

1 **Diffusion or advection? Mass transfer and diffusive boundary**
2 **layer landscapes of the brown alga *Fucus vesiculosus***

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15 Running title: DBL landscapes around *Fucus vesiculosus*

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23 **Abstract**

24 The role of hyaline hairs on the thallus of brown-algae in the genus *Fucus* is long debated
25 and several functions have been proposed. We used a novel motorized setup for 2D- and
26 3D-mapping with O₂-microsensors to investigate the spatial heterogeneity of the diffusive
27 boundary layer (DBL) and O₂ flux around single and multiple tufts of hyaline hairs on
28 the thallus of *Fucus vesiculosus*. Flow was a major determinant of DBL thickness, where
29 higher flow decreased DBL thickness and increased O₂ flux between algal thallus and the
30 surrounding seawater. However, the topography of the DBL varied and did not directly
31 follow the contour of the underlying thallus. Areas around single tufts of hyaline hairs
32 exhibited both increased and decreased DBL thickness as compared to areas over smooth
33 thallus surfaces. Over thallus areas with several hyaline hair tufts, the overall effect was
34 a local increase in the DBL thickness. We also found indications for advective O₂
35 transport driven by pressure gradients or vortex-shedding downstream from dense tufts
36 of hyaline hairs alleviating local mass-transfer resistance imposed by thickened DBL.
37 Mass-transfer dynamics around hyaline hair tufts are thus more complex than hitherto
38 assumed and may have important implications for algal physiology and plant-microbe
39 interactions.

40

41 **Keywords:** biological fluid mechanics, diffusive boundary layer, macroalgae, mass
42 transfer, oxygen, topography.

43

44 **Introduction**

45 Aquatic macrophytes are limited compared to their terrestrial counterparts by the $\sim 10^4$
46 slower diffusion and a much lower solubility of gases in water than in air [1, 2]. The
47 efficient exchange of nutrients and gases is further exacerbated by the diffusive boundary
48 layer (DBL) surrounding all submersed surfaces [3]. The thickness and topography of
49 diffusive boundary layers around submerged impermeable objects is affected by flow
50 velocity and surface topography [3]. Higher flow velocities decrease the DBL thickness
51 by exerting a higher shear stress on the viscous sublayer near the surface. The effect of
52 surface roughness on DBL thickness is variable, where angled planes facing the flow
53 generally will have decreased boundary layers, while the DBL downstream of protruding
54 structures will have increased boundary layers. At the same time, the effect of surface
55 roughness on mass transfer across a boundary layer is dichotomous, where a thicker DBL
56 will decrease mass transfer, while surface roughness tends to increase the overall surface
57 area increasing mass transfer [3]. In microsensor-based studies of the DBL, it is important
58 to note that the presence of the microsensor tip in itself can affect the local DBL thickness
59 [4], where flow acceleration around the microsensor shaft will compress the local DBL
60 thickness leading to locally enhanced O_2 fluxes in the order of 10%. However, this effect
61 is only significant when investigated on smooth surfaces, while a clear effect is apparently
62 undetectable when measured over tufts in e.g. a cyanobacterial mat [5].

63 It has been estimated that the DBL accounts for up to 90% of the resistance to carbon
64 fixation in freshwater plants [6], and structural- and biochemical regulations to alleviate
65 such mass transfer resistance have evolved across lineages. Some aquatic macrophytes
66 have e.g. developed i) thinner leaves and a reduced cuticle that decreases the diffusion
67 path length to chloroplasts, ii) carbon concentrating mechanisms that increase internal
68 carbon concentration, and iii) the ability to utilize HCO_3^- which constitutes the largest
69 fraction of dissolved inorganic carbon at ocean pH ([7], [8] and references therein). In
70 photosymbiotic corals it has also been proposed that vortical ciliary flow can actively
71 enhance mass transfer between the coral tissue and the surrounding water in stagnant or
72 very low flow regimes with concomitant thick diffusive boundary layers [9].

73 Hyaline hairs, i.e., colourless, filamentous multicellular structures, are often observed as
74 whitish tufts on the thallus of brown macroalgae in the genus *Fucus*. The hairs originate
75 and are anchored in cryptostomatal cavities on the apical- and mid-regions of the thallus

76 [10]. It is recognized that hyaline hairs aid in the uptake of nutrients [11, 12] e.g. during
77 springtime, when photosynthetic potential is higher due to increased light levels and the
78 need for nutrients apparently triggers growth of hyaline hairs [10].
79 How hyaline hairs affect solute exchange and nutrient acquisition in *Fucus* is still debated,
80 although a number of suggestions currently exist in the literature suggesting that: i) the
81 hairs increase the algal surface area available for nutrient uptake, albeit this is probably
82 not the major limitation on nutrient uptake [11]; ii) the hyaline hairs might decrease the
83 diffusive boundary layer (DBL) due to turbulence created by the hairs as water flows
84 across them, decreasing the mass resistance imposed by the DBL [11]; iii) the thin cell
85 walls of the hairs relative to the thallus could have less resistance to the passage of ions
86 [13, 14]; iv) the hyaline hairs increase DBL thickness, thereby retaining the products of
87 thallus surface-active enzymes such as extracellular phosphatases and ensuring more
88 efficient nutrient uptake [14].
89 There can be no doubt, however, that the DBL has great importance for macroalgal
90 growth rates. The mass resistance imposed by the DBL has been correlated with nutrient
91 limitation for the giant kelp *Macrocystis pyrifera* [15], and a considerable spatial variation
92 of the DBL over thallus and cryptostomata of *Fucus vesiculosus* has been observed [16].
93 However, current knowledge of the DBL characteristics of aquatic plants is largely based
94 on point measurements with O₂ microsensors [14], while it is known from boundary layer
95 studies in biofilms [17], corals [18-20] and sediments [17, 21-23] that the DBL exhibits
96 a spatio-temporal heterogeneity, which is modulated by both flow velocity and surface
97 topography. Similar studies of DBL topography are very limited in aquatic plant science
98 [24] and the aim of this study was to explore how the DBL thickness and the local O₂ flux
99 varied spatially over the thallus of *F. vesiculosus* with and without tufts of hyaline hairs.
100 Such first time exploration of the 3D DBL topography was done with O₂ microsensors
101 mounted in a fully automated motorized micromanipulator system that allowed
102 measurements of DBL transects and grids over the thallus of *F. vesiculosus* (Fig. S1). Our
103 results reveal a complex DBL landscape over the algal thallus, where mass transfer across
104 the DBL apparently can be supplemented by advective processes due to the presence of
105 hyaline hairs.
106

107 **Materials and Methods**

108 Sampling and experimental setup

109 Specimens of *Fucus vesiculosus* and seawater used in the experimental setup were
110 sampled on the day of usage at <1 m depth at Kronborg, Helsingør, Denmark from May
111 through August. If the influence of a single tuft of hyaline hairs on the DBL was
112 measured, the surface of the thallus surrounding the tuft was carefully shaved off any
113 additional tufts with a scalpel and observation under a dissection microscope to avoid
114 thallus damage. When the influence of multiple tufts was analysed, the thallus was left
115 intact. Prior to measurements, a piece of *F. vesiculosus* thallus with hyaline hairs was
116 fixed on a slab of agar (~1.5% w/w in seawater) in a small flow chamber creating a
117 defined unidirectional flow of seawater across the thallus surface [16]. The flow chamber
118 was connected via tubing to a submersible water pump in a continuously aerated seawater
119 reservoir tank underneath the flow chamber. Flow velocities were adjusted by restricting
120 water flow to the flow chamber, and flow velocity was determined by collecting water
121 from the flow chamber outlet for one minute, where after the sampled volume per time
122 was divided by the cross sectional area of the flow chamber to obtain an estimate of the
123 mean free flow velocity. All measurements were done with flow velocities of either 1.65
124 cm s⁻¹ or 4.88 cm s⁻¹.

125 The sample was illuminated from above with light from a halogen lamp (Schott KL-
126 2500LCD) equipped with a collimating lens yielding a downwelling photon irradiance
127 (400-700 nm) of ~350 μmol photons m⁻² s⁻¹, as measured with a quantum irradiance meter
128 (LI-250, LiCor Inc., USA).

129

130 Microsensor measurements

131 Measurements of O₂ concentration above the thallus of *F. vesiculosus* were done with
132 Clark type O₂ microelectrodes (tip diameter 10 μm, OX10, Unisense A/S, Denmark;
133 Revsbech 1989) with a response time of 1-3 seconds and low stirring sensitivity (<2%).
134 The microelectrode was connected to a pA meter (PA2000, Unisense A/S, Denmark) and
135 sensor signals from the pA meter were acquired on a PC via a parallel port -connected
136 A/D converter (ADC-101, Pico Technologies Ltd., England). The O₂ microsensor was
137 mounted in a custom built micromanipulator setup enabling motorized positioning at
138 defined x, y and z coordinates at ~1 μm resolution by use of 3 interconnected motorized

139 positioners (VT-80, Micos GmbH, Germany) and controllers (MoCo DC, Micos GmbH,
140 Germany). Data acquisition and positioning was controlled by a custom made software
141 (Volfix) programmed in LabView (National Instruments, Japan).
142 The O₂ microelectrode signal was linearly calibrated at experimental temperature
143 (~17°C) and salinity (S=16) from measurements in air saturated seawater and in seawater
144 made anoxic by addition of sodium dithionite. The O₂ concentration in air saturated
145 seawater, C₀, and the molecular diffusion coefficient of O₂, D₀, in seawater at
146 experimental temperature and salinity was taken from tabulated values (Unisense A/S,
147 Denmark) as C₀ = 274 μmol O₂ L⁻¹ and D₀ = 1.87·10⁻⁵ cm² s⁻¹.

148

149 Mapping of diffusive boundary layers

150 The diffusive boundary layer (DBL) around tufts of hyaline hairs anchored in
151 cryptostomata of *F. vesiculosus* was mapped as 2D transects and 3D grid measurements
152 of O₂ concentration profiles. The Volfix software enabled us to specify a measuring
153 grid/transect with any number of sampling points in the x, y and z-directions. In this study,
154 the y-direction corresponds to the direction of flow (where negative values indicate
155 distance behind a single tuft of hyaline hair), the x-direction corresponds to the width of
156 the flow chamber, and the z-direction corresponds to the height above the thallus surface
157 (Fig. S1). The approximate height and radius of the hyaline hairs was approximated by
158 manual manipulation of the microelectrode tip relative to the structures as observed under
159 a stereomicroscope (SV6, Zeiss, Germany). Thallus samples were placed in the flow
160 chamber with the length of the thallus oriented along the direction of flow, i.e., the y-
161 direction.

162 *2D DBL transects.* For 2D transect measurements, the O₂ microelectrode tip was
163 positioned manually as close to the centre of the selected cryptostomata with hyaline hairs
164 as possible using a dissection microscope for observation; this position was set to y=0 in
165 the Volfix measuring software. The transect measurements started 2 mm upstream (y=2
166 mm) from the cryptostomata and 1 mm above the hyaline hairs (Point A in Fig. S1D),
167 and ended 4 mm downstream (y=-4 mm). Transects of O₂ concentration profiles were
168 measured at a lateral resolution of 0.5 mm in the y-direction, with vertical O₂
169 concentration profiles done at each transect point in steps of 0.1 mm in the z-direction.
170 All profile measurements started at the same z-position and were finished in the upper

171 thallus layer, where a characteristic jump in the O₂ concentration, due to the physical
172 impact of the O₂ microsensor and the solid thallus surface, enabled precise determination
173 of the thallus surface.

174 *3D DBL grids.* For measuring 3D grids of O₂ concentration profiles over thallus areas
175 with only one tuft of hyaline hairs, 9 transects covering a 24 mm² sampling grid area
176 around a central tuft of hairs was set up (Fig. S1E) for O₂ measurements at 0.5 mm lateral
177 resolution (x and y direction) and 0.1 mm vertical resolution (z-direction). Measurements
178 started ~1 mm above the hyaline hairs (Fig. S1E).

179 For measurements of 3D grids of O₂ concentration profiles over larger thallus areas with
180 multiple tuft of hyaline hairs, an oblong 12 mm by 2 mm measuring grid was set up.
181 Oxygen measurements were done at a lateral resolution of 0.5 mm (x and y directions)
182 and a vertical resolution of 0.2 mm (z direction). Tufts of hyaline hair were scattered
183 across the thallus, and the starting point of the grid measurement was set randomly, but
184 with the same starting point for both the 1.65 cm s⁻¹ and 4.88 cm s⁻¹ measurements.

185 Due to the length of measurements different thallus samples were used for individual
186 experiments.

187

188 Diffusive boundary layer thickness and calculations

189 There are different ways of determining the effective thickness of the diffusive boundary
190 layer from O₂ microsensor measurements [17]. The DBL thickness is often found by
191 extrapolating the linear O₂ gradient in the DBL to the bulk concentration of the free-flow
192 region. The distance from the surface to the intersection of the extrapolated linear gradient
193 and the bulk concentration is denoted the effective diffusive boundary layer thickness, Z_δ
194 [3]. However, analysing large numbers of O₂ profiles in this manner is very time
195 consuming, and a somewhat faster determination can be done by defining Z_δ as the
196 distance between the surface and the vertical position above the surface where the O₂
197 concentration has changed 10% relative to the O₂ concentration in the bulk water.
198 Estimations of Z_δ via this method were found to differ <10% from more precise
199 determinations [17].

200 The diffusive flux of O₂ across the DBL, J (in units of nmol O₂ cm⁻² s⁻¹) was calculated
201 from steady state O₂ concentration profiles using Fick's 1st law:

202

203 $J = D_0(C_\infty - C_0)/Z_\delta$ (Eq. 1)

204

205 where C_∞ is the O_2 concentration in the free-flow region ($\mu\text{mol } O_2 \text{ L}^{-1} = \text{nmol } O_2 \text{ cm}^{-3}$),
206 C_0 is the O_2 concentration at the thallus surface ($\mu\text{mol } O_2 \text{ L}^{-1}$), Z_δ is the effective DBL
207 thickness (cm), and D_0 is the molecular diffusion coefficient of O_2 in seawater ($\text{cm}^2 \text{ s}^{-1}$).
208 To compensate for the uneven surface of thalli, measurements below the thallus surface
209 are not shown on final transects. The depth axis on transects is therefore denoted $z' = z -$
210 z_0 , where z is the z -coordinate from the sample data, and z_0 is the z -coordinate of the
211 thallus surface. The thallus surface position was determined from the intermittent sudden
212 drop in O_2 concentration when the microsensor pushed against the thallus surface cortex.
213 Maps of O_2 concentration, Z_δ , and J were generated from measured transects and grids
214 using data interpolation software (Kriging gridding method using default settings, i.e.
215 Linear Variogram and Point Kriging, Surfer v.8, Golden Software Inc., USA).

216

217 Statistics

218 Two-way ANOVAs were applied to test differences in the mean Z_δ over *F. vesiculosus*
219 between flow rates and light conditions (light/dark). For significant main effects (flow
220 rate and/or light condition) and interaction effects, Tukey's multiple comparisons post
221 hoc test was applied. Two-way ANOVAs were applied to test differences between mean
222 O_2 flux values between flow rates and thallus condition (single or multiple tufts). For
223 significant main effects (flow rate and/or thallus condition) and interaction effects,
224 Tukey's multiple comparisons post hoc test was applied. Statistical analysis was
225 performed using Rstudio (Rstudio version 0.99.491, 2016) with the level of significance
226 set to $p < 0.05$.

227

228 **Results**

229 The DBL around single tufts of hyaline hairs

230 Isopleths of O_2 concentration in the water column above the thalli showed a local increase
231 in effective DBL thickness, Z_δ associated with the hyaline hair tuft (Fig. 1), with highest
232 Z_δ values located downstream from the hyaline hairs, either directly behind the tuft or
233 even within the expanse of the hyaline hairs. In light ($350 \mu\text{mol photons m}^{-2} \text{ s}^{-1}$), and at a
234 flow of 1.65 cm s^{-1} , Z_δ reached a maximum thickness of 1.2 mm downstream relative to

235 the tuft at $y = -0.5$ mm (Fig. S2, S3A), while under a flow of 4.88 cm s^{-1} , the maximum
236 Z_{δ} was reduced to 0.6 mm at $y = -2.5$ mm (Fig. S3A). In darkness, the maximal Z_{δ} values
237 were 0.9 mm and 0.4 mm at $y = -2$ mm and $y = 0$ mm under flows of 1.65 and 4.88 cm s^{-1} ,
238 respectively. The mean Z_{δ} did not change significantly ($p < 0.05$) between measurements
239 in light and darkness (Fig. S3B) under low flow ($Z_{\delta(\text{Light})} = 0.72$ mm, $Z_{\delta(\text{Dark})} = 0.58$ mm;
240 $p \text{ adj} = 0.26$) or high flow ($Z_{\delta(\text{Light})} = 0.36$ mm, $Z_{\delta(\text{Dark})} = 0.18$ mm; $p \text{ adj} = 0.13$). However,
241 flow velocity had a significant effect on the mean Z_{δ} that was significantly thinner under
242 high flow (4.88 cm s^{-1}) as compared to the low flow (1.65 cm s^{-1}) ($p \text{ adj} < 0.001$ for both
243 main effects).

244 The hyaline hairs affected Z_{δ} downstream from the hair tuft (Fig. 1A,B) causing a
245 thickening of the DBL that also expanded perpendicular to the flow direction (Fig. 1C,D),
246 reaching a maximum expansion at $y = -2$ mm at both flows ($Z_{\delta\text{Max}} = 1.8$ mm for 1.65 cm s^{-1}
247 and $Z_{\delta\text{Max}} = 0.9$ mm for 4.88 cm s^{-1}). Beyond the local peak in boundary layer thickness,
248 the DBL closely followed the contours of the thallus surface topography. Two transects
249 measured at higher resolution along the x-axis at $y = -2$ mm showed that the increase in Z_{δ}
250 was roughly identical and extended ~ 1 mm on both sides of the hyaline hair tuft (Fig.
251 1C,D). At distances > 1 mm away from the local maximum, the DBL thickness
252 approached a lower more homogeneous DBL thickness over thallus areas unaffected by
253 the hair tuft.

254 Unexpectedly, a transect of O_2 concentrations above the illuminated thallus at $y = -2$ mm
255 in the x direction, i.e., perpendicular to the flow, showed a local O_2 increase followed by
256 a decrease in oxygen concentration that was apparently independent of Z_{δ} (Fig. 1C). A
257 longitudinal transect at $x = -0.5$ mm along the y-direction, i.e., the flow direction, showed
258 further indications of an apparent local “upwelling” of O_2 into the transition zone between
259 the DBL and the fully mixed water column protruding downstream from the hyaline hair
260 tuft (Fig. 2). We found such “upwelling” zones most pronounced under low flow located
261 ~ 2 mm downstream from the centre of the tuft and extending several mm into the water
262 column with O_2 concentrations reaching up to > 2 times air saturation in some cases (Fig.
263 2A).

264

265 The DBL around multiple tufts of hyaline hairs

266 3D grid measurements of O₂ concentration over a *F. vesiculosus* thallus with several tufts
267 of hyaline hairs spaced at approximately 2-5 mm distance were done to investigate
268 potential combined effects of multiple tufts on the DBL. Such measurements showed that
269 the smooth local thickening of the DBL around a single hyaline tuft relative to the DBL
270 of the smooth thallus was altered in the presence of multiple tufts (Fig. 3). The DBL
271 topography was more heterogeneous with Z_δ varying >1 mm reaching a maximum
272 thickness of >2.5 mm under low flow and >1.5 mm under high flow, respectively. The
273 DBL topography was apparently largely determined by the interaction between flow and
274 the tufts of hyaline hairs under low flow, while the smooth thallus surface topography led
275 to local minima in Z_δ in-between individual tufts at higher flow velocity (Fig. 3B).

276 Transects of O₂ concentrations at x=1 mm (extracted from the 3D grids in Fig. 3A,B)
277 gave a detailed information on how O₂ concentration varied over the thallus with distance
278 along the thallus in the flow direction (Fig. 3C,D). In light, the thallus surface O₂
279 concentration reached >900 μM in both flows, while the O₂ concentration in the transient
280 zone of the DBL (z=0.7 mm) varied between 350 and 750 μM under low flow and
281 between 300 and 550 μM in high flow, respectively, clearly demonstrating a compression
282 of the boundary layer and more effective O₂ exchange between thallus and water under
283 higher flow.

284 However, the thickening of the 300-350 μM O₂ contour areas e.g. at y=-11 mm and y=-
285 6 mm (Fig. 3C) was due to gradual increasing O₂ concentrations from the bulk water
286 towards the upper part of the DBL (data not shown). This creates an artefact in the precise
287 determination of Z_δ by the method proposed by Jørgensen and Des Marais [17] that will
288 overestimate the local DBL thickness e.g. compared to the local profile in y=-1 mm where
289 a more steady O₂ increase was measured.

290

291 Diffusive O₂ fluxes over the *Fucus* thallus with single and multiple tufts

292 Although inconsistencies were found (e.g. in the area around x = -1.5 mm, y = 1.5mm in
293 Fig. 4A,B), increases in DBL thickness generally correlated with a decrease in O₂ flux,
294 and the flow-dependent DBL topography strongly affected the O₂ flux from the
295 illuminated *F. vesiculosus* thallus.

296 Comparing the O₂ fluxes in transects over the *F. vesiculosus* thallus with single and
297 multiple tufts of hyaline hairs (Fig. 5A,B) showed an increased O₂ flux just upstream to
298 the position of the hyaline hair tufts independent of the flow velocity. The flux values
299 generally correlated with the DBL thickness and the flux gradually decreased downstream
300 relative to the hair tuft. However, some local variations were seen in areas exhibiting less
301 uniform increases in O₂ concentration towards the thallus surface.

302 The average O₂ flux calculated from transects over *F. vesiculosus* thalli with single and
303 multiple hyaline hair tufts (Fig. 5C,D) showed that flow was the major determinant of gas
304 exchange between macroalga and the surrounding seawater. Both in measurements over
305 single and multiple hair tufts, the O₂ flux values were higher in high flow treatments as
306 compared to low flow (p adj <0.001). The O₂ fluxes measured around a single hair tuft
307 under high flow were higher than the corresponding measurements over multiple tufts
308 treatment (Fig. 5C,D; p adj <0.001). However, the flux values in the multiple tuft
309 treatments were averaged over a two times larger distance (Fig. 5C; 12 mm), and thus
310 include the combined effect of multiple tufts and DBL variation over these, while the
311 values of the single tuft treatments (Fig. 5D; 6mm) only reflect the DBL effects on O₂
312 flux immediately downstream from the hair tuft.

313

314 **Discussion**

315 In our measurements around single tufts of hyaline hairs, the DBL followed the contour
316 of the smooth thallus surface, except around the hair tufts where a thickening occurred
317 downstream of tufts and perpendicular to the flow direction, with a concomitant decrease
318 in the thickness 1-2 mm away from the hair tufts, depending on the flow-regime.

319 In measurements over multiple tufts, the smooth thickening of the DBL observed around
320 isolated single tufts was absent. This more dynamic DBL landscape is probably caused
321 by the close vicinity of neighbouring hyaline hair tufts, where the DBL thickness between
322 tufts never reach 'normal' conditions over a smooth thallus. Interestingly, this suggests
323 that the effect of multiple hair tufts overall increased the DBL thickness across the thallus.
324 Intuitively, a thin boundary layer would create physical conditions that could better avoid
325 high detrimental O₂ concentrations and inorganic carbon limitations in light and O₂
326 limitation in darkness. So why does *Fucus* spend metabolic energy on production of
327 hyaline hairs? In early work by Raven [11], it was suggested that hyaline hairs aid in

328 nutrient uptake by having a highly decreased diffusion resistance over the plasmalemma
329 compared to the thick algal thallus, and in addition the hairs could protrude through the
330 viscous sublayer and into the mainstream flow with better nutrient access. However, as
331 pointed out by Hurd [14], the thin and flexible hairs are considered unlikely to disrupt the
332 viscous sublayer and create turbulence themselves. Here we show that across a thallus
333 with multiple hair tufts, the overall DBL thickness is increased, which has a functional
334 significance similar to observed DBL effects of epiphytes on submerged macrophytes
335 [24, 25]. The thickening of the boundary layer thickness creates a mass transfer limitation
336 that in light can lead to high O₂ concentrations [24, 26] potentially inducing
337 photorespiration [27] and limiting the inorganic carbon supply [28] to the thallus.
338 However, such mass transfer limitation would also maintain higher nutrient
339 concentrations due to surface-associated enzyme activity that can aid in the uptake of e.g.
340 phosphorous and other nutrients [14, 29].

341 A thicker diffusive boundary layer over thalli with tufts of hyaline hairs could also create
342 a niche for epibiotic bacteria, and the presence of bacteria on algal thalli is well known
343 [30-32]. In light of the recently developed ‘holobiont’ concept [32, 33] a physical
344 structure that would keep the algal associated bacteria more protected and provide them
345 with metabolic compounds, could present a competitive advantage. Studies of the role of
346 bacteria in algal life-cycle and metabolism have shown that a strong host specificity of
347 epiphytic bacterial communities exists, possibly shaped by the algal metabolites as the
348 primary selective force [34]. Previous studies have e.g. demonstrated the presence of N₂
349 fixing cyanobacteria as part of the algal microbiome, and it has also been shown that
350 native bacteria are required for normal morphological development in some algae [35].
351 However, the actual distribution and ecological niches of such macroalgae-associated
352 microbes are not well studied. Spilling et al. (2010) found more pronounced O₂ dynamics,
353 reaching anoxia during darkness, in the cryptostomata cavities of *F. vesiculosus*, wherein
354 the hyaline hairs are anchored. Cryptostomata could thus represent potential niches for
355 bacterial aerobic and anaerobic degradation of organic substrates or O₂-sensitive N₂
356 fixation that warrant further exploration.

357 In some DBL transects measured on light exposed *Fucus* thalli, we observed areas of
358 enhanced O₂ concentration detached from the boundary layer (Fig. 2). In a previous study,
359 it was shown that nutrient uptake rates could increase 10-fold when the boundary layer

360 was periodically stripped by passing waves [36]. However, in our case the flow upstream
361 from the tuft was laminar and no waves or DBL stripping occurred. The observed
362 phenomenon of enhanced O₂ above the DBL could be explained by a combination of
363 factors. As flow is obstructed by a physical object a differential pressure field is created
364 where a local drop in pressure is created around the hyaline hairs due to the locally smaller
365 cross section of unobstructed flow. Such pressure gradient could create a local advective
366 upwelling around the area of low pressure thus affecting the O₂ transport. The phenomena
367 is well described in e.g. sediment transport [37] and plumes of O₂ release have also been
368 observed before in coral-reef-associated algae *Chaetomorpha sp.* using planar optodes
369 [38].

370 In addition, so-called vortex shedding (von Kármán vortex sheets) could also be a factor
371 influencing the observed O₂ release. Shedding of vortices can occur at certain Reynold
372 numbers at the transition between laminar and turbulent flow when the pressure increases
373 in the direction of the flow, i.e., in the presence of a so-called adverse pressure gradient
374 [39]. In our study, the flow upstream from the hyaline hairs was laminar but a transition
375 to turbulent flow can occur, even at low Reynold numbers, when a certain surface
376 roughness is present and vortex shedding can initiate at Reynold numbers of ~50 [40].
377 Using characteristic scales from this study (hyaline hair tuft diameter = 2 mm; free-stream
378 velocity = 1.65 or 4.88 cm s⁻¹; fluid density = 1 kg L⁻¹ and a dynamic fluid viscosity of
379 1.08 × 10⁻³ Pa s [41], we calculated Reynold numbers of ~30-90. Von Kármán vortices
380 have previously been connected to flow patterns on the lee side of plant parts [42] and
381 based on the calculated Reynold numbers the theoretical basis for generation of vortex
382 shedding [40] due to tufts of hyaline hairs is present in our experimental setup.

383 We thus speculate that a combination of pressure gradient mediated upwelling of O₂ and
384 vortex shedding (Fig. 6) could explain the observed phenomena in Fig. 2. The local mass
385 transfer related to the presence of hyaline hair tufts on Furoid macroalgae may thus be
386 more complex. A more detailed investigation of such phenomena was beyond the scope
387 of the present study and would clearly require more detailed characterization of the
388 hydrodynamic regimes over thalli with and without tufts of hyaline hairs.

389 In conclusion, our study of the chemical boundary layer landscape over the thallus of *F.*
390 *vesiculosus* revealed a strong local effect on the DBL over and around tufts of hyaline
391 hairs anchored in cryptostomata. Single tufts showed a thickening of the DBL

392 downstream and horizontally relative to the thinner DBL over the smooth thallus surface,
393 while areas with multiple tufts exhibited a consistently thickened DBL that may affect
394 gas and nutrient exchanges between the alga and seawater. Furthermore, we also observed
395 more complex solute exchange phenomena that were apparently driven by pressure
396 gradients and/or vortex shedding over the hyaline hair tufts. Altogether, this study
397 demonstrates that interactions between flow and distinct macroalgal surface topography
398 gives rise to local heterogeneity in the chemical landscape and solute exchange that may
399 allow for microenvironmental niches harbouring microbial epiphytes facilitating a
400 diversity of aerobic and anaerobic processes. Further microscale studies of such niches in
401 combination with e.g. microscopy and molecular detection of microbes in relation to
402 hyaline hairs and cryptostomata thus seem an important next step to reveal further insights
403 to the presence and role of the microbiome of Furoid algae.

404

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412

413 **Author contributions**

414 RDN and MK designed the research, RDN and ML performed the research, ML, RDN
415 and MK analysed data, ML wrote the paper with editorial assistance from RDN and MK.
416 All authors gave final approval for publication.

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Figure legends

Figure 1. Fine-scale mapping of DBL around a single tuft of hyaline hairs on an illuminated *Fucus vesiculosus* thallus ($350 \mu\text{mol photons m}^{-2} \text{s}^{-1}$). (A) and (B) 3D plots of *F. vesiculosus* thallus surface (grey area) and upper extension of DBL (coloured area) around a single tuft of hyaline hairs, at flow velocities of 1.65 (left panels) and 4.88 cm s^{-1} (right panels). Colour bars depict the effective DBL thickness, Z_δ (mm), and arrows indicate the direction of flow. (C) and (D) Transects in the x-direction (perpendicular to the flow), at position $y=-2$ mm from Fig. 1 A,B, respectively, normalized to thallus surface showing the local O_2 concentration. The zero position (0,0) indicates the position of the cryptostomata. Colour bars denote O_2 concentration (in $\mu\text{mol O}_2 \text{L}^{-1}$). (E) and (F) Transects of O_2 concentration (in $\mu\text{mol L}^{-1}$) measured across a single tuft of hyaline hairs in *Fucus vesiculosus* measured at flow velocities of 1.65 (E) and 4.88 cm s^{-1} (F), in light ($350 \mu\text{mol photons m}^{-2} \text{s}^{-1}$). The arrows indicate flow direction. The zero position (0,0) indicates the position of the cryptostomata, and transects were adjusted to the thallus surface. Colour bars denote O_2 concentration (in $\mu\text{mol O}_2 \text{L}^{-1}$).

Figure 2. Local transects of O_2 concentration around single hair tufts from two different measurement series over an illuminated *Fucus vesiculosus* thallus ($350 \mu\text{mol photons m}^{-2} \text{s}^{-1}$). (A) shows a transect taken from Fig. 1A at $x=-0.5$ under a flow velocity of 1.65 cm s^{-1} , while (B) was measured similarly as Fig. 1E, also at a flow velocity of 1.65 cm s^{-1} . The hair tufts were 2.5-3 mm in diameter and protruded 3-3.5 mm from the thallus. Both transects were adjusted to the thallus surface. The black arrow indicates the flow direction. Colour bars denote O_2 concentration (in $\mu\text{mol O}_2 \text{L}^{-1}$).

Figure 3. Mapping of DBL over several tufts of hyaline hairs on an illuminated *Fucus vesiculosus* thallus ($350 \mu\text{mol photons m}^{-2} \text{s}^{-1}$). (A) and (B) 3D plot of *F. vesiculosus* thallus surface (grey area) and the upper extension of the DBL (coloured area) of multiple tufts of hyaline hairs under a flow velocity 1.65 (left panels) and 4.88 cm s^{-1} (right panels). Colour bars depict the effective DBL thickness, Z_δ (mm). (C) and (D) Transects of O_2 concentration at position $x=1$ (along the y-axis direction) normalized to thallus surface. Colour bars denote O_2 concentration (in $\mu\text{mol O}_2 \text{L}^{-1}$).

Figure 4. Isoleths of O₂ flux (in nmol O₂ cm⁻² s⁻¹) (A) and (C) and effective DBL thickness, Z_δ (in mm) (B) and (D) measured over an illuminated *Fucus vesiculosus* thallus (350 μmol photons m⁻² s⁻¹) around a single tuft of hyaline hairs at flow velocities of 1.65 cm s⁻¹ (A, B) and 4.88 cm s⁻¹ (C, D). The hyaline hairs were rooted in the cryptostomata located at the (0,0) coordinate, as indicated by the black cross. Black arrows indicate the flow direction.

Figure 5. Comparison of O₂ flux values (in nmol O₂ cm⁻² s⁻¹) calculated from transects of O₂ concentration profiles measured over an illuminated (350 μmol photons m⁻² s⁻¹) intact *Fucus vesiculosus* thallus with several tufts of hyaline hairs (A) and a thallus with only a single hair tuft (B) measured under flow velocities of 1.65 cm s⁻¹ and 4.88 cm s⁻¹. Note the difference in x-scale. The black arrow indicates the flow direction. The individual position of the multiple hairtufts in panel (A) were not mapped and the zero position on the x-axis thus only reflects the starting point of the transect. In panel (B), the zero position indicates the centre of the cryptostomata. Panel (C) and (D) show the average O₂ flux (±SEM) across (C) the intact thallus and (D) the thallus with a single hair tuft protruding.

Figure 6. Conceptual drawing showing possible scenarios for the observed upwelling of O₂ downstream of the hyaline hairs. Flow velocity (straight black lines) decreases from the free stream velocity as the thallus surface is approached through the diffusive boundary layer (DBL). The hyaline hair tuft protruding from cryptostomata alters the DBL thickness locally and creates a differential pressure field (shown in gradient blue and red colours) due to the smaller cross section of unobstructed flow. This creates a flow acceleration in the areas of relative low pressure which could result in advective upwelling. In addition, an adverse pressure gradient is generated downstream from the hair tuft potentially resulting in vortex shedding.

Figures

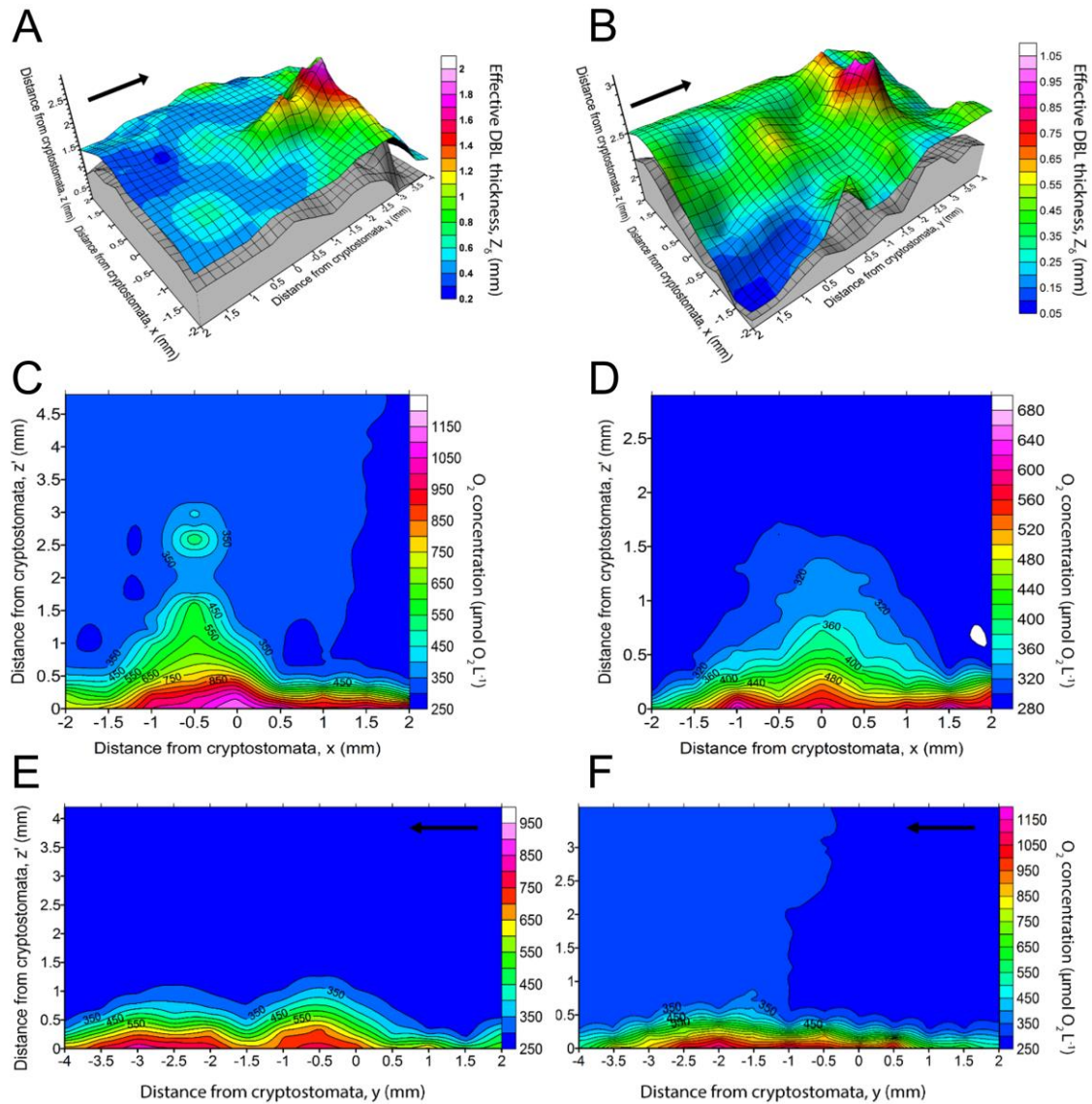


Figure 1

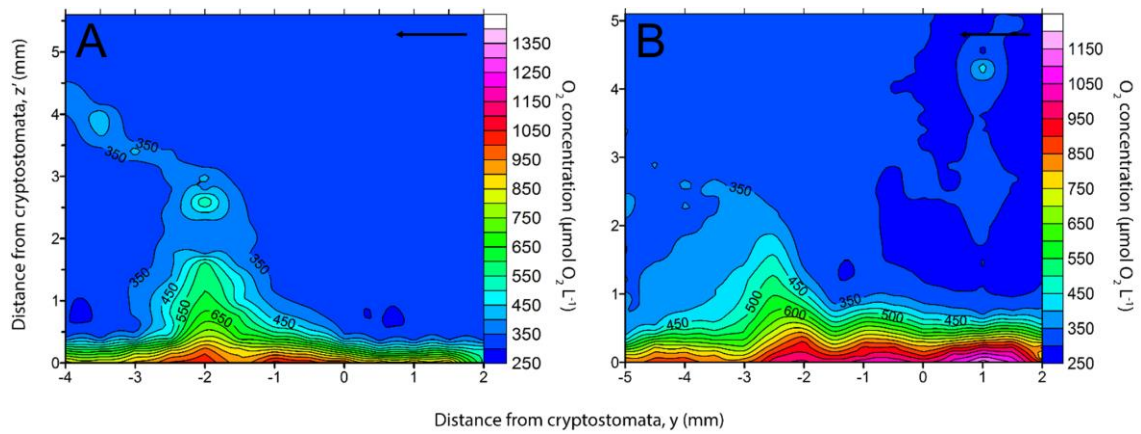


Figure 2

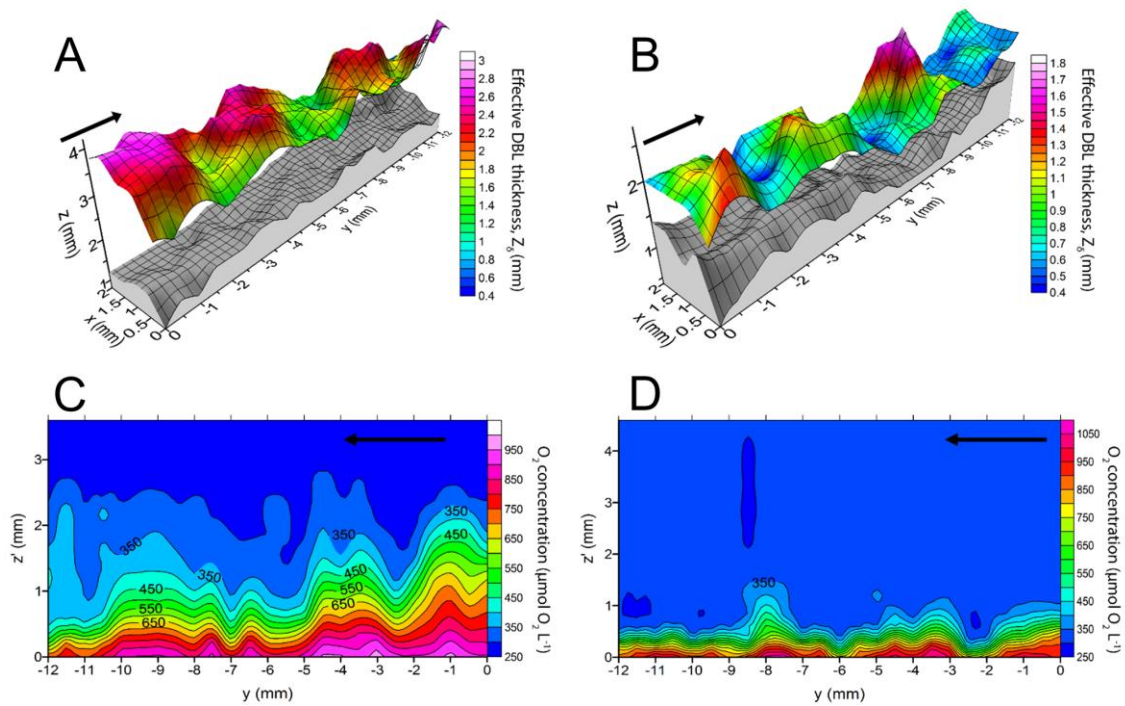


Figure 3

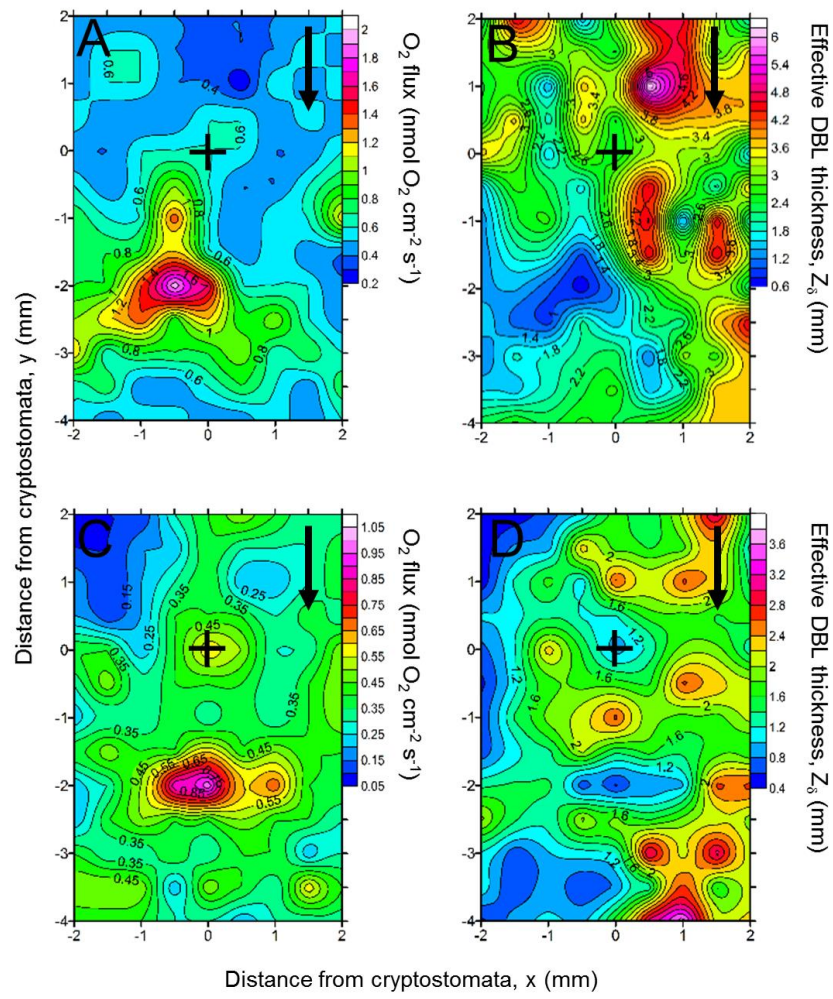


Figure 4

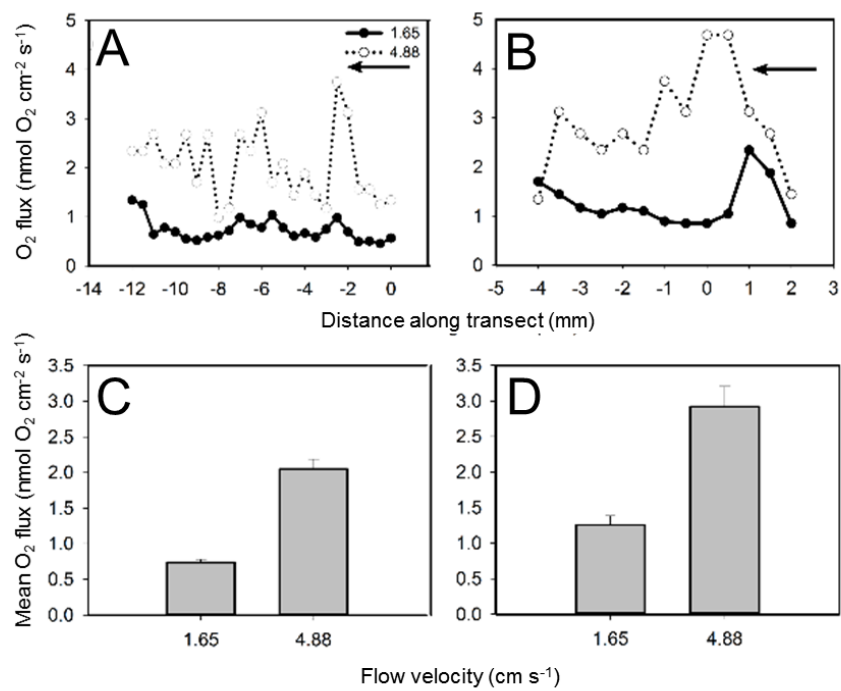


Figure 5

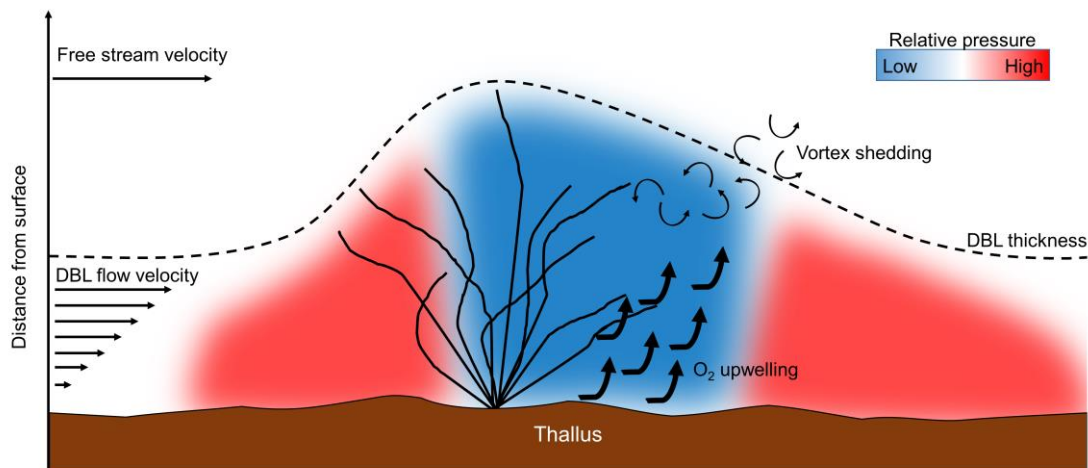


Figure 6