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RESEARCH ARTICLE

Summer shifts of bacterial communities associated with the invasive brown seaweed *Sargassum muticum* are location and tissue dependent

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Abstract

Seaweed-associated microbiota experience spatial and temporal shifts in response to changing environmental conditions and seaweed physiology. These shifts may result in structural, functional and behavioral changes in the host with potential consequences for its fitness. They, thus, may help the host to adapt to changing environmental conditions. The current knowledge of seasonal variation of seaweed-associated microbiota is however still limited. In this study, we explored temporal and spatial variation of microbial communities associated with the invasive brown seaweed *S. muticum*. We sampled in northern and southern Portugal, in September, March and July-August (summer). In addition, as (pseudo-)perennial seaweeds display seasonal reproductive phenology, we sampled various parts of the individuals to disentangle the effect of temporal changes from those due to structural development variations. The diversity and structure of associated microbial communities were determined using next generation sequencing of the variable regions V5-7 of the 16S rDNA. We expected to find differentiation in associated microbial communities between regions and sampling months, but with differences depending on the seaweed structure examined. As expected, the study revealed substantial temporal shifts in *S. muticum* microbiome, for instance with large abundance of *Rhodobacteraceae* and *Loktanella* in September-March but prevalence of *Pirellulales* during the summer months. Variations between regions and tissues were also observed: in northern Portugal and on basal structures, bacterial diversity was higher as compared to the South and apical parts. All examined seaweed structures showed temporal differences in associated microbial community structure over time, except for holdfasts between September and March. Bacteria contributing to these changes varied spatially. Conversely to all other structures, the holdfast also did not show differences in associated community structure between southern and northern regions. Our study highlights the importance of structural microscale differentiations within seaweeds hosts with regard to their associated microbial communities and their importance across temporal and spatial dimensions.

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Introduction

Seaweeds live in association with abundant and diverse microbiota, which plays an important role in the life of its hosts in nature [1, 2, 3]. Seaweeds rely on associated microbial communities for diverse functions including morphological development [4, 5, 6, 7, 8], consumption of organic matter and nitrogen source [9], defense [10, 11, 12, 13, 14, 15, 16], or provision of vitamins [17]. Microbial host-specificity has been reported (e.g., in fucoid seaweed *Phyllospora comosa* [18]) but most studies documented changes of seaweed associated microbial communities in response to various factors and conditions [11, 18].

Structure and composition of the seaweed-associated microbiota are known to change with the host conditions, as well as in space and time [11, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27]. Regarding host condition, for example, healthy vs. stressed *Ecklonia radiata* exhibit different microbial communities [28]. Host traits may therefore be critical in determining the community structure of associated microbiota or *vice versa*. Other studies documented spatial variation [29], for instance, high within-species variability was observed among microbial communities associated with specimens of *Ulva australis* from different rock pools [30]. However, most studies documented shifts in microbial communities over time, varying from short-term (i.e. less than a day) to long-term (i.e. inter-annual) scales, with changes most often associated with seasonal variability [21, 31]. Some bacterial taxa seem more representative of a given season than others [19, 21, 22, 25]. For instance, bacterial communities associated with *Fucus vesiculosus* showed persistent seasonal variation, at the phyla level, over two consecutive years [21]. Recent research conducted on the Mediterranean seaweed *Cystoseira compressa* also revealed rather dynamic associated bacterial communities [22]. Bacterial communities associated with thalli of *C. compressa* displayed a clear successional pattern over time as well as an increase in abundance of pathogenic bacteria, associated with natural degradation of thalli at the end of the annual life cycle [22].

Seasonal variations are thought to be related to the combined effect of biotic (i.e. seaweed growth cycle, the age of the algal tissue) and abiotic factors (i.e. seawater temperature) [19, 22, 25]. *Planctomycetes*, *Verrucomicrobia* and *Alphaproteobacteria*, for example, were among the early colonizers of young thalli in *Laminaria hyperborea* [19]. As the algal tissue aged and biofilms matured, communities associated with *L. hyperborea* were complemented by *Gamma-proteobacteria*, *Bacteroidetes*, and *Cyanobacteria* [19]. Other biotic factors include interactions (including competition) among bacterial taxa (i.e. both internal and from the surrounding water; [19]), consequences of biological interactions with organisms from other trophic levels (i.e. grazing, cross-feeding) as well as the loss of certain functions by bacteria (which results in the dependence on services provided by other microorganisms, [32]). Concerning abiotic factors, seasonal shifts in microbial community composition generally mirror seasonal environmental changes [31]. In particular, the summer increase of water temperature is likely to modify bacterial community composition [19, 22]. During summer, microbial communities on *L. hyperborea* were for instance characterized by a high diversity and such conditions were thought to be beneficial for seaweed-associated bacteria populating algal exudates [19]. On the other hand, some bacteria, such as *Betaproteobacteria* and *Verrucomicrobia* associated with *L. hyperborea*, were only observed in months when the seawater temperature was below 10°C, suggesting a preference for colder seawater temperatures [19]. Climate change and ocean acidification are also expected to intensify these changes [33, 34, 35].

Apart from the seasonality, host-associated microbiota is documented to be tissue-specific [24]. While microbial tissue-specificity was studied previously in, for example, corals [36], the number of studies conducted on seaweeds is very limited, but suggest that different seaweed tissues are populated by specific bacteria [24]. For instance, *Laminaria saccharina* had specific

bacteria within its young and undisturbed tissues regardless of seasonality or geographic location [24]. It still remains to be explored to what extent tissue differentiation is affected seasonally. In addition, although many studies explored the seasonal differences of seaweed-associated microbiota and surrounding water column [19, 21, 22, 30, 37], no studies looked at similarities between the seaweed microbiome and surrounding water over time and, more importantly, at tissue or structure specificity. Differences in microbial communities of a seaweed between summer and winter can be directly related to its development and the structures sampled during each time. In winter, for example, pseudo-perennial seaweeds (as *S. muticum*) might consist of only a basal holdfast whereas in early summer they might be several meters long with various different structures present including blades, floatation vesicles and reproductive structures (receptacles), which each might have a different associated microbiome.

The present study addresses the disentanglement of spatial and temporal shifts in seaweed associated bacterial communities among different seaweed structures using the pseudo-perennial invasive brown seaweed *Sargassum muticum* as a case study. This brown alga is one of the most invasive macroalgae in the northern hemisphere, but its invasive success is not yet entirely understood [38]. Microbiota might play a key role in the acclimatization of this non-native seaweed, but it has been so far poorly examined. As detailed before, several studies indeed showed that bacterial seasonal variations are driven by environmental factors, notably seawater temperature. Seawater temperature data from southern to northern Portugal clearly demonstrate that there is a much larger variation in temperature between seasons in the North as compared to the South [39]. We thus hypothesized that microbial communities associated to *S. muticum* would show combined effects of spatio-temporal differences, but that these differences also depend on the seaweed structure/part examined. Next-generation sequencing of the variable regions V5-V7 of bacterial 16S rDNA genes was applied to characterize the diversity of associated microbiota and describe differences in microbial community structure.

Methodology

Study area and samples collection

Samples were collected in northern Portugal at Praia Norte (Viana do Castelo) and in southern Portugal at Praia do Queimado (Porto Covo) in September 2013, as well as in March, July (Porto Covo) & August (Viana do Castelo) 2014. The tissues, pieces of 1–2 cm, collected constituted: the tip, basal blades and holdfast, as well as reproductive structures (receptacles) (Fig 1) collected in July in Porto Covo and in September & August in Viana do Castelo. Each tissue was sampled in triplicate, separated by several meters to cover local variation, at each location and month. Sampled individuals were haphazardly chosen from the more complete developed individuals of the population to assure the presence of as many different structures as available and standardize the developmental status among individuals. In addition, sediments and water samples were collected. For sediments, three replicates were sampled in each month apart from September in Viana do Castelo (only 2 replicates). Seawater samples were also represented by three replicates, except September (in both locations), when only 2 replicates were available, and August in Viana do Castelo, where only 1 sample was collected. In the field, seawater microbiome was sampled by filtering 0.5 L of seawater over a 0.2 µm filter. All samples, seaweed and environmental, were preserved in the field directly upon collection in Xpedition lysis buffer (Zymo Research, California, USA). Overall, 94 samples were collected across both locations: 47 from Porto Covo (Southern) and 47 from Viana do Castelo (Northern). No collection permits were required as no individuals were sampled. The model species is not a protected or endangered species.

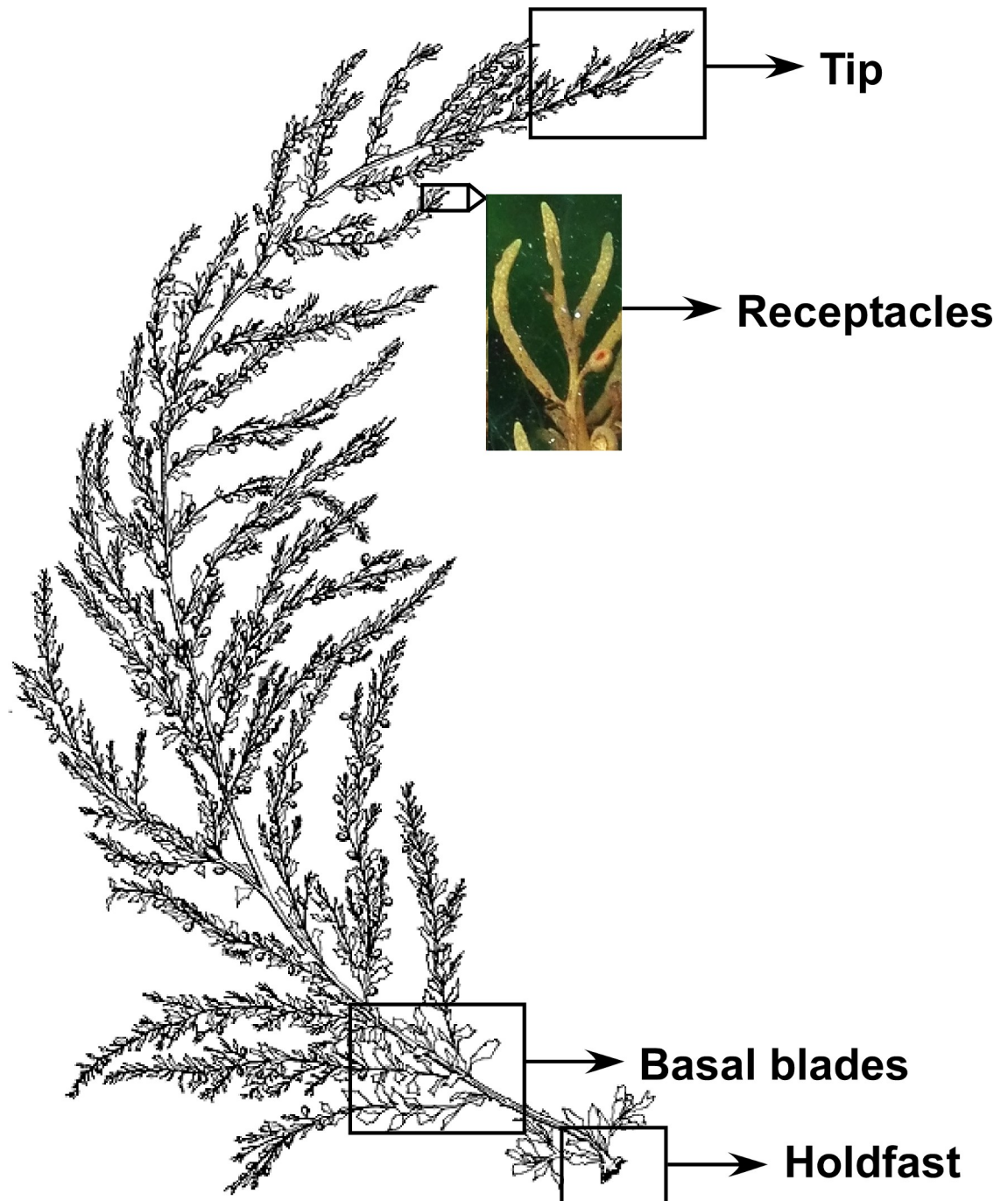


Fig 1. Schematic representation of *S. muticum* identifying the differentiated tissues considered in this study: Tip, receptacle, basal blades and holdfast.

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High-throughput sequencing of the microbiome

Epiphytic and endophytic bacteria were extracted from differentiated structures (holdfast, basal blades, tip blades and receptacles) of *S. muticum* (Fig 1), seawater and sediments using MoBio PowerSoil DNA Isolation Kit in following the manufacturer protocol. The total 16S rRNA was amplified using the universal primers 27F and 1492r with the following changes to

the original protocol [40]: following the initial denaturation step at 95°C for 2min, conditions constituted 35 cycles of denaturation at 95°C for 20s, annealing at 55°C for 20s, and extension at 72°C for 90s. The final extension was at 72°C for 3min. The 25 µl reaction mixture contained 250 µM dNTPs, 0.6 µM of each primer, 1 × 2PCR buffer mix, 2 µl of template DNA (with a final concentration of about 10 ng µl⁻¹), and 0.3 µl of Taq polymerase (Advantage R2 Clontech). PCR products were cleaned using the ExoFastAP enzyme according to the manufacturer protocol (Thermo Scientific) and amplified DNA was sent to Molecular Research (MR DNA), Shallowater, Texas where a nested-PCR was implemented before the sequencing. Modified 8 bp key-tagged primer 799F along with the reverse primer 1193R (fragment ~ 400 bp), which avoid chloroplast cross amplification [41], were applied and PCR conditions constituted: 95°C for 3 min, 10 cycles of 95°C for 20 s, 50°C for 30 s, 72°C for 30 s, and a final elongation of 72°C for 3 min. Samples were pooled together in equal proportions based on their molecular weight and DNA concentrations and purified using calibrated Ampure XP beads. DNA libraries were prepared applying the Illumina TruSeq DNA library preparation protocol and paired-end (2 x 250 bp) sequencing performed at MR DNA (www.mrdnab.com, Shallowater, TX, USA) on a MiSeq following the manufacturer's guidelines.

Sequence analysis and bioinformatics

The microbial community analysis was implemented using the Quantitative Insights into Microbial Ecology (QIIME version 1.8.0) program [42]. Sequences were screened and filtered for a minimum read length of 350 bp and less than 2 undetermined nucleotides. The filtered dataset, comprising only high-quality sequences, was applied to a conservative chimera detection filter using the ChimeraSlayer method [43]. Selected high quality chimera-free sequences were clustered into Operational Taxonomic Units (OTUs) within reads using the UCLUST algorithm [44] with a pairwise identity threshold of 0.97. Representative sequences for each OTU were selected using the “most-abundant” method and OTU sequence alignment was implemented with the Pynast tool [42]. The Ribosomal Database Project (RDP) classifier [45] was applied for taxonomic assignment with a 95% confidence threshold. To assign each OTU to the closest matching described taxon, the search was performed against the Greengenes reference database (version 12_10) [46] with a maximum e-value to record an assignment of 0.001. Sequences with the best match for eukaryotes (i.e. chloroplasts & mitochondria), rare OTUs (i.e. singletons & doubletons), and unassigned sequences were removed from the OTU table in the downstream analysis. The filtered rarefied OTU table was applied to calculate alpha diversity statistics, including the Chao I richness estimates [47], the observed number of OTUs, and the Shannon index, using QIIME software [42]. Permutational multivariate analysis of variance (PERMANOVA) was conducted to test for spatial and temporal differences in the microbiomes of *S. muticum*. To visualize dissimilarity between samples, Canonical Analysis of Principal coordinates (CAP) plots were constructed using the interaction among locationXtissueXseason as *a priori* factors. Bacterial contributions to similarity and dissimilarity between microbial communities was assessed by SIMPER analysis. All statistical analysis mentioned above were implemented using the software PRIMER-E+PERMANOVA v.6 [48, 49]. For bacterial groups, which showed notable differences in abundance between seasons, two-way analyses of variance (ANOVA) were implemented (with the preliminary tests for normality and homogeneity of variances).

Results

Rarefaction to 2719 high quality sequences per sample based on the lowest number of available reads of a sample, resulted in 255,586 sequences in total that were used and

corresponded to 52,109 OTUs. Overall, diversity was 1.2 times higher in northern compared to southern Portugal when expressed as Chao1 (PERMANOVA, $P = 0.001$; Fig 2A) and OTU Richness (PERMANOVA, $P = 0.001$, Fig 2B), except for Chao1 in March, and all months for the Shannon index (Fig 2C). Differences among tissues were also revealed for Chao1 (PERMANOVA, $P = 0.001$, Fig 2C), OTU Richness (PERMANOVA, $P = 0.001$, Fig 2B), and Shannon index (PERMANOVA, $P = 0.009$, Fig 2C). However, an interaction was observed between month and seaweed structure for OTU Richness (PERMANOVA, $P = 0.035$; Fig 2B). Overall, holdfasts were characterized by the highest bacterial diversity among all seaweed structures examined, after sediments (Fig 2B and 2C). Bacterial diversity tended to decrease towards the apical seaweed tissues with the lowest diversity associated to the tips (Fig 2B and 2C).

Overall, 794 bacterial genera were distributed across 137 classes from 52 phyla. In the South, 623 genera were distributed across 104 classes from 40 phyla, while in the North 682 genera were distributed across 124 classes from 52 phyla. September-March bacterial communities associated to *S. muticum* were dominated by *Proteobacteria* (60.5%; Alpha 36%, Gamma 17.9% & Delta 4.9%), *Bacteroidetes* (29.9%; *Flavobacteriia* 21.1% & *Saprospirae* 4.7%), and *Actinobacteria* (7.3%; *Acidimicrobiia* 7.2%) (Fig 3). Although *Proteobacteria* (40%; Alpha 18.3% & Gamma 17.5%) and *Bacteroidetes* (31.4%; *Flavobacteriia* 15.5% & *Saprospirae* 11.7%) are still major contributors to the bacterial communities in summer (July-August), a major change was reported with a high prevalence of *Planctomycetes* (14.5%; *Planctomycetia* 11.4%) and *Spirochaetes* (6.5%; *Spirochaetes* 6.5%) (Fig 3).

Bacterial community structure associated with different *S. muticum* structures, sediments and seawater differed structurally in the two regions (North and South), among September, March and July-August (PERMANOVA, $p = 0.006$, Table A in S1 File), as shown by the canonical analysis of principle coordinates (Fig 4). Overall, *S. muticum* structures, sediment and seawater had different communities associated to them. Across structures, receptacles and basal blades had marginally more similar communities (PERMANOVA pairwise comparison $P < 0.053$) as compared to holdfasts which harbored the most distinct associated community (Fig 4).

Microbial communities associated with seaweed structures showed temporal differences among all sampling times in the North (PERMANOVA $P = 0.0001$), but not in the South between September and March (PERMANOVA $P = 0.1766$; Fig 4). In contrast, environmental (sediments and seawater) microbial communities did not show temporal variations depending on location (PERMANOVA $P = 0.070$). Temporal variation in community structure was dependent on the seaweed structure examined (PERMANOVA $P = 0.001$), only holdfast associated communities did not differ among all sampling times ($P = 0.188$).

The most overall differences in community structure between summer and March—September constituted a decrease in *Proteobacteria* by 20.5% and *Actinobacteria* by 7.4%, as well as an increase in *Planctomycetes* by 13.8% (Fig 3). The lower abundances of *Proteobacteria* in summer, compared to September, occurred mostly due to the significant decline in abundance of an unidentified *Rhodobacteraceae* (within the tip, receptacles & basal blades) and *Loktanella* (within the tip, receptacles, basal blades and holdfast) both belonging to the order *Rhodobacterales*. Strongly increased abundance of *Planctomycetes* in summer, compared to September and March, occurred mostly due to an increase of an unidentified *Pirellulaceae* associated with receptacles, basal blades and holdfast.

Overall, in September-March, the most abundant bacteria associated with *S. muticum* were unidentified *Rhodobacteraceae* (9.9%), *Loktanella* (8.6%), unidentified *Flavobacteriaceae* (8.5%) and unidentified *Hyphomonadaceae* (5.4%). In contrast, in July-August, the most abundant bacteria were unidentified *Saprospiraceae* (10.2%), unidentified *Pirellulaceae* (10%), unidentified *Gammaproteobacteria* (9.4%), unidentified *Flavobacteriaceae* (7%), *Spirochaeta* (6.4%), and unidentified *Rhodobacteraceae* (5.5%).

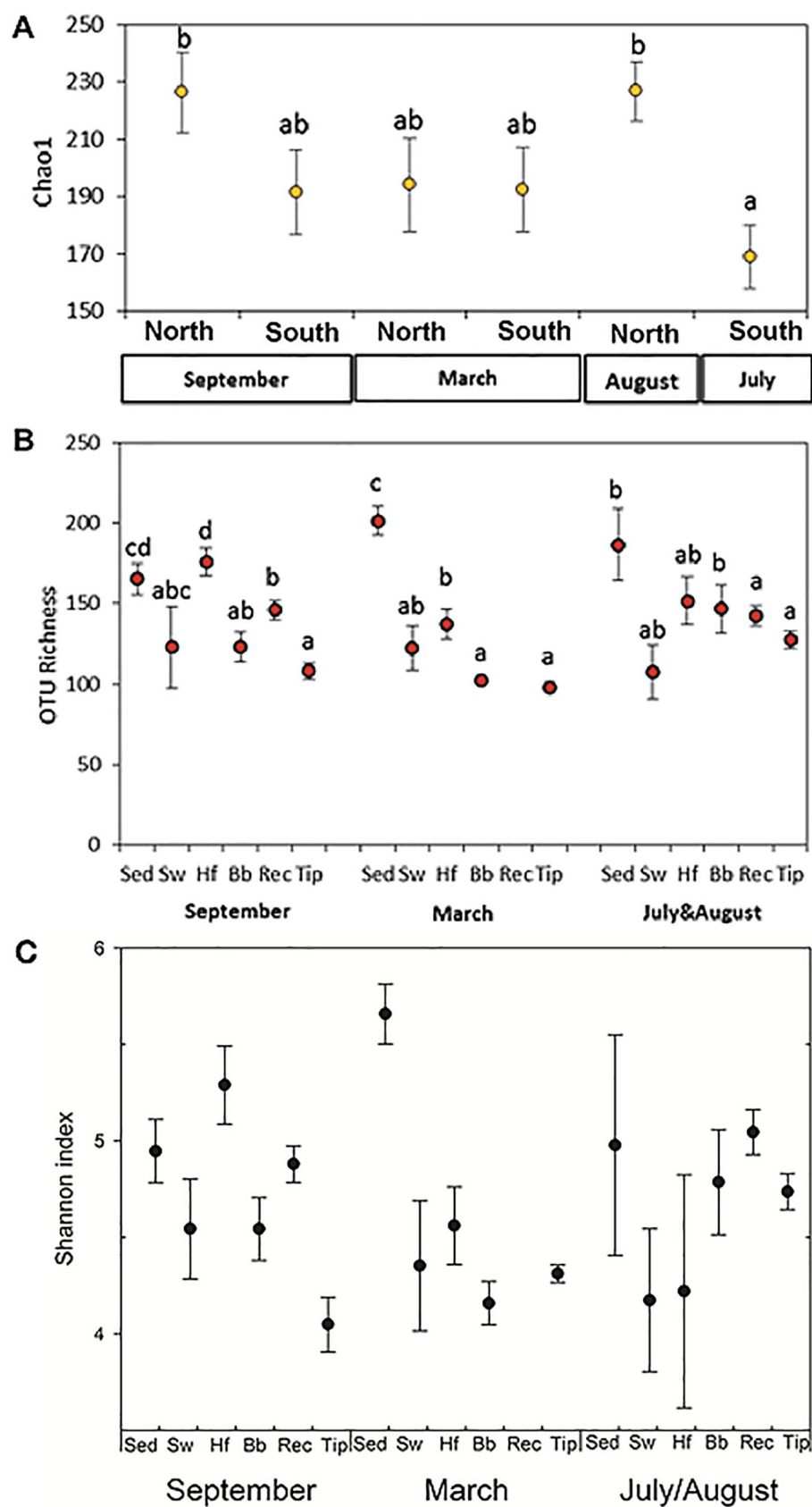


Fig 2. Alpha-diversity of bacterial communities. Chao 1 (A), OTU richness (B) and Shannon index (C) alpha diversity measures of bacterial communities associated to *S. muticum* across seasons at its northern (VC: Viana do Castelo) and southern distribution in Portugal (PC: Porto Covo). Differentiated are the environmental samples of sediments (color circle) and seawater (empty square), and the seaweed structures holdfast (rombe), basal blades (color square), receptacles (color triangle), and tip (empty circle). Values are means $\pm$ standard error (n = 3).

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Community structure associated with *S. muticum* was different between southern and northern Portugal (PERMANOVA, $P = 0.001$). The main differences were due to the prevalence of unidentified *Pirellulaceae* (*Pirellulales*) and unidentified *Gammaproteobacteria* in the South (contributing 1.71% and 1.67% to dissimilarity respectively; SIMPER), and higher abundance of *Loktanella* (*Rhodobacterales*) and unidentified *Saprospiraceae* (*Saprospirales*) in the North (contributing 1.65% and 1.62% to the dissimilarity respectively; SIMPER). The North-South differences in community was reflected in the communities associated with sediment ($P = 0.0012$) and each seaweed structure ($P = 0.0004$ – 0.0014), except holdfast ($P = 0.4647$) and seawater ($P = 0.0754$), where no such difference was detected.

Overall, 18 and 19 bacterial genera in southern and northern Portugal, respectively, were present in all tissues at all times, constituting *Alphaproteobacteria*, *Gammaproteobacteria*, *Flavobacteria*, *Saprospirae* and *Acidimicrobia* (Fig 5). These communities were very similar between the locations (Table B in S1 File). *Proteobacteria* and *Bacteroidetes* constituted the seasonally-independent bacterial phyla observed in all months sampled, *Planctomycetes* were unique to *S. muticum* only in Summer.

Temporal differences in communities for each seaweed structure were seldom at the southern (2 out of 9 pairwise comparisons), but common at the northern location (7 out of 10 pairwise comparisons; Table C in S1 File).

Tip associated communities changed over all sampled seasons, but differently so between the northern and southern *S. muticum*, with in the South only temporal differences in Summer, while in the north temporal differences occurred across all sampling times (Table C in S1 File). Between September and March, an unidentified *Acidimicrobiales* (*Acidimicrobiia*) and *Loktanella* (*Alphaproteobacteria*) contributed most (5.68% and 4.6% respectively) to the dissimilarity in the North (43.04% average dissimilarity, SIMPER), while an unidentified *Rhodobacteraceae* (*Alphaproteobacteria*) contributed most (3.66%) to the dissimilarity in the South (40.98% average dissimilarity, SIMPER) (Fig 6). Between March and summer, an unidentified *Acidimicrobiales* (*Acidimicrobia*) and *Tenacibaculum* (*Flavobacteria*) contributed most to the dissimilarity (4.77% and 3.81% respectively) in the North (47.74% average dissimilarity, SIMPER), while an unidentified *Pirellulaceae* (*Planctomycetia*) contributed most (4.58%) to the dissimilarity in the South (52.10%) (see Fig 6 for relative abundances). Between summer and September, *Loktanella* (*Alphaproteobacteria*) and an unidentified *Saprospiraceae* the North (45.64% average dissimilarity, SIMPER), while *Glaciecola* (*Gammaproteobacteria*) and an unidentified *Pirellulaceae* (*Planctomycetia*) contributed most to the dissimilarity (4.34% and 4.22% respectively) in the South (55.64% average dissimilarity, SIMPER) (Fig 6).

Receptacles were only found during August (North) and September (North and South). In the North, differences were detected between summer and September ($P = 0.019$, average dissimilarity 39.65%, SIMPER) with an unidentified *Pirellulaceae* (*Planctomycetia*) and *Loktanella* (*Alphaproteobacteria*) contributing most to the dissimilarity, 3.48% and 3.41% respectively (Fig 6).

Microbial communities associated with basal blades showed differences between months only at the northern location, across all sampling times (Table C in S1 File). Between Summer and September (average dissimilarity 39.45%, SIMPER) an unidentified *Pirellulaceae* (*Planctomycetia*) contributed most (3.45%) to the dissimilarity, while between March and Summer

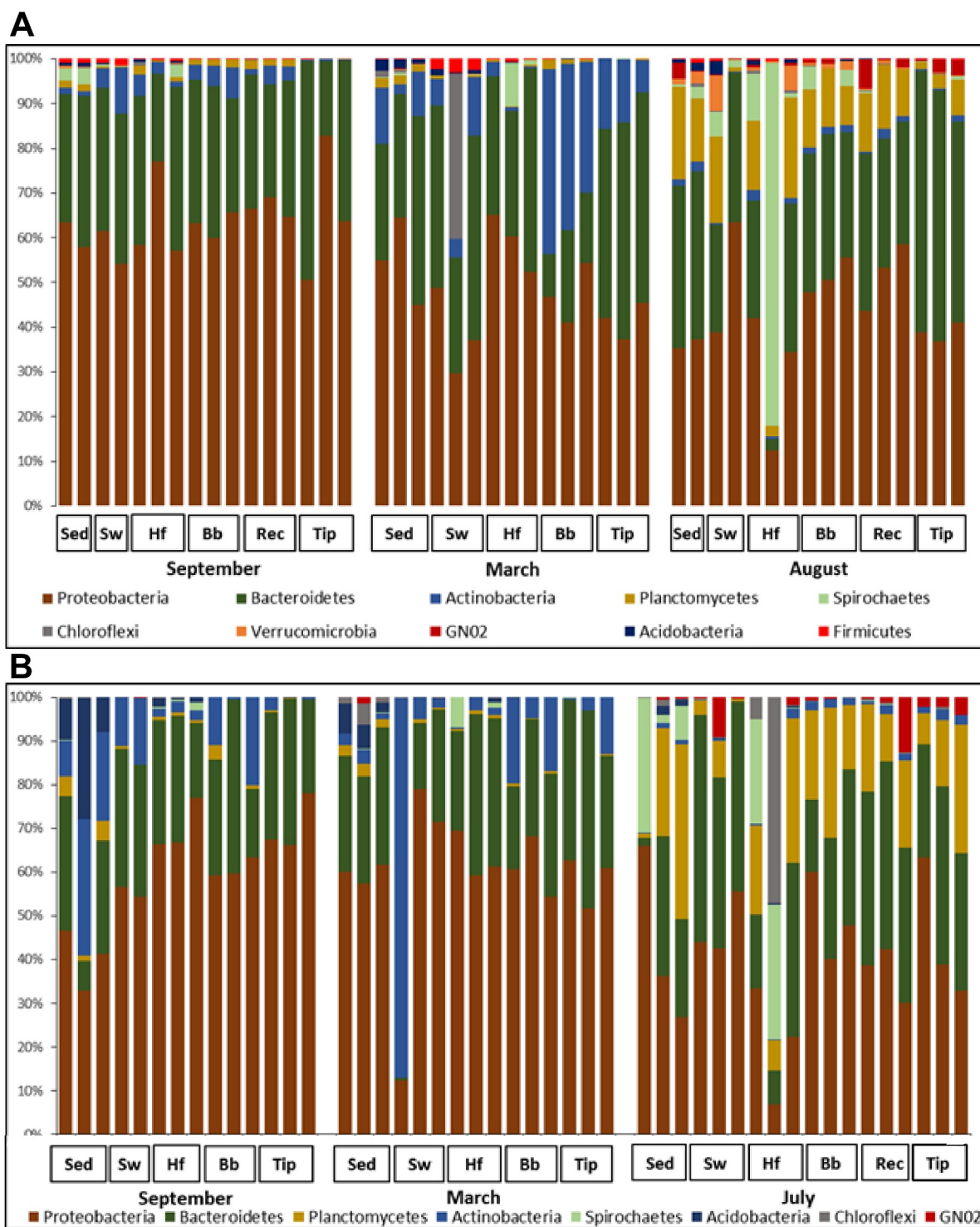


Fig 3. Distribution of bacteria phyla associated with *Sargassum muticum*. Relative distribution of the bacteria phyla associated with different structures of the brown seaweed *Sargassum muticum*, sediments (Sed) and surrounding seawater (SW) in northern (Viana do Castelo (A)) and southern (Porto Covo (B)) Portugal in September, March and July/August. Tissues are holdfast (Hf), basal blades (Bb), receptacle (Rec) and tip (Tip).

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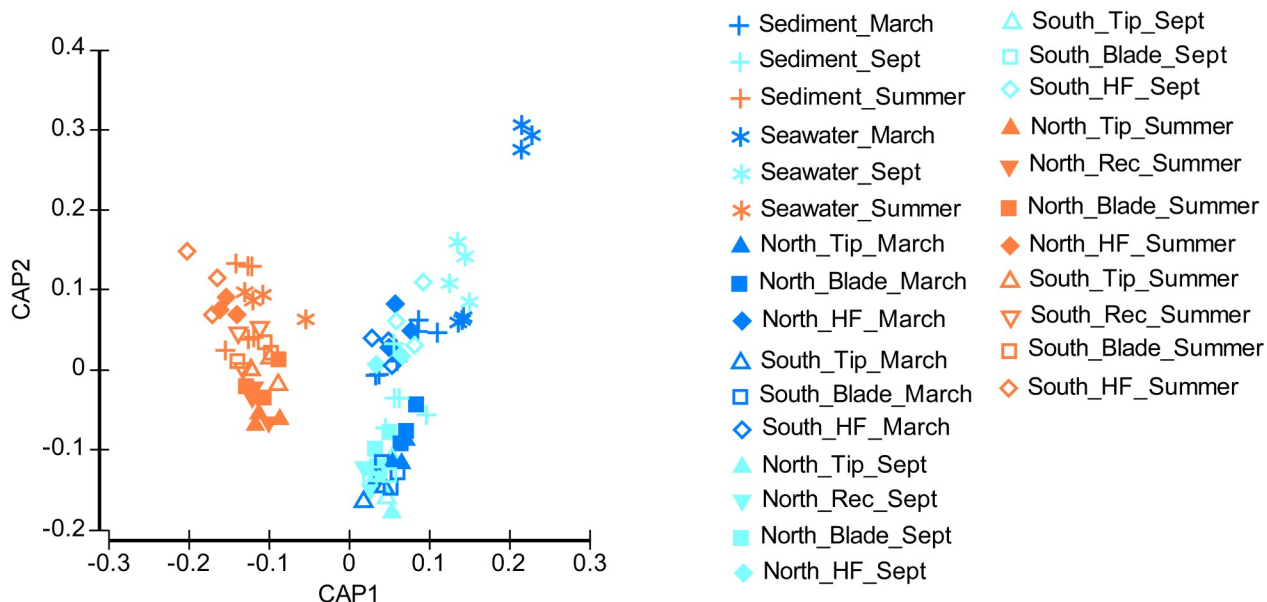


Fig 4. Structure of microbial communities associated with *Sargassum muticum*. Canonical Analysis of Principal coordinates of the bacterial communities associated with *Sargassum muticum*, sediments (+) and seawater (*) samples collected in southern (Porto Covo) and northern (Viana do Castelo) Portugal in September (light blue), March (dark blue) and July-August (orange). Distinguished structures are holdfast (HF) (diamond), basal blades (square), receptacle (Rec) (down facing triangle) and tip (up facing triangle).

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(average dissimilarity 54.01%, SIMPER) most contribution to the dissimilarity was done by an unidentified *Acidimicrobiales* (4.38%) and an unidentified *JdFBGBact* (3.06%) both belonging to the class *Acidimicrobiia* (Fig 6). Between September and March (average dissimilarity 42.59%, SIMPER), an unidentified *Acidimicrobiales* contributed most (5.12%) to dissimilarity (Fig 6). No significant temporal differences in bacterial structure were revealed within the holdfast tissues.

Environmental communities showed temporal variation in community structure, but whereas in sediments structure differed among September, March and summer ($P = 0.002$ – 0.007), in seawater differences were restricted to between March and summer ($P = 0.039$).

Discussion

Our results show that microbial communities associated to *Sargassum muticum* show temporal differences between September/March and July/August, but with different dynamics in northern and southern Portugal. In addition, these temporal and spatial differences depend on the seaweed structure examined.

The most pronounced temporal changes in microbial community were the significantly decreased abundance of *Proteobacteria* (by 20.5%) and *Actinobacteria* (by 7.4%) and increased abundance of *Planctomycetes* (by 13.8%) in summer. The summer decrease in *Proteobacteria* occurred mostly due to the significant decline in of unidentified *Rhodobacteraceae* (within the tip, receptacles & basal blades) and *Loktanella* (within the tip, receptacles, basal blades and holdfast). *Rhodobacterales* have been previously isolated from the seagrasses *Thalassia hemprichii* [50] and *Zostera marina* [51] and are primary colonizers of in marine surfaces with a known ability to fix nitrogen [52, 53]. Although not all members of the *Rhodobacteraceae* are considered pathogens, certain members are known to cause infections and disease (with the potential of getting more severe at increased temperatures) in *Fucus vesiculosus* [54] and *Deli-sea pulchra* [55]. *Loktanella* are known from various macroalgal species [2, 56, 57], including

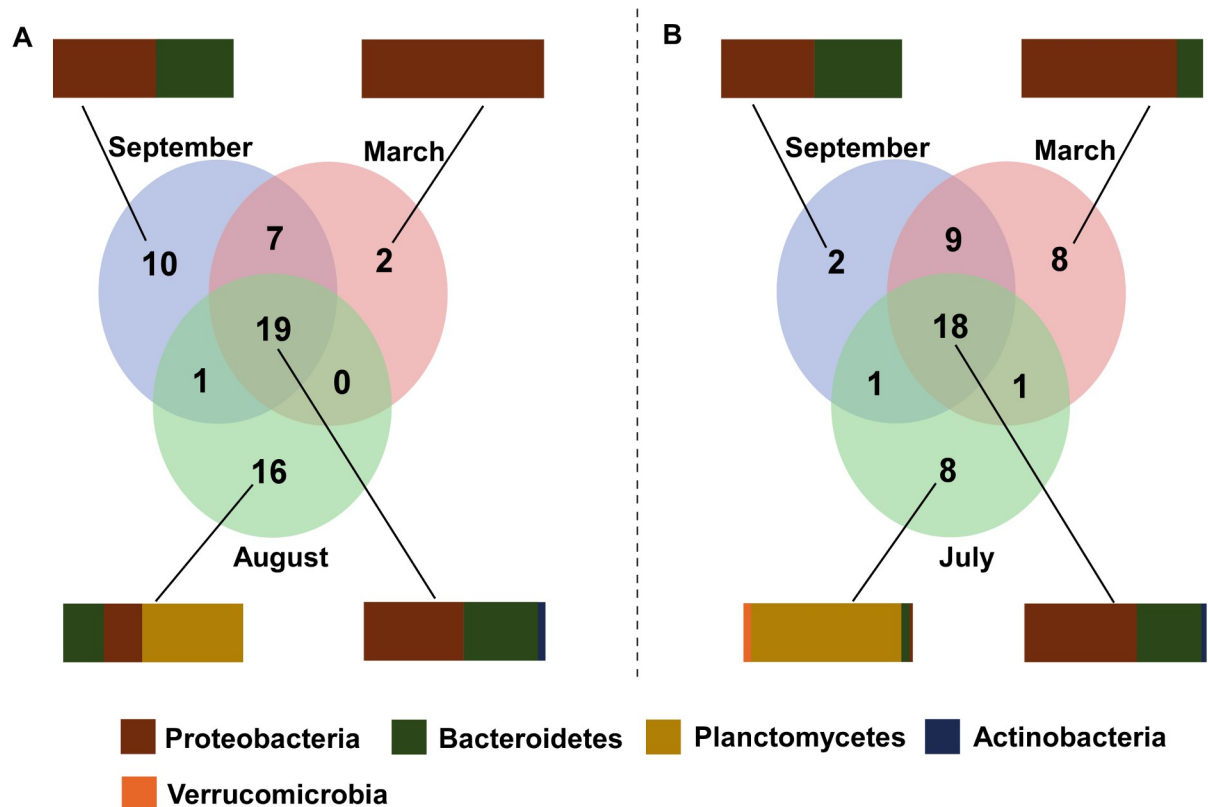


Fig 5. *Sargassum muticum* consistent bacterial genera across structures. Venn diagram representing the number of bacterial genera present across all *S. muticum* structure samples in North (A) and South (B) Portugal in different months. Vertical bars show phylum composition of each month (non-shared genera) and shared among all months.

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Fucus vesiculosus [21, 54], *Ulva australis* [30] and *Cystoseira compressa* [22]. Bacteria from this genus are highly adaptive and known for their capacity to utilize and rapidly metabolize organic carbon sources from seaweed exudates [20, 58, 59]. A significant increase of these bacteria was associated with the natural degradation of aged thalli in *Cystoseira compressa* during annual shedding in October [22]. Decrease in abundance of *Rhodobacteraceae* in response to elevated temperature levels was also observed in association with *Fucus vesiculosus* forma *mytili* [60]. The *S. muticum* material used in this study all seemed in good health and where not degrading and without signs of infection or disease.

Another important change was the summer increase of *Planctomycetes*, which occurred mostly due to the increase of unidentified *Pirellulaceae* from the order *Pirellulales* (within the holdfast, basal blades & receptacles). *Planctomycetes* are frequent associates of macroalgae [2, 9, 19, 30, 61]. They were reported in association with *Laminaria hyperborea* [62, 63], *Macrocystis pyrifera* [64], *Porphyra umbilicalis* [57], *Ulva australis* and *Delisea pulchra* [29], *Fucus vesiculosus*, *Gracilaria vermicuphylla* and *Ulva intestinalis* [21]. *Planctomycetes* are known for their ability to mineralize organic into inorganic compounds matching nutritional requirements of macroalgae [61]. *Planctomycetes* are also proposed to degrade algal polymers and important contributors to the global nitrogen cycle [65]. Because *Planctomycetes* contain a high number of sulfatases genes, they could participate in the degradation of sulfated polysaccharides produced by *S. muticum* [66, 67]. Although *Planctomycetes* are known to be abundant on macroalgae [61, 68], their relative abundances vary substantially between seasons and seaweed species [21]. The summer increase of *Planctomycetes* is in line with the study on

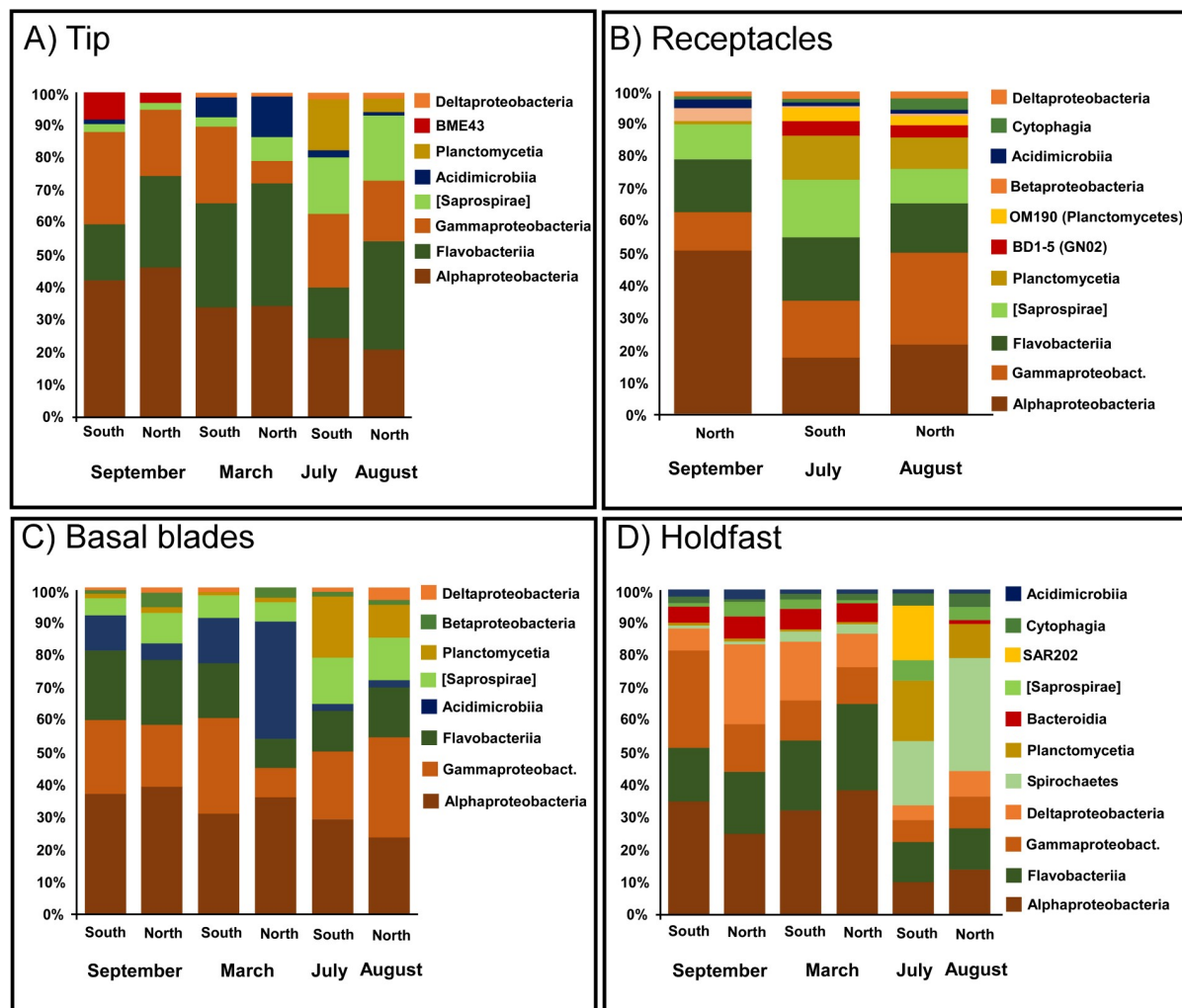


Fig 6. Most abundant bacterial classes associated with the different types of *S. muticum* tissues across seasons between North and South Portugal. A) Tips; B) Receptacles; C) Basal blades; D) Holdfasts.

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Laminaria hyperborea, where the abundance of this bacterial phylum was minimal in September and maximal in July [19]. It has been proposed that without the seaweed chemical defense, *Planctomycetes* could lose their competitiveness over other bacteria, resulting in the low abundance observed in September [62].

Temporal independent bacterial communities

Overall, *Proteobacteria* and *Bacteroidetes* constituted the most abundant bacterial phyla associated with *S. muticum* which was consistent with previous studies in other seaweeds [19, 21, 22, 24, 25, 30]. *Proteobacteria* (Alpha, Gamma & Delta), *Bacteroidetes* and *Actinobacteria* were prevalent in association with Baltic and North Sea *Fucus vesiculosus* [54, 60, 69], *Cystoseira compressa* [22] and *Macrocystis pyrifera* [64]. Prevalence of *Proteobacteria*, *Bacteroidetes* and *Actinobacteria* over the year indicates of their seasonal independence (consistent with results of [21]) and temporal adaptation, as well as an important role towards *S. muticum* and functioning of its associated bacterial community.

Differences between North and South

Loktanella (Alphaproteobacteria) and unidentified *Saprospiraceae* (Bacteroidetes) were more prevalent at the northern, while unidentified *Pirellulaceae* (Planctomycetes) and unidentified *Gammaproteobacteria* were more abundant at the southern location. Between September–March and summer, *Loktanella* and unidentified *Gammaproteobacteria* exhibited significant decrease in abundance, while unidentified *Saprospiraceae* and unidentified *Pirellulaceae* showed significant increase. Overall, the effect of season on *S. muticum* associated microbiota was more important than the effect of geographic location (Fig 4). A possible explanation for these results might be that despite the temperature gradient between the two locations throughout the year, the difference in seawater temperature might not be sufficient to significantly re-organize and re-structure associated microbial communities.

Temporal differences by tissue

The highest temporal changes in *S. muticum* microbiota were detected within the tip tissues. This could be due to the fact that tips are made of newly developed (annual) tissues and, as such, recently colonized as compared to older perennial holdfast tissues. Another major change were the significant differences within the basal blades among the sampled months and within the receptacles between September and summer in the North, but not in the South. This could be due to the higher variation in water temperature in the North of Portugal compared to the South. In addition, for receptacles, reproductive activity of *S. muticum* in northern Portugal is finished by September and re-organization of associated microbiota is likely to have taken place before (notably when reproduction occurs between April and August).

Possible sources of bacteria

Variation of bacterial communities associated with *S. muticum* could be explained by the variation in abundance of bacteria in the environment, from where they could be acquired, at the particular season [70]. The differences in composition between bacterial communities associated with *S. muticum* and seawater maybe due to the influence of the chemistry of the seaweed surface tissue (i.e. effect of seaweed metabolites on bacterial growth and attachment), which selectively attracts specific bacteria forming microbial composition driven by seaweed exudates [56, 64]. Pre-existing bacterial communities on the seaweed surface may affect the ability of settling bacteria to attach [70].

In this study, the abundance of unidentified *Rhodobacteraceae* was generally the highest in the seawater (followed by *S. muticum*) and the lowest in the sediments, indicating that these bacteria could be acquired by the seaweed from the surrounding water. Similarity in patterns between unidentified *Rhodobacteraceae* observed within *S. muticum* and in the seawater suggest that these bacteria could follow the seasonal pattern of its availability in the water column in response to variations in temperature, light availability and other factors. In contrast, *Loktanella* was more abundant in association with *S. muticum* than in the seawater across all seasons and locations, and least abundant in the sediments. An increased abundance of *Loktanella* during September–March could be linked to the fact that these bacteria could be attracted by seasonal changes in algal exudates at these months. The abundance of unidentified *Pirellulaceae* was higher in the sediments than within *S. muticum* and the lowest in the seawater across all seasons and locations, indicating that these bacteria could also be acquired by the seaweed from the sediments.

Conclusion

In this study, we demonstrated that bacterial communities associated with *S. muticum* experience significant temporal shifts as well as variation between geographic locations. The

temporal effect was reflected in significant abundance of unidentified *Rhodobacterales* and *Laktonella* in September-March and substantial prevalence of unidentified *Pirellulales* in summer. Such changes within *S. muticum* microbiota could be related to the seaweed productivity as temporally changing algal exudates attract different bacteria, which degrade algal polysaccharides and cell walls among other functions [62]. The temporal changes occurred mostly within the tip tissues and less within the basal blades and, possibly, receptacles. This could be related to the tips being younger compared to the other seaweed tissues and possibly under more direct colonization by bacteria from the surrounding environment. To what extent these re-organization and re-structuring of microbiota associated to *S. muticum* have potential consequences for the seaweed fitness and adaptation to environmental changes resulting in increased invasiveness [71] remains to be resolved.

Supporting information

S1 File. Table A. PERMANOVA results: Structure of bacterial communities associated with *S. muticum*, seawater and sediments (ti) at two locations (lo) and three months (mo). Table B. PERMANOVA Pair-wise tests between locations by month and seaweed structure of the interaction term 'location x month x tissue' of bacterial community structure associated with *S. muticum*, seawater and sediments at two locations and three months. Table C. PERMANOVA Pair-wise tests among months by location and seaweed structure of the interaction term 'location x month x tissue' of bacterial community structure associated with *S. muticum*, seawater and sediments at two locations and three months. (DOCX)

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References

1. Egan S, Thomas T, Kjelleberg S. Unlocking the diversity and biotechnological potential of marine surface-associated microbial communities. *Curr Opin Microbiol*. 2008; 11: 219–225. <https://doi.org/10.1016/j.mib.2008.04.001> PMID: 18524668
2. Hollants J, Leliaert F, De Clerck O, Willems A. What we can learn from sushi: A review on seaweed-bacterial associations. *FEMS Microbiol Ecol*. 2013; 83: 1–16. <https://doi.org/10.1111/j.1574-6941.2012.01446.x> PMID: 22775757
3. Whitman WB, Coleman DC, Wiebe WJ. Prokaryotes: the unseen majority. *Proc Natl Acad Sci*. 1998; 95(12): 6578–6583. PMID: 9618454
4. Marshall K, Joint I, Callow M, Callow J. Effect of marine bacterial isolates on the growth and morphology of axenic plantlets of the green alga *Ulva linza*. *Microb Ecol*. 2006; 52: 302–310. <https://doi.org/10.1007/s00248-006-9060-x> PMID: 16897307
5. Matsuo Y, Suzuki M, Kasai H, Shizuri Y, Harayama S. Isolation and phylogenetic characterization of bacteria capable of inducing differentiation in the green alga *Monostroma oxyspermum*. *Environ Microbiol*. 2003; 5(1): 25–35. <https://doi.org/10.1046/j.1462-2920.2003.00382.x> PMID: 12542710
6. Nakanishi K, Nishijima M, Nishimura M, Kuwano K, Saga N. Bacteria that induce morphogenesis in *Ulva pertusa* (Chlorophyta) grown under axenic conditions. *J Phycol*. 1996; 32(3): 479–482. <https://doi.org/10.1111/j.0022-3646.1996.00479.x>
7. Singh RP, Mantri VA, Reddy CRK, Jha B. Isolation of seaweed-associated bacteria and their morphogenesis-inducing capability in axenic cultures of the green alga *Ulva fasciata*. *Aquat Biol*. 2011; 12(1): 13–21. <https://doi.org/10.3354/ab00312>
8. Spoerner M, Wichard T, Bachhuber T, Stratmann J, Oertel W. Growth and thallus morphogenesis of *Ulva mutabilis* (Chlorophyta) depends on a combination of two bacterial species excreting regulatory factors. *J Phycol*. 2012; 48(6): 1433–1447. <https://doi.org/10.1111/j.1529-8817.2012.01231.x> PMID: 27009994
9. de Oliveira LS, Gregoracci GB, Silva GGZ, Salgado LT, Filho GA, Alves-Ferreira M, et al. Transcriptomic analysis of the red seaweed *Laurencia dendroidea* (Florideophyceae, Rhodophyta) and its microbiome. *BMC Genomics*. 2012; 13: 487. <https://doi.org/10.1186/1471-2164-13-487> PMID: 22985125
10. Ballestrero F, Thomas T, Burke C, Egan S, Kjelleberg S. Identification of compounds with bioactivity against the nematode *Caenorhabditis elegans* by a screen based on the functional genomics of the marine bacterium *Pseudoalteromonas tunicata* D2. *Appl Environ Microb*. 2010; 76(17): 5710–5717. <https://doi.org/10.1128/AEM.00695-10>
11. Campbell AH, Harder T, Nielsen S, Kjelleberg S, Steinberg P.D. Climate change and disease: bleaching in a chemically-defended seaweed. *Glob Chang Biol*. 2011; 17: 2958–2970. <https://doi.org/10.1111/j.1365-2486.2011.02456.x>
12. Kubanek J, Jensen PR, Keifer PA, Sullards MC, Collins DO, Fenical W. Seaweed resistance to microbial attack: a targeted chemical defense against marine fungi. *Proc Natl Acad Sci*. 2003; 100(12): 6916–6921. <https://doi.org/10.1073/pnas.1131855100> PMID: 12756301
13. Lemos ML, Toranzo AE, Barja JL. Antibiotic activity of epiphytic bacteria isolated from intertidal seaweeds. *Microb Ecol*. 1985; 11(2): 149–163. <https://doi.org/10.1007/BF02010487> PMID: 24221303
14. Paul N, de Nys R, Steinberg P. Chemical defence against bacteria in the red alga *Asparagopsis armata*: linking structure with function. *Mar Ecol Prog Ser*. 2006; 306: 87–101. <https://doi.org/10.3354/meps306087>
15. Weinberger F. Pathogen-induced defense and innate immunity in macroalgae. *Biol Bullet*. 2007; 213(3): 290–302. <https://doi.org/10.2307/25066646>
16. Wiese J, Thiel V, Nagel K, Staufenberg T, Imhoff J. Diversity of antibiotic-active bacteria associated with the brown alga *Laminaria saccharina* from the Baltic Sea. *Mar Biotechnol*. 2009; 11: 287–300. <https://doi.org/10.1007/s10126-008-9143-4> PMID: 18855068
17. Croft MT, Lawrence AD, Raux-Deery E, Warren MJ, Smith AG. Algae acquire vitamin B12 through a symbiotic relationship with bacteria. *Nature*. 2005; 438(7064): 90–93. <https://doi.org/10.1038/nature04056> PMID: 16267554
18. Campbell AH, Marzinelli EM, Gelber J, Steinberg PD. Spatial variability of microbial assemblages associated with a dominant habitat forming seaweed. *Front Microbiol*. 2015; 6: 230. <https://doi.org/10.3389/fmicb.2015.00230> PMID: 25859245

19. Bengtsson MM, Sjøtun K, Øvreås L. Seasonal dynamics of bacterial biofilms on the kelp *Laminaria hyperborea*. *Aquat Microb Ecol*. 2010; 60: 71–83. <https://doi.org/10.3354/ame01409>
20. Goecke F, Labes A, Wiese J, Imhoff JF. Chemical interactions between marine macroalgae and bacteria. *Mar Ecol Prog Ser*. 2010; 409: 267–299. <https://doi.org/10.3354/meps08607>
21. Lachnit T, Meske D, Wahl M, Harder T, Schmitz R. Epibacterial community patterns on marine macroalgae are host-specific but temporally variable. *Environ Microbiol*. 2011; 13(3): 655–665. <https://doi.org/10.1111/j.1462-2920.2010.02371.x> PMID: 21078035
22. Mancuso FP, D'hondt S, Willems A, Airoldi L, De Clerck O. Diversity and Temporal Dynamics of the Epiphytic Bacterial Communities Associated with the Canopy-Forming Seaweed *Cystoseira compressa* (Esper) Gerloff and Nizamuddin. *Front Microbiol*. 2016; 7. <https://doi.org/10.3389/fmicb.2016.00476>
23. Martin M, Portetelle D, Michel G, Vandenbol M. Microorganisms living on macroalgae: diversity, interactions, and biotechnological applications. *Appl Microbiol Biotechnol*. 2014; 98: 2917–2935. <https://doi.org/10.1007/s00253-014-5557-2> PMID: 24562178
24. Staufenberger T, Thiel V, Wiese J, Imhoff JF. Phylogenetic analysis of bacteria associated with *Laminaria saccharina*. *FEMS Microbiol Ecol*. 2008; 64: 65–77. <https://doi.org/10.1111/j.1574-6941.2008.00445.x> PMID: 18328081
25. Tujula NA, Crocetti GR, Burke C, Thomas T, Holmstrom C, Kjelleberg S. Variability and abundance of the epiphytic bacterial community associated with a green marine *Ulvean* alga. *ISME J*. 2010; 4: 301–311. <https://doi.org/10.1038/ismej.2009.107> PMID: 19829319
26. Fernandes N, Steinberg PD, Rusch D, Kjelleberg S, Thomas T. Community structure and functional gene profile of bacteria on healthy and diseased thalli of the red seaweed *Delisea pulchra*. *PLoS ONE*. 2012; 7: e50854. <https://doi.org/10.1371/journal.pone.0050854> PMID: 23226544
27. Sato Y, Willis BL, Bourne DG. Successional changes in bacterial communities during the development of black band disease on the reef coral, *Montipora hispida*. *The ISME J*. 2010; 4(2): 203–214. <https://doi.org/10.1038/ismej.2009.103> PMID: 19776765
28. Marzinelli EM, Campbell AH, Zozaya Valdes E, Vergés A, Nielsen S, Wernberg T, et al. Continental-scale variation in seaweed host associated bacterial communities is a function of host condition, not geography. *Environ Microbiol*. 2015; 17: 4078–4088. <https://doi.org/10.1111/1462-2920.12972> PMID: 26148974
29. Longford SR, Tujula NA, Crocetti GR, Holmes AJ, Holmstrom C, Kjelleberg S, et al. Comparisons of diversity of bacterial communities associated with three sessile marine eukaryotes. *Aquat Microb Ecol*. 2007; 48: 217–229. <https://doi.org/10.3354/ame048217>
30. Burke C, Thomas T, Lewis M, Steinberg P, Kjelleberg S. Composition, uniqueness and variability of the epiphytic bacterial community of the green alga *Ulva australis*. *ISME J*. 2011; 5: 590–600. <https://doi.org/10.1038/ismej.2010.164> PMID: 21048801
31. Fuhrman JA, Cram JA, Needham DM. Marine microbial community dynamics and their ecological interpretation. *Nat Rev Microbiol*. 2015; 13: 133–146. <https://doi.org/10.1038/nrmicro3417> PMID: 25659323
32. Morris JJ, Lenski RE, Zinser ER. The Black Queen Hypothesis: evolution of dependencies through adaptive gene loss. *MBio*. 2012; 3. <https://doi.org/10.1128/mBio.00036-12>
33. O'Brien PA, Morrow KM, Willis B, Bourne D. Implications of Ocean Acidification for Marine Microorganisms from the Free-Living to the Host-Associated. *Front Mar Sci*. 2016; 3(47). <https://doi.org/10.3389/fmars.2016.00047>
34. Joint I, Doney SC, Karl DM. Will ocean acidification affect marine microbes? *The ISME J*. 2011; 5(1): 1–7. <https://doi.org/10.1038/ismej.2010.79> PMID: 20535222
35. Liu J, Weinbauer MG, Maier C, Dai M, Gattuso JP. Effect of ocean acidification on microbial diversity and on microbe-driven biogeochemistry and ecosystem functioning. *Aquat Microb Ecol*. 2010; 61: 291–305. <https://doi.org/10.3354/ame01446>
36. Li J, Chen Q, Long LJ, Dong J, Yang J, Zhang S. Bacterial dynamics within the mucus, tissue and skeleton of the coral *Porites lutea* during different seasons. *Sci Rep*. 2014; 4:7320. <https://doi.org/10.1038/srep07320> PMID: 25475855
37. Taylor MW, Schupp PJ, Dahllöf I, Kjelleberg S, Steinberg PD. Host specificity in marine sponge-associated bacteria and potential implications for marine microbial diversity. *Environ Microbiol*. 2004; 6: 121–130. <https://doi.org/10.1046/j.1462-2920.2003.00545.x> PMID: 14756877
38. Engelen AH, Serebryakova A, Ang P, Britton-Simmons K, Mineur F, Pedersen MF, et al. Circumglobal invasion by the brown seaweed *Sargassum muticum*. *Oceanography and Marine Biology: An Annual Review*. 2015; 53: 81–126. <https://doi.org/10.1201/b18733-4>
39. Lima FP, Ribeiro PA, Queiroz N, Hawkins SJ, Santos AM. Do distributional shifts of northern and Southern species of algae match the warming pattern? *Global Change Biology*. 2007; 13: 2592–2604. <https://doi.org/10.1111/j.1365-2486.2007.01451.x>

40. Lane DJ. 1991. 16S/23S rRNA sequencing. *Nucleic acid techniques in bacterial systematics*: 125–175.
41. Bodenhausen N, Horton MW, Bergelson J. 2013. Bacterial Communities Associated with the Leaves and the Roots of *Arabidopsis thaliana*. *PLoS One* 8(2):e56329. <https://doi.org/10.1371/journal.pone.0056329> PMID: 23457551
42. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, et al. 2010. QIIME allows analysis of high-throughput community sequencing data. *Nature Methods* 7(5):335–6. <https://doi.org/10.1038/nmeth.f.303> PMID: 20383131
43. Haas BJ, Gevers D, Earl AM, Feldgarden M, Ward DV, Giannoukos G. et al. Chimeric 16S rRNA sequence formation and detection in Sanger and 454-pyrosequenced PCR amplicons. *Genome research*. 2011; 21(3): 494–504. <https://doi.org/10.1101/gr.112730.110> PMID: 21212162
44. Edgar RC. Search and clustering orders of magnitude faster than BLAST. *Bioinformatics* 2010; 26(19): 2460–2461. <https://doi.org/10.1093/bioinformatics/btq461> PMID: 20709691
45. Wang G, Shuai L, Li Y, Lin W, Zhao X, Duan D. Phylogenetic analysis of epiphytic marine bacteria on hole-rotten diseased sporophytes of *Laminaria japonica*. *J Appl Phycol*. 2008; 20: 403–409.
46. McDonald D, Price MN, Goodrich J, Nawrocki EP, DeSantis TZ, Probst A. An improved Greengenes taxonomy with explicit ranks for ecological and evolutionary analyses of bacteria and archaea. *The ISME J*. 2012; 6(3): 610. <https://doi.org/10.1038/ismej.2011.139> PMID: 22134646
47. Chao A. 1984. Nonparametric estimation of the number of Classes in a population. *Scandinavian Journal of Statistics* 11(4):265–270. <https://doi.org/10.2307/4615964>
48. Clarke KR, Warwick RM. 1994. Change in Marine Communities: An Approach to Statistical Analysis and Interpretation. Plymouth: Plymouth Marine Laboratory, Natural Environment Research Council.
49. Clarke KR, Gurley RN. 2006. PRIMER v6: user manual/tutorial (Plymouth Routines in Multivariate Ecological Research). Plymouth: Primer-E Ltd.
50. Jiang YF, Ling J, Dong JD, Chen B, Zhang YY, Zhang YZ, et al. Illumina-based analysis the microbial diversity associated with *Thalassia hemprichii* in Xincun Bay, South China Sea. *Ecotoxicology*. 2015; 24(7–8): 1548–1556. <https://doi.org/10.1007/s10646-015-1511-z> PMID: 26092035
51. Ettinger CL, Voerman SE, Lang JM, Stachowicz JJ, Eisen JA. Microbial communities in sediment from *Zostera marina* patches, but not the *Z. marina* leaf or root microbiomes, vary in relation to distance from patch edge. *PeerJ* 2017; 5: e3246. <https://doi.org/10.7717/peerj.3246> PMID: 28462046
52. Palacios R, Newton WE. Genomes and genomics of nitrogen-fixing organisms. Dordrecht: Springer; 2005.
53. Dang H, Li T, Chen M, Huang G. Cross-ocean distribution of *Rhodobacterales* bacteria as primary surface colonizers in temperate coastal marine waters. *Appl Environ Microb*. 2008; 74: 52–60. <https://doi.org/10.1128/AEM.01400-07>
54. Stratil SB, Neulinger SC, Knecht H, Friedrichs AK, Wahl M. Temperature-driven shifts in the epibiotic bacterial community composition of the brown macroalga *Fucus vesiculosus*. *Microbiology open*. 2013; 2: 338–349. <https://doi.org/10.1002/mbo3.79> PMID: 23568841
55. Zozaya-Valdes E, Egan S, Thomas T. A comprehensive analysis of the microbial communities of healthy and diseased marine macroalgae and the detection of known and potential bacterial pathogens. *Front Microbiol*. 2015; 6: 146. <https://doi.org/10.3389/fmicb.2015.00146> PMID: 25759688
56. Egan S, Harder T, Burke C, Steinberg P, Kjelleberg S, Thomas T. The seaweed holobiont: understanding seaweed–bacteria interactions. *FEMS Microbiol Rev*. 2013; 37(3): 462–476. <https://doi.org/10.1111/1574-6976.12011> PMID: 23157386
57. Miranda LN, Hutchison K, Grossman AR, Brawley SH. Diversity and abundance of the bacterial community of the red macroalga *Porphyra umbilicalis*: did bacterial farmers produce macroalgae? *PLoS One*. 2013; 8: e58269. <https://doi.org/10.1371/journal.pone.0058269> PMID: 23526971
58. Bengtsson MM, Sjøtun K, Storesund JE, Øvreås L. Utilization of kelp-derived carbon sources by kelp-surface-associated bacteria. *Aquat Microb Ecol*. 2011; 62: 191–199. <https://doi.org/10.3354/ame01477>
59. Wahl M, Goecke F, Labes A, Dobretsov S, Weinberger F. The second skin: ecological role of epibiotic biofilms on marine organisms. *Front Microbiol*. 2012; 3. <https://doi.org/10.3389/fmicb.2012.00292>
60. Mensch B, Neulinger SC, Graiff A, Pansch A, Künzel S, Fischer MA, et al. Restructuring of epibacterial communities on *Fucus vesiculosus* forma *mytili* in response to elevated pCO₂ and increased temperature levels. *Front Microbiol*. 2016; 7. <https://doi.org/10.3389/fmicb.2016.00434>
61. Lage OM, Bondoso J. *Planctomycetes* and macroalgae, a striking association. *Front Microbiol*. 2014; 5: 267. <https://doi.org/10.3389/fmicb.2014.00267> PMID: 24917860
62. Bengtsson MM, Øvreås L. 2010. *Planctomycetes* dominate biofilms on surfaces of the kelp *Laminaria hyperborea*. *BMC Microbiology* 10:261 <https://doi.org/10.1186/1471-2180-10-261> PMID: 20950420

63. Bengtsson MM, Sjøtun K, Lanzén A, Øvreås L. Bacterial diversity in relation to secondary production and succession on surfaces of the kelp *Laminaria hyperborea*. ISME J. 2012; 2188–2198. <https://doi.org/10.1038/ismej.2012.67> PMID: 22763650
64. Michelou VK, Caporaso JG, Knight R & Palumbi SR. The ecology of microbial communities associated with *Macrocystis pyrifera*. PLoS One. 2013; 8: e67480. <https://doi.org/10.1371/journal.pone.0067480> PMID: 23840715
65. Fuerst JA. *Planctomycetes: cell structure, origins and biology*. New York: Springer; 2013.
66. Glöckner FO, Kube M, Bauer M, Teeling H, Lombardot T, Ludwig W, et al: Complete genome sequence of the marine planctomycete *Pirellula* sp. strain 1. Proc Natl Acad Sci USA. 2003, 100 (14): 8298–8303. <https://doi.org/10.1073/pnas.1431443100> PMID: 12835416
67. Wegner CE, Richter-Heitmann T, Klindworth A, Klockow C, Richter M, Achstetter T, et al. Expression of sulfatases in *Rhodopirellula baltica* and the diversity of sulfatases in the genus *Rhodopirellula*. Mar Genom. 2013; 9: 51–61. <https://doi.org/10.1016/j.margen.2012.12.001>
68. Bondoso J, Balagué V, Gasol JM, Lage OM. Community composition of the *Planctomycetes* associated with different macroalgae. FEMS Microbiol Ecol. 2014; 88(3): 445–456. <https://doi.org/10.1111/1574-6941.12258> PMID: 24266389
69. Lachnit T, Fischer M, Künzel S, Baines JF, Harder T. Compounds associated with algal surfaces mediate epiphytic colonization of the marine macroalga *Fucus vesiculosus*. FEMS Microbiol Ecol. 2013; 84 (2): 411–420. <https://doi.org/10.1111/1574-6941.12071> PMID: 23311942
70. Singh RP, Reddy CRK. Seaweed–microbial interactions: key functions of seaweed-associated bacteria. FEMS Microbiol Ecol. 2014; 88(2): 213–230. <https://doi.org/10.1111/1574-6941.12297> PMID: 24512602
71. Aires T, Serebryakova A, Viard F, Serrão EA, Engelen AH. Acidification increases abundances of *Vibrionales* and *Planctomycetia* associated to a seaweed- grazer system: potential consequences for disease and prey digestion efficiency. PeerJ. 2018; 6: e4377. <https://doi.org/10.7717/peerj.4377> PMID: 29610702