# 1 Anti-epiphyte defenses in the red seaweed Gracilaria vermiculophylla: non-

# 2 native algae are better defended than their native conspecifics

- 3 Running headline: Epiphytes on non-native *Gracilaria vermiculophylla*
- 4 Shasha Wang<sup>1\*¶</sup>, Gaoge Wang<sup>2¶</sup>, Florian Weinberger<sup>1</sup>, Dapeng Bian<sup>3</sup>, Masahiro Nakaoka<sup>4</sup> and
- 5 Mark Lenz<sup>1</sup>
- 6 <sup>1</sup> Department of Benthic Ecology, GEOMAR Helmholtz-Zentrum für Ozeanforschung Kiel,
- 7 24105 Kiel, Germany
- <sup>8</sup> <sup>2</sup> College of Marine Life Sciences, Ocean University of China, Qingdao 266003, China
- <sup>3</sup> Xunshan Group Co., Ltd, Rongcheng 264316, China
- <sup>4</sup> Akkeshi Marine Station, Field Science Center for Northern Biosphere, Hokkaido University,
- 11 Akkeshi, Hokkaido 088-1113, Japan

## 12

- 13 <sup>\*</sup>Corresponding author: Shasha Wang
- 14 E-mail: <u>swang@geomar.de</u>
- 15 Fax: +49 431 600 1671

## 16

<sup>17</sup> <sup>¶</sup>These authors are both first authors and contributed equally to this work.

## 19 Summary

1. Epibiosis in the marine environment is a stressor that may determine invasion success in introduced species. Previous comparisons showed resistance to epibionts can be higher in non-native than in resident seaweed species, but we do not know whether it is an intrinsic trait of the non-natives or it has been acquired during the invasion process. To elucidate this question, a comparison between native and non-native populations of the same species is needed.

25 2. Resistance against two groups of epiphytes was assessed in living thalli and in artificial 26 substrata coated with surface extracts, both gained from four Asian (native) and four European (non-native) populations of the red alga Gracilaria vermiculophylla. Two diatom species and 27 two filamentous macroalgae were used as micro- and macro-epiphytes, and one of each type was 28 collected in Asia, while the other came from Europe. Laboratory assays were done in both 29 30 distributional ranges of G. vermiculophylla and in different seasons. We used G. vermiculophylla 31 from four populations in each range and used a fully-crossed design with the factors a) 'Origin of 32 Gracilaria', b) 'Origin of epiphytes', c) 'Season' and d) 'Solvent used for extraction'.

33 3. Both groups of epiphytes, regardless of their origin, attached less to living thalli and to
34 surface extracts from non-native *G. vermiculophylla*. Fewer diatoms attached to hexane-based
35 extracts, while fewer *Ceramium* filaments settled on extracts gained with dichloromethane.

4. Synthesis. Our results show for the first time that non-native individuals of a marine organism are better defended against epiphytes than native conspecifics. Furthermore, we found evidence that at least a part of the defense is based on extractable secondary metabolites. We discuss several mechanisms that could explain the increased resistance to epiphytes in non-native individuals, including the release from enemies in the non-native range, which could lead to an

41	increase in algal performance during the invasion process. We suggest that an enhanced defense
42	against epiphytes after introduction is one reason for G. vermiculophylla's invasion success. Our
43	observation may also apply to other basibiont-epibiont and host-enemy systems, including plant-
44	plant, plant-animal and animal-animal interactions, in aquatic environments and could be a key
45	feature of bioinvasions.
46	
47	
48	
49	
50	
51	
52	
53	
54	
55	

56	Key-words: anti-fouling, biological invasions, chemical defense, Enemy Release Hypothesis,
57	epiphytes, fouling, Gracilaria vermiculophylla, invasion ecology, non-native seaweeds
58	
59	
60	
61	
62	
63	
64	
65	
66	
67	
מט	

## 69 Introduction

Biological invasions are an important component of global change (Mack et al. 2000; Ricciardi 70 2007) and can cause severe ecological or economic problems by altering local biodiversity and 71 72 affecting the services of ecosystems (Pimentel, Zuniga & Morrison 2005; Williams & Smith 73 2007; Vilà et al. 2011; Newton et al. 2013; Paini et al. 2016). Therefore, bioinvasions in the 74 aquatic and terrestrial realm are receiving attention by scientists, authorities and environmental managers worldwide (Torchin & Mitchell 2004; Olenin et al. 2014). This growing awareness is 75 needful, since the number of successful invasions is rising rapidly. In the marine environment, 76 for instance, it increases exponentially and about 20% of the invasions are due to the spread of 77 78 macroalgae (Schaffelke, Smith & Hewitt 2006).

To manage invasive species and to alleviate their negative effects, it is necessary to understand 79 the mechanisms that determine the success or failure of invasions: Why do certain species invade 80 new habitats successfully while others fail? Which factors allow the establishment and spread of 81 introduced species? Currently, several hypotheses suggest explanatory concepts for these 82 83 questions. Among them, the Enemy Release Hypothesis (ERH) is one of the most well-known (Keane & Crawley 2002). It states that when species are introduced into new habitats they are 84 85 confronted with an abiotic and biotic environment that can be substantially different from the one 86 they adapted to over evolutionary time scales. If co-evolved antagonists are absent in the new habitat and resident predators or parasites cannot recognise the newly introduced species as a 87 resource, a release of the introduced species results from enemy control (Vermeij et al. 2009; 88 Cacabelos et al. 2010). Furthermore, the Evolution of Increased Competitive Ability Hypothesis 89 (EICA), which is an extension of the ERH, argues that non-native species that are released from 90 their native enemies do not need to defend at all or, at least, less than in their home range. This 91

92 release would allow them to invest more energy into growth, reproduction or into tolerating 93 environmental stress (Hierro, Maron & Callaway 2005; Joshi & Vrieling 2005; Lenz *et al.* 2011), 94 what, in turn, should make them competitively superior to native species (Blossey & Nötzold 95 1995; Müller-Schärer, Schaffner & Steinger 2004). Finally, the "Novel Weapons"-hypothesis by 96 Callaway and Ridenour (2004) suggests that non-native species should be less susceptible to 97 resident enemies than native ones, because they possess biochemical defense mechanisms to 98 which native antagonists have not adapted to so far.

Most if not all marine organisms are prone to colonization by sessile life forms (epibionts), 99 100 including bacteria, protists, microalgae, macroalgae and invertebrates. This phenomenon is known as epibiosis and it can severely impair the performance of the host organisms (basibionts) 101 (Wahl 2008; Thomsen et al. 2012). It has been reported that epibionts can affect the growth and 102 survival of macroalgal hosts by a) limiting the uptake of oxygen, carbon dioxide and nutrients 103 through the thallus surface, b) reducing the amount of light available for photosynthesis, c) 104 physically inhibiting sporulation, d) decreasing thallus flexibility, and/or e) increasing the 105 106 palatability of the thallus (Wahl, Hay & Enderlein 1997; Hemmi et al. 2005). Macroalgae therefore need physical or chemical defenses to minimize colonization of their surfaces. 107

All the hypotheses described above should also hold true for host-epibiont interactions, but have so far been widely neglected as a factor determining invasion success. So far, studies that tried to elucidate the relevance of host-enemy interactions for biological invasions mainly focused on non-native spermatophytes and their antagonists as well as their competitors in terrestrial systems (Keane & Crawley 2002; Verhoeven *et al.* 2009), while less research has been done on marine species (Wikström *et al.* 2006). Furthermore, most studies on non-native species resident enemy interactions in marine systems considered herbivory (Wikström *et al.* 2006; 115 Forslund, Wikström & Pavia 2010; Engelen et al. 2011; Hammann et al. 2013), while little is 116 known about the interactions between non-native basibionts and resident epibionts (Strong, Maggs & Johnson 2009; Baer & Stengel 2014). A comparison between the non-native brown 117 alga Fucus evanescens and the native Fucus vesiculosus in Swedish waters revealed that 118 resistance to epibiosis was higher in the non-native species (Wikström & Kautsky 2004; 119 Wikström & Pavia 2004). However, we lack knowledge whether this difference is based on 120 species-specific traits or whether resistance to epibiosis can be gained during the invasion 121 process. This could, for instance, be due to the release from other enemies that would allow the 122 123 allocation of energy to defenses against epibionts.

To elucidate whether non-native seaweed species are better defended against epibionts than their 124 native conspecifics, we compared the susceptibility to epiphytism by microalgae and macroalgae 125 126 between native and non-native populations of the red macroalga Gracilaria vermiculophylla. This perennial seaweed originates from the Northwest Pacific but during the last four decades it 127 invaded many coastal habitats in the eastern Pacific (Bellorin, Oliveira & Oliveira 2004), the 128 129 eastern Atlantic (Rueness 2005) and the western Atlantic (Freshwater et al. 2006; Thomsen et al. 2006) and the Mediterranean Sea (Sfriso et al. 2012). Gracilaria vermiculophylla has proven to 130 131 be a particularly suitable marine model organism for the testing of theoretical concepts that predict the causes or consequences of biological invasions. It has, for example, been 132 demonstrated that non-native populations of this species have a much lower genetic diversity 133 than native populations (Kim, Weinberger & Boo 2010), while, nonetheless, the former proved 134 to be more resistant towards herbivory (Weinberger et al. 2008; Rempt et al. 2012; Hammann et 135 al. 2013; Hammann et al. 2016) and heat stress (Hammann 2014). These findings shed light on 136

potential reasons for the invasion success of *G. vermiculophylla* and stimulated curiosity abouthow this seaweed interacts with epibionts in its non-native range.

Putative differences in the resistance against micro-epiphytes between native and non-native populations of *G. vermiculophylla* have been studied for bacteria (Saha *et al.* 2016), while we are not aware of studies that were done with eukaryote micro-epiphytes and macro-epiphytes. We conducted common garden experiments, i.e. individuals of *G. vermiculophylla* from different geographic ranges were transplanted into a common environment, with *G. vermiculophylla* from Asia and Europe and with micro- and macro-epiphytes coming from the macroalga's native and non-native range.

## 160 Materials and methods

#### 161 Collection of *Gracilaria vermiculophylla* and of epiphytes

Algal individuals were collected from four native and four non-native populations (Table 1). 162 163 Sampling was conducted three times: from May to June and again from August to September 164 2014 as well as from June to July 2015. Laboratory experiments were conducted in June (summer) and in October (autumn) 2014 at the Helmholtz Centre for Ocean Research at Kiel, 165 166 Germany, and in September (autumn) 2014 at the Xunshan Group Co., Ltd, Rongcheng, China, as well as in July (summer) 2015 at the Akkeshi Marine Station, Akkeshi, Japan (Table 2). Prior 167 to experiments, living algal specimens from all sampling sites were transferred to the respective 168 169 laboratory and kept under laboratory conditions for at least one week to allow them to recover from the transport (see Appendix S1 in Supporting Information for details). 170

Two pennate diatom species of the genus *Stauroneis* were isolated from individuals of *G. vermiculophylla* that were collected in Rongcheng and in the Kiel Fjord, respectively.
Individuals of *Ceramium tenerrimum* were collected in Rongcheng, while specimens of *Ceramium virgatum* stem from Kiel Fjord (see Appendix S2 for details).

### 175 Algal surface extraction

Before the extraction procedure, adherent water was removed from the algal thalli in a centrifuge (Eppendorf 5810 R) with 200 rpm for 30 s. Surface associated metabolites were extracted from 80 g algal wet mass using the 'dipping technique' (de Nys, Dworjanyn & Steinberg 1998; Nylund *et al.* 2007), by dipping the algal individuals into a stirred mixture of dichloromethane (DCM) and hexane 1:4 (v/v) for 5 s. This procedure was benign and was chosen after different mixtures of solvents and dipping times were tested with regard to their effects on the survival of 182 epidermal cells which was verified with Evan's blue (Figure S1). This was done to make sure that any damaging of cell walls, which could have led to the leaching of non-surface compounds, 183 was avoided. The resulting solution was immediately filtered through a paper filter (Macherey 184 Nagel, 185 mm in diameter) to remove particles and the solvents were then evaporated under 185 vacuum at 30 °C. The residue was re-dissolved in hexane to exclude non-polar compounds and 186 this step was repeated until the hexane appeared colourless. The residue that remained after this 187 first extraction step was then re-dissolved in DCM to extract existing polar components. Finally, 188 4 ml of both extracts were collected and stored at -20 °C. 189

#### 190 Extracted surface area

To identify the extracted surface area, the relationship between algal surface area and algal wet 191 weight was determined. Ten algal fragments, taken haphazardly across all populations, were 192 carefully dried with paper and then scanned and weighed. The imaging software Image J 193 (National Institute of Health, Bethesda, Maryland, USA) was used to analyze the surface area of 194 195 each fragment. The algal thallus was viewed as a cylinder, so the projection area = thallus diameter  $\times$  thallus length. The surface would then be =  $\pi \times$  thallus diameter  $\times$  thallus length =  $\pi \times$ 196 projection area. We identified the average surface area per g algal material across all ten 197 fragments as  $46.06 \pm 2.8 \text{ cm}^2 \text{g}^{-1}$  (mean  $\pm$  SD). The total extracted surface area was 80 g \* 46.06 198  $\pm 2.8 \text{ cm}^2 \text{g}^{-1} = 3684.8 \pm 224 \text{ cm}^2$ 199

## 200 Defense capacity against diatoms in living Gracilaria vermiculophylla thalli

We combined living thalli of *G. vermiculophylla* from both ranges (native and non-native) with diatoms from Kiel as well as Rongcheng, respectively. For each basibiont-epibiont combination, six fragments of 2 cm were cut from six randomly chosen algal individuals (n = 6) and were 204 placed separately into the wells of a 6-well plate (Standard, Sarstedt AG & Co.) together with 3 205 ml of a homogenized diatom suspension per well. The covered plate was then incubated for 3 h. A pilot study was done with different incubation periods (3 h, 5 h and 6 h), but diatom 206 207 attachment did not increase any further after 3 h (Table S1). During incubation the plates were placed on a rotary shaker (100 rpm) that served to gently homogenize the suspension. 208 209 Afterwards, each algal fragment was rinsed with 3 ml of sterile seawater. Algal pieces were then transferred to tubes containing 50 ml of sterile seawater and all attached diatoms were extricated 210 by shaking the tube with a vortex shaker for 3 min. The shaken-off diatoms were collected on 211 212 polycarbonate filters (0.2 µm pore size, 25 mm in diameter), which were then inspected under a fluorescence microscope and photographed. The photos were later used to assess the number of 213 diatoms per algal fragment. The tested algal area was determined by scanning the fragments 214 afterwards and quantifying their surface area with Image J. 215

#### 216 Defense capacity against *Ceramium* sp. in living *Gracilaria vermiculophylla* thalli

217 The majority of studies about the attachment of epiphytes or epizoans to hosts focused on colonization by microscopic propagules and spores, which are the mobile stages of the otherwise 218 219 sessile epibionts. However, vegetative thalli of *Ceramium* sp. can directly attach or penetrate into 220 the host by the formation of hapteria (Leonardi et al. 2006; Lion et al. 2006; Michetti et al. 221 2016). We therefore conducted assays with C. tenerrimum and C. virgatum with filaments of these algae and organized them in the same way as the diatom trials described in the previous 222 paragraph. For this, ten algal individuals per population of G. vermiculophylla were used (i.e. n = 223 224 10 per basibiont-epibiont combination). From each of these ten replicates we cut a fragment of 2 225 cm, while a *Ceramium* filament of the same length was then bound to *G. vermiculophylla* using colored paper clips. This was done to shorten the distance between the fragments and by this to 226

increase the likelihood of attachment. These pairs were put into Petri dishes containing 30 ml of
the modified culture medium of Provasoli's enriched seawater (PES) (Bold & Wynne 1978). The
covered Petri dishes were incubated for two weeks and attachment rates were quantified after
this time.

#### 231 Chemical defense capacity against diatoms in *Gracilaria vermiculophylla* surface extracts

232 These assays were organized in the same way as the ones with living thalli. For the assays with 233 extracts we used an extract concentration that was five times higher than the natural surface concentration. This was done to compensate for the possible degradation and incomplete 234 extraction of active compounds. In one cylindrical well of a 96-well plate (flat bottom, Greiner 235 bio-one), 100 µl of both, DCM and hexane, cover a total surface area of 94 mm<sup>2</sup>. Thus, 5.1 µl of 236 237 surface extracts and 94.9 µl of pure solvent were then pipetted into each well to cover the aspired wall area. Wells loaded with pure DCM and hexane were later used as controls. Solvents were 238 then evaporated overnight in a freeze-dryer. After this, 100 µl of the homogenized diatom 239 240 suspension were transferred into the wells. Four wells of each experimental group received extracts and diatoms, while four wells received extracts only to check for extract background 241 fluorescence. Control wells were treated in the same way. Afterwards, the covered 96-well plate 242 was incubated for 3 h and then each well was rinsed with 200 µl of sterile seawater. Finally, 243 244 fluorescence intensity per well was measured and the number of diatoms per well was calculated from fluorescence intensity by using the linear function that was established in a pilot study 245 (Appendix S3, Figure S2). 246

247 Chemical defense capacity against *Ceramium* sp. in *Gracilaria vermiculophylla* surface
248 extracts

249 These assays were organized in the same way as the ones with living thalli. In a 6-well plate, 120 250 µl of solvent can cover the total surface area of the bottom of one well. A paper filter (Carl Roth, 251 3.5 cm in diameter) was put into each well to avoid erosion by solvents. For applying a fivefold 252 natural surface concentration, 52  $\mu$ l of surface extracts and 68  $\mu$ l of pure solvent were then pipetted into each well, while we had five wells per population. Some wells received pure DCM 253 254 or hexane and served as controls. The solvent was then evaporated overnight in a freeze-dryer. After that, 5 ml of PES medium and ten Ceramium filaments (1 cm) were transferred to each 255 well. The covered 6-well plate was then incubated for two weeks. Afterwards, the proportion of 256 257 *Ceramium* filaments that attached to the paper filter was quantified.

#### 258 Statistical analyses

259 All statistical and graphical analyses were done using the free statistical computing software R (R Development Core Team 2014). We used mixed effect-modelling to analyze the data from 260 our multifactorial experimental approach. In case of the assays with living thalli, our 261 262 experimental design included three fixed factors: 1) 'Origin' with the levels 'Native' and 'Nonnative' (i.e. origin of Gracilaria), 2) 'Diatom' / 'Ceramium' with the levels 'China' and 263 'Germany' (i.e. origin of epiphytes), and 3) 'Season' with the levels 'Summer' and 'Autumn' 264 (i.e. the time of the experiment). In the assays with surface extracts we had one more fixed 265 factor: 'Solvent' with the levels 'DCM' and 'hexane'. In all analyses, the algal sampling sites 266 were included as a random factor, while the two types of epiphytes (diatoms/Ceramium) were 267 analyzed separately. To achieve homogeneity of variances and normality of errors, data from the 268 assays with diatoms and living thalli were square root transformed and data from the assays with 269 270 diatoms and surface extracts were log-transformed. However, homogeneity of variances could not be achieved for all factors. We therefore included weights for 'Season' and 'Diatom' to 271

account for the differences in the variance structures between their factor levels in the modelling. For this we used the varIdent function of the nlme package in R (Zuur et al. 2009). Test assumptions were checked graphically with residual plots (Zuur, Ieno & Elphick 2010). In addition, a mixed effect-modelling, with the factors 'Origin', 'Diatom' / 'Ceramium', 'Season' and 'Material'(i.e. thalli and surface extracts), was used to analyze the data from the two assays with diatoms and from the two assays with *Ceramium* sp., respectively, in a common approach (see results in Table S2, S3). To achieve homogeneity of variances and normality of errors, data from the two assays with diatoms were square root transformed. 

## 290 **Results**

#### 291 Defense capacity against diatoms in living Gracilaria vermiculophylla thalli

292 After 3 h of exposure to colonization by diatoms, a three-way-interaction among the factors 293 'Diatom', 'Origin' and 'Season' was observed: Fewer diatoms from Rongcheng attached to non-294 native than to native G. vermiculophylla in autumn and this difference was less pronounced in 295 Summer and less observed with diatoms from Kiel in both seasons (Fig. 1, Table 3). For both 296 diatom species fewer cells (by 60% less) attached to non-native than to native G. vermiculophylla 297 individuals (Fig. 1, Table 3) and only 4% of the unexplained variation was found to be covered by the random factor 'Site'. Furthermore, for both diatom species, settlement rates were on 298 average by 66% lower in summer than in autumn. Averaged across the two seasons, diatoms 299 from Kiel settled by 21% less often on G. vermiculophylla thalli than their congeners from 300 Rongcheng (Fig. 1, Table 3). 301

#### 302 Defense capacity against *Ceramium* sp. in living *Gracilaria vermiculophylla* thalli

After two weeks of colonization by *Ceramium* filaments there was an interaction between 303 304 'Ceramium' and 'Origin' (Fig. 2, Table 4). Ceramium virgatum (from Kiel, Germany) attached less to non-native than to native G. vermiculophylla, while this difference was less pronounced in 305 C. tenerrimum (from Rongcheng, China). An interaction between 'Ceramium' and 'Season' also 306 307 emerged since attachment rates of C. tenerrimum differed between autumn and summer, while 308 this was not the case for C. virgatum (Fig. 2, Table 4). Filaments of both Ceramium species attached, on average, by 33% less often to non-native than to native G. vermiculophylla 309 310 specimens (Fig. 2, Table 4). In this case, 37% of the unexplained variation was covered by 'Site'. 311 Furthermore, attachment rates, averaged across both Ceramium species, were by 10% lower in autumn than in summer, while they, when averaged across both seasons, did not differ between
the two *Ceramium* species (Fig. 2, Table 4).

#### 314 Chemical defense capacity against diatoms in *Gracilaria vermiculophylla* surface extracts

The results from this assay are generally in accordance with those of the diatom trials with living 315 thalli. After exposing the surface extracts to diatom settlement for 3 h, an interaction among the 316 317 factors 'Solvent', 'Diatom' and 'Season' was observed: Attachment rates of diatoms from Kiel on surfaces coated with DCM-based extracts were lower in summer than in autumn, but no such 318 difference was observed on surfaces coated with hexane-based extracts or with diatoms from 319 Rongcheng on any coated surfaces (Table 5). Additionally, fewer diatoms attached (by 9% less) 320 321 to surfaces coated with extracts from non-native than from native G. vermiculophylla (Figs 3 and 322 4, Table 5) and only 2% of the unexplained variation was covered by 'Site'. In general, diatom settlement rates were again by 22% lower in summer than in autumn (Figs 3 and 4, Table 5). 323 Interestingly, different from the assays with living thalli, diatoms from Kiel settled two times 324 325 more often than diatoms from Rongcheng (Fig. 3, Table 5). Moreover, we found fewer diatoms (by 4% less) attached to surfaces covered with non-polar compounds (extracted with hexane) 326 than to those coated with polar compounds (extracted with DCM) (Fig. 4, Table 5). 327

# 328 Chemical defense capacity against *Ceramium* sp. in *Gracilaria vermiculophylla* surface 329 extracts

In autumn we found, after two weeks of exposing surface extracts to colonization by *Ceramium* filaments that fewer (by 13% less) *Ceramium* filaments attached to surfaces coated with moderately polar compounds than on such covered with non-polar compounds. This was not the case in summer and this difference led to an interaction between 'Solvent' and 'Season' (Fig. 6,

334	Table 6). In addition, there was an interaction between 'Ceramium' and 'Season': Attachment
335	rates of C. virgatum were lower in autumn than in summer, while this difference was less
336	pronounced in C. tenerrimum (Fig. 5, Table 6). Furthermore, fewer Ceramium filaments (by 10%
337	less) attached to surfaces coated with extracts gained from non-native than to surfaces with
338	extracts from native G. vermiculophylla (Figs 5 and 6, Table 6) and only 0.5% of the
339	unexplained variation was covered by 'Site'. In general, Ceramium filaments attached 22% less
340	often in autumn than in summer (Figs 5 and 6, Table 6), but attachment rates never differed
341	between the two Ceramium species (Fig. 5, Table 6).
342	
343	
545	
344	
345	
346	
347	
348	
349	
545	
350	
351	
352	

## 353 **Discussion**

For this intra-specific comparison, we sampled individuals of G. vermiculophylla at different 354 sampling locations in either the native or the non-native range of the species. Within the 355 356 respective ranges, the different sampling sites were located in nearly the same biogeographical 357 region according to the Marine Ecoregions of the World (MEOW) system suggested by Spalding et al. (2007). The sites in the native range were located in the Cold Temperate Northwest 358 359 Pacific/Yellow Sea (China) and in Northeastern Honshu (Japan) as well as in the Warm Temperate Northwest Pacific/Central Kuroshio Current (Japan), which is adjacent to 360 Northeastern Honshu. The sites in the non-native range were located in the Northern European 361 362 Sea/Baltic & North Sea (Germany) and in the Celtic Sea as well as the Lusitanian/South European Atlantic Shelf (France), which is adjacent to the Celtic Sea. Furthermore, we took care 363 that the distances between the various sampling sites in the native as well as in the non-native 364 range were similar, in order to have the same degree of between-site variability within the 365 ranges. We therefore assumed that the within-range variability, which could be attributed to 366 367 potential differences in the diversity and composition of the resident flora and fauna as well as to climate conditions, would be low. This assumption was confirmed by the low amount of 368 unexplained variation (0.5% to 4%) that was actually covered by the random factor 'Site' in most 369 370 of our statistical modellings. This was true for diatom attachment rates on both living thalli and on extracts and for *Ceramium* attachment to extracts. In contrast to this, *Ceramium* attachment 371 rates on living thalli varied considerably (37% of the unexplained variation) among sites within 372 both ranges and we cannot plausibly explain this deviation from the otherwise consistent picture. 373

In our study we used two types of epiphytes to test for inter-population differences in *G*. *vermiculophylla* with regard to its susceptibility to epiphytes: Diatoms as a common type of

376 micro-epiphytes and *Ceramium* filaments as a macro-epiphyte. We found that both of them, 377 independent of their actual origin, attached by 60% and 33%, respectively, less to the living thalli of European G. vermiculophylla than to those of Asian conspecifics. In general, such a difference 378 379 could either due to the fact that non-native G. vermiculophylla individuals are better defended and therefore attract fewer/repel more epiphytes or it could be due to lower settlement rates of 380 381 native colonizers on the non-native macroalga. However, our experimental design excluded the latter option, since we exposed non-native macroalgae to epiphytes from the native as well as 382 from the non-native range and both combinations showed the same trend. This finding indicates 383 384 that non-native G. vermiculophylla are better defended against epiphytes than those that stem from the native range. 385

In macroalgae, resistance to epibiosis can be mediated by a) the thallus surface structure 386 (Schumacher et al. 2007; Chapman et al. 2014), b) surface associated bacterial communities that 387 388 repel epibionts (Boyd, Adams & Burgess 1999; Dobretsov, Dahms & Qian 2006), and c) 389 surface-bound secondary metabolites that have anti-epibiont activities (Nylund et al. 2007; Saha et al. 2011; Thabard et al. 2011). The question is now whether one or more of these mechanisms 390 changed with regard to their mode of action or with regard to their efficiency during the invasion 391 process. Such a change could, *inter alia*, be caused by a directional selection of genotypes that 392 exhibit a low susceptibility to epibionts during transport or after release into the new habitat. 393 However, we cannot think of a scenario during these phases that would specifically select for 394 resistance to epibiosis. Furthermore, since the non-native gene pool is a reduced subset of the 395 396 gene pool in the donor region, it is possible, although presumably not very likely, that by chance 397 an epibiosis-resistant genotype was highly frequent among the introduced individuals. Finally, a change in the quality of anti-epiphyte defenses could be attributed to increased energy resources, 398

399 which are a consequence of the release from abiotic and biotic pressures in the new environment 400 (Joshi & Vrieling 2005). Under such conditions, non-native seaweeds may reduce specific defenses they developed against enemies in their native range and shift energy resources towards 401 402 more general anti-enemy defenses. An observation made at our study site in the native range hints at the potential relevance of the last mechanism: In Rongcheng, China, an amphipod 403 species, Caprella sp., is the main grazer of G. vermiculophylla in many habitats and it can 404 consume substantial parts of this local alga during summer (S. Wang, pers. obs.). In Kiel, 405 Germany, so far no herbivore makes use of this alga to such an extent and it seems that the 406 407 grazing pressure on G. vermiculophylla is generally lower than in Rongcheng (Hammann et al. 2013). However, we do not have information whether the picture is the same in the other non-408 native habitats that we sampled in Europe. If G. vermiculophylla is mostly ungrazed in coastal 409 410 habitats in Europe, this could have allowed the non-native G. vermiculophylla to allocate a larger part of their energy budget to anti-epibiont defenses and this possibly caused their lower 411 attractiveness for colonizers. 412

An important aspect of our study was to identify properties of G. vermiculophylla that mediate 413 its anti-epiphyte defenses. For this we compared epiphyte attachment rates on living thalli to 414 those on extract coated surfaces. Here we observed that the general trend in epiphyte attachment 415 was the same for living thalli and extract coated substrata. However, the effect size, i.e. the 416 difference in the susceptibility to epiphytes in native and non-native G. vermiculophylla, was 417 consistently smaller in the latter assays: Fewer diatoms (on average 9% for extracts, 60% for 418 living thalli) and fewer *Ceramium* filaments (on average 10% for extracts, 33% for living thalli) 419 420 attached to substrata that were covered with extracts from European G. vermiculophylla than to those with extracts from Asian specimens. This, first of all, confirms that resistance to epiphytes 421

422 in G. vermiculophylla has, at least partly, a chemical component. If the lower susceptibility in 423 non-native G. vermiculophylla is due to this chemical component, it could either be based on an increased synthesis of active compounds (Forslund, Wikström & Pavia 2010) or due to the 424 presence of some chemical compounds that are novel to resident enemies in these individuals 425 (Enge et al. 2012). Overall, extracts exhibited a lower inhibitory activity against diatom and 426 Ceramium settlement than living algae. This difference could be due to the fact that active 427 metabolites were insufficiently captured by the extraction process or degraded after extraction. 428 Alternatively, other non-chemical components such as surface properties - which were of course 429 430 excluded in the assays with extracts - could also have contributed to the overall deterrence. Finally, the compounds which were responsible for the anti-epiphyte activity we observed in 431 living G. vermiculophylla may not only have stemmed from the thallus surface but also from the 432 inside of algal cells. This reason could have been relevant since some epibionts, including 433 species of the genus Ceramium, penetrate into algal thalli and therefore also get in contact with 434 their interior (e.g. Leonardi et al. 2006). We have no data that could elucidate which of the three 435 436 scenarios was responsible for the picture we observed. However, since we used only two solvents for the extractions (i.e. hexane and DCM) that cover a limited part of the polarity 437 438 spectrum, it is at least likely that we missed relevant compounds and thereby underestimated the potential of chemical defenses in G. vermiculophylla. 439

So far, no tests have been made to investigate whether the surface texture, microtopography or consistency of *G. vermiculophylla* thalli mediates a defense against epibionts. Such effects are known from *Saccharina* species (Chapman *et al.* 2014; da Gama, Plouguerné & Pereira 2014) that belong to the brown macroalgae and possess an outer cell wall with a mucilage consisting of 444 alginic acid with traces of sulphated fucoidan that could, theoretically, act as a low-adhesion,445 gelatinous covering.

Whatever the mechanism is, a low susceptibility to epiphytes in non-native populations of G. 446 *vermiculophylla* can, at least partly, explain the invasion success of the species. It has been 447 448 proposed that marine algal invaders have more effective anti-epibiont defenses than comparable resident species, e.g. in its non-native habitats in northern Europe the brown alga Fucus 449 evanescens is known to get less colonized by filamentous algae and sessile invertebrates than its 450 native congener Fucus vesiculosus (Wikström & Pavia 2004). When their surface is free of 451 epibionts, macroalgae can take up more oxygen, carbon dioxide and nutrients. Furthermore, they 452 receive more light for photosynthesis and are less prone to dislodgement caused by 453 454 biomechanical drag. Furthermore, they may be less attractive to grazers (Wahl, Hay & Enderlein 1997). Therefore, algae, which are free of epibionts or show low degrees of epiphyte or epizoan 455 456 cover, should have more energy available for reproduction and growth, as well as for tolerating adverse environmental conditions – what in turn should increase their potential to establish and 457 spread in new environments. 458

459 The Enemy Release Hypothesis proposes that non-indigenous species are commonly released from biotic pressures, e.g. grazing, in their recipient habitat, because they leave their co-evolved 460 461 antagonists behind while, at the same time, resident enemies fail to recognize the new species as a food source (Keane & Crawley 2002). In this context, Hammann et al. (2013) found that the 462 periwinkle species Littorina brevicula, which lives in the native range of G. vermiculophylla, 463 consumes more of this seaweed, regardless from which distributional range the algal material 464 465 stems, than *Littorina littorea*, which is from its non-native range. This finding is presumably due to the fact that L. brevicula coevolved with G. vermiculophylla and can make better use of it as a 466

467 food source. A somewhat comparable preference of native over non-native G. vermiculophylla was also shown by the diatoms we used in the assays with living thalli: diatoms from Rongcheng 468 generally attached in higher numbers to G. vermiculophylla from both distributional ranges than 469 470 diatoms from Kiel. This finding may also be due to the fact that these diatoms recognize G. vermiculophylla more readily as a suitable settlement substratum than diatoms from the non-471 472 native range. Surprisingly, we observed the opposite picture when we tested the surface extracts. This mismatch suggests that, besides a chemical defense, other properties of the thallus surface, 473 such as its structure or the presence of biofilms, play a role in the anti-diatom defense in living 474 475 thalli. However, we did not find a difference between the attachment rates exhibited by the two *Ceramium* species for both substrata. However, it is not clear why this epiphyte did not show a 476 preference. It is possible that C. virgatum – with regard to G. vermiculophylla as a settlement 477 substratum - generally has the same settlement capacity as C. tenerrimum. The absence of a 478 difference is somewhat surprising, because the way the Ceramium filaments attach to algal 479 surfaces constitutes a very intimate connection of the two organisms. It is most often 480 481 characterized as an infection of the basibiont, because it is mediated by the formation of hapteria that first attach to the thalli of the host and then penetrate into its tissue (Lion et al. 2006). Lion 482 483 et al. (2006) found that after wounding Gracilaria chilensis released oxylipins, which suppressed the development of hapteria in Ceramium rubrum. This fact indicates that co-evolution occurs 484 between Ceramium species and their hosts and hence a difference in settlement rates between C. 485 486 *tenerrimum* and *C. virgatum* would be likely.

Previous studies have suggested that season (Culioli *et al.* 2002; Hellio *et al.* 2004) can influence the capacity of a seaweed to defend itself against epibionts. In accordance with this finding, we found differences in epiphyte settlement rates between summer and autumn of the same year of 490 which we assume that they are attributed to differences in the anti-epiphyte activity of G. 491 vermiculophylla. Fewer diatoms attached to both living fragments and extract-coated surfaces in summer, while fewer Ceramium filaments attached to those substrates in autumn. Such inter-492 493 seasonal differences in anti-epiphyte defenses presumably correlate with natural fluctuations in the overall propagule abundance in the colonizer pool (Steinberg & Vanaltena 1992; Amade & 494 495 Lemée 1998; Wahl et al. 2010; Rickert et al. 2015), which means that marine macroalgae can adjust their anti-epibiont activities to quantitative or qualitative changes in colonization pressure. 496 In both regions where we collected algae, diatoms are more abundant from April to June than 497 498 from August to October (Trimonis, Vaikutiene & Gulbinskas 2010; Wang et al. 2014), while 499 *Ceramium* is more abundant during the latter time span (Weinberger *et al.* 2014; S. Wang, pers. 500 obs.). However, we collected our data only during the course of one year and we therefore do not 501 have robust evidence for seasonality in the defense capacity of G. vermiculophylla. To establish 502 such a pattern, assays would need to be repeated over several years.

A further interesting observation that we made was that fewer diatoms attached to surfaces 503 covered with non-polar compounds than to those coated with polar compounds, while the 504 opposite was true for *Ceramium* filaments. This indicates that the defenses against these two 505 506 epiphytes are mediated by compounds that differ in polarity. A similar observation has been reported earlier: surface compounds extracted with a mixture of hexane and DCM from 507 508 *Caulerpa filiformis* significantly inhibited spore settlement of *Polysiphonia* sp., while more polar compounds, which were extracted with DCM from surfaces of the same species, inhibited 509 510 settlement and germling development of gametes of Ulva australis (Nylund et al. 2007).

511 Our study is the second biogeographical comparison of defense capacities against epibionts 512 between native and non-native populations of *G. vermiculophylla* (Saha *et al.* 2016), which is 513 now invasive in many coastal areas worldwide. However, it gives the first evidence that the capacity to defend against epibionts is higher in non-native individuals than in native -514 regardless of whether the epibionts originate from the native of the non-native range of G. 515 *vermiculophylla*. Our findings therefore seemingly contradict the observations made by Saha *et* 516 517 al. (2016), who focused on seaweed-bacteria interactions and showed that non-native G. 518 *vermiculophylla* are better defended against bacterial epibionts from the non-native range but, at the same time, have reduced their capacity to defend themselves against epibionts from their 519 home range. The contradiction may be due to the use of different micro-epibionts. Bacteria are 520 521 the first colonizers of bare substrata in the marine environment (Wahl 1989) and can regulate the production of bioactive compounds, motility, and biofilm formation by Quorum Sensing (QS), 522 which is a density-dependent cell-cell signaling communication among bacteria (da Gama, 523 524 Plouguerné & Pereira 2014). Furthermore, it is known that bacterial biofilm formation can mediate further colonization by eukaryote micro- and macro-epibionts. The differences between 525 epibacteria and other epibionts could have led to the evolution of different defense strategies 526 527 against them in seaweeds. Unlike compounds that function against eukaryote micro- and macroepibionts through growth inhibition or lethality, most antimicrobial settlement and attachment 528 529 defenses impact the behavior of bacteria, such as swarming (Rasmussen & Givskov 2006).

Even though our study focused on macrophyte-epiphyte interactions in the marine environment, our findings should be applicable to all basibiont-epibiont interactions in aquatic systems, including plant-plant, plant-animal and animal-animal combinations, since epibionts are widespread and most of them are generalists (Wahl & Mark 1999). Additionally, our findings may also be applicable to host-herbivore interactions in aquatic systems, since, similar to epibionts, many herbivores are generalists and an increased chemical resistance to herbivory has already been documented in non-native plants and seaweeds (Forslund, Wikström & Pavia2010).

We conclude that the lower susceptibility to epiphytes that we observed in non-native *G*. *vermiculophylla* cannot be explained by a lower epibiont pressure experienced by the non-native individuals, but is due to an elevated resistance to epibiosis that, at least partly, is linked to an enhanced chemical defense capacity. Our study therefore provides the first evidence of an increased resistance to epibiosis in introduced populations of a widely distributed marine species. This change in its performance during the invasion process may be critical for the invasion success of the macroalga.

- 545
- 546
- 547
- 548
- 549

550

551

552

# 554 Acknowledgments

S. Wang is supported by a scholarship from the China Scholarship Council (CSC) at GEOMAR -Helmholtz-Zentrum für Ozeanforschung in Kiel. We would like to thank Prof. Dr. Martin Wahl for his valuable support and technical advices for bioassay design and methods. We are grateful to Inken Kruse, Takehisa Yamakita, Haruka Yamaguchi, Carola Schuller, Hiromi Sugai, Myriam Perschke for collecting and sending algal samples, as well as to Nadja Stärck for her technical advices and help with algal surface extraction. We are also very grateful to two anonymous reviewers for their very valuable comments on the first version of the manuscript. 

# 573 Author contributions

574	M.L., F.W. and S.W. conceived and designed the experiments. S.W. performed the experiments
575	and collected algal material. G.W., D.B. and M.N. contributed to algal collection and the labs in
576	China and Japan for bioassays. M.L. and S.W. performed statistical analyses. S.W. wrote the
577	manuscript, and M.L., F.W. and S.W. contributed to revisions.
578	
579	
580	
581	
582	
583	
584	
585	
586	
587	
588	
589	
590	

# 591 Data accessibility

592	The authors confirm that all data underlying the findings are fully available without restriction.
593	Data are available at https://doi.pangaea.de/10.1594/PANGAEA.865230 (Wang et al. 2016). R
594	scripts: uploaded as online supporting information.
595	
596	
597	
598	
599	
600	
601	
602	
603	
604	
605	
606	
607	

## 608 **References**

- Amade, P. & Lemée, R. (1998) Chemical defence of the Mediterranean alga *Caulerpa taxifolia*:
   variations in caulerpenyne production. *Aquatic toxicology*, 43, 287-300.
- Baer, J. & Stengel, D.B. (2014) Can native epiphytes affect establishment success of the alien
   seaweed Sargassum muticum (Phaeophyceae)? Biology and Environment: Proceedings of
   the Royal Irish Academy, 114B, 41-52.
- Bellorin, A.M., Oliveira, M.C. & Oliveira, E.C. (2004) *Gracilaria vermiculophylla*: A western
   Pacific species of Gracilariaceae (Rhodophyta) first recorded from the eastern Pacific.
   *Phycological Research*, 52, 69-79.
- Blossey, B. & Nötzold, R. (1995) Evolution of increased competitive ability in invasive
  nonindigenous plants: a hypothesis. *Journal of Ecology*, 83, 887-889.
- Bold, H.C. & Wynne, M.J. (1978) *Introduction to the Algae: Structure and Reproduction*.
  Prentice-Hall, Englewood Cliffs, New Jersey.
- Boyd, K.G., Adams, D.R. & Burgess, J.G. (1999) Antibacterial and repellent activities of marine
  bacteria associated with algal surfaces. *Biofouling*, 14, 227-236.
- Cacabelos, E., Olabarria, C., Incera, M. & Troncoso, J.S. (2010) Do grazers prefer invasive
  seaweeds? *Journal of Experimental Marine Biology and Ecology*, **393**, 182-187.
- Callaway, R.M. & Ridenour, W.M. (2004) Novel weapons: invasive success and the evolution of
   increased competitive ability. *Frontiers in Ecology and the Environment*, 2, 436-443.
- Chapman, J., Hellio, C., Sullivan, T., Brown, R., Russell, S., Kiterringham, E., Le Nor, L. &
   Regan, F. (2014) Bioinspired synthetic macroalgae: examples from nature for antifouling
   applications. *International Biodeterioration & Biodegradation*, 86, 6-13.
- Culioli, G., Ortalo-Magné, A., Richou, M., Valls, R. & Piovetti, L. (2002) Seasonal variations in
   the chemical composition of *Bifurcaria bifurcata* (Cystoseiraceae). *Biochemical systematics and ecology*, **30**, 61-64.
- da Gama, B.A.P., Plouguerné, E. & Pereira, R.C. (2014) The Antifouling Defence Mechanisms
  of Marine Macroalgae. *Advances in Botanical Research* (ed. N. Bourgougnon), pp. 413440. Academic Press, Oxford.
- de Nys, R., Dworjanyn, S.A. & Steinberg, P.D. (1998) A new method for determining surface
   concentrations of marine natural products on seaweeds. *Marine Ecology Progress Series*,
   162, 79-87.
- Dobretsov, S., Dahms, H.U. & Qian, P.Y. (2006) Inhibition of biofouling by marine
   microorganisms and their metabolites. *Biofouling*, 22, 43-54.
- Enge, S., Nylund, G.M., Harder, T. & Pavia, H. (2012) An exotic chemical weapon explains low
  herbivore damage in an invasive alga. *Ecology*, 93, 2736-2745.
- Engelen, A.H., Henriques, N., Monteiro, C. & Santos, R. (2011) Mesograzers prefer mostly
  native seaweeds over the invasive brown seaweed *Sargassum muticum*. *Hydrobiologia*,
  645 669, 157-165.
- Forslund, H., Wikström, S.A. & Pavia, H. (2010) Higher resistance to herbivory in introduced
   compared to native populations of a seaweed. *Oecologia*, 164, 833-840.
- Freshwater, D.W., Montgomery, F., Greene, J.K., Hamner, R.M., Williams, M. & Whitfield, P.E.
  (2006) Distribution and identification of an invasive *Gracilaria* species that is hampering
  commercial fishing operations in southeastern North Carolina, USA. *Biological Invasions*, 8, 631-637.

- Hammann, M. (2014) Invasion ecology of marine macroalgae: the relevance of stress resistance
   for the invasion success of *Gracilaria vermiculophylla* and consequences of its spread.
   Christian-Albrechts-University Kiel.
- Hammann, M., Rempt, M., Pohnert, G., Wang, G.G., Boo, S.M. & Weinberger, F. (2016)
  Increased potential for wound activated production of Prostaglandin E-2 and related toxic
  compounds in non-native populations of *Gracilaria vermiculophylla*. *Harmful Algae*, 51, 81-88.
- Hammann, M., Wang, G.G., Rickert, E., Boo, S.M. & Weinberger, F. (2013) Invasion success of
   the seaweed *Gracilaria vermiculophylla* correlates with low palatibility. *Marine Ecology Progress Series*, 486, 93-103.
- Hellio, C., Marechal, J.P., Véron, B., Bremer, G., Clare, A.S. & Le Gal, Y. (2004) Seasonal
  variation of antifouling activities of marine algae from the Brittany coast (France). *Marine Biotechnology*, 6, 67-82.
- Hemmi, A., Mäkinen, A., Jormalainen, V. & Honkanen, T. (2005) Responses of growth and
   phlorotannins in *Fucus vesiculosus* to nutrient enrichment and herbivory. *Aquatic Ecology*, 39, 201-211.
- Hierro, J.L., Maron, J.L. & Callaway, R.M. (2005) A biogeographical approach to plant
   invasions: the importance of studying exotics in their introduced and native range.
   *Journal of Ecology*, 93, 5-15.
- Joshi, J. & Vrieling, K. (2005) The enemy release and EICA hypothesis revisited: incorporating
   the fundamental difference between specialist and generalist herbivores. *Ecology Letters*,
   8, 704-714.
- Keane, R.M. & Crawley, M.J. (2002) Exotic plant invasions and the enemy release hypothesis.
   *Trends in Ecology & Evolution*, 17, 164-170.
- Kim, S.Y., Weinberger, F. & Boo, S.M. (2010) Genetic data hint at a common donor region for
  invasive Atlantic and Pacific populations of *Gracilaria Vermiculophylla* (Gracilariales,
  Rhodophyta). *Journal of Phycology*, 46, 1346-1349.
- Lenz, M., da Gama, B.A.P., Gerner, N.V., Gobin, J., Gröner, F., Harry, A., Jenkins, S.R.,
  Kraufvelin, P., Mummelthei, C., Sareyka, J., Xavier, E.A. & Wahl, M. (2011) Non-native
  marine invertebrates are more tolerant towards environmental stress than taxonomically
  related native species: results from a globally replicated study. *Environmental research*,
  111, 943-952.
- Leonardi, P.I., Miravalles, A.B., Faugeron, S., Flores, V., Beltrán, J. & Correa, J.A. (2006)
   Diversity, phenomenology and epidemiology of epiphytism in farmed *Gracilaria chilensis* (Rhodophyta) in northern Chile. *European Journal of Phycology*, 41, 247-257.
- Lion, U., Wiesemeier, T., Weinberger, F., Beltrán, J., Flores, V., Faugeron, S., Correa, J. &
   Pohnert, G. (2006) Phospholipases and galactolipases trigger oxylipin-mediated wound activated defence in the red alga *Gracilaria chilensis* against epiphytes. *Chembiochem*, 7,
   457-462.
- Mack, R.N., Simberloff, D., Lonsdale, W.M., Evans, H., Clout, M. & Bazzaz, F.A. (2000) Biotic
   invasions: causes, epidemiology, global consequences, and control. *Ecological applications*, 10, 689-710.
- Michetti, K.M., Miravalles, A.B., Hughes, M.H. & Leonardi, P.I. (2016) Infection process of
   *Ceramium rubrum* (Rhodophyta, Ceramiales) on the agarophyte *Gracilaria chilensis* (Rhodophyta, Gracilariales). *Botanica Marina*, **59**, 51-61.

- Müller-Schärer, H., Schaffner, U. & Steinger, T. (2004) Evolution in invasive plants:
   implications for biological control. *Trends in Ecology & Evolution*, 19, 417-422.
- Newton, C., Bracken, M.E.S., McConville, M., Rodrigue, K. & Thornber, C.S. (2013) Invasion
   of the red seaweed *Heterosiphonia japonica* spans biogeographic provinces in the
   western North Atlantic Ocean. *PLoS One*, 8, e62261.
- Nylund, G.M., Gribben, P.E., de Nys, R., Steinberg, P.D. & Pavia, H. (2007) Surface chemistry
   versus whole-cell extracts: antifouling tests with seaweed metabolites. *Marine Ecology Progress Series*, 329, 73-84.
- Olenin, S., Narščius, A., Minchin, D., David, M., Galil, B., Gollasch, S., Marchini, A.,
   Occhipinti-Ambrogi, A., Ojaveer, H. & Zaiko, A. (2014) Making non-indigenous species
   information systems practical for management and useful for research: An aquatic
   perspective. *Biological Conservation*, **173**, 98-107.
- Paini, D.R., Sheppard, A.W., Cook, D.C., De Barro, P.J., Worner, S.P. & Thomas, M.B. (2016)
  Global threat to agriculture from invasive species. *Proceedings of the National Academy*of Sciences of the United States of America, **113**, 7575-7579.
- Pimentel, D., Zuniga, R. & Morrison, D. (2005) Update on the environmental and economic costs associated with alien-invasive species in the United States. *Ecological economics*, 52, 273-288.
- Rasmussen, T.B. & Givskov, M. (2006) Quorum sensing inhibitors: a bargain of effects.
   *Microbiology-Sgm*, 152, 895-904.
- Rempt, M., Weinberger, F., Grosser, K. & Pohnert, G. (2012) Conserved and species-specific
   oxylipin pathways in the wound-activated chemical defense of the noninvasive red alga
   *Gracilaria chilensis* and the invasive *Gracilaria vermiculophylla*. *Beilstein Journal of Organic Chemistry*, 8, 283-289.
- Ricciardi, A. (2007) Are modern biological invasions an unprecedented form of global change?
   *Conservation Biology*, 21, 329-336.
- Rickert, E., Karsten, U., Pohnert, G. & Wahl, M. (2015) Seasonal fluctuations in chemical
   defenses against macrofouling in *Fucus vesiculosus* and *Fucus serratus* from the Baltic
   Sea. *Biofouling*, **31**, 363-377.
- Rueness, J. (2005) Life history and molecular sequences of *Gracilaria vermiculophylla* (Gracilariales, Rhodophyta), a new introduction to European waters. *Phycologia*, 44, 120-128.
- Saha, M., Rempt, M., Grosser, K., Pohnert, G. & Weinberger, F. (2011) Surface-associated
   fucoxanthin mediates settlement of bacterial epiphytes on the rockweed *Fucus vesiculosus*. *Biofouling*, 27, 423-433.
- Saha, M., Wiese, J., Weinberger, F. & Wahl, M. (2016) Rapid adaptation to controlling new microbial epibionts in the invaded range promotes invasiveness of an exotic seaweed.
   *Journal of Ecology*, **104**, 969-978.
- Schaffelke, B., Smith, J.E. & Hewitt, C.L. (2006) Introduced macroalgae a growing concern.
   *Journal of Applied Phycology*, 18, 529-541.
- Schumacher, J.F., Carman, M.L., Estes, T.G., Feinberg, A.W., Wilson, L.H., Callow, M.E.,
  Callow, J.A., Finlay, J.A. & Brennan, A.B. (2007) Engineered antifouling
  microtopographies effect of feature size, geometry, and roughness on settlement of
  zoospores of the green alga *Ulva*. *Biofouling*, 23, 55-62.
- Sfriso, A., Wolf, M.A., Maistro, S., Sciuto, K. & Moro, I. (2012) Spreading and autoecology of
   the invasive species *Gracilaria vermiculophylla* (Gracilariales, Rhodophyta) in the

- lagoons of the north-western Adriatic Sea (Mediterranean Sea, Italy). *Estuarine, Coastal and Shelf Science*, **114**, 192-198.
- Spalding, M.D., Fox, H.E., Halpern, B.S., McManus, M.A., Molnar, J., Allen, G.R., Davidson,
  N., Jorge, Z.A., Lombana, A.L., Lourie, S.A., Martin, K.D., McManus, E., Molnar, J.,
  Recchia, C.A. & Robertson, J. (2007) Marine ecoregions of the world: A
  bioregionalization of coastal and shelf areas. *Bioscience*, 57, 573-583.
- Steinberg, P.D. & Vanaltena, I. (1992) Tolerance of marine invertebrate herbivores to brown
   algal phlorotannins in temperate Australasia. *Ecological Monographs*, 62, 189-222.
- Strong, J.A., Maggs, C.A. & Johnson, M.P. (2009) The extent of grazing release from
   epiphytism for *Sargassum muticum* (Phaeophyceae) within the invaded range. *Journal of the Marine Biological Association of the United Kingdom*, **89**, 303-314.
- Thabard, M., Gros, O., Hellio, C. & Maréchal, J.P. (2011) Sargassum polyceratium
  (Phaeophyceae, Fucaceae) surface molecule activity towards fouling organisms and embryonic development of benthic species. *Botanica Marina*, 54, 147-157.
- Thomsen, M.S., Gurgel, C.F.D., Fredericq, S. & McGlathery, K.J. (2006) *Gracilaria vermiculophylla* (Rhodophyta, Gracilariales) in Hog Island Bay, Virginia: A cryptic alien
   and invasive macroalga and taxonomic correction. *Journal of Phycology*, 42, 139-141.
- Thomsen, M.S., Wernberg, T., Engelen, A.H., Tuya, F., Vanderklift, M.A., Holmer, M.,
  McGlathery, K.J., Arenas, F., Kotta, J. & Sillimann, B.R. (2012) A Meta-Analysis of
  Seaweed Impacts on Seagrasses: Generalities and Knowledge Gaps. *PLoS One*, 7, 21-28.
- Torchin, M.E. & Mitchell, C.E. (2004) Parasites, pathogens, and invasions by plants and animals. *Frontiers in Ecology and the Environment*, 2, 183-190.
- Trimonis, E., Vaikutiene, G. & Gulbinskas, S. (2010) Seasonal and spatial variations of
   sedimentary matter and diatom transport in the Klaipeda Strait (Eastern Baltic). *Baltica*,
   23, 127-134.
- Verhoeven, K.J.F., Biere, A., Harvey, J.A. & Van Der Putten, W.H. (2009) Plant invaders and
   their novel natural enemies: who is naive? *Ecology Letters*, 12, 107-117.
- Vermeij, M.J.A., Smith, T.B., Dailer, M.L. & Smith, C.M. (2009) Release from native
   herbivores facilitates the persistence of invasive marine algae: a biogeographical
   comparison of the relative contribution of nutrients and herbivory to invasion success.
   *Biological Invasions*, 11, 1463-1474.
- Vilà, M., Espinar, J.L., Hejda, M., Hulme, P.E., Jarošík, V., Maron, J.L., Pergl, J., Schaffner, U.,
   Sun, Y. & Pyšek, P. (2011) Ecological impacts of invasive alien plants: a meta-analysis
   of their effects on species, communities and ecosystems. *Ecology Letters*, 14, 702-708.
- Wahl, M. (1989) Marine epibiosis. I. Fouling and antifouling: some basic aspects. *Marine Ecology Progress Series*, 58, 175-189.
- Wahl, M. (2008) Ecological lever and interface ecology: epibiosis modulates the interactions
  between host and environment. *Biofouling*, 24, 427-438.
- Wahl, M., Hay, M.E. & Enderlein, P. (1997) Effects of epibiosis on consumer–prey interactions.
   *Hydrobiologia*, 355, 49-59.
- Wahl, M. & Mark, O. (1999) The predominantly facultative nature of epibiosis: experimental and observational evidence. *Marine Ecology Progress Series*, **187**, 59-66.
- Wahl, M., Shahnaz, L., Dobretsov, S., Saha, M., Symanowski, F., David, K., Lachnit, T., Vasel,
  M. & Weinberger, F. (2010) Ecology of antifouling resistance in the bladder wrack *Fucus vesiculosus*: patterns of microfouling and antimicrobial protection. *Marine Ecology Progress Series*, **411**, 33-48.

- Wang, D., Huang, B.Q., Liu, X., Liu, G.M. & Wang, H. (2014) Seasonal variations of
   phytoplankton phosphorus stress in the Yellow Sea Cold Water Mass. *Acta Oceanologica Sinica*, 33, 124-135.
- Weinberger, F., Buchholz, B., Karez, R. & Wahl, M. (2008) The invasive red alga *Gracilaria vermiculophylla* in the Baltic Sea: adaptation to brackish water may compensate for light
   limitation. *Aquatic Biology*, 3, 251-264.
- Weinberger, F., Hammann, M., Griem, M. & Siedentopp, D. (2014) Ostsee-Makroalgenblüten.
   Helmholtz-Zentrum für Ozeanforschung Kiel, Germany.
- Wikström, S.A. & Kautsky, L. (2004) Invasion of a habitat-forming seaweed: effects on associated biota. *Biological Invasions*, 6, 141-150.
- Wikström, S.A. & Pavia, H. (2004) Chemical settlement inhibition versus post-settlement
   mortality as an explanation for differential fouling of two congeneric seaweeds.
   *Oecologia*, 138, 223-230.
- Wikström, S.A., Steinarsdóttir, M.B., Kautsky, L. & Pavia, H. (2006) Increased chemical
   resistance explains low herbivore colonization of introduced seaweed. *Oecologia*, 148, 593-601.
- Williams, S.L. & Smith, J.E. (2007) A global review of the distribution, taxonomy, and impacts
  of introduced seaweeds. *Annual Review of Ecology, Evolution, and Systematics*, 38, 327359.
- Zuur, A.F., Ieno, E.N., Walker, N.J., Saveliev, A.A. & Smith, G.M. (2009) *Mixed Effects Model and Extensions in Ecology with R.* Springer, New York.
- Zuur, A.F., Ieno, E.N. & Elphick, C.S. (2010) A protocol for data exploration to avoid common
   statistical problems. *Methods in Ecology and Evolution*, 1, 3-14.
- 812
- 813

Origin	Collection site	Geographic	1 <sup>st</sup>	2 <sup>nd</sup>	3 <sup>rd</sup>	Salinity	Attachm-	Morphology	Epibionts
		Coordinate	sampling	sampling	sampling		ent status		(incomplete
									information)
Native	Rongcheng,	37°9'4.29"N,	21.05.2014	23.09.2014	06.07.2015	28-33	Holdfast	Brown,	Diatoms, Ceramium
	China, Yellow	122°33'35.60"E						relatively more	tenerrimum, Ulva
	Sea							branches	sp., Polysiphonia
									sp., <i>Folliculina</i> sp.
	Qingdao, China,	36°3'0.6"N,	21.05.2014	21.09.2014	06.07.2015	28-33	Holdfast	Brown,	Diatoms, Ceramium
	Yellow Sea	120°20'59.1"E						relatively more	tenerrimum, Ulva
								branches	sp., Cladophora sp.
	Akkeshi, Japan,	43°1'25.80"N,	01.05.2014	19.08.2014	16.07.2015	29-33	Holdfast	Brown,	Diatoms, Ceramium
	Northeastern	144°52'47.20"E						relatively more	kondoi, Circeis
	Honshu							branches	spirillum
	Tokyo, Japan,	35°19'25.72"N,	01.06.2014	02.08.2014	10.07.2015	29-33	Holdfast	Brown,	Diatoms, Ulva sp.
	Central	139°38'8.30"E						relatively more	
	Kuroshio							branches	
	Current								

# **Table 1.** Sampling information of the four native and four non-native populations of *Gracilaria vermiculophylla*.

Non-	Kiel, Germany,	54°23'57.03"N,	02.06.2014	12.08.2014	12.07.2015	15-20	Free-	dark	brown,	Diatoms, Ceramium
native	Baltic Sea	10°12'53.12"E					floating	relativel	y fewer	<i>tenuicorne</i> , Ulva sp.,
								branche	8	Pilayella sp.,
										Amphibalanus
										improvises, Mytilus
										edulis
	Nordstrand,	54°29'10.0"N	25.05.2014	04.09.2014	08.07.2015	30-32	Free-	dark	brown,	Diatoms, Ulva sp.
	Germany, North	8°48'44.8"E					floating	relativel	y fewer	Chaetomorpha sp.,
	Sea							branche	S	Cladophora sp.,
										Amphibalanus
										improvises, Mytilus
										edulis
	Belon, France,	47°49'35.80"N,	18.05.2014	08.09.2014		2-33	Free-	dark	brown,	Diatoms
	South European	3°40'20.50"W					floating	relativel	y fewer	
	Atlantic Shelf							branche	S	
	Pouldouran.	48°45'55.90"N.	18.05.2014	08.09.2014	18.06.2015	2-33	Free-	dark	brown.	Diatoms
	France, Celtic	3°12'1.40"W					floating	relativel	v fewer	
	Seas							branche	s	
	~~~~							Siunene	0	

816 Holdfast: thalli attached by a holdfast to the substratum; Free-floating: thalli not attached by a holdfast;

817 Table 2. Overview of the locations and timing over the attachment assays with *Gracilaria*818 *vermiculophylla* and different epiphytes.

Epiphytes	Assays after 1 <sup>st</sup>	Assays after 2 <sup>nd</sup>	Assays after 3 <sup>rd</sup>
	sampling	sampling	sampling
Diatoms from	Kiel,	Kiel,	Akkeshi,
Rongcheng	11.06.2014	24.10.2014	20.07.2015
Diatoms from Kiel	Kiel,	Kiel,	
	11.06.2014	24.10.2014	
Ceramium tenerrimum	Rongcheng,	Rongcheng,	
from Rongcheng	30.09.2014	30.09.2014	
Ceramium virgatum	Kiel,	Kiel,	
from Kiel	10.06.2014	23.10.2014	

**Table 3.** Influence of season, origin of *Gracilaria vermiculophylla* and origin of diatoms on the

	Source of variation	numDF	denDF	F - value	p - value
	Season	1	220	283.691	<.0001
	Origin	1	6	138.724	<.0001
	Diatom	1	220	24.636	<.0001
	Diatom:Origin	1	220	4.429	0.0365
	Diatom:Season	1	220	2.047	0.1540
	Origin:Season	1	220	40.930	<.0001
	Diatom:Origin:Season	1	220	11.439	0.0009
828	Results from mixed-effect	et modelling.			
829					
830					
831					
832					
833					
834					
835					
836					

827 attachment rates of diatoms on living thalli.

838 Table 4. Influence of season, origin of *Gracilaria vermiculophylla* and origin of *Ceramium* on

839	filament attachment rates	on living thalli.
-----	---------------------------	-------------------

	Source of variation	numDF	denDF	F - value	p - value
	Season	1	20	8.4325	0.0088
	Origin	1	6	30.5134	0.0015
	Ceramium	1	20	2.1081	0.1620
	Ceramium:Origin	1	20	13.1757	0.0017
	Ceramium:Season	1	20	75.8923	<.0001
	Origin:Season	1	18	2.2500	0.1510
	Ceramium:Origin:Season	1	18	0.5625	0.4629
840	Results from mixed effect-m	odelling.			
841					
842					
843					

850	Table 5. Influence of season, solvent, origin of Gracilaria vermiculophylla and origin of diatoms
851	on the attachment rates of diatoms on surface extracts.

Source of variation	numDF	denDF	F - value	p - value
Season	1	238	283.0	<.0001
Solvent	1	238	17.4	<.0001
Origin	1	6	11.8	0.0139
Diatom	1	238	1772.5	<.0001
Solvent:Season	1	238	21.9	<.0001
Diatom:Season	1	238	648.9	<.0001
Solvent:Diatom	1	238	0.2	0.6686
Solvent:Origin	1	238	1.3	0.2479
Diatom:Origin	1	238	0.1	0.7242
Origin:Season	1	238	0.0	0.9599
Solvent:Diatom:Season	1	238	6.0	0.0154
Solvent:Diatom:Origin	1	234	0.1	0.8000
Solvent:Origin:Season	1	234	0.7	0.4059
Diatom:Origin:Season	1	234	0.3	0.6021
Solvent:Diatom:Origin:Season	1	234	0.0	0.9073

852 Results from mixed effect-modelling.

856	Table6.	Influence	of	season,	solvent,	origin	of	Gracilaria	vermiculophylla	and	origin	of
857	<i>Ceramium</i> on filament attachment rates on surface extracts.											

Source of variation	numDF	denDF	F - value	p - value
Season	1	307	124.418	<.0001
Solvent	1	307	11.090	0.0010
Origin	1	6	27.095	0.0020
Ceramium	1	307	0.126	0.7224
Solvent:Season	1	307	9.835	0.0019
Ceramium:Season	1	307	20.782	<.0001
Solvent:Ceramium	1	298	2.308	0.1297
Solvent:Origin	1	298	0.126	0.7224
Ceramium:Origin	1	298	0.051	0.8211
Origin:Season	1	298	0.026	0.8717
Solvent:Ceramium:Origin	1	298	0.001	0.9742
Solvent:Ceramium:Season	1	298	3.395	0.0664
Solvent:Origin:Season	1	298	0.235	0.6281
Ceramium:Origin:Season	1	298	0.026	0.8717
Solvent:Ceramium:Origin:Season	1	298	2.718	0.1003

858 Results from mixed effect-modelling.



Fig. 1. Colonization of living thalli of native and non-native *Gracilaria vermiculophylla* by
diatoms from both origins. Assays were run in summer and in autumn 2014 and in summer 2015.
Means and 95% CIs. n =24 to 48 in each group.



Fig. 2. Colonization of living thalli of native and non-native *Gracilaria vermiculophylla* by *Ceramium* from both origins, *C. virgatum* from Germany and *C. tenerrimum* from China. Assays
were run in summer and in autumn 2014. Means and 95% CIs. n = 40 in each group.



Fig. 3. Colonization of surface extracts from native and non-native *Gracilaria vermiculophylla* by diatoms from both origins. Assays were run in summer and in autumn 2014. Means and 95% CIs. n = 32 in each group. The horizontal lines indicate mean colonization rate on controls, which were without extracts (n = 8).



Fig. 4. Colonization of DCM and hexane surface extracts from native and non-native *Gracilaria vermiculophylla* by diatoms from both origins. Assays were run in summer and in autumn 2014.
Means and 95% CIs. n = 32 in each group. The horizontal lines indicate mean colonization rate
on controls, which were without extracts (n = 8).
903
904
905
906
907
908
909



Fig. 5. Colonization of surface extracts from native and non-native *Gracilaria vermiculophylla*by *Ceramium* from both origins, *C. virgatum* from Germany and *C. tenerrimum* from China.
Assays were run in summer and in autumn 2014. Means and 95% CIs. n = 40 in each group. The
horizontal lines indicate mean colonization rate on controls, which were without extracts (n = 10).
915
916
917
918
919



Fig. 6. Colonization of DCM and hexane surface extracts from native and non-native *Gracilaria vermiculophylla* by *Ceramium* from both origins, *C. virgatum* from Germany and *C. tenerrimum*from China. Assays were run in summer and in autumn 2014. Means and 95% CIs. n = 40 in each
group. The horizontal lines indicate mean colonization rate on controls, which were without
extracts (n = 10).

# 936 Supporting Information

937 Additional Supporting Information may be found in the online version of this article:

938 Appendix S1. Transport and cultivation of *Gracilaria vermiculophylla*.

939 Appendix S2. Collection, isolation, transport and cultivation of epiphytes.

940 Appendix S3. Establishing the relationship between diatom density and fluorescence intensity.

**Table S1.** Change of fluorescence intensities of attached diatoms with concentration of diatoms

942 in suspensions after different incubation periods.

Table S2. Influence of season, material (extracts vs. thalli), origin of *Gracilaria vermiculophylla*and origin of diatoms on the attachment rates of diatoms on *G. vermiculophylla*.

Table S3. Influence of season, material (extracts vs. thalli), origin of *Gracilaria vermiculophylla*and origin of *Ceramium* on filament attachment rates on *G. vermiculophylla*.

**Figure S1.** Determination of solvents and dipping times for surface extraction of *Gracilaria vermiculophylla*. (a) Healthy algal cells. The alga was extracted by: (b) Methanol-hexane mixture 1:9 (v/v) for 5 s. (c) Methanol-hexane mixture 1:19 (v/v) for 5 s. (d) Propanol-hexane mixture 1:9 (v/v) for 5 s. (e) Propanol-hexane mixture 1:19 (v/v) for 5 s. (f) dichloromethane (DCM)-hexane mixture 1:3 (v/v) for 5 s. (g) DCM-hexane mixture 1:4 (v/v) for 10 s. (h) DCM-hexane mixture 1:4 (v/v) for 7 s. (i) DCM-hexane mixture 1:4 (v/v) for 5 s. Scale bars: 10  $\mu$ m.

Figure S2. The relationship between fluorescence intensity and diatom density. (a) diatom fromRongcheng, China. (b) diatom from Kiel, Germany.