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Accumulation of detached kelp biomass in a subtidal temperate coastal ecosystem induces succession of epiphytic and sediment bacterial communities

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Summary
Kelps are dominant primary producers in temperate coastal ecosystems. Large amounts of kelp biomass can be exported to the seafloor during the algal growth cycle or following storms, creating new ecological niches for the associated microbiota. Here, we investigated the bacterial community associated with the kelp Laminaria hyperborea during its accumulation and degradation on the seafloor. Kelp tissue, seawater and sediment were sampled during a 6-month in situ experiment simulating kelp detritus accumulation. Evaluation of the epiphytic bacterial community abundance, structure, taxonomic composition and predicted functional profiles evidenced a biphasic succession. Initially, dominant genera (Hellea, Litorimonas, Granulosicoccus) showed a rapid and drastic decrease in sequence abundance, probably outcompeted by algal polysaccharide-degraders such as Bacteroidia members which responded within 4 weeks. Acidimicrobia, especially members of the Sva0996 marine group, colonized the degrading kelp biomass after 11 weeks. These secondary colonizers could act as opportunistic scavenger bacteria assimilating substrates exposed by early degraders. In parallel, kelp accumulation modified bacterial communities in the underlying sediment, notably favouring anaerobic taxa potentially involved in the sulfur and nitrogen cycles. Overall, this study provides insights into the bacterial degradation of algal biomass in situ, an important link in coastal trophic chains.

Introduction
Brown macroalgae of the order Laminariales, collectively known as kelps, are crucial foundation species that thrive in coastal regions of temperate and Arctic seas worldwide. Kelps feature some of the highest growth rates and primary productivity on the planet (Mann, 1973). Kelp biomass thus represents an abundant standing stock of organic matter, primarily composed of polysaccharides and proteins that account for ca. 50% and 3%–15% of the dry weight, respectively (Kloareg and Quatrano, 1988; Fleurence, 1999). Up to 82% of the kelp primary production can be exported as dissolved or particulate organic matter to the shores or neighbouring intertidal, subtidal and benthic zones (Krumhansl and Scheibling, 2012). This may be due to the drift of naturally detached old blades as well as the fragmentation or dislodging of kelps caused by hydrodynamic forces such as strong storm events (Krumhansl and Scheibling, 2012; de Bettignies et al., 2015; Krause-Jensen and Duarte, 2016). This massive input of organic matter can strongly influence the stranding ecosystems, creating new niches for meiofauna and microorganisms (Duarte and Cebrián, 1996). This is particularly relevant in benthic subtidal habitats where sediments are otherwise devoid of kelps. The turnover of algal biomass largely depends on bacterial degradation processes (Mann, 1982; Newell et al., 1982) and is influenced by several physico-chemical parameters. For example, small fragments are degraded faster, and degradation is favoured when temperature and hydrodynamics increase (Krumhansl and Scheibling, 2012).
Live macroalgae host diverse and abundant bacterial communities (Burke et al., 2011), with about $10^6$ to $10^9$ bacterial cells cm$^{-2}$ on algal surfaces (Martin et al., 2014). Bacteria play an important role in algal health and physiology: they can provide vitamins to the host, defend the host against pathogens via antibiotic production, or degrade algal tissues (Egan et al., 2013, 2014; Minich et al., 2018). Epiphytic bacterial communities have been thoroughly characterized for an increasing number of kelp species and generally comprise large proportions of Alphaproteobacteria, Gammaproteobacteria, Bacteroidetes, Verrucomicrobia and Planctomycetes (Bengtsson and Øvreås, 2010; Lemay et al., 2018). Studies further revealed that kelp epiphytic communities are stable over large spatial scales (Marzinelli et al., 2015), can be species-specific (Lemay et al., 2018), depend on the age of algal tissues (Weigel and Pfister, 2019; Ramirez-Puebla et al., 2020) and follow reproducible seasonal successions that may be explained by water temperature (Bengtsson et al., 2010; Minich et al., 2018). In addition, physiological parameters varying at the algal surface may impact communities. Oxygen release by the alga can promote the growth of aerobic bacteria but also harm epiphytic bacteria by producing reactive oxygen species (ROS) (Goecke et al., 2010; Egan et al., 2013). Kelps may also control resident or pathogenic microorganisms by secreting secondary metabolites such as halogenated compounds (Egan et al., 2014). In contrast to epiphytic communities on intact macroalgae, the specific bacterial communities colonizing detached kelps and their succession are still understudied. Total microflora has been determined essentially from kelp detritus stranded on intertidal areas (Koop et al., 1982; Bouvy et al., 1986; Delille and Perret, 1991). Even less information is available on the taxonomy, function and succession of bacterial communities colonizing large submersed accumulations of kelp fragments on benthic subtidal habitats. Furthermore, kelp accumulation and degradation processes are also expected to influence sediment communities, which are known to harbour contrasted and more diverse microbial communities compared to the overlying water column (Gobet et al., 2012).

We previously reported an experimental approach to mimic dislodgement and accumulation of the kelp *Laminaria hyperborea* on a subtidal sandy bottom ecosystem of the north-western coast of Brittany, France (de Bettignies et al., 2020). We showed that algal biomass decomposition started after 2 weeks and reached a critical step after 11 weeks, with an increase in respiration rate and phlorotannin content. After 24 weeks, 85% of the algal biomass was lost. Using samples collected during the same experiment, we investigated here the changes in bacterial community abundance and composition and predicted functions associated with the *in situ* degradation of the kelp biomass over 6 months, in four compartments: *L. hyperborea*’s tissues, sediments directly underneath the experimental cages (hereafter ‘underlying sediment’), control sediments 2 m away from the cages and devoid of kelp tissues (hereafter ‘external sediment’), and the overlying seawater 1 m above the cages. We tested the following hypotheses: (i) the bacterial diversity and community structure are specific to a given compartment; (ii) there is a succession of different bacterial groups associated with the degradation of *L. hyperborea*’s tissues; and (iii) bacterial communities in the underlying sediment are affected by the degrading kelp tissues and different from the external sediment.

**Results**

**Fluctuations and diversity patterns of the bacterial community during kelp degradation**

The characteristics of the 92 samples analysed in this study are listed in the Supporting Information Table S1. Total bacterial abundance was estimated by quantitative PCR in each compartment using 16S rRNA gene copy number as a proxy. The average number of 16S rRNA gene copies on kelp surfaces was initially $2.27 \times 10^8$ copies cm$^{-2}$. Considering an average of four copies of 16S rRNA gene per bacterial cell (Větrovský and Baldrian, 2013), it corresponds to ca. $6 \times 10^9$ cells cm$^{-2}$. It decreased significantly to $5.65 \times 10^7$ copies cm$^{-2}$ (ca. $1.4 \times 10^7$ cells cm$^{-2}$) during the first 2 weeks and remained stable afterwards [analysis of variance (ANOVA), $F_{7,16} = 7.37$, $P < 0.001$; Fig. 1A]. Over the 6 months, 16S rRNA gene copy number significantly increased in the surrounding seawater (ANOVA, $F_{7,16} = 4.02$, $P = 0.010$), the underlying sediment (ANOVA $F_{7,16} = 3.83$, $P = 0.012$) and to a lesser extent in the external sediment (ANOVA $F_{7,14} = 4.32$, $P = 0.010$) (Fig. 1B and C). Notably, the number of 16S rRNA gene copies was similar between the two sediment compartments the first 2 weeks, but the increase tended to be stronger in the underlying sediment (4.6-fold change between 0 and 24 weeks) than in the external sediment (1.7-fold change).

Alpha-diversity indices were two to three times higher in sediment samples compared to kelp-associated and water column samples (Fig. 1D and F; Supporting Information Fig. S1). The OTU richness and the Shannon index from kelp-associated samples increased significantly in the first 11 weeks (OTU richness from 351–493 to 2090–2312, Shannon index from 2.4–3.1 to 5.4–5.6; ANOVA, $F_{7,15} = 44.68$, $P < 0.001$ and $F_{7,15} = 54.78$, $P < 0.001$, respectively) before reaching a plateau. Bacterial diversity slightly increased in the water column throughout the experiment (ANOVA, $F_{7,16} = 48.73$, $P = 0.001$ for OTU richness and $F_{7,16} = 62.50$, $P = 0.001$).
for Shannon index) while it stayed stable in the underlying and the external sediment samples ($P > 0.05$). Further, sample compartment showed a strong impact on the bacterial community structure (permutational analysis of variance PERMANOVA, $F_{3,84} = 35.77$, $P < 0.001$, Fig. 2A). Pairwise comparisons showed significant dissimilarities between each compartment ($P = 0.001$), although less pronounced between external and underlying sediments (Supporting Information Table S2). A higher dispersion was visible on the principal coordinates analysis (PCoA) within the algal samples (mean dispersion: 0.60) compared to the other compartments (seawater 0.55, underlying sediment 0.56, external sediment 0.53; $P < 0.001$). Only 1 946 OTUs, i.e. 2.2% of the total number of OTUs were shared between the four compartments (Fig. 2B). The sequences contained in this core set of OTUs are present in different proportions of the complete dataset in each compartment, they represented 12% of the sequence abundance in kelp-associated samples, 17% in seawater, 12% in underlying sediment and 7% in external sediment. In contrast, 58.2% (51 344 OTUs) of the total number of OTUs were specific to a given compartment, but they represented only 10% of the total number of sequences in the dataset.

**Succession of epiphytic bacteria on kelp tissues**

The structure of the kelp-associated bacterial community was impacted by the elapsed degradation time (PERMANOVA, $F_{7,15} = 6.33$, $P = 0.001$), showing distinct groups for freshly detached kelps (week 0) and samples collected between weeks 2–6 or 11–24 (Fig. 3A). Pairwise ANOVA comparisons confirmed that samples from weeks 2–6 significantly differed from samples from weeks 11–24 ($F_{1,19} = 13.46$, $P = 0.003$). Although visible on the ordination plot, the difference between samples from week 0 and the other groups could not be reliably tested due to one missing sample. Constrained ordination using distance-based redundancy analysis (db-RDA) showed that four environmental parameters were significant explanatory variables, namely remaining biomass...
To further understand these patterns of the community structure, we investigated fluctuations of bacterial taxonomic groups at the kelp surface. The epiphytic community of freshly detached kelps was dominated by Alphaproteobacteria and Gammaproteobacteria (ca. 62% and 32% of all sequences at week 0, respectively), while other classes represented less than 3% (Fig. 3B). In particular, 83% of the sequences retrieved from kelp-associated samples at week 0 belonged to only 10 OTUs, with four of them assigned to Litorimonas and three to Hellea (Supporting Information Fig. S2). 

In situ degradation of kelp biomass was associated with a succession pattern in the taxonomic composition of the epiphytic community (Fig. 3B; Supporting Information Fig. S2). Alphaproteobacteria remained the most abundant class throughout the experiment, although their contribution started decreasing after 4 weeks and reached 40% after 24 weeks. The proportion of Bacteroidia increased early on from 2.5% at week 2 to 14.6% after

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Kelp-degrading bacterial communities

Fig 3. Fluctuations of kelp-associated bacterial communities during the degradation of detached *L. hyperborea*. A. db-RDA plot with vectors representing significant contextual predictors. Samples from the same sampling date are connected by coloured segments. B. Relative abundance of bacterial classes in kelp-associated samples during the 6 month-experiment. Sequence relative abundances of classes representing less than 0.5% of all sequences at each sampling time were summed into the ‘Other’ category. Values are mean ± standard deviation (n = 3, except for T0 n = 2). C. Relative abundance of differentially abundant kelp-associated OTUs during the experiment. OTUs showing significant differential abundance (P < 0.05) and representing at least 1.5% of the sequences in at least one sample are shown. OTUs were annotated at the genus level. For unclassified genera, taxa were assigned to the lowest taxonomic level identified. OTUs were grouped by class and order. Pl: *Plactomycetacia*; Acidi: *Acidimicrobia*; Bact: *Bacteroidia*.

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24 weeks. *Acidimicrobia* were rare (<0.5%) until week 6 but increased afterwards, peaking at 22% relative sequence abundance at week 20. The contribution of *Deltaproteobacteria* increased slightly from the initial sampling date (0.6%) to week 11 (3.2%) and stayed stable afterwards. This pattern differed from the one observed in the surrounding seawater (Supporting Information Fig. S3A), where a seasonal increase of *Gammaproteobacteria* during summer (weeks 11–15) coincided with a decrease of *Alphaproteobacteria* and *Bacteroidia*.

A differential abundance test detected 249 OTUs (i.e. 1.8% of the total OTUs) for which the relative sequence abundance significantly changed with kelp degradation time (*P* < 0.05 (Supporting Information Table S3)). For clarity, we only display in Fig. 3C the 54 differential OTUs that represent ≥1.5% of the total community in at least one sample. Among these, seven OTUs affiliated to *Granulosicoccus* (*Gammaproteobacteria*) and *Litorimonas* and *Hellea* (*Alphaproteobacteria*) were abundant at the beginning of the experiment and decreased over time (*Tukey;* *P* < 0.001 between week 0 and 2). This decrease was particularly drastic for OTU10 (*Granulosicoccus*) and OTU14 (*Hellea*), which accounted for 25.4% and 22.5% at week 0 and only 0.03% and 1.1% after 24 weeks respectively (*P* < 0.001). The sequence abundance of 18 *Alphaproteobacteria* OTUs out of the 32 represented in Fig. 3C increased during the first 4 weeks of the experiment and then decreased. The sequence abundance of 8 out of 14 *Gammaproteobacteria* OTUs, including members of *Cellvibionales*, *Ectothiorhodospirales* and the KI89A clade, increased after 11 weeks of degradation. In particular, one OTU related to *Thiohalophilus* (OTU22) peaked at 7% mean relative sequence abundance at week 20. The sequence abundance of four OTUs related to the Sva0996 marine group (*Microtrichales*, *Acidimicrobia*) showed an important increase between 6 and 11 weeks of experiment (*P* = 0.003). The most abundant one (OTU16) represented 0.1% of the communities after 6 weeks vs. 5.6% after 6 months. Finally, 14% (35/249) of the differential OTUs belonged to *Bacteroidia* (Supporting Information Table S3) but are not represented in Fig. 3C since they did not pass the 1.5% abundance threshold we chose for clarity.

We built a correlation network of the kelp-associated bacterial communities to search for potential associations (Fig. 4). The final network comprised 134 nodes and 572 edges. Sixty-seven percent of the associations were positive, including those with a maximum delay of one sampling date (384 and 188 positive and negative associations, respectively). Among the 10 contextual variables included in the dataset, eight showed significant associations within the network. The variables with the most associations were ‘Temperature’ (23 significant edges), ‘C/N’ (22 significant edges) and ‘Respiration’ (19 significant edges) (Supporting Information Fig. S4). Most positive associations of OTU relative sequence abundance to respiration and C/N ratio were delayed. Community respiration was negatively correlated with algal biomass, indicating that the bulk of oxygen consumption did not come from algal respiration. Instead, respiration was positively correlated with temperature, which is expected to increase bacterial activity. Eight clusters were detected in the network (Supporting Information Table S4), which might represent strong microbial associations, i.e. nodes that are highly connected to each other but weakly connected to nodes outside of their module. In particular, the two strongest clusters 1 and 2 comprised 66% of the nodes in the network (Fig. 4, inset) and were almost mutually exclusive, suggesting they denote two different communities during the bacterial succession. Most OTUs from cluster 2 showed a major change in their relative sequence abundance during the first 6 weeks (Supporting Information Fig. S5). This suggests that cluster 2 represents associations occurring early during the bacterial succession. Seventy-five percent of these OTUs from cluster 2 belonged to *Alphaproteobacteria* (Supporting Information Table S5). None of these OTUs had significant associations to environmental parameters (Fig. 4). In contrast, OTUs from cluster 1 were more diverse (Supporting Information Table S5), significantly associated with a number of environmental parameters (Fig. 4) and showed a later response (Supporting Information Fig. S6). Only four OTUs were shared between clusters 1 and 2 (Fig. 4, inset, nodes in crimson colour), including OTU923 (*Arenicella, Gammaproteobacteria*), OTU190 (*Saprospiraceae, Bacteroidia*), OTU5587 (*Hellea, Alphaproteobacteria*) and OTU361 (*Litorimonas, Alphaproteobacteria*). These shared OTUs showed the highest closeness centrality (a measure indicating the distance of a node to all other nodes) of the entire network (CC > 0.4) and might therefore act as hubs connecting the two types of communities found in clusters 1 and 2.

Putative functional profiles were inferred from 16S rRNA data using FAPROTAX for the kelp-associated communities (Fig. 5), leading to the assignation of 3 753 OTUs to at least one functional group (31.4% of all OTUs). The predicted abundance of functional groups involved in aerobic chemoheterotrophy and the oxidation of sulfur compounds increased from 2 weeks on before a slight decrease at 11 weeks. Moreover, predicted groups involved in fermentation and nitrate reduction were virtually absent initially (<18 and 28 estimated sequence read counts, respectively) and more represented during the degradation (55–997 and 177–899 estimated sequence read counts, respectively). Since sugars constitute a major part of algal biomass, we specifically assessed the
predicted relative sequence abundance of glycoside hydrolases (GH) and polysaccharide lyases (PL) with PICRUSt2. The mean Nearest Sequenced Taxon Index (NSTI) was 0.21 ± 0.03. NSTI estimates the extent to which microorganisms in a given sample are related to sequenced genomes used to infer metagenomes and ranges from 0 (high similarity) to 1 (no similarity). The relatively high NSTI values obtained here denote that few genomes of kelp-associated bacteria are available yet, and that predictions should be interpreted carefully. The
predicted relative sequence abundance of endolytic enzymes such as cellulase and endo-1,4-β-xylanase decreased over time (Supporting Information Fig. S7). In contrast, the predicted relative sequence abundance of several exolytic GHs increased significantly with elapsed time of degradation, either at an early stage, from 2 weeks (i.e. α-glucosidase, α-galactosidase and β-mannosidase) or later, after 6 weeks (i.e. α-amylase, 4-α-D-(1- > 4)-α-D-Glc)trehalose trehalohydrolase, α-mannosidase).

Effect of algal degradation on sediment bacterial communities

We further focused on the effect of kelp accumulation on sediment bacterial communities of the receiving ecosystem. Beta-diversity analysis of the underlying sediment samples showed an effect of the elapsed degradation time on the bacterial community structure (Fig. 6A) (PERMANOVA, $F_{7,16} = 2.10$, $P < 0.001$). Constrained ordination using db-RDA showed that three environmental parameters were significant explanatory variables, namely remaining biomass ($F_{1,16} = 4.53$, $P = 0.001$), respiration ($F_{1,16} = 1.97$, $P = 0.009$) and temperature ($F_{1,16} = 1.95$, $P = 0.012$). Sediment communities were initially dominated by Gammaproteobacteria and Bacteroidia (about 36% and 23% of all sequences at week 0, respectively) (Fig. 6B). Only ca. 9.5% of OTUs present in the underlying or external sediments were shared with the algal epiphytic microbiota (Fig. 2B). The abundance of these OTUs in the sediments and in the kelp-associated compartments represented 22.3% and 20.8% of the overall sequence abundance, respectively. Bacterial communities in external sediments were stable with a slight increase of Bacteroidia during kelp degradation (Supporting Information Fig. S3B), while changes were more pronounced in the underlying sediment (Fig. 6B). Furthermore, about 40% of the OTUs detected in the underlying sediments (containing 13% of the number of sequences in the underlying sediments) were not found in the external sediments (Fig. 2B). In underlying sediments, the sequence relative abundance of Gammaproteobacteria decreased.

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until week 6, reaching 22% before an increase at 31% and stabilization around 27%. This was different from external sediments, where Gammaproteobacteria gradually decreased to 27% until week 15. The sequence relative abundance of Bacteroidia increased rapidly in underlying sediments, peaking at about 40% after 6 weeks before stabilization around 31%. This increase was less pronounced in external sediments. The relative sequence abundance of Alphaproteobacteria in underlying sediments slightly decreased from about 14% at week 0 to
10% at week 24, while it stayed stable in external sediments. Deltaproteobacteria were enriched along the time course, representing about 9% of the total sequence relative abundance at week 0 and reaching 15% and 11% at week 24 in underlying and external sediments, respectively. Differential abundance analysis detected 177 OTUs (i.e. 0.3% of the total number of OTUs) for which the relative sequence abundance significantly changed in the underlying sediment during kelp degradation ($P < 0.05$), while 27 were detected in external sediments (Supporting Information Table S6 and S7). Among these, only 11 OTUs affiliated with Bacteroidia, Deltaproteobacteria and Gammaproteobacteria were found to significantly vary with time in both underlying and external sediments (Supporting Information Fig. S8). For consistency, we applied the same filter to differential OTUs in underlying sediment than that used for kelp-associated samples (>1.5% relative abundance in at least one sample), but only seven OTUs passed this criteria. Consequently, we lowered the cut-off at 0.5% to show 35 differential OTUs (Fig. 6C). The sequence abundance of the 35 represented OTUs stayed generally stable during the first 2 weeks. The sequence abundance of 16 members of Bacteroidales and Flavobacteriales (Bacteroidia) as well as two Clostridiales (Clostridia) and two Rhodobacterales (Alphaproteobacteria) increased between 0 and 4 weeks (Tukey; $P = 0.015$, $P = 0.005$, $P = 0.010$, respectively) and decreased afterwards ($P = 0.001$, $P = 0.044$, $P = 0.037$ between 6 and 24 weeks, respectively). Five Desulfovobacterales OTUs (Deltaproteobacteria) increased between 0 and 6 weeks ($P = 0.015$) and two Sulfurovum OTUs (Campylobacteria) increased between 0 and 15 weeks ($P = 0.011$). Putative functional profiles were inferred for underlying sediment communities (Fig. 5). The 10 657 OTUs were assigned to at least one functional group (24.5%). The predicted abundance of functional groups involved in aerobic chemoheterotrophy, nitrogen cycle and fermentation increased during the first 4 weeks and remained stable afterwards. In contrast, sulfate-reducing functional groups increased throughout the degradation process. The predicted sequence abundance of the sulfate respiration group increased 10 times over 24 weeks.

Discussion

In temperate coastal ecosystems, large amounts of kelp tissues are exported to the seafloor naturally or following seasonal storms. To investigate the degradation dynamics and the effect of kelp tissue accumulation on receiving ecosystems, a 6-month experiment was deployed on a low subtidal sandy floor. Here, we investigated the bacterial community succession associated with the degradation of L. hyperborea tissue accumulated on the seafloor in four compartments: the degrading kelp, the seawater, the underlying and the external sediment.

Microbial communities are specific to a given compartment

Bacterial diversity patterns showed specific communities and succession in the four compartments. In particular, half of the kelp-associated OTUs were not detected in any other compartment. This observation was expected as each compartment features contrasting biotic and abiotic conditions. The kelp surface is an interface between the alga and its surrounding environment where specific metabolic exchanges occur (e.g. defence, biofilm control). This is an organic-rich environment, ideal for copiotrophs but with high competition between microorganisms for surface colonization, which might select for K-strategists (Andrews and Harris, 1986; Fierer et al., 2007). In contrast, the water column is more substrate-depleted, which may select for oligotrophs and K-strategists. Finally, sediments concentrate decaying organic matter, available for r-strategist copiotrophs, and oxygen availability as well as chemical conditions varying with depth offer multiple microniches. This likely explains the higher diversity with various microbial life strategy-types in this compartment compared to kelp-associated and seawater samples. Overall, bacterial diversity was not strongly impacted in the seawater and external sediments during the 6-month period, pointing to a local effect of kelp accumulation. Indeed, constant water renewal due to tides and currents likely dilutes the compounds liberated upon degradation (Masson, 2002). In the following paragraphs, we focus on the major changes observed in kelp-associated and underlying sediment communities.

Bacterial communities associated with healthy kelp tissue

Throughout the experiment, kelp-associated bacterial communities were dominated by Alphaproteobacteria and Gammaproteobacteria. These bacterial classes are regularly found among the most abundant on brown algal tissue (Lachnit et al., 2011; Stratil et al., 2013; Martin et al., 2015; Ramirez-Puebla et al., 2020). At the initial sampling date, in April 2017, the surface of fresh L. hyperborea hosted an abundant resident bacterial community, concurring with previous studies showing $10^6$–$10^9$ cells.cm$^{-2}$ on kelp surfaces (Corre and Prieur, 1990; Martin et al., 2014). The initial community had a low diversity and was dominated by a few OTUs affiliated to Granulosicoccus (Gammaproteobacteria), Hellea and Litorimonas (Alphaproteobacteria). These genera were always present in the epiphytic community of L. hyperborea and were previously detected on other brown algae such as Macrocystis pyrifera (Florez et al., 2019), Taonia atomaria (Paix et al., 2019), Fucus vesiculosus (Lachnit et al., 2011; Stratil et al., 2013), and
L. digitata (Gobet, personal communication). We detected < 1% Planctomycetia and Verrucomicrobia in the initial communities. This contrasts with previous studies using fluorescent in situ hybridization and denaturing gel gradient electrophoresis where these taxa contributed up to 50% of epiphytic bacteria on L. hyperborea at the Norwegian coast (Bengtsson and Øvreås, 2010; Bengtsson et al., 2010). This discrepancy may be due to (i) ecological differences between Norway and Brittany coasts, (ii) seasonal variations and (iii) the technique used to survey the microbial community (Zinger et al., 2012). The primer used for our metabarcoding approach, designed to minimize amplification of plastid sequences, may partly miss Planctomycetia and Verrucomicrobia (Thomas et al., 2020).

Dynamics of bacterial communities at the surface of degrading kelp tissues

The kelp-associated bacterial community shifted after 2 weeks. 16S rRNA gene copies decreased fourfold and stayed stable during the 6 months. This could reflect both a decrease in the total number of bacteria attached to the kelp surface and a shift towards taxa featuring fewer 16S rRNA gene copies per genome such as Acidobacteria (Vetrovsky and Baldrian, 2013). Analysis of kelp-associated community composition further showed a biphasic succession, with changes during the first 11 weeks followed by stabilization of the bacterial community composition until 24 weeks. This corroborates the kinetics of algal biomass degradation (de Bettignies et al., 2020), which showed a rapid loss of algal biomass until week 11 that slowed down until week 24. During the 6-month experiment, oxygen consumption increased in cages while algal biomass decreased. Although this could partially be due to colonization by fauna (de Bettignies et al., 2020), it likely indicates the important contribution of kelp-degrading bacteria to the holobiont respiration (here, the kelp host and its associated fauna and microbiota) (de Bettignies et al., 2020). Overall, we suggest that the observed changes in kelp-associated bacterial communities result from a succession of heterotrophic taxa with different substrate niches, as previously observed for the degradation of terrestrial plant biomass (Jiménez et al., 2017) and marine polysaccharide particles (Enke et al., 2019). During the first phase, kelp biomass was colonized by bacteria specialized in the degradation of complex algal polysaccharides and exudates. This corroborates results we previously obtained, showing an increase of mannitol- and alginate-utilizing bacteria between 0 and 2 weeks (de Bettignies et al., 2020). The activity of these early degraders likely opened new substrate niches and increased the accessibility for opportunistic scavenger bacteria that responded during the second phase. Some of these changes might be partly due to seasonal variations that would also occur on intact, non-detached algal tissue. This is suggested by the significant effect of temperature and light found with db-RDA and the number of associations between individual OTUs and temperature in the network. Previous studies showed that seasonal parameters influence brown algal microbiomes, notably increasing temperature that can favour the growth of Flavobacteriales, Rhodobacteriales and Rhizobiales (Stratil et al., 2013; Minich et al., 2018). In addition, the observed increase in OTU richness is stronger than what was previously measured on kelps due to seasons (Weigel and Pfister, 2019) tissue aging (Bengtsson et al., 2012) or temperature elevation (Minich et al., 2018), suggesting a combined effect of degradation and seasonal variations. Furthermore, kelp defence reactions in early time points (e.g. emission of reactive oxygen species and halogenated compounds) that occur in stress situations such as blade detachment, cutting, displacement and degradation (Cosse et al., 2007) may select for a resistant fraction of the resident bacterial community. The control exerted by living kelps on biofilm formation through surface metabolites (Bengtsson et al., 2012; Salaün et al., 2012) might also be relaxed during algal decay, possibly explaining some of the observed changes.

During the first phase of the succession (0–6 weeks), the diversity of the kelp-associated community increased, due to a decrease in the abundance of initially dominant OTUs affiliated to Litorimonas, Hellea and Granulosicoccus. These genera were reported as primary surface colonizers (Dang and Lovell, 2016; Ramirez-Puebla et al., 2020) but might be outcompeted by specialist algae-degraders. In parallel, a number of OTUs that were initially absent or rare at the kelp surface showed a transient increase in abundance during the first 6 weeks of degradation, including Alphaproteobacteria (e.g. Fretibacter, Hellea, unclassified Rhizobiaceae, Octadecabacter, Sulfitobacter, Tateyamaria, Altererythrobacter) and Gammaproteobacteria (e.g. Paraglacieola, unclassified Ectothiorhodospiraceae). The same changes were not detected in the surrounding seawater, suggesting that they result more from an enrichment of initially rare taxa at the kelp surface than from a colonization by free-living bacteria. The rapid and transient enrichment of these bacteria on detached blades might be due to their capacity to assimilate low-molecular weight algal compounds. For instance, a study on the Fucus spiralis microbiota showed that Sulfitobacter and Octadecabacter isolates were able to grow on mannitol (Dogs et al., 2017), a brown algal polyol whose exudation is enhanced upon stress such as blade damage. Members of the genus Tateyamaria, previously found on kelp (Bengtsson et al., 2012), can use the brown algal storage compound laminarin (Sass et al., 2010). The first phase of the succession was also...
characterized by a global increase in the relative sequence abundance of numerous Bacteroidia OTUs which individually represented a low proportion of the microbiota, and stayed stable afterwards. Bacteroidia are commonly known as particle-associated taxa (Thiele et al., 2015) and positively respond to an input of marine polysaccharides (Wietz et al., 2015; Balmonte et al., 2018, 2019; Enke et al., 2019). In particular, members of the family Flavobacteriaceae are specialized in the degradation of algal biomass (Thomas et al., 2011; Teeling et al., 2012). Flavobacteriaceae are known to colonize algal surfaces and form biofilms (Mann et al., 2013) and to possess numerous polysaccharide utilization loci (PULs) (Grondin et al., 2017) that encode a great number and diversity of carbohydrate-active enzymes for the breakdown of algal polysaccharides, together with regulators and transporters. Genomes of marine flavobacteria frequently feature PULs predicted to be involved in the degradation of brown algal polysaccharides such as alginate, laminarin and fucoidan (Barbeyron et al., 2016; Kappelmann et al., 2019). Notably, the PUL dedicated to alginate, the most abundant polysaccharide in kelps, was biochemically characterized and shown to account for the full substrate degradation into simple sugars (Thomas et al., 2012; Dudek et al., 2020). Therefore, the Bacteroidia detected here on detached kelp likely contributed to the decomposition of the algal biomass.

During the second phase of the succession (11–24 weeks), kelp-associated bacterial community was the most diverse, suggesting that new niches were available for bacterial colonization. The most striking observation is the enrichment in Acidimicrobiia, mainly due to OTUs related to the Sva0996 marine group. This group may be ubiquitous as it was previously detected on the brown macroalgae Taonia atomaria (Paix et al., 2019) and L. digitata (Gobet, personal communication), as well as sponges and ascidians (Steinert et al., 2015; Verhoeven et al., 2017; Dat et al., 2018), in deep marine sediments (Chen et al., 2016) and in the water column of oligotrophic environments (Reintjes et al., 2019). Little is known about the Sva0996 marine group, except its ability to use phytoplankton-derived dissolved proteins and potential importance in the dissolved organic nitrogen cycle (Orsi et al., 2001; de Beer et al., 2005; Ettinger et al., 2017). Comparisons of underlying and external sediments indicate that algal degradation had mostly a local effect directly underneath the accumulated kelps. The low proportion of OTUs shared between kelp-associated and underlying sediment communities suggests that this change in community composition is not due to a direct transfer of OTUs from algae, but to a change of environmental conditions linked to kelp accumulation and degradation. Indeed, in the underlying sediment, there was a shift in abundance between the two most abundant bacterial classes, with the Gammaproteobacteria decreasing in sequence relative abundance until week 6 and the Bacteroidia increasing and remaining the most abundant group until 24 weeks. As stated in the section above, bacteria from the class Bacteroidia such as the Flavobacteriaceae are involved in degradation of complex molecules such as algal polysaccharides. This taxonomic group is likely enriched in the underlying sediment through the experiment as fragments of degrading algal tissue are continuously deposited onto the seafloor. Both the limited oxygen diffusion due to kelp accumulation and the activity of aerobic degrading bacteria are expected to rapidly deplete O2 in underlying sediments. Accordingly, differential sequence abundance analyses and functional predictions indicated an increase of taxa involved in sulfur and nitrogen cycles, as well as fermentation. Although some of these changes might partially be due to seasonal variations (Gobet et al., 2012; Marchant et al., 2014), our data showed a significant effect of algal accumulation since early changes (2–4 weeks) were not seen in external sediments. Further, during tissue degradation, there was an increase in Deltaproteobacteria, a group regularly found in...
temperate coastal sediment and known for sulfate and nitrate reduction (Mußmann et al., 2005). Sulfate reduction is one of the most common anaerobic degradation pathways for organic matter in marine sediments (Wasmund et al., 2017). Subsequently, several OTUs related to Desulfovibrio and Clostridiales were enriched during the degradation. Members of these orders are well-known sulfate reducers and are frequently detected in temperate marine sediments rich in organic matter (Muyzer and Stams, 2008; Gobet et al., 2012; Mahmoudi et al., 2015; Ettinger et al., 2017). Desulfovibrio feature metabolically versatile and larger genomes than other sulfate-reducing bacteria (SRB), which might allow a faster response to organic matter inputs (Strittmatter et al., 2009). OTUs related to Arcobacter and Sulfurovum were also enriched in underlying sediments during the degradation. These genera can oxidize reduced sulfur compounds produced by SRB (e.g. H₂S, thiosulfate) back to sulfate, therefore completing the sulfur cycle (Wasmund et al., 2017). Furthermore, FAPROTAX analysis showed a functional enrichment of nitrate reduction pathways. Nitrate reducers use either denitrification to N₂ or dissimilatory nitrate reduction to ammonium (DNRA). Here, the massive input of organic matter and probable increase in sediment sulfide concentrations due to SRB enrichment would be expected to favour DNRA (Burgin and Hamilton, 2007; Hardison et al., 2015). Therefore, we can speculate that kelp degradation allows retaining nitrogen inputs as bio-available ammonium instead of loss as N₂. Similar to these results, a recent study (Aires et al., 2019) also showed that additions of red or green algal biomass (Gracilaria vermiculophylla or Ulva rigida, respectively) strongly favoured Desulfobacteriales, Bacteroidales, Clostridiales and Campylobacteriales. Therefore, at the order level changes in sediment bacterial communities due to seaweed accumulation do not appear to be specific to the type of algae.

Conclusion

In conclusion, following kelp detachment and accumulation on the seafloor, the structure and composition of the epiphytic bacterial community shifted, showing a biphasic succession of early degraders and secondary colonizers that might exploit different substrate niches. Besides this direct effect on kelp-associated bacteria, we also evidenced changes in the underlying sediments, showing that kelp accumulation likely has a global effect on the microbial communities of receiving ecosystems. This work paves the way for future studies investigating the functional determinants, mechanisms and bacterial interactions of the epiphytic microbiota during the kelp breakdown process.

Experimental procedures

The in situ experiment was conducted from April 2017 to October 2017 in the bay of Morlaix (48°42’33.78” N, 003°57’12.36” W) on the north-western coast of Brittany (France) and detailed elsewhere (de Bettignies et al., 2020). Briefly, cages were randomly filled with 1 kg of new blades from young healthy adult L. hyperborea sporophytes (140–200 cm). A total of 35 cages were set at 9 m depth on a sandy floor to simulate an accumulation of kelp fragments. Three cages were randomly selected at each of eight sampling times (0, 2, 4, 6, 11, 15, 20, 24 weeks) to sample algal surfaces, underlying and external sediments and overlying water as follows. Ten algal pieces of 1.3 cm diameter were subsampled from each cage using a sterilized stainless steel punch. The surface sediments (0–4 cm depth) directly below each cage (‘underlying sediment’) or 2 m away from the cages (‘external sediment’) were sampled in sterile plastic containers. Initial sediment was collected immediately prior to the cages set-up. Three litres of the overlying water (1 m above the cages) were taken using Niskin bottles. Triplicates of 1 l were prefiltered at 3 μm to remove large particles and then filtered through a 0.2 μm Sterivex filter (Merck, Darmstadt, Germany). All samples were stored at −80°C before DNA extraction. Measurements of contextual parameters related to physico-chemical conditions (temperature, light, percent organic matter in sediment) and state of the kelp tissue (remaining biomass, oxygen consumption, photosynthetic capacity, carbon/nitrogen ratio, phlorotannin content) were detailed elsewhere (de Bettignies et al., 2020) and summarized in the Supporting Information text. The list of samples and contextual parameters is available in the Supporting Information Table S1.

DNA extraction. We adapted a protocol for DNA extraction from the algal surface. Algal pieces were immersed in a bacterial lysis buffer, without previous grinding of the tissues to prevent extraction of endophytic bacteria and algal plastids while efficiently lysing epiphytic bacteria. After incubation, algal pieces were discarded and DNA was extracted only from the soluble lysate using a phenol:chloroform:isoamyl alcohol extraction combined with filtration steps of the Nucleospin PlantII kit (Macherey Nagel, Hoerdt, France). For water samples, DNA was extracted from the 0.2 μm filters following a similar protocol as used for the algal samples. For sediment samples, DNA was extracted from 0.5 g of sediment using the DNeasy PowerLyzer PowerSoil Kit (Qiagen, Courtaboeuf, France). For further details, please see the Supporting Information text.

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16S rRNA gene copies quantification. The total number of bacterial 16S rRNA gene copies in each sample was assessed by quantitative real-time PCR as described previously (Bacchetti De Gregoris et al., 2011), using the universal primers 926F/1062R. This primer pair has a predicted coverage of 55% for chloroplast-derived 16S rRNA genes (Silva Testprime on SSU r138.1). Therefore, we cannot exclude that a small proportion of quantified 16S rRNA gene copies originates from chloroplasts. Amplification and detection were performed in a LightCycler 480 Instrument II. For each sample, triplicate reactions were prepared as described in the Supporting Information text. Serial dilutions of purified bacterial genomic DNA ranging from $10^3$ to $10^8$ 16S rRNA gene copies were prepared as described in the Supporting Information text. The total number of bacterial 16S rRNA gene copies quantified in each sample was assessed by quantitative real-time PCR as described previously (Bacchetti De Gregoris et al., 2011), using the universal primers 926F/1062R. This primer pair has a predicted coverage of 55% for chloroplast-derived 16S rRNA genes (Silva Testprime on SSU r138.1). Therefore, we cannot exclude that a small proportion of quantified 16S rRNA gene copies originates from chloroplasts. Amplification and detection were performed in a LightCycler 480 Instrument II. For each sample, triplicate reactions were prepared as described in the Supporting Information text. Serial dilutions of purified bacterial genomic DNA ranging from $10^3$ to $10^8$ 16S rRNA gene copies equivalent were used as a standard curve and were amplified in triplicate in the same run as the environmental samples. A non-template control was included in the run. Results were analysed using the LightCycler 480 Software v1.5. One-way ANOVA analyses followed by pairwise post-hoc Tukey HSD were conducted with time as fixed factor, individually for each compartment. Details on qPCR assays are given in the Supporting Information Table S8, following the MIQE guidelines (Bustin et al., 2009).

Library preparation, sequencing and sequence processing. As a positive control, we constructed a mock community based on the genomic DNA of 32 pure marine bacterial isolates as reported previously (Thomas et al., 2020). Libraries were prepared for the 94 samples (24 algal samples, 24 water samples, 3 initial sediment samples, 21 underlying sediment samples, 20 external sediment samples, one mock community and one negative control (PCR molecular grade water)). Libraries were prepared using the primers S-D-Bact-0341-b-S-17 (5’ CCTACGGGNGGCWGCAG 3’) and 799F_rc (5’ CMGGGTATCTAATCCGCTTT 3’) targeting the V3-V4 region of the 16S rRNA gene and sequenced on a MiSeq paired-end sequencing run (300 cycles × 2, Illumina, San Diego, CA, USA) as described previously (Thomas et al., 2020). 16S rRNA gene amplicon sequences are available at NCBI under BioProject ID PRJNA667331. Raw Illumina sequence reads were trimmed to remove low-quality bases and Illumina adapters using Trimmomatic v0.38 (Bolger, Lohse and Usadel, 2014, for details see Supporting Information text). Overlapping paired-end sequence reads were assembled using PANDAseq v2.11 and assembled sequences between 400 and 500 bp long were kept (Masella et al., 2012). All the following steps for sequence processing were performed using the sequence processing pipeline FROGS (Find Rapidly OTU with Galaxy Solution) developed for the Galaxy platform (http://galaxy3.sb-roscot.fr, Escudie et al., 2018). Sequences were clustered into OTUs using SWARM v2 (Mahé et al., 2014). PCR chimera and singletons were filtered out and OTUs were taxonomically assigned using the RDP classifier (Wang et al., 2007) on the Silva 16S rRNA (v132) database.

Multivariate and statistical analyses. Prior to further analysis, one kelp-associated sample from week 0 (AA-T0-C2 in the Supporting Information Table S1) was removed from the dataset due to its poor sequencing depth (3 302 sequences) compared to other samples (range 24 482–123 368 sequences). Samples from the bacterial community dataset were rarefied (100 permutations) to 24 482 sequences prior to OTU richness and Shannon diversity index calculation. Dissimilarities in community structure were calculated using the Bray–Curtis dissimilarity index (Bray and Curtis, 1957) before principal coordinate analysis (PCoA). Permutational analysis of variance (PERMANOVA, 999 permutations) was applied using adonis to discriminate groups of samples according to time or type of compartment (Anderson, 2001), followed by multivariate pairwise comparisons using pairwise.perm.manova. Mean dispersions within the compartments were calculated using betadisper (999 permutations). The effect of environmental factors on kelp-associated and underlying sediment communities were assessed using distance-based redundancy analysis (db-RDA) (Legendre and Anderson, 1999) with capscale function followed by ANOVA. Environmental factors were log10-transformed prior to this analysis. Only significant environmental factors were selected for db-RDA visualization (ANOVA, $P < 0.05$). The above multivariate and statistical analyses were performed in R v3.5.0 with the packages phyloseq, vegan, ggplot and EcolUtils (Wickham, 2009; McMurdie and Holmes, 2013; Oksanen et al., 2013; R Core Team, 2018). Differential sequence abundance analysis through time was performed at the OTU level for kelp-associated and underlying sediment communities using the GLM test with the ALDEx2 R package (Fernandes et al., 2014). The variation of selected differential OTUs through time was further tested using ANOVA followed by Tukey HSD post hoc test. An association network was built for kelp-associated bacterial communities. Extended Local Similarity Analysis (eLSA v1.0.4) (Xia et al., 2011) was performed on a dataset comprising 10 contextual parameters (eight environmental factors and two bacterial diversity measures) and the relative sequence abundance of OTUs found in at least 19 samples (135 OTUs). This prevalence filter was set according to recommendations in Rötjers and Faust (2018). A delay of one sampling date was allowed for temporal associations. P-values were estimated using the permutation method, with 1 000 permutations. Q-values were estimated to control for false positives. Default eLSA parameters were used otherwise. The
network was visualized in Cytoscape v3.3.0 (Shannon et al., 2003). Only associations with \( |L| \geq 0.5, Q \leq 0.05 \) and \( P \leq 0.001 \) were represented with the ‘edge-weighted, spring-embedded layout’ (nodes are strongly repelled or attracted as a function of their LS value; spring strength = 50, spring rest length = 100, strength of disconnected spring = 0.05, rest length of disconnected spring = 500, and strength to avoid collisions = 500). Clusters were detected using the MCODE algorithm (Bader and Hogue, 2003) with ‘haircut’ and ‘fluff’ options and otherwise default parameters.

Functional profiles for kelp-associated and underlying sediment communities were inferred using the FAPROTAX v1.2.2 database and script (Louca et al., 2016) against the rarefied OTU dataset with the assigned taxonomy. In addition, PICRUS2 v2.1.4 (Douglas et al., 2020) was run with the default parameters (except minreads set to 5) to get metagenome predictions for EC numbers. Differences in sequence abundance of functional groups and enzymes over time were assessed using the GLM test with ALDEx2.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

Appendix S1: Supporting Information