1	The diversity and functional capacity of microbes associated with coastal phototrophs
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17	
18	Abstract
19	Coastal marine phototrophs exhibit some of the highest rates of primary productivity in the
20	world. They have been found to host a diverse set of microbes, many of which may impact the
21	biology of their phototroph hosts through metabolisms that are unique to microbial taxa. Here we
22	characterized the metabolic functions of phototroph-associated microbial communities using
23	metagenomes collected from 2 species of kelp (Laminaria setchellii and Nereocystis luetkeana)

24 and 3 marine angiosperms (Phyllospadix scouleri, P. serrulatus and Zostera marina), including 25 the rhizomes of two surfgrass species (*Phyllospadix* spp.) and the seagrass Zostera marina, and 26 the sediments surrounding *P. scouleri* and *Z. marina*. Using metagenomic sequencing, we 27 describe 72 metagenome assembled genomes (MAGs) that potentially benefit from being 28 associated with macrophytes and may contribute to macrophyte fitness through their metabolic 29 gene content. All host-associated metagenomes contained genes for the use of dissolved organic 30 matter from hosts and vitamin (B_1, B_2, B_7, B_{12}) biosynthesis. Additionally, we found a range of 31 nitrogen metabolism genes that transform dissolved inorganic nitrogen into forms that may be 32 more available to the host. The rhizosphere of surfgrass and seagrass contained genes for 33 anaerobic microbial metabolisms, including *nifH* genes associated with nitrogen fixation, despite 34 residing in a well-mixed and oxygenated environment. The range of oxygen environments 35 engineered by macrophytes likely explains the diversity of both oxidizing and reducing microbial 36 metabolisms, and contributes to the functional capabilities of microbes and their influence on 37 carbon and nitrogen cycling in nearshore ecosystems.

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Importance

Kelps, seagrasses and surfgrasses are ecosystem engineers on rocky shorelines where they show remarkably high levels of primary production. Through analysis of their associated microbial communities, we found a variety of microbial metabolisms that may benefit the host, including nitrogen metabolisms and the production of B vitamins. In turn, these microbes have the genetic capability to assimilate the dissolved organic compounds released by their phototroph hosts. We describe a range of oxygen environments associated with surfgrass, including low-oxygen microhabitats in their rhizomes that host genes for nitrogen fixation. The tremendous

47 productivity of coastal phototrophs is likely due in part to the activities of associated microbes48 and an increased understanding of these associations is needed.

- 49
- 50

Introduction

51 We are experiencing a paradigm shift in biology with the recognition that many species exist as a 52 consortium with microbes (1). These microbial associations are nearly ubiquitous, spanning a 53 diversity of hosts across ecosystems. In coastal marine environments, phototrophic microbial 54 hosts are diverse and range from marine angiosperms to large eukaryotic protists (macroalgae). 55 Different macroalgal host species (2, 3) and different phototroph tissues (4, 5) host distinct microbial communities numbering in the millions per cm^2 of host tissue (6), yet we still know 56 57 little about the functional role the microbiome plays in host fitness or how the host influences the 58 microbiome. The microbiome of phototroph species has been shown to have metabolisms that 59 provide nitrogen to the host (7, 8). Bacteria also supply B vitamins (9) and affect development of 60 their host (10). Further, the contributions that marine phototrophs make to host carbon and 61 nitrogen cycling have largely ignored the role that microbes play. Even as we begin to describe 62 their microbiome, we are discovering that environmental change affects these communities (11). 63 For many of the foundational phototrophic species in the coastal ocean, our understanding of the 64 diversity and role of their microbiome is nascent.

65

A unique aspect of host-associated microbes are the strong gradients in oxygen that they
experience due to the biological activities of the host. The photosynthetic and respiratory
activities of the host can generate a 'phycosphere' (12) where the host influences the physical
environment experienced by microbes, sometimes over micron or mm scales. For example, the

basal leaf meristem of the seagrass *Zostera* ranges from oxic to anoxic conditions over a scale of
300 microns when measured with oxygen microsensors (13). This range of oxygen
concentrations likely selects for a diversity of microbial metabolisms in association with
macrophytes.

74

75 Another factor important to the microbial metabolisms associated with coastal macrophytes is 76 nutrient availability. In coastal systems, nitrogen can limit primary production and microbial 77 associates that aid in accessing nitrogen might be selected. Microbial metabolisms that can 78 increase the available dissolved inorganic nitrogen (DIN) for the host (14) include pathways that 79 cleave carbon-nitrogen bonds to generate ammonium. This ammonification in biological systems 80 can result from a diversity of hydrolases, including ureases and other enzymes that cleave C-N 81 bonds (15). Further, microbes that fix atmospheric nitrogen have been discovered in an 82 increasing number of taxa (8, 16), now recognized to include heterotrophic as well as 83 phototrophic taxa (17–19). Nitrogen fixation was previously assumed to be restricted to nitrogen-84 poor environments, but has been quantified recently in systems thought to be nitrogen-rich (8, 85 20), an enigmatic finding given that nitrogen fixation is a costly metabolic process that consumes 86 16 ATPs per N₂ fixed (21). Sediments where oxygen is low and nutrients can be depleted by 87 macrophytes, such as the rhizosphere of seagrasses have provided evidence of nitrogen fixation 88 (22–26). The recent discovery that nitrogen fixation takes place on particles in the coastal ocean 89 where nitrate is relatively abundant (8, 20) suggests that *nifH* genes could be abundant in other 90 nearshore systems.

92	Microbial metabolisms that synthesize compounds and vitamins needed by seaweeds and
93	seagrasses may also underlie host-microbe exchanges. The active form of Vitamin B1 (thiamin)
94	is essential for all organisms and is involved in carbohydrate and amino acid metabolisms.
95	Vitamin B2 (riboflavin)-binding proteins are co-enzymes in various oxidases and are involved in
96	photosynthesis and phototropism (27). Vitamin B7 (biotin) is a cofactor for acetyl coenzyme A
97	(coA) which is essential for fatty acid synthesis. Vitamin B12 (cobalamin) is required as a
98	coenzyme in the mitochondria for many algae, yet they depend upon prokaryotes to produce it
99	(9, 28). Thus, marine macrophytes may be auxotrophic for key vitamins, and their production by
100	host-associated bacteria may be another basis for phototroph-microbiome interactions in nature.
101	
102	Hosts might reciprocally benefit microbes, especially if heterotrophic microbes benefit from the
102 103	Hosts might reciprocally benefit microbes, especially if heterotrophic microbes benefit from the dissolved organic carbon that is released by their hosts. Of the carbon that is fixed, kelp have
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103 104 105 106 107 108	dissolved organic carbon that is released by their hosts. Of the carbon that is fixed, kelp have been demonstrated to release 15-16% of it as dissolved organic matter (29, 30), and seagrasses too provide a constant source of dissolved organic carbon (31, 32), likely stimulating heterotrophic bacterial processes (33). These rates of organic carbon release, often involving highly labile organic carbon compounds (34), could provide the basis for reciprocal benefits

112 reciprocally benefit host and microbes. We analyzed the surface microbiome on the blade of two

113 kelp species (Laminaria setchellii and Nereocystis luetkeana) and the surfgrass Phyllospadix

114 scouleri, the rhizomes of P. scouleri, P. serrulatus, and the seagrass species, Zostera marina, and

115 the sediment surrounding the rhizomes of P. scouleri and Z. marina. We quantified the variable 116 oxygen environment in the rhizomes of *Phyllospadix spp.* to determine if they allow for aerobic 117 as well as anaerobic metabolisms. We analyzed the microbial taxa present and examined their 118 gene content to estimate their functional and metabolic capacities. We hypothesized that 119 microbial partners: 1) enhance host access to dissolved inorganic nitrogen through nitrogen 120 recycling, ammonification and nitrogen fixation, 2) provision vitamins B₁, B₂, B₇, B₁₂, and 3) use 121 a diversity of abundant dissolved organic carbon exudates from the host. We tested whether 122 microbial taxonomy and function differed across hosts and host tissue types, and whether 123 anaerobic metabolisms were present in low-O₂ environments (e.g., rhizomes and sediment). 124 Through this study, we find that the range of oxygen environments engineered by host 125 phototrophs likely explains the diversity of both oxidizing and reducing microbial metabolisms, 126 and contributes to the functional capabilities of microbes and their influence on carbon and 127 nitrogen cycling in nearshore ecosystems. 128 129 Methods 130 Sampling and DNA Extraction 131 We collected metagenome samples from the surfaces of 5 different phototroph species (Table 132 S1). The surface of *Phyllospadix scouleri* blades, *Laminaria setchellii* fronds and the inner bulbs 133 of Nereocystis luetkeana were swabbed with a sterile swab and brushed with an interdental brush 134 (GUM Proxabrush Go-Betweens). We preserved sections of the rhizomes of *Phyllospadix* 135 scouleri, P. serrulatus and Zostera marina. Sediment surrounding P. scouleri and Z. marina was 136 also collected. All samples were collected from Tatoosh Island, WA, USA (48.393679, -137 124.734617) on 16-17 Jul 2019, except for Z. marina samples which were sampled from West

138	Falmouth Bay, MA, USA (41.60708333, -70.64527778) on 19 Sept 2019. We included samples
139	from the rhizosphere of Z. marina from the Atlantic Ocean as a known positive control for
140	nitrogen fixation (22, 23). Swabs, tissue and sediment were immediately frozen at 20° C and
141	shipped to storage at -80° C. DNA from these collections was extracted with a Qiagen PowerSoil
142	Kit and multiple samples were pooled for each metagenome sample to increase DNA quantity
143	and possible discovery: P. scouleri blade, rhizome and sediment (3 pooled individuals each), P.
144	serrulatus rhizome (3 individuals), L. setchellii blade (3 individuals), N. luetkeana interior bulb
145	(4 individuals), Z. Marina rhizomes and sediment (2 individuals).
146	
147	Shotgun metagenomic sequencing, assembly, and read recruitment
148	The above 8 samples were run over 2 lanes on a HiSeq 2500 (2x150) with TruSeq DNA library
149	preps at Argonne National Laboratory. For each sample, resulting DNA sequences were first
150	quality filtered (35)(Minoche et al. 2011), then assembled with IDBA-UD v1.1.3 (36) (Peng et
151	al. 2012) with a minimum scaffold length of 1 kbp. Metagenomic short reads from each sample
152	were then recruited back to their corresponding assembled contigs using Bowtie2 (37). Samtools
153	(38) was used to generate sorted and indexed BAM files. Anvi'o v7.0 (39) was used as the
154	command line environment for all downstream analyses. 'anvi-gen-contigs-database' was used
155	to generate anvi'o contigs databases, during which Prodigal v2.6.3 (40) identified open reading
156	frames, and 'anvi-run-hmms' was used to identify genes matching to archaeal and bacterial
157	single-copy core gene collections using HMMER (41).

158

159 *Reconstructing metagenome-assembled genomes (MAGs)*

To reconstruct genomes from the assembled metagenomes, we used a combination of automatic binning via CONCOCT v1.1.0 (42), followed by a manual curation of each MAG as outlined by Shaiber et al. 2020 (43). Genome taxonomy was determined using GTDB v.1.3.0 (44), and 'anvirun-scg-taxonomy'. We also inferred gene-level taxonomy using Centrifuge v1.0.4 (45) to aid manual curation.

165

166 Phylogenomic analysis of MAGs

167 To perform a phylogenomic analysis of our MAGs, we recovered amino acid sequences for

168 bacterial single-copy core genes (SCGs) from each genome (except the only archaeal genome in

169 our collection) using the program 'anvi-get-sequences-for-hmm-hits' with the parameter `--

170 hmm-source 'Bacteria_71' on the ribosomal gene set 'Ribosomal L1-L6' and the flag '--

171 concatenate`, which independently aligned each SCG independently using Muscle v3.8.1 (46)

before concatenating them into a final superalignment. We then refined the alignment using

trimAl v1.4.rev15 (47) to remove any position in the alignment if more than 50% of the residues

174 were gap characters. A maximum-likelihood phylogeny was inferred using IQTree (48) with

175 1,000 bootstrap replicates, and a LG+R6 model best fit our data using ModelFinder (49).

176

177 Functional analysis of microbial communities

178 To address the metabolic capabilities of host-associated microbes, we annotated genes in each

179 anvi'o contigs database with 3 different databases using 'anvi-run-kegg-kofams', 'anvi-run-ncbi-

180 cogs', and 'anvi-run-pfams', which used the databases of Kyoto Encyclopedia of Genes and

181 Genomes (KEGG) (50), NCBI's Clusters of Orthologous Genes (COGs) (51) and EBI's Pfam

182 database (52) respectively. We used these annotated genes to test for 1) nitrogen cycling

183	metabolisms, especially those within the nitrogen-fixation pathway, 2) hydrolases, including
184	ureases, as well as ammonia-lyases, to cleave the C-N bonds in amino acids and make
185	ammonium available to the host, 3) vitamin production, namely vitamins B_1 , B_2 , B_7 and B_{12} and
186	4) a set of dissolved organic matter (DOM) transporter genes identified by Poretsky et al. (34)
187	that indicate the ability of the microbial community to assimilate DOM exudates from kelps and
188	surfgrasses. The list of genes used is indicated in Table S4. We additionally developed and used
189	a graph-based algorithm on KEGG definitions for vitamins B1, B2, B7 and B12 to detect the
190	presence of these biosynthetic pathways (Supplementary Code 1). To expand our functional
191	analysis of kelp blade genes, we included 32 MAGs from the surface of <i>N. luetkeana</i> blades that
192	were collected from the same location at the same time using similar methods as those described
193	above (53).

194

195 Phylogenetic analysis of nifH genes

196 To search for *nifH* amino acid sequences in our environmental samples, we identified 9 MAGs 197 which contained *nifH* genes using the KEGG identifier K02588 with e-value < 1e-20. We 198 aligned the AA sequences for these genes against 89 well-characterized reference nifH AA 199 sequences (Table S6) using Muscle v3.8.1 (46) and refined the alignment using trimAl (gap-200 threshold: 0.5) and 'anvi-script-reformat-fasta' (max-percentage-gap: 50%). A maximum-201 likelihood phylogeny was inferred using IQTree (48) with 1,000 bootstrap replicates, and a 202 LG+R5 model best fit our data using ModelFinder (49). nifH genes from the Zostera samples 203 served as positive controls to detect nitrogen fixation genes in other samples. Figures 2, 3, 4 were 204 generated using iTol v5 (54), R v4.0.3 and FigTree respectively. We additionally took tissue 205 samples from *P. scouleri* rhizome (n = 16), basal meristematic region just distal to the sheath

206 (n = 12) and blade 35 cm above the rhizome (n = 12) to quantify stable isotopes of $\delta 15N$ and 207 $\delta 13C$ to look for signatures of nitrogen fixation (methods described in Appendix 1).

208

209 Quantifying the Oxygen Environment

210 We quantified the oxygen concentrations in proximity to *Phyllospadix spp.* rhizomes by

211 comparing dissolved oxygen (DO) concentrations in the surrounding seawater and in the

sediment around the rhizome. We used a Pyro Science Robust Oxygen Probe (OXROB10,

213 FirestingTM, Pyroscience), and repeated measurements around 0900h across 4 days (7-9 June

214 2019, 13 June 2021) within P. scouleri (n = 18) and P. serrulatus (n = 11) rhizomes. Each

215 reading first measured the surrounding seawater after which we gently pushed the tip of the

216 oxygen probe into the sediment and rhizome mass to a depth of 1-3 mm, the typical thickness

217 (*pers. observation*). We let the probe equilibrate and took a reading at 150 sec. This allowed the

218 rhizome oxygen environment to equilibrate after we disturbed the intact rhizome. We compared

219 surrounding water and within-rhizome oxygen using paired t-tests in R.

220

221 Data Availability

In addition to the code available on GitHub (____), the final MAG database files generated in

anvi'o are available on the FigShare repository: (_____). Metagenomic sequence data are

available at the NCBI's Sequence Read Archive under accession no. (submission in progress).

225

226

Results

227 Surfgrass rhizomes have lower oxygen concentrations than surrounding seawater

228	The oxygen environment in the rhizomes differed significantly from that of the surrounding
229	seawater (Fig. 1). Rhizomes maintained a lower dissolved oxygen (DO) concentration than the
230	surrounding seawater for both <i>P. scouleri</i> (n=18, pairwise t-test: <i>p</i> <0.001) and <i>P. serrulatus</i>
231	(n=11, pairwise t-test: p<0.001). P. serrulatus maintained a slightly lower DO concentration in
232	the rhizome at 2.11 mg l^{-1} , compared to 5.61 mg l^{-1} for <i>P. scouleri</i> . However, the nature of
233	sampling likely introduced more oxygenated water from the surrounding water column to the
234	rhizome-sediment microenvironment, suggesting that the actual DO concentration within the
235	sediment is lower than the value reported.
236	
237	Diversity of MAGs assembled across hosts
238	Following filtering, we obtained an average of 41 million reads per sample (range 6.48 to 67.73
239	million), with 79.8% of raw reads retained on average. When these reads were assembled into
240	contigs of at least 1000 nucleotides, a mean of 42,026 contigs and a mean of 110,054 genes were
241	present across samples (Table S1).
242	
243	Across 8 metagenomes we manually binned 33 high quality MAGs, defined as having a
244	completion score >90% and contamination (or redundancy) < 10% (Table 2). We also identified
245	39 lower quality MAGs that had completion scores between 38 and 93% and redundancy scores
246	between 0 and 21% (Table S3). All MAGs were bacterial except for a single archaeal MAG on
247	the rhizome of P. scouleri. The bacterial MAGs spanned 7 phyla, including Proteobacteria
248	(n=34), Bacteroidota (n=19), Verrumicrobia (n=2), Campylobacterota (n=3), Desulfobacterota
249	(n=5), and a single MAG in each of <i>Desulfomonadota</i> , <i>Acidobacteriota</i> , and <i>Spirochaetota</i> . The
250	Archaea belonged to the phylum Chrenarchaeota. There were 46 MAGs resolved to the species

level, with 8 to the genus level, 9 to family, 2 to order, and 2 to class level. Five MAGs wereresolved only as Bacteria (Table S2).

253

254 The 72 MAGs belong to diverse microbial phyla, which were distributed across the 5 host

species and tissue types (Fig. 2). In some cases, bacterial taxa from kelp blade tissues were most

256 closely related to bacteria collected from the rhizome or sediment of a seagrass, suggesting that

257 closely related bacterial taxa can associate with diverse hosts. Known anaerobic sulfur cyclers

258 like Desulfobulbia, Desulfobacteria, Desulfuromonadia and Campylobacteria (Sulfurovum

259 sp000296775 and Sulfurimonas autotrophica) were exclusively found in the low oxygen rhizome

and sediment samples of Zostera marina and Phyllospadix spp. Conversely,

261 *Alphaproteobacteria*, were exclusively found on surfaces exposed to the water column.

262 Gammaproteobacteria was the only class found across the range of tissue types (6 out of 8 host

263 environments). We did not include the only well-resolved archaeal taxon found in our samples,

264 *Crenarchaea* (*P. scouleri* rhizome), as our analysis compared single-copy core genes specific to

bacterial phyla.

266

267 Host-associated microbial genomes contain pathways to synthesize vitamins, recycle nitrogen,

268 and use host-generated carbon

269 We found evidence for a number of metabolic pathways that are likely important for exchanges

between host phototrophs and their microbial partners (Fig. 3). Microbes on hosts had genes for

diverse carbohydrate and carboxylic acid assimilation via cell membrane transport proteins.

272 Host-associated microbes also had genes for a diversity of nitrogen metabolisms, including

273 ureases and hydrolases that could regenerate ammonium. Nitrogen metabolisms were most

- diverse in rhizome and sediment samples where we identified both oxidizing (nitrification) and
- 275 reducing (nitrate reduction, nitrogen fixation, denitrification) metabolisms, as well as
- 276 metabolisms that both oxidize and reduce (annamox).
- 277
- 278 Every sample had at least one gene from B-vitamins biosynthesis pathways. Using a simple-path
- 279 based algorithm on KEGG definitions (Supplementary Code 1), we determined that all microbial
- 280 communities had the metabolic pathways to synthesize vitamins B_1 (with the exception of the *P*.
- 281 scouleri rhizome), B_2 and B_7 (except inside the bulb of N. luetkeana). The Vitamin B_{12} anaerobic
- biosynthesis pathway, however, was only present in MAGs found on the blades of L. setchellii
- 283 (2) and *P. scouleri* (3) and the rhizomes of *P. serrulatus* (2) and *Z. marina* (1). Additionally, all
- three MAGs on the blade of *P. scouleri* that had this anaerobic pathway had the genes necessary
- to synthesize Vitamin B_{12} aerobically as well.
- 286

287 Novel detection of nifH genes in surfgrass

288 We identified the nitrogenase gene (*nifH*) in 9 MAGs with e-value support < 1.3e-120 (KEGG) 289 and < 1.1e-135 (COG). These 9 MAGs were assembled from *P. serrulatus* rhizomes (n = 2) and 290 Z. marina rhizomes (n = 3) and the surrounding sediment (n = 4). Of these 9 MAGs, 5 were 291 resolved to the genus level, while others were resolved to the order and family level, including 292 *Campylobacterales*, *Desulfobacterales* and 2 *Flavobacteriaceae* (Fig. 4, Table S5). *nifH* genes 293 identified in the rhizomes of *P. serrulatus* and *Z. marina* belonging to the class *Desulfobacteria* 294 and family *Flavobacteriaceae*, clustered within Cluster III: anaerobic nitrogen-fixers that are 295 often coupled with sulfate-reduction metabolisms. Samples from Z. marina sediment and 296 rhizome also contained 3 nifH genes in Campylobacterial MAGs that clustered together in a

297	sister clade to the aerobic nitrogen-fixers of Cluster I. The COG gene identified as <i>nifH</i>
298	(COG1348) also includes the homologous protochlorophyllides, which are involved in
299	photosynthetic pigment synthesis but have high sequence similarity to the <i>nifH</i> gene (21, 55).
300	Instead, we used the KEGG gene (K02588) that does not detect these homologs. When we
301	inspected genes on the same contig with <i>nifH</i> , we found a number of genes related to nitrogen
302	fixation (Table S5), including nifD (COG 2710) in 7 of the 9 contigs, nitrogen regulatory protein
303	PII (COG 347), nifB (COG 535), and multiple iron containing proteins including ferrodoxin and
304	Fe-Mo cluster-binding proteins (Table S5).
305	
306	Discussion
307	Phototroph tissues and sediment host distinct microbial taxa and functions
308	The phototroph species we sampled in this study are foundational in coastal ecosystems (56–59),
309	yet a description of the diversity and function of their microbiomes have been lacking. All
310	MAGs were bacterial, except for a single archaeal MAG (Crenarchaeota) in the rhizome of
311	Phyllospadix scouleri, which was identified as Nitrosopumulis, a genus associated with
312	nitrification (Table S3). Together, these 5 phototrophs hosted bacteria from 9 phyla. The only
313	low diversity sample was the interior of the bulb of Nereocystis, where we assembled only a
314	single MAG (UBA7415 sp002470515) suggesting that this environment of high carbon
315	monoxide and nitrogen gas (60) may inhibit microbial activity or pose a highly selective
316	environment. Blades of kelp and surfgrass, in contrast, were a locus of microbial diversity and
317	function, a finding that is similar to many recent studies of macroalgal and seagrass microbiomes
318	reporting high microbial diversity (2, 4, 5, 61–63). The functional attributes of microbial taxa
319	associated with marine macrophytes include pathogen resistance (64), the ability to provision the

host with B vitamins (9), and enhanced host algal fitness (65), perhaps through some of thenitrogen metabolisms we documented here (14, 66).

322

323 Host-microbe interactions in a dynamic oxygen microenvironment

Grouping MAGs by microbial metabolisms (Fig. 3) showed key functional differences among 324 325 phototroph hosts. Blade tissues that interacted directly with the water column were associated 326 with microbial nitrogen metabolisms that were mostly oxidizing. The abundance of dissolved 327 organic carbon from phototroph hosts (29-31, 59) might select for heterotrophic metabolisms. 328 Indeed, we found an abundance of genes for dissolved organic matter assimilation and transport 329 in all metagenomes, suggesting that hosts may stimulate heterotrophy in their associated 330 microbial community similar to findings by Poretsky et al. (34). Improved characterization of the 331 components of dissolved organic matter and the genomes of hosts will allow us to better assess 332 complementarity in resource supply by hosts and resource use by microbes.

333

334 The host tissue types in this study differed in surface oxygen concentrations. Blade tissue 335 interacts with the water column and is likely more oxygenated than rhizome tissue or sediments, 336 though a previous study suggests there can also be a 60% reduction in oxygen along the immediate surface of kelp blades (67), and along the mucus layer where some kelp-associated 337 338 bacteria reside (6). Over two-thirds of the bacterial taxa on blades of N. luetkeana belonged to 339 families associated with obligately aerobic metabolisms, demonstrating the role of oxygen in 340 shaping phototroph-associated microbial communities (68). The sediment surrounding the 341 rhizomes of Phyllospadix spp. contained low oxygen microenvironments (Fig. 1) likely 342 maintained by macroinvertebrate respiration (69)(Moulton and Hacker 2011), similar to the

343	biological processes in the anaerobic sediment surrounding Zostera (13). Low rhizosphere
344	oxygen concentrations likely structured the taxonomic composition of Z. marina to include
345	anaerobic taxa such as Campylobacteria, Desulfatitalea and Desulfobulbus. The presence of
346	anaerobes like Desulfuromonadia, Desulfobaceria, Spirochaeta and Aminicenantia in P.
347	serrulatus rhizomes suggests sulfate reduction also occurs, possibly coupled to dissolved organic
348	carbon use as an energy source (e.g. (70) Howarth & Hobbie 1982). Additionally,
349	Campylobacteria and the genus Thiodiazotropha were associated with Z. marina and may
350	remove detrimental sulfide accumulation through sulfur oxidation (71, 72).
351	
352	Nitrogen metabolisms that were both oxidizing and reducing were found in MAGs associated
353	with rhizomes of both Z. marina and Phyllospadix (Fig. 3), suggesting the potential for temporal
354	niches when, for example, ammonium oxidation to nitrate occurs during high-O2 daylight
355	periods, followed by nitrate reduction or nitrogen fixation during O ₂ -depleted nighttime hours.
356	Additionally, all MAGs in this study contained hydrolases that cleave carbon-nitrogen bonds to
357	produce ammonium (14), recycling nitrogen compounds for host uptake. Oxidizing and reducing
358	metabolisms are likely separated only by microns in the hosts studied here.
359	
360	We detected biosynthetic pathways for vitamins B1, B2, B7 and B12 that are required by the
361	auxotrophic phototroph hosts in this study $(9, 10, 73)$. We found that only the blades of <i>P</i> .
362	scouleri harbored MAGs with both anaerobic and aerobic biosynthetic pathways for Vitamin
363	B ₁₂ , suggesting that the variable oxygen environment driven by host-metabolism creates diverse

364 metabolic niches for associated microbes. Strong gradients in oxygen and metabolically diverse

365 microbial metabolisms are present in a diversity of animal hosts such as corals and sponges as a

result of host metabolism (74–76). Fluctuating oxygen microenvironments might also promote
cross-feeding, where microbial taxa produce a metabolite that can be consumed by other taxa.
Cross-feeding is potentially important for nitrogen (77) and carbon metabolisms (78, 79) in
microbial communities.

370

371 Characteristics of previously undescribed nitrogen fixation in surfgrass

372 Building on recent studies that illustrate the association of nitrogen fixing microbes with a 373 diversity of macroalgae (80) and seagrasses (22, 23, 81, 82), we found a previously undescribed 374 diversity of nitrogenase genes associated with the surfgrass Phyllospadix. We detected nifH 375 genes in *P. serrulatus* rhizomes that resolved into the Cluster I group of nifH genes, which are 376 characterized by aerobic nitrogen fixers. P. serrulatus, in comparison to P. scouleri, is found 377 higher up in the intertidal zone and often in sheltered tidepools that tend to undergo dramatic 378 daily fluctuations in oxygen, possibly allowing for a temporal low-O₂ niche during the night 379 (83). Conversely, we did not detect nitrogenase genes in the microbiome of *P. scouleri*, which 380 inhabits more wave-exposed and thus better oxygenated environments (Fig. 1). However, stable 381 isotope analyses across *P. scouleri* samples show a lower nitrogen isotopic signature in the 382 rhizome compared to the rest of the plant, a possible indication of nitrogen from an atmospheric 383 source (Fig. S1), though *in situ* experiments with stable isotope tracers are needed to confirm the 384 presence of nitrogen fixation.

385

Nitrogen fixation by microbial associates provides a key means of increasing the availability of
ammonium, possibly supporting primary productivity. *P. scouleri* biomass reaches 12.7 kg of
wet mass per square meter of shore and exudes 0.93 mg C per hour per gram dry mass as

389	dissolved organic carbon that may fuel microbial activity (59). There is evidence that nitrogen
390	fixation can contribute to seagrass productivity (66, 84), a possible adaptation to low nitrogen
391	environments. Our finding that nitrogen fixing microbes are associated with a rocky intertidal
392	surfgrass is especially surprising given that Tatoosh Island is in an area of upwelling and high
393	DIN (86) at the more northerly end of the California Current Large Marine Ecosystem. Whether
394	nitrogen fixation forms the basis for reciprocal host-microbe exchange is still unknown.
395	
396	The metagenomic analyses we present here suggest that phototroph-associated microbiomes may
397	be involved in carbon, nitrogen and vitamin metabolisms important to their hosts, likely
398	generating commensal or mutualistic interactions. Future experiments should test these
399	hypothesized interactions between host and microbiome. The importance of seaweeds and
400	seagrasses to coastal productivity, and the demonstrated sensitivity of both host and microbes to
401	increasing temperatures and pH (11, 62, 85), pathogens (61), and other anthropogenic stressors,
402	underline the importance of further studying phototroph-microbiome interactions.
403	
404	Acknowledgements
405	Our gratitude to the Makah Tribal Nation for access to Tatoosh Island. We thank The University
406	of Chicago's Microbiome Center for pilot award funding, and Washington Department of
407	Natural Resources grants 93099282, 93100399 (CAP) and NSF-DEB grant (#1556874) awarded
408	to JT Wootton. We appreciate the work of C Sauceda in the isotope analysis, and A Wootton, A
409	Wood and K Foreman in the field sampling. KM was supported by an EE Fellowship from The
410	University of Chicago. S Owens and S Greenwald at Argonne National Lab provided expertise
411	in sequencing.

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643

Table 1. Summary of the features of 8 metagenomes. More information is in Table S1 and the

645 taxonomy based on single copy genes is in Table S2.

Phyllospadix			Phyllospadix	Laminaria	Nereocystis	Zostera	
scouleri			serrulatus	setchellii	luetkeana	marina	
Sediment	Rhizome	Blade	Rhizome	Blade	Inner bulb	Sediment	Rhizome
# quality read	# quality reads (in millions)						
43.68	67.73	38.41	37.99	48.58	6.48	19.37	65.76
Bacteria							
63.7%	58.3%	63.6%	63.0%	63.9%	62.2%	33.6%	60.6%
Archaea							
34.2%	38.3%	33.1%	33.3%	32.7%	35.9%	62.1%	34.2%

646

648 Table 2. Metagenome assembled genomes across all samples and their representation across

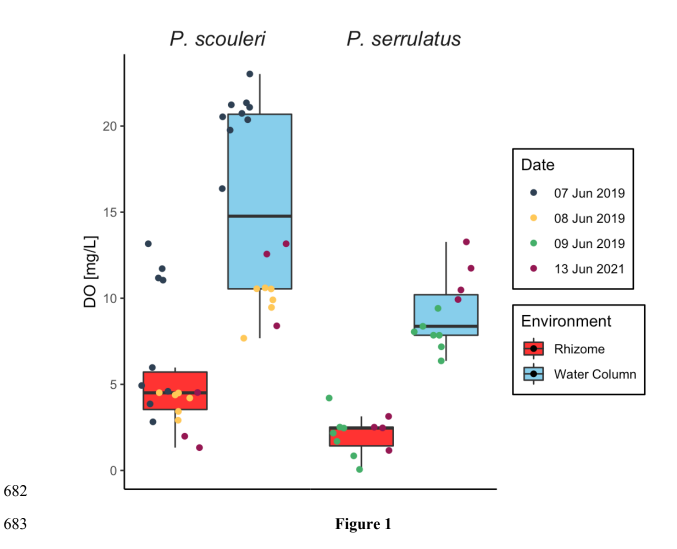
649 phyla. More detailed information on the MAGs can be found in Table S3.

Phyllospadix scouleri			Phyllospadix	Laminaria setchellii	Nereocystis luetkeana	Zostera marina	
			serrulatus				
Sediment	Rhizome	Blade	Rhizome	Blade	Inner bulb	Sediment	Rhizome
High Quality MA	AGs						
3	1	7	6	9	1	2	5
Other MAGs							
2	2	8	7	7	0	5	7
Proteobacteria							
2	-	9	2	10	1	5	5
Bacteroidota							
3	2	5	4	4	-	-	1
Verrucomicrobia	1						
-	-	-	-	2	-	-	-
Campylobactero	ta						
-	-	-	-	-	-	2	1
Desulfobacterota	1						
-	-	-	2	-	-	-	3
Desulfuromonad	lota						
-	-	-	1	-	-	-	-
Acidobacteriota							
-	-	-	1	-	-	-	-
Spirochaetota							
-	-	-	1	-	-	-	-
No ID							
-	-	1	2	-	-	-	2
Crenarchaeota							
-	1	_	_	_	_	_	_

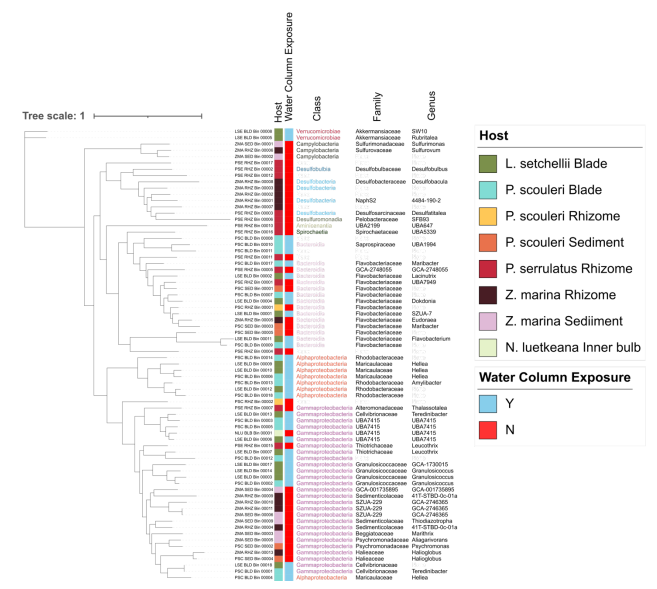
650

652	Figure Captions
653	Figure 1. Boxplot comparing the dissolved oxygen concentrations of water column (blue) and
654	the sediment-rhizome environment (red) of <i>P. scouleri</i> (pairwise t-test: p<0.001) and <i>P.</i>
655	serrulatus (pairwise t-test: p<0.001). Sampling dates are represented by different colors.
656	
657	Figure 2. A phylogenomic tree of 6 concatenated bacterial single-copy core ribosomal genes
658	from 71 bacterial MAGS across 8 samples, showing the results from 33 high quality MAGs and
659	38 lower quality ones. One MAG, PSC_RHZ_Bin_00003, from the rhizome of <i>P. scouleri</i> , was
660	identified as an archaeal genome and was thus omitted from this tree. Gaps in class, family and
661	genus indicate the level to which taxonomic classification was resolved in each MAG. All blade
662	tissues have 'water column exposure', while rhizome and sediment samples do not.
663	
664	Figure 3. Microbial Metabolisms in the MAGs reported in Fig. 2 and Table S3 across all hosts
665	and grouped as those that might benefit the host ("hosts benefit") and microbial metabolisms that
666	might utilize host provisioned metabolites ("microbes benefit"). Each tick along the x-axis
667	corresponds to a MAG. N. luetkeana blade MAGs are from Weigel et al. (in review). The
668	metabolisms for DOC Uptake that benefit microbes are shown as a heatmap of the count of the
669	number of genes that can metabolize Compatible Solutes, Carboxylic Acids, Carbohydrate
670	Pentoses and General Carbohydrates. Microbial metabolisms that benefit the host are
671	Ammonification Hydrolases, where the heatmap provides a count of the hydrolases acting on
672	C-N bonds other than peptide bonds, Nitrogen Metabolisms and Vitamin Synthesis, both
673	shown as the presence or absence of a gene in a pathway. The genes used in this are in Table S4.
674	

- 675 Figure 4. A phylogenomic tree of *nifH* genes found on the rhizomes of *P. serrulatus* (PSE,
- n = 3 and the rhizomes and surrounding sediment of Z. marina (ZMA, n = 2 and 5,
- 677 respectively). Some *nifH* genes group into Cluster I, including a sulfur oxidizing taxon on the
- 678 rhizome of Z. marina, and other taxa in Campylobacterota, including Sulfurovum. Cluster III
- 679 contains taxa associated with rhizomes including rhizomes including Desulfobulbus
- 680 *mediterraneus* on *P. serrulatus* and a Desulfobacterales associated with *Z. marina* rhizomes.

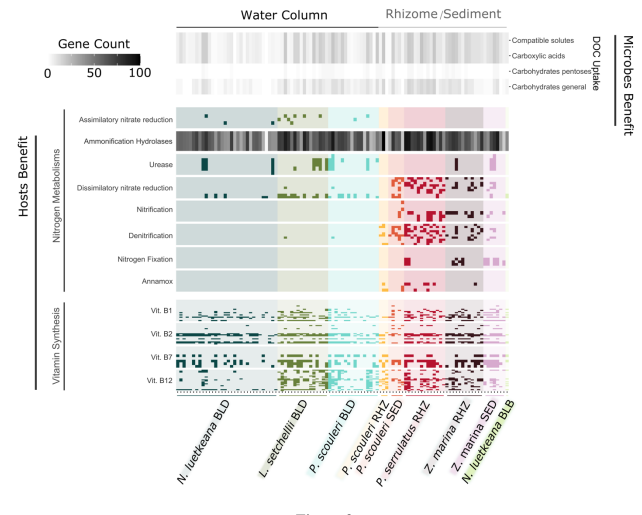


33



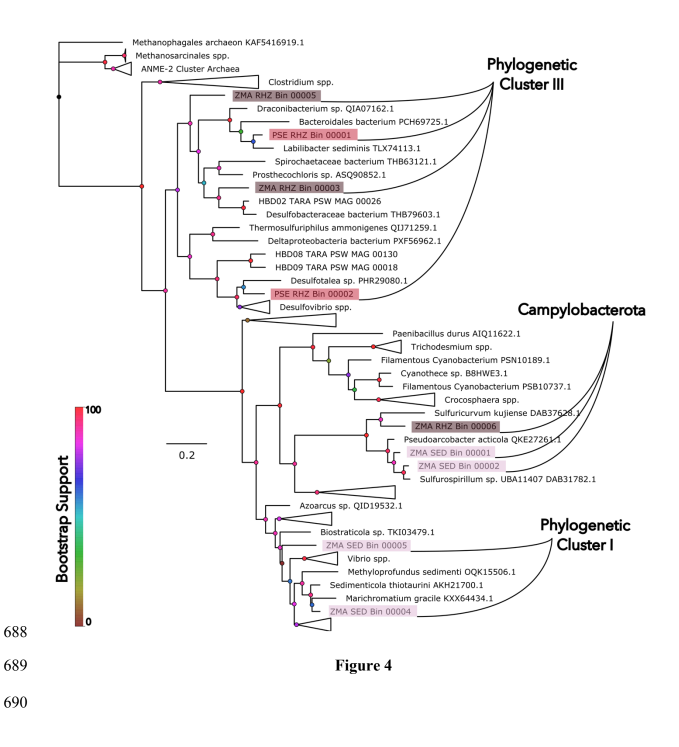
684

Figure 2



687

Figure 3



691	Supplementary Files
692	Table S1. A summary of eight metagenomes from five macrophyte taxa.
693	Table S2. A summary of taxonomy of MAGs
694	Table S3. The features of 72 metagenome assembled genomes (MAGs).
695	Table S4. Genes used to generate Fig. 3
696	Table S5. The features of <i>nifH</i> genes found in MAGs.
697	Table S6. nifH reference amino acid sequences
698	
699	Appendix 1. Additional methods to quantify carbon and nitrogen stable isotopes in <i>P. scouleri</i>
700	Figure S1. Stable isotope analysis of delC13 and delN15 at blade tip, meristem, rhizome of P.
701	scouleri. From blade tip to rhizome, water flow and thus elemental mixing reduces due to
702	attenuation and boundary layer effects of surfgrass canopy. Assuming elemental uptake occurs
703	from the same pools of C and N, the lower the extent of mixing, the heavier the isotopic
704	signature should be at that point of the plant. This is observed with delC13 which gets heavier
705	from the tip to the blade. This is observed with delN15 till the meristem after which it lightens.
706	This is probably occurring as nitrogen is taken up from a different pool of nitrogen from that
707	around the blade/meristem. This different pool is probably made available through n-fixation