 384 disturb and weaken any protective components of the algal microbiome. The circumstance that bleaching occurred 'spontaneously' at a low rate but could be prevented by an 385 important percentage of all tested isolates suggests that some opportunistic pathogens 386 387 survived the treatment with antibiotics but could then not become virulent when protective bacteria were inoculated - similar as previously reported for bacteria that had been isolated 388 389 from Gracilaria 'conferta' and prevented thallus tip bleaching in this alga (Weinberger et al., 390 1997). The protective effect of various isolates on *Agarophyton* was further confirmed when all the 19 protective strains (nine 'significantly protective' and ten 'potentially protective') 391 392 were tested together in combination with the 3 pathogenic strains (one 'significant 393 pathogen' and two 'potential pathogens') and a bleaching reduction was still documented. Alltogether, our observations strongly hint at the presence of protective epibacteria on the 394 395 surface of *Agarophyton*. They could (again estimated from the number of identified isolates 396 in our tested strain collection) comprise at least 15% of all taxa present in this microbiome. The presence of such beneficial bacteria has been previously demonstrated not only for 397 other Gracilarioids (Weinberger et al., 1997), but also for Delisea pulchra (Longford et al., 398 2019), corals (Rosenberg et al., 2007) and other seaweeds like the brown alga Fucus 399

*vesiculosus,* in which surface associated bacteria were found to inhibit the settlement of
macrofoulers (Nasrolahi *et al.,* 2012).

402 The biofilms on seaweed surfaces represent highly competitive environments, with microbes competing for refuge, nutrients and substratum, and interspecific antagonistic 403 404 effects of bacterial strains are not rare. For example, such effects have been previously 405 demonstrated for the brown alga Saccharina latissima (Wiese et al., 2009), the red alga Delisea pulchra and the green alga Ulva australis (Penesyan et al., 2009). Release of 406 407 inhibitory components like antibiotics (Wiese et al., 2009) and/or quorum sensing inhibitors 408 (Romero et al., 2010) has been observed and could also explain the 'protective' effect observed by us. Interestingly, one of the significant protective strains, *Pseudoalteromonas* 409 410 *piscicida*, belongs to a genus which comprises several species that are known to produce 411 antibacterial products to outcompete other bacteria for space and nutrients (Holmström & Kjelleberg, 1999). Also Pseudoalteromonas piscicida has been recently demonstrated to 412 413 inhibit and/or kill competing bacteria - including several marine pathogens, such as Vibrio 414 vulnificus, Vibrio parahaemolyticus, Vibrio cholerae, Photobacterium damselae, and Shewanella algae - through secretion of antimicrobial substances and the direct transfer of 415 416 digestive vesicles to competing bacteria (Richards et al., 2017). On the other hand, 417 Shewanella aquimarina exhibited a strong protective effect against Aqarophyton tip bleaching in the current study and the same was observed with two other potentially 418 419 protective species of the genus Shewanella, i.e. S. marisflavi and S. loihica. These 420 observations contrast with the findings that S. marisflavi is a pathogen of sea cucumbers (Li et al., 2010) and other bacteria of the genus are pathogenic towards humans. The 421 422 mechanisms behind the protective effects on Agarophyton deserve further investigation. 423 The exact (additive or synergistic) contributions of the active epibacterial players in the

424 cross-infection experiment with all 19 'protectors' combined with 3 'pathogens' are not425 known yet.

426 Beneficial roles of certain components in the bacterial communities around the rhizosphere 427 of terrestrial plants are well known. They can not only facilitate nutrient acquisition, but also 428 support plant growth under biotic and abiotic plant stress (Lareen et al., 2016; Mendes et 429 al., 2013). Seaweed-associated bacteria may also facilitate nutrient acquisition and provide essential vitamins and growth factors (Wahl et al., 2012), and – as confirmed in the present 430 431 study – mediate biotic stress. However, while we have started to understand these benefits and to gather evidence of a selective recruitment of bacteria both in terrestrial (Lebeis et al., 432 2015) and aquatic environments, empirical links between this selective recruitment of 433 communities and a health benefit for the host are still very rare in the aquatic realm. Kessler 434 435 et al., 2018 recently demonstrated for the marine macroalga Ulva mutabilis (Chlorophyta) a mediating role of the algal osmolyte DMSP (dimethylsulfoniopropionate) in the attraction of 436 437 the beneficial bacterium Roseovarius sp. MS2, responsible for release of morphogenetic compounds that ensure proper algal morphogenesis. In absence of these morphogenetic 438 compounds under axenic conditions, Ulva mutabilis develops into callus-like colonies 439 440 consisting of undifferentiated cells and abnormal cell walls. While microbial 'gardening' via use of chemicals has been documented in terms of growth and development of seaweeds 441 (Kessler et al., 2018), our study demonstrates for the first time such a link between the 442 443 disease resistance capacity of a seaweed and beneficial selective gardening of 'protective' bacteria based upon surface chemistry. 444

445 Metabolites present on the surface of seaweeds or in the rhizosphere are a cocktail of 446 metabolites originating both from the algal or plant host and from associated surface

microbiota. Such surface associated metabolites from the algal holobiont are also known to 447 function as a defence against fouling by microfoulers (e.g. bacteria, diatoms) and 448 macrofoulers (e.g. barnacle larvae, mussels) (reviewed by Da Gama, 2014; Saha et al., 2017). 449 Also epibacteria from seaweeds are well known to have inhibitory activities against other 450 451 fouling organisms (Singh & Reddy, 2014). Thus, it is possible that beneficial bacteria 452 recruited by Agarophyton will not only act as a defence against pathogens but also against 453 other foulers, like filamentous algae. Using a transcriptomic approach, de Oliviera et al., 454 2012 demonstrated that the red seaweed host Laurencia dendriodea (rather the surface associated bacteria) is involved in the biosynthesis of terpenoids (chemical defence 455 456 compounds against bacterial colonization and infection) through the mevalonate independent pathway. For the Agarophyton holobiont, we do not know yet the identity of 457 surface associated bioactive compounds. Thus, it was not possible for us to distinguish the 458 459 relative contributions of surface metabolites originating from the algal host Agarophyton 460 and from surface associated microbiota. Mutants of Arabidopsis thaliana with suppressed salicylic acid signalling pathways formed abnormal root microbiomes when compared to the 461 wild plants (Lebeis et al., 2015), which could suggest that the role of the host was more 462 important in this specific case. The contribution of seaweed microbiome metabolites 463 depends on the community composition, abundance and metabolic activity (Wahl et al., 464 465 2010) and may be expected to be more variable than that of the host. Selective effects observed with surface associated metabolites coming from different algal individuals varied 466 relatively little in our study, which could suggest that metabolites generated by the host 467 have more importance. However, our knowledge of the species-species interactions of 468 469 cultivatable and non-cultivatible taxa associated with Agarophyton or other plants is 470 rudimentary at best. The involvement of multiple protective microorganisms in our and

471 several other cases (see above) strongly supports the view that the traditional conceptual
472 model emphasizing direct interactions of hosts and single microbes needs to be expanded to
473 a holobiont concept if seaweed- or plant-microbe interactions are to be understood (Egan *et*474 *al.*, 2013).

In conclusion, our study demonstrates for the first-time selective chemical 'gardening' of 475 476 protective epibacterial strains by a seaweed holobiont with regard to disease resistance capacity. The combined effect of metabolites generated by the host alga and the protection 477 478 offered by associated microbial partners determines the virulence of harmful opportunistic 479 bacterial pathogens. A major component of the epibacterial community appears capable of contributing to this protection against co-occurring pathogens, which suggests that 480 microbiota of very different taxonomic groups may provide the holobiont with the same 481 482 ecological function, which could be pivotal for the establishment of Agarophyton in new environments. Thus, absence of protective microbiota in new environments might not be a 483 484 factor limiting the invasion success of Agarophyton.

As known for other seaweeds like the brown alga *Fucus vesiculosus*, bioactive surface metabolites often act in synergism or additively and/or antagonistically, producing an overall defensive or prebiotic effect on bacterial recruitment (Saha *et al.*, 2011; Saha *et al.*, 2012). The identification of metabolites responsible for such chemical 'gardening' effects via classical bioassay guided fractionation techniques in the near future may allow us to manipulate algal thallus microbiomes to enhance seaweed health, prevent bleaching diseases and ensure production and sustainability in *Agarophyton* aquaculture.

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497	Author contributions
498	M.S. isolated and identified the bacterial isolates, designed and performed the anti-
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500	F.W. analysed the data. M.S. wrote the paper and F.W. contributed to the editing.
501	Author Declaration
502	The authors declare no conflict of interest. Data underlying this publication are freely
503	accessible and can be downloaded from the DRYAD data repository (Provisional DOI:
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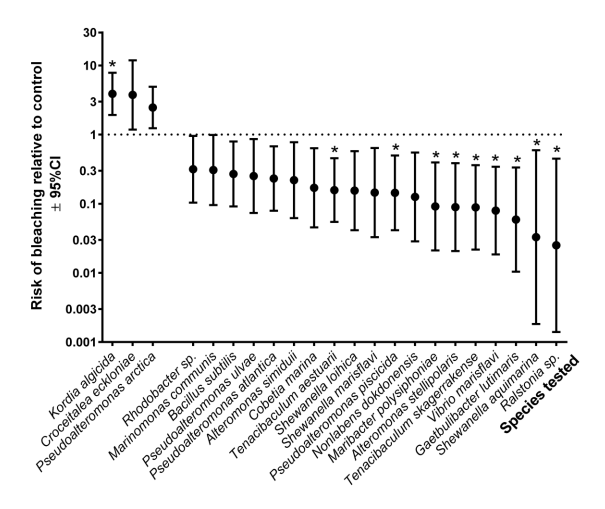
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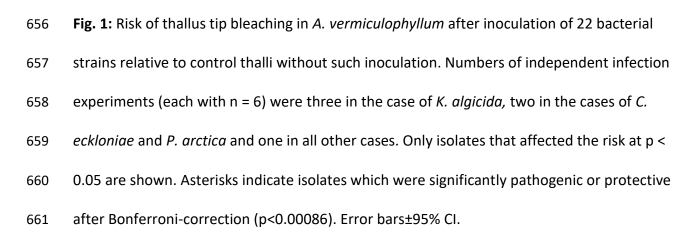
#### 646 Tables

Table 1. Epibacterial strains [isolated from *Agarophyton vermiculophyllum* (GV) populations]
that were tested in tip bleaching assays. IR=bacteria isolated from the invasive populations
of GV; NR=bacteria isolated from the native populations of GV. Significantly protective
strains are bacteria that were still protective after Bonferroni correction. Asterisks indicate
strains that were tested in anti-settlement assays with surface associated compounds of GV.

Isolate code, Range of	Closest match on RDF	Designation	
isolation			
G-NY6, IR	Ralstonia sp.*	Significantly Protective	
G-JI1, NR	Shewanella aquimarina*	Significantly Protective	
G-NORD3, IR	Gaetbulibacter lutimaris	Significantly Protective	
G-G2, NR	Vibrio marisflavi	Significantly Protective	
G-MAN7, IR	Tenacibaculum skagerrakense*	Significantly Protective	
G-HO9, IR	Alteromonas stellipolaris*	Significantly Protective	
G-ODO3, NR	Maribacter polysiphoniae	Significantly Protective	
G-FALK1, IR	Nonlabens dokdonensis*	Potentially Protective	
G-NORD11, IR	Pseudoalteromonas piscicida	Significantly Protective	
G-G4, NR	Shewanella marisflavi	Potentially Protective	
G-DA3, NR	Shewanella loihica	Potentially Protective	
G-NORD6, IR	Tenacibaculum aestuarii*	Significantly Protective	
G-JI4, NR	Cobetia marina*	Potentially Protective	
G-NY1, IR	Alteromonas simiduii	Potentially Protective	

G-DA5, NR	Pseudoalteromonas atlantica	Potentially Protective
G-FALK2, IR	Pseudoalteromonas ulvae	Potentially Protective
G-MAN5, IR	Bacillus amyloliquefaciens	Potentially Protective
G-JI5, NR	Marinomonas communis	Potentially Protective
G-HO8, IR	Rhodobacter sp.	Potentially Protective
G-MAN6, IR	Pseudoalteromonas arctica*	Potentially Pathogenic
G-NORD9, IR	Croceitalea eckloniae*	Potentially Pathogenic
G-MAN4, IR	Kordia algicida*	Significantly Pathogenic





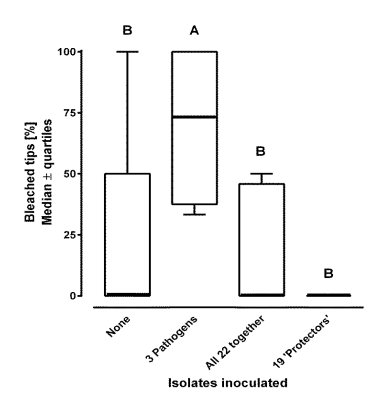
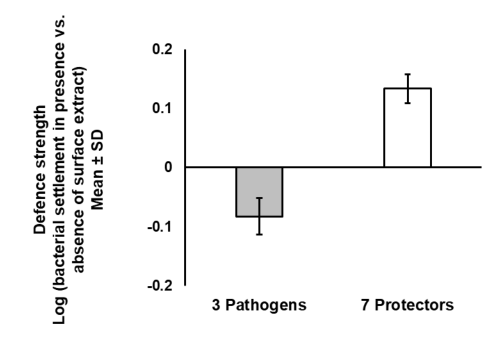




Fig. 2: Relative amounts of bleached thallus tips in *A. vermiculophyllum* after inoculation
with 3 pathogenic bacterial isolates with 19 protective bacterial isolates, with all 22 isolates
together and in controls without any inoculation. Different letters indicate treatments that
are significantly different (n = 12; Kruskal-Wallis-ANOVA (p < 0.0001) and Dunn's post hoc</li>
test (p < 0.05)). Pathogenic strains include both 'significant pathogens' and 'potential</li>
pathogens'. Protective strains include both 'significant protectors' and 'potential
protectors'. Median ± quartiles.



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**Fig. 3:** Mean anti-settlement activity of *Agarophyton* surface metabolites against three pathogenic (one 'significant pathogen' and two 'potential pathogen') and seven protective (five 'significant protector' and two 'potential protector') strains. Error bars  $\pm$  SD (n=5); Welch-corrected t-test, p < 0.0001.