

Microorganisms living on macroalgae: diversity, interactions, and biotechnological applications.

Marjolaine Martin, Daniel Portetelle, Gurvan Michel, Micheline Vandenbol

▶ To cite this version:

Marjolaine Martin, Daniel Portetelle, Gurvan Michel, Micheline Vandenbol. Microorganisms living on macroalgae: diversity, interactions, and biotechnological applications. Applied Microbiology and Biotechnology, 2014, 98 (7), pp.2917-35. 10.1007/s00253-014-5557-2. hal-00999506

HAL Id: hal-00999506 https://hal.sorbonne-universite.fr/hal-00999506

Submitted on 3 Jun 2014

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Microorganisms living on macroalgae:

Diversity, interactions, and biotechnological applications

Marjolaine Martin^{1*}, Daniel Portetelle¹, Gurvan Michel ^{2,3}, Micheline Vandenbol¹

- Gembloux Agro-Bio Tech, Microbiology and Genomics Unit, Passage des déportés 2, 5030 Gembloux, Belgique
- Sorbonne Universités, UPMC Univ Paris 06, UMR 8227, Integrative Biology of Marine Models, Station Biologique de Roscoff, CS 90074, F-29688, Roscoff cedex, Bretagne, France
- CNRS, UMR 8227, Integrative Biology of Marine Models, Station Biologique de Roscoff, CS 90074, F-29688, Roscoff cedex, Bretagne, France

Abstract: Marine microorganisms play key roles in every marine ecological process, hence the growing interest in studying their populations and functions. Microbial communities on algae remain underexplored, however, despite their huge biodiversity and the fact that they differ markedly from those living freely in seawater. The study of this microbiota and of its relationships with algal hosts should provide crucial information for ecological investigations on algae and aquatic ecosystems. Furthermore, because these microorganisms interact with algae in multiple, complex ways, they constitute an interesting source of novel bioactive compounds with biotechnological potential, such as dehalogenases, antimicrobials and alga-specific polysaccharidases (e.g. agarases, carrageenases, alginate lyases). Here, to demonstrate the huge potential of alga-associated organisms and their metabolites in developing future biotechnological applications, we first describe the immense diversity and density of these microbial biofilms. We further describe their complex interactions with algae, leading to the production of specific bioactive compounds and hydrolytic enzymes of biotechnological interest. We end with a glance at their potential use in medical and industrial applications.

Keywords: microbial biofilms; algal symbiosis; glycoside hydrolases; algal polysaccharidases; antimicrobial compounds.

*Corresponding author: Marjolaine Martin; E-Mail: marjolaine.martin@ulg.ac.be

Tel: +3281622355 Fax: +3281611555

Introduction: surface-associated marine microorganisms

Marine microorganisms live freely in seawater (planktonic microorganisms) or attached to biotic or abiotic surfaces. Microorganisms on a surface commonly form a biofilm, defined as "an assemblage of microbial cells that is irreversibly associated with a surface and enclosed in a matrix of primarily polysaccharide material" (Donlan 2002). This matrix, called the exopolysaccharide layer, allows close spatial proximity, enhancing communication and interactions among bacteria and between bacteria and their host (Kilian et al. 1985; Pasmore and Costerton 2003; Wilson et al. 2011). In the marine environment, where competition for space and nutrients is intense, the surfaces of marine eukaryotes such as invertebrates and algae offer a nutrient-rich habitat uniquely suited for microbial colonization and biofilm formation (Egan et al. 2008; Goecke et al. 2010). As such surfaces are highly complex and differentiated, marine microbial biofilms should constitute a huge source of diversity, and the bacterial communities forming them should differ considerably in composition from populations of pelagic bacteria. Oddly, although bacteria in marine environments are most often surfaceassociated, previous investigations have preponderantly focused on the diversity of planktonic microorganisms rather than on microbial epibionts. Yet although investigators are increasingly using both culture-dependent and -independent methods to zoom in on microbial symbionts living on the surfaces of organisms such as corals, sponges, tunicates, and macroalgae (Erwin et al. 2011; Rohwer et al. 2002; Taylor et al. 2003; Wegley et al. 2007; Wilson et al. 2010), the biotechnological potential of these symbionts remains little discussed. In this review we focus on microbial biofilms on marine macroalgae, including their potential importance in developing future biotechnological applications. We discuss the diversity and density of these biofilms and the factors influencing the microbial communities that live on diverse algal species. We further outline interactions between algae and their epibionts leading to the production of metabolites of biotechnological interest. Particularly, we review the state of the art on algal-specific polysaccharidases from seaweed-associated bacteria. Finally we draw attention to the potential importance of these microorganisms and their metabolites, such as secondary bioactive compounds and specific hydrolytic enzymes, for biotechnological applications in diverse industrial fields.

Diversity of microorganisms on algae

Microorganisms are very abundant on the surfaces of marine organisms (> 1.1x10⁸ microorganisms/cm²) (Cundell et al. 1977). Although the microorganisms observed and identified on the surfaces of diverse algae include yeasts, fungi, and protists (Armstrong et al. 2000; Cundell et al. 1977; Genilloud et al. 1994; Schaumann and Weide 1995; Uchida and Murata 2004), most available reports on alga-associated microbial populations concern bacteria. Therefore this review focuses mainly on alga-associated bacterial communities.

The density of bacteria on algal surfaces has been estimated by cell counts under the microscope (Cundell et al. 1977), by culture-based methods (Mazure and Field 1980), and by molecular approaches (Armstrong et al. 2000). Mean densities between 10⁶ and 10⁹ bacteria/cm² algal surface have been recorded. There is some controversy regarding the composition of bacterial communities on algae. Bacterioplankton studies have shown most marine bacteria to be gram negative, but recent studies on marine-sediment-associated bacteria have revealed a large proportion of gram-positive bacteria, too (Gontang et al. 2007). Table 1 shows the most abundantly represented phyla (and classes or orders) of bacteria identified on diverse algal species, with the sampling location and season. Gram-negative bacteria of the phyla *Bacteroidetes* and *Proteobacteria* emerge as the most abundant, having been found on practically all the listed algal species (Table 1). Although gram-negative bacteria appear to preponderate, gram-positive species are also present. In particular, gram-positive bacteria of the phyla *Actinobacteria* and *Firmicutes* have been observed on most algae (Table 1). On some species or in a particular season, other bacterial phyla can also be abundant. For instance, Bengtsson et al. found peptidoglycan-less *Planctomycetes* species to dominate the bacterial biofilm on the kelp *Laminaria hyperborea* for long periods of the year (Bengtsson et al. 2010; Bengtsson et al. 2013; Bengtsson and Øvreås 2010). Ocean surface water shows a phylum distribution quite similar to that of algal surface bacteria, the most abundantly represented phylum being the *Proteobacteria*

(particularly the class Alpha-proteobacteria) (Morris et al. 2002), followed by the Bacteroidetes, Actinobacteria, Planctomycetes, and Chloroflexi (Longford et al. 2007). At the bacterial genus and species levels, however, recent investigations have revealed notable differences in composition between epibacterial communities and the surrounding bacterioplankton communities (Burke et al. 2011b; Lachnit et al. 2011; Longford et al. 2007). This suggests that colonization patterns are host-specific and strongly influenced by the seaweed, because of physicochemical constraints, such as cell wall component diversity (Michel et al. 2010a; Popper et al. 2011), and/or active defense mechanisms (Cosse et al. 2007; Potin et al. 2002). Table 1 further shows that the sampling season and region and the algal species or phylum can influence community composition. In fact, diverse factors shape the composition of alga-associated bacterial populations: (i) Recent studies on bacterial biofilm composition have shown it to vary considerably with the algal phylum (green, brown, or red algae) and, to a lesser extent, with the algal species (Lachnit et al. 2009; Longford et al. 2007). Longford et al. compared the bacterial beta-(between host) and alpha-(within host) diversity of the marine sponge Cymbastela concentrica and two co-habiting algae species, the red alga Delisea pulchra and the green alga Ulva australis (Longford et al. 2007). Between the two algal species, the community patterns were very similar at bacterial phylum level, but at bacterial species level little overlap was observed. Lachnit et al. focused on compositional variability among the bacterial communities associated with diverse species of the three algal phyla Rhodohpyta, Chlorophyta and Phaeophyta (Lachnit et al. 2009). They found that host phylum seems to contribute more than host species to dissimilarity in epibacterial composition, explaining this dissimilarity on the basis of different physico-chemical properties and metabolite compositions and more or less effective defense mechanisms (Potin et al. 2002) and/or attractants (Pasmore and Costerton 2003). For instance, brown algae produce and secrete large amounts of mannitol (Gravot et al. 2010), a main carbon storage compound (Michel et al. 2010b). This organic exudate was recently shown to affect the formation of biofilms of marine

(ii) The part of the thallus sampled and its age also influence both the composition and the specificity of the bacterial population. On the brown alga *Laminaria saccharina*, for example, Staufenberger et al. (2008) found a greater bacterial diversity on the old phyloid than on any other part of the alga, explaining it on the basis of tissue age/mechanical stress: this tissue should contain more damaged cells vulnerable to bacterial decomposition, enhancing bacterial colonization. Furthermore, the association appeared most specific (i. e., between-specimen variability was lowest) on the meristem (where new tissue is formed) and cauloid. On all parts of the alga, however, the bacterial communities differed markedly from those of the surrounding seawater. The authors also point out that the composition of the bacterial community present on the substratum-anchored rhizoid is likely to reflect the presence, in the substratum, of other marine organisms with their own surface communities.

bacteria such as *Pseudolatermonas* spp. 3J6 and D41 and *Zobellia galactanivorans* (Salaün et al. 2012). This latter microorganism, which was isolated from the red alga *Delesseria sanguinea* (Barbeyron et al. 2001), is a model bacterium for the study of bacteria-seaweed interactions and particularly the bioconversion of algal polysaccharides (Michel and

- (iii) **Seasonal changes** in the composition of alga-associated bacterial populations have also been recorded. Mazure and Field (1980) observed on the brown alga *Laminaria saccharina* a predominance of mesophilic bacteria in summer, with a switch to a more psychrophilic population in winter. A similar seasonal shift was observed on *Laminaria digitata* (Corre and Prieur 1990; Salaün 2009). Furthermore, bacterial abundance can be two to three times greater in summer, likely because the higher temperature favors enhanced microbial metabolism (Rao 2010). Moreover, Stratil et al. (Stratil et al. 2013) studied the shift in diversity and density of bacterial populations on *F. vesiculosus* when cultured at different temperatures. They found 20% of the bacterial diversity variation between host groups to be due to temperature, but bacterial density was not affected by this factor.
- (iv) Rapid changes in bacterial community composition and abundance have also been observed **between healthy and bleached (diseased) algal tissues.** On diseased macroalgae, the density of bacteria and other microfoulers can be as much

Czjzek 2013).

as 400 times that found on healthy tissues (Weinberger et al. 1994). Furthermore, comparative metagenomics applied to healthy and bleached tissues of *D. pulchra* has evidenced differences in bacterial taxa and functional genes (Fernandes et al. 2012). These shifts have been explained by reduced defenses in stressed thalli (due to high summer temperature), leading to colonization by opportunistic and pathogenic bacteria.

(v) Finally, Burke et al. (2011a), studying *Ulva australis*, observed **intraspecies differences** in bacterial community composition. They noted similar functional profiles for the communities found on different specimens, suggesting that (functional) genes, rather than bacterial species, may explain the diversity of bacterial epibionts on algae. These intraspecies differences were also observed on *F. vesiculosus* (Stratil et al. 2013). As only 20% of the community variation could be explained by temperature changes, a large proportion of variation between hosts is left unexplained. This strengthens the 'functional profile' theory of Burke et al.

Alga-associated microorganisms produce specific enzymes and bioactive compounds

Microorganisms on algae, through their complex and numerous interactions with the host, constitute an immense source of bioactive compounds and specific polysaccharidases. Therefore, before discussing the biotechnological potential of algal epibionts and their metabolites, we will have a glance at microorganism-alga interactions and at the biotechnologically useful bioactive compounds and enzymes produced by alga-associated microorganisms.

Seaweed-associated bacteria produce alga-specific polysaccharidases

It is generally assumed that microorganisms benefit from the ready availability of a range of organic carbon sources produced by the host alga. Green, red, and brown algae produce a wide diversity of complex polysaccharides which are essential components of their cell walls (Popper et al. 2011). These polysaccharides constitute a crucial biomass in coastal ecosystems. Interestingly, in contrast to the polysaccharides of terrestrial plants, most algal polysaccharides are non-lignocellulosic and sulfated (Popper et al. 2011). Whereas lignocellulosic biomass consists of cellulose, lignin, and hemicelluloses, macroalgal biomass is much more complex. About ten different polysaccharides (e.g. agars, carrageenans, ulvans) and as many monosaccharides (e.g. glucose, mannose, xylose), are found over the three algal phyla (Jung et al. 2013). Accordingly, alongside common polysaccharidases (e.g. cellulases, beta-glucosidases and amylases), very specific carbohydrate-active enzymes are found in microorganisms living on algae. Here we present the current state of knowledge on these enzymes (see http://www.cazy.org/, Cantarel et al., 2009), focusing solely on those characterized at both the molecular and biochemical levels, and particularly on those whose 3D structure has been determined (Table 2).

Carrageenases

Carrageenans and agars are sulfated galactans. They are the main cell wall components of red macroalgae (Popper et al, 2011). Carrageeenases are currently divided into three classes according to the number of sulfate substituents per disaccharide repeating unit which are specifically recognized: kappa- (1 sulfate, EC 3.2.1.83), iota- (2 sulfates, EC 3.2.1.157) and lambda-carrageenases (3 sulfates, EC 3.2.1.-). All these enzymes cleave β -1,4 glycosidic bonds in carrageenans.

Kappa-carrageenase genes have been cloned from several *Pseudoalteromonas* species (Barbeyron et al. 1994; Kobayashi et al. 2012; Liu et al. 2011), from *Zobellia* species (Barbeyron et al. 1998; Liu et al. 2013), and from *Cellulophaga lytica* strain N5-2 (Yao et al. 2013). The corresponding enzymes belong to glycoside hydrolase family 16 (GH16) (Barbeyron et al. 1994). The kappa-carrageenase of *P. carrageenovora* adopts a β jelly-roll fold and displays a tunnel active site (Figure 1A). These features suggest that this enzyme has an endo-processive mode of action (Michel et al. 2001a), and this prediction has been biochemically confirmed (Lemoine et al. 2009).

The first cloned iota-carrageenase genes originated from the marine bacterium *Alteromonas fortis* and from *Z. galactanivorans*, and their products defined the GH82 family (Barbeyron et al. 2000). Additional iota-carrageenase genes have been cloned from *Cellulophaga* sp. QY3, a flavobacterium isolated from the red alga *Grateloupia livida* (Ma et al. 2013),and from *Microbulbifer thermotolerans* JAMB-A94T, a deep-sea bacterium (Hatada et al. 2011). The iota-carrageenase CgiA of *A. fortis* adopts a right-handed β -helix fold with two additional domains (A and B) in the C-terminal region (Michel et al. 2001b). Upon substrate binding, the (α/β)-fold domain A shifts towards the β -helix cleft, forming a tunnel that encloses the iota-carrageenan chain (Figure 1B), thus explaining the highly processive character of CgiA (Michel et al. 2003). A mechanistic study has demonstrated that CgiA is chloride ion dependent and that its catalytic residues are Glu245 and Asp247 (Rebuffet et al. 2010).

Lambda-carrageenases constitute a new GH family, unrelated to kappa- and iota-carrageenases (Guibet et al. 2007). Only two genes have been cloned so far, one from the seaweed-associated bacterium *P. carrageenovora* (Guibet et al. 2007) and one from the deep-sea bacterium *Pseudoalteromonas* sp. strain CL19 (Ohta and Hatada 2006). The products of these genes are highly similar (98% sequence identity), explaining why no CAZY family number has yet been attributed (Cantarel et al. 2009). These large enzymes (~105 kDa) feature a low-complexity linker connecting two independent modules, an N-terminal domain predicted to fold as a β-propeller and a C-terminal domain of unknown function (Guibet et al. 2007).

Agarases

Agarases are divided into two classes, alpha-agarases (EC 3.2.1.158) and beta-agarases (EC 3.2.1.81), which respectively hydrolyze α -1,3 and β -1,4 linkages between neutral agarose motifs in agar chains. The first alpha-agarase activity was purified and characterized from *Alteromonas agarlyticus* twenty years ago (Potin et al. 1993). The gene was later cloned, revealing a large enzyme (154 kDa) with a complex modular architecture including five calcium-binding thrombospondin type 3 repeats, three family-6 carbohydrate-binding modules (CBM6s), and a C-terminal catalytic module defining a novel GH family (GH96) (Flament et al. 2007). Bioinformatic studies suggest that the CBM6s specifically bind agars and were acquired from modular GH16 beta-agarases (Michel et al. 2009). A highly similar alpha-agarase (72% sequence identity) has also been cloned from *Thalassomonas* sp. JAMB-A33, a strain isolated from marine sediment (Hatada et al. 2006).

Beta-agarases are found in four unrelated CAZY families: GH16, GH50, GH86, and GH118 (Cantarel et al. 2009). The first beta-agarases to be both structurally and biochemically characterized were the GH16 beta-agarases ZgAgaA and ZgAgaB of Z. galactanivorans (Allouch et al. 2003; Jam et al. 2005). ZgAgaA is an extracellular monomeric enzyme with a GH16 module appended to a putative CBM and a PorSS secretion domain, while ZgAgaB is a dimeric lipoprotein anchored to the outer membrane (Jam et al. 2005). In both enzymes, the GH16 module displays a β jelly-roll fold with an open catalytic groove (Allouch et al. 2003). Two agar-binding sites have been identified in the structure of $ZgAgaA_{GH16}$ complexed with oligo-agars: one in the active site cleft and one at the external surface of the protein, explaining the high agar-fiber-degrading efficiency of this enzyme (Allouch et al. 2004). The crystal structure of a third beta-agarase from Z galactanivorans has been solved recently. ZgAgaD has a longer catalytic groove with 8 subsites (Figure 1C) and is specific for unsubstituted agarose motifs (Hehemann et al. 2012a). Numerous GH16 beta-agarases have been cloned from bacteria isolated from seawater or marine sediments, but relatively few from seaweed-associated bacteria (Kim and Hong 2012; Oh et al. 2010; Schroeder 2003; Yang et al. 2011).

The first GH50 beta-agarase was cloned from *Vibrio* sp. JTO107, isolated from seawater in Japan (Sugano et al. 1993). So far, however, no GH50 gene has been cloned from an alga-associated microorganism. The first structure of a GH50 beta-agarase was determined last year: Aga50D from *Saccharophagus degradans* (Pluvinage et al. 2013). This bacterium was isolated from a halotolerant land plant in a salt marsh, and is thus not a genuine marine microorganism (Andrykovitch and Marx 1988). Aga50D features two domains, a $(\beta/\alpha)_8$ -barrel connected to a small β -sandwich domain reminiscent of a CBM

(Figure 1D). The putative catalytic residues (Glu534 and Glu695) are located in an active site with a tunnel topology, in keeping with the exo-lytic mode of action of this beta-agarase (Pluvinage et al. 2013).

One of the first characterized beta-agarases (AgrA) was purified from *Pseudoalteromonas atlantica* Tc6, a gammaproteobacterium isolated in Canada from the red alga *Rhodymedia palmata* (Yaphe 1957). Its gene remained an orphan sequence for a long time (Belas 1989), before defining the GH86 family (Cantarel et al. 2009). No other GH86 beta-agarase has been characterized from alga-associated bacteria.

The GH118 family includes only 8 sequences from marine bacteria, and none of them was isolated from a seaweed-associated bacterium. The first GH118 beta-agarase was cloned from *Vibrio* sp. PO-303 (Dong et al. 2006). The beta-agarase of *Pseudoalteromonas* sp. CY24 has also been extensively characterized, revealing a large binding site with 12 subsites. This GH118 enzyme proceeds according to a mechanism of inversion of the anomeric configuration (Ma et al. 2007), in contrast to GH16 beta-agarases, which act via a retaining mechanism (Jam et al. 2005). The families GH50 and GH86 are also predicted to encompass retaining enzymes (Pluvinage et al. 2013). Currently there is no GH86 or GH118 beta-agarase of known 3D structure, although a note mentions the crystallization of a beta-agarase from *Pseudoalteromonas* sp. CY24 (Ren et al. 2010).

Porphyranases

Porphyran is the usual name of the agar extracted from red algae of the genus Porphyra. The porphyran backbone is composed of ~30% agarose repetition moieties (LA-G), the remaining moieties being essentially L-galactopyranose-6sulfate (L6S) linked via an α-1,3 bond to a beta-D-galactopyranose (G) residue. A porphyran repetition moiety (L6S-G) is linked via a β-1,4 linkage to either another porphyran moiety or to an agarose moiety (Correc et al. 2011). Such a hybrid structure is usual for agars, and the number of porphyran motifs varies according to the red algal species (Popper et al. 2011). Recently, a new class of enzymes has been discovered in the genome of Z. galactanivorans: β-porphyranases, which specifically hydrolyze the β-1,4 linkage between porphyran motifs in agars. These enzymes define a new subfamily within the GH16 family. The crystal structures of ZgPorA (Figure 1E) and ZgPorB reveal a porphyran binding mode involving conserved basic amino acids (Hehemann et al, 2010). The fine differences in substrate specificity between the β-agarases and β-porphyranases of Z. galactanivorans have been further studied, and a comprehensive model for this complex agarolytic system has been proposed (Hehemann et al. 2012a). Fascinatingly, β-porphyranase genes from algal epibionts have been found in human gut bacteria isolated from Japanese individuals, suggesting that edible seaweeds with their associated marine bacteria were the route through which the gut bacteria acquired these novel polysaccharidases (Hehemann et al. 2010). This hypothesis is strengthened by the experimental demonstration that the Japanese gut bacterium Bacteroides plebeius can grow on porphyran (Hehemann et al. 2012b). Moreover, the putative glycoside hydrolases BpGH16B and BpGH86A have been characterized as active β-porphyranases. The structure of BpGH86A in a complex with an oligo-porphyran has also been solved (Figure 1F), revealing a TIM barrel domain with an extended substratebinding cleft and two accessory β-sandwich domains (Hehemann et al. 2012b). Thus, GH86 enzymes constitute a polyspecific family including both β -agarases and β -porphyranases.

α -1,3-(3,6-Anhydro)-L-galactosidases

Z. galactanivorans has also been pivotal in the discovery of a third class of enzymes involved in the catabolism of agars: the hypothetical protein Zg4663, distantly related to GH43 enzymes, has emerged as a specific α-1,3-galactosidase catalyzing the removal of 3,6-anhydro-L-galactose residues from the non-reducing ends of oligo-agars released by β-agarases, hence the name α-1,3-(3,6-anhydro)-L-galactosidase (ZgAhgA, also known as α-1,3-L-neoagarooligosaccharide hydrolase). It defines a new family of glycoside hydrolases, the GH117 family (Rebuffet et al. 2011). AhgA features a helix-turn-helix (HTH) domain connected to a five-bladed β-propeller domain and forms a dimer by swapping of the HTH

domain (Figure 1G). The putative catalytic residues, partially conserved with GH43 enzymes, are located at the bottom of the funnel-like active site. The mechanism of ZgAhgA is cation dependent, and a zinc ion has been identified in the active site, with an unusual coordination sphere occupied by water molecules. The amino acids binding these water molecules (and thus indirectly this cation) are strictly conserved with the GH117 family(Rebuffet et al. 2011). Two homologs of ZgAhgA have been characterized more recently, SdNABH from S. degradans and BpGH117 from B. plebeius. While no cation was found in the structure of SdNABH (Ha et al. 2011), BpGH117 features a magnesium ion at the position conserved with ZgAhgA (Hehemann et al. 2012c), suggesting a degree of plasticity for this cation-binding site. The structure of an inactive mutant of BpGH117 has also been determined, in a complex with neoagarobiose, identifying key residues for substrate recognition and catalysis. A mutagenesis approach has confirmed the involvement of five residues in catalysis: Asp90, Asp245, and Glu303 (conserved in family GH43), Glu167 (involved in the cation-binding site), and His302 (Hehemann et al. 2012c).

Alginate lyases

Alginate is a polymer of D-mannuronate and of its C5-epimer L-guluronate. It is an expolysaccharide in some bacteria and also the main cell wall compound of brown algae (Popper et al. 2011). Interestingly, genomic analysis has provided evidence that the common ancestor of brown algae acquired the alginate biosynthesis pathway from actinobacteria (Michel et al. 2010a). This highlights the importance of associated bacteria in the evolution of macroalgae. Alginate lyases (EC 4.2.2.3) are the key enzymes in alginate degradation and remodeling, to be found in seven polysaccharide lyase families: PL5, PL6, PL7, PL14, PL15, PL17, and PL18. Despite the importance of algal alginate as renewable biomass, most of the characterized alginate lyases originate from alginate-producing bacteria and from terrestrial bacteria feeding on bacterial alginate (Cantarel et al. 2009). Only five genes have been cloned from seaweed-associated bacteria: three PL7 genes (Han et al. 2004; Kim et al. 2009; Malissard et al. 1993) and two PL18 genes (Li et al. 2011; Sawabe et al. 2001). Knowledge in this field has recently advanced with the characterization of the alginolytic system of Z. galactanivorans (Thomas et al. 2012). This flavobacterium possesses seven alginate lyase genes (two PL6, three PL7, one PL14, and one PL17 gene) and a PL15 gene of uncertain specificity. Five of these genes are organized in clusters: a small cluster (alyA4, alyA5, alyA6) and a large cluster including alyA2, alyA3, and numerous carbohydrate-related genes predicted to be involved in alginate uptake and assimilation and in transcriptional regulation. These clusters have been shown to be genuine operons induced by alginate. ZgAlyA1, ZgAlyA4, ZgAlyA5, and ZgAlyA7 have been overexpressed in Escherichia coli and confirmed to be active alginate lyases. Zg2622 and Zg2614 are, respectively, a dehydrogenase and a kinase, further converting the terminal unsaturated monosaccharides released by alginate lyases to 2-keto-3-deoxy-6-phosphogluconate (Thomas et al. 2012). An in-depth study has demonstrated that ZgAlyA1 (PL7) is an endolytic guluronate lyase (EC 4.2.2.11), and ZgAlyA5 (PL7) cleaves unsaturated units, α-L-guluronate, or β-D-manuronate residues at the nonreducing ends of oligo-alginates in an exolytic fashion (EC 4.2.2.-). Despite a common jelly-roll fold, these striking differences in mode of action are due to different active site topologies: an open cleft in ZgAlyA1 (Figure 1H), whereas ZgAlyA5 displays a pocket topology due to the presence of additional loops partially obstructing the catalytic groove (Figure 1I). Lastly, in contrast to PL7 alginate lyases from terrestrial bacteria, both enzymes proceed according to a calcium-dependent mechanism, suggesting an exquisite adaptation to their natural substrate in the context of brown algal cell walls (Thomas et al. 2013).

Fucoidanases

Fucoidans are sulfated polysaccharides containing α -L-fucose residues and present in the cell wall of brown algae. They encompass a continuous spectrum of highly ramified polysaccharides, ranging from high-uronic-acid, low-sulfate polymers with significant proportions of D-xylose, D-galactose, and D-mannose to highly sulfated homofucan molecules (Popper et al, 2011). Only one fucanolytic gene has been cloned to date: the fucoidanase *fncA* from *Mariniflexile fucanivorans* SW5

(Colin et al. 2006). This marine flavobacterium was isolated from a water-treatment facility that recycles the effluent from an algal alginate extraction plant (Barbeyron et al. 2008; Descamps et al. 2006). FcnA encompasses an N-terminal catalytic module (~400 residues), three immunoglobulin-like modules, and a PorSS secretion module. A recombinant protein including the N-terminal module and the immunoglobulin-like modules has been overexpressed in *E. coli*, purified, and shown to retain the same activity as the wild-type enzyme. This fucoidanase releases as end products a tetrasaccharide and a hexasaccharide, and cleaves the α -1,4 glycosidic bonds between L-fucose-2,3-disulfate- α -1,3-L-fucose-2-sulfate repeating units. The N-terminal catalytic module displays ~25% identity to two patented fucoidanases from the bacterial strain SN-1009, and together these three proteins define a novel family of glycoside hydrolases, family GH107 (Colin et al. 2006).

Laminarinases

Laminarin, the storage polysaccharide of brown algae, is a small vacuolar beta-1,3-glucan containing ~25 glucosyl residues and some occasional β-1,6-linked branches. It includes two series, the minor G-series, containing only glucose residues, and the more abundant M-series, displaying a D-mannitol residue at the reducing end (Read et al. 1996). The unique presence of mannitol in laminarin is also explained by the horizontal gene transfer event involving the common ancestor of brown algae and an ancestral actinobacterium (Michel et al. 2010b). Laminarinases (EC 3.2.1.6 and 3.2.1.39) are found in several GH families (GH16, GH17, GH55, GH64, GH81, and GH128). Numerous beta-1,3-glucanases of terrestrial bacteria have been characterized in the context of the degradation of cell-wall beta-1,3-glucans of fungi, oomycetes, and land plants. Amazingly, however, among all the characterized beta-1,3-glucanases reported in the CAZY database (Cantarel et al. 2009), only one laminarinase gene has been cloned from a seaweed-associated bacterium: the GH16 laminarinase ZgLamA of Z. galactanivorans (Labourel et al. 2014). The 3D structure of ZgLamA_{GH16} and of two enzyme-substrate complexes, one with laminaritetraose and one with a trisaccharide of 1,3-1,4-β-D-glucan, have been determined this year. Compared to other GH16 laminarinases, ZgLamA_{GH16} contains a unique additional loop which gives a bent shape to the active-site cleft of the enzyme. This particular topology is perfectly adapted to the U-shaped conformation of laminarin chains in solution, and thus explains the predominant specificity of ZgLamAGH16 for this substrate (Labourel et al. 2014).

Ulvan lyases

Ulvans are the main cell-wall components of green algae of the genus *Ulva* (Popper et al. 2011). These complex sulfated polysaccharides are composed mainly of sulfated L-rhamnose, D-glucuronic acid and its C5-epimer L-iduronic acid, and a minor fraction of D-xylose (Lahaye and Robic 2007). The first described ulvanolytic bacterium was isolated at a "green tide" site in the Saint-Brieuc Bay (Brittany). This bacterium was not further characterized, but a semi-purified enzyme was shown to cleave the β-(1,4) linkage between L-rhamnose-3-sulfate (Rha3S) and D-glucuronic acid (GlcA), releasing an oligosaccharide with an unsaturated uronic acid at the non-reducing end. This enzyme was thus a polysaccharide lyase, referred to as an ulvan lyase (Lahaye et al. 1997). The only ulvan lyase gene to have been cloned was obtained from *Persicivirga ulvanivorans* (Nyvall Collén et al. 2011) a flavobacterium isolated from the faeces of the mollusc *Aplysia punctata* having fed on *Ulva* sp. (Barbeyron et al. 2011). This enzyme is endolytic and cleaves the glycosidic bond between the sulfated rhamnose and a glucuronic or iduronic acid. The sequence of this ulvan lyase has no similarity to known proteins (Nyvall Collén et al. 2011) and is currently an unclassified polysaccharide lyase in the CAZY database.

Microorganisms enhance algal defense, growth, and nutrient uptake

An increasing number of reviews discuss the beneficial contribution of microorganisms to algae, and notably their role in improving algal defense and nutrient uptake and in stimulating algal morphology and algal spore germination (Barott et al. 2011; Egan et al. 2013; Goecke et al. 2010; Harder et al. 2012). Alga-associated bacteria contribute to algal defense by

producing antimicrobial and antifouling compounds (Wilson et al. 2011). Table 3 shows diverse algal species on which bacteria with antimicrobial activities have been identified. Some 27% of isolated strains, on the average, show antimicrobial/antibacterial activity. The percentage is much lower for planktonic strains isolated from seawater (only 7% show antimicrobial activity) and even lower in terrestrial samples (Penesyan et al. 2009). The most represented bacterial genera are *Bacillus*, *Pseudoalteromonas*, *Pseudomonas*, and *Streptomyces*. Gram-positive *Bacillus* and *Streptomyces* and gram-negative *Pseudomonas* and *Pseudoalteromonas* are genera known for their ability to produce bioactive compounds (Bhatnagar and Kim 2010).

Prokaryotes have also been observed to synthesize necessary vitamins (Croft et al. 2006; Croft et al. 2005) and growth factors (Dimitrieva et al. 2006; Tsavkelova et al. 2006) and to improve algal growth by making these compounds accessible in sufficient amount.

Lastly, microbial epibionts produce common hydrolytic enzymes that improve algal nutrient uptake and development. A bacterial strain isolated from the red alga *Sargassum serratifolium*, for example, was shown to contain, in addition to agarase activities, diverse other hydrolytic activities such as amylase, alkaline phosphatase, esterase and lipase (C14), β-galactosidase, and urease activities (Kim and Hong 2012). In other bacterial strains also found on *Sargassum sp.*, amylase, carboxymethylcellulase, and protease activities were found (Mohapatra et al. 2003). An alkaline serine protease with potential use in the laundry industry was found in *Bacillus megaterium* RRM2, isolated from a red alga species (Rajkumar et al. 2011). Furthermore, bacterial enzymes such as lipases and esterases (Rajkumar et al. 2011), cellulases (Dong et al. 2010; Fu et al. 2010; Gibbs et al. 1992), proteases (Cristóbal et al. 2011; Yang et al. 2013), amylases (Both et al. 1993; Liu et al. 2012), laccases (Fang et al. 2012; Ge et al. 2011), and beta-glucosidases (Cristóbal et al. 2009; Mai et al. 2013), distantly related to terrestrial ones and displaying original biochemistry, are increasingly being isolated from the marine environment.

All these interesting bioactive compounds and enzymes produced by microorganisms in interaction with algae might predictably be very useful in diverse medical and industrial applications, as described in the next section.

Interest of macroalga-associated microorganisms in biotechnological applications

Medical and pharmaceutical applications

Microbial pathogens are becoming increasingly resistant to antibiotics, making some human infections untreatable. Hence, new antimicrobial compounds of natural origin, specifically targeting certain pathogens, are urgently needed. The marine environment is increasingly explored for such compounds. Although marine macroorganisms, including algae, are known to produce many interesting antimicrobial, antifungal, and potentially therapeutic compounds (Engel et al. 2002; Kubanek et al. 2003; Mayer and Gustafson 2003; Paul and Puglisi 2004; Steinberg and de Nys 2002; Takamatsu et al. 2003), ensuring a continued supply of eukaryotic compounds seems quite impossible. Producing such compounds would require growing macroorganisms in large quantity, and this would require much time and space (Dobretsov et al. 2006). Furthermore, the chemical synthesis of complex eukaryotic compounds is difficult. Therefore as microorganisms on algae release many bioactive compounds that prevent extensive colonization by other microorganisms, larvae, or algae, they could represent an interesting source of new antimicrobials (see Table 2), easily exploitable as they produce compounds faster in large quantity and are easier to culture. Moreover, marine microorganisms seem extremely productive of secondary metabolites: in addition to antimicrobial metabolites, they have been found to produce antitumor, anticancer, cytotoxic, and photoprotective compounds (Bhatnagar and Kim 2010). For example, it has recently been shown that phloroglucinol, a precursor of brown algal phlorotannins used in medicine to treat abdominal pain (Chassany et al. 2007), is synthesized by a polyketide synthase acquired through horizontal gene transfer (HGT) from an ancestral actinobacterium (Meslet-Cladiere et al. 2013). This highlights the importance of bacterial epibionts both in algal evolution and as a source of interesting bioactive compounds (Meslet-Cladiere et al, 2013). Lastly, algal-polysaccharide-degrading enzymes have a wide range of medical and pharmaceutical applications because they produce remarkable biologically **active oligosaccharides** with properties useful in maintaining human health, such as anticoagulant (Pereira et al. 1999; Pushpamali et al. 2008), anti-inflammatory (Berteau and Mulloy 2003), antioxidant (Hatada et al. 2006; Jiao et al. 2012), or immunostimulating activity (Bhattacharyya et al. 2010). Furthermore, oligosaccharides derived from ulvans, agars, carrageenans, alginates, and other less known algal polysaccharides are explored for their potential use as prebiotics favoring gut health in humans and animals (O'Sullivan et al. 2010). To obtain active oligosaccharides with the desired properties, enzymatic production with specific algal-polysaccharide-degrading enzymes is required. As microorganisms on algae are main producers of such specific enzymes, they represent a great source of them.

Production of biofuels

The need to preserve fossil fuels has prompted increasing efforts to produce biofuels. Initial efforts focused on producing biofuels from plant biomass. Unfortunately, using this biomass requires complex extraction methods due to the presence of recalcitrant polysaccharides such as lignocellulose. Furthermore, to obtain plant biomass one needs land for cultivation, in competition with human and animal food. Therefore, non-lignocellulosic macroalgal biomass, requiring no land for cultivation and possessing a high carbohydrate content, seems an interesting alternative for biofuel production. Promising results have been obtained in studies aiming to produce bioethanol from brown algae (Enquist-Newman et al. 2013; Wargacki et al. 2012) or red algae (Kim et al. 2012). All of these studies used microbial enzymes, directly or indirectly, to degrade specific algal polysaccharides. For example, Wargacki et al. (2012) used *Escherichia coli* strains transformed with DNA encoding enzymes involved in alginate transport and metabolism, in combination with an extracellular depolymerization system, to metabolize alginate and synthesize ethanol. Kim et al. (2012) used several microbial agarases to saccharify agarose to monosugars for further fermentation to ethanol. Lastly, the very recent study of Enquist-Newman et al. (2013) used bacterial alginate and mannitol catabolism genes in *Saccharomyces cerevisiae* to metabolize alginate monomers (4-deoxy-L-erythro-5-hexoseulose uronates) and mannitol from brown seaweeds, for further fermentation of sugar to ethanol. These recent promising works demonstrate the advantage of identifying algal polysaccharide-degrading enzymes and the encoding genes for the production of green energy.

Industrial applications

The diversity of non-lignocellulosic, sulfated poly- and monosaccharides makes algal hydrocarbons interesting for diverse industrial and biotechnological applications (see Table 4). The most used and studied alga-specific polysaccharides are agars and carrageenans (red algae), ulvans (green algae), alginates, laminarin, and sulfated fucoidans (brown algae) (Popper et al. 2011). Polysaccharide biotechnology uses enzymes or enzyme systems to convert carbohydrate polymers to addedvalue new polysaccharides (De Ruiter and Rudolph 1997) and thus requires hydrolytic enzymes such as agarases, carrageenases, alginate lyases, fucoidanases, porphyranases, and sulfatases to modify useful algal polysaccharides, improve their structures, and enhance their functionalities. Moreover, enzymatic hydrolysis is increasingly viewed as a promising alternative to current chemical extraction methods (Gavrilescu and Chisti 2005). Therefore, industrialists also seek hydrolytic enzymes with commonly exploited activities, e.g. proteases, cellulases, amylases, beta-glucosidases and laccases, but with original properties making them suitable for new applications. To date, most enzymes used in industry have been isolated from microorganisms living in terrestrial environments, mainly soils. Marine microorganisms, being exposed to extreme temperature, pressure, salinity, and nutrient availability conditions, should provide new enzymes with original biochemistry and characteristics (Kennedy et al. 2011). Lastly, alga-associated microbial communities respond to their exposure to alga-derived metabolites by producing a range of specific compounds, potentially of biotechnological interest. For example, various industries use halogenated compounds that become a hazard when they end up in the environment. Many studies therefore focus on optimizing their biodegradation by microbial dehalogenases (Swanson 1999). Alga-associated microorganisms constitute a potential source of dehalogenases, as they appear to resist the halogenated metabolites that algae produce as a defense mechanism (Potin et al. 1999).

Prospects for exploiting algal epibionts

As shown in the first part of this review, microorganisms living on algae are highly diverse but underexplored. The composition of alga-associated microbial communities varies, for example, according to the alga phylum and species, the season, and the age of the thalli. Furthermore, as these microorganisms constantly metabolize algal products, they produce numerous specific enzymes and secondary metabolites. From their immense diversity and their constant activity stems their great potential as a source of novel and original enzymes and metabolites. Furthermore, specific hydrolytic enzymes with novel biochemistry are increasingly sought for biotechnological applications in biomass and biofuel production, medicine, and wide-ranging industrial applications. Algal polysaccharidases identified to date (such as agarases, carrageenases, and alginate lyases) display very specific structures and biochemistry, related only distantly to those of known terrestrial glycoside hydrolases. This highlights their huge potential for new and original biotechnological uses and the importance of investigating these interesting enzymes.

Most published investigations on algal epibionts and their metabolites have relied on cultivation methods. To our knowledge, indeed, all specific enzymes isolated from algal epibionts have been obtained from cultivable microbial strains. Some high-throughput screens of algal microbial communities have been performed, but functional metagenomics has not been used to identify new microbial enzymes and metabolites. Functional metagenomics and techniques such as high-throughput sequencing are powerful means of gaining knowledge on the microorganisms composing these underexplored communities and of identifying novel metabolites and specific enzymes produced by alga-associated microbes.

Acknowledgments

This project was funded by Wallonie-Bruxelles International (WBI) and the Fonds Scientifique de la Recherche (F.R.S-F.N.R.S) in the framework of the Collaboration Program Hubert Curien. GM was also supported by the French National Research Agency with regard to the Investment Expenditure Program IDEALG (http://www.idealg.ueb.eu/, grant agreement No. ANR-10-BTBR-04).

Conflicts of Interest

The authors declare no conflict of interest.

References

- Allouch J, Helbert W, Henrissat B, Czjzek M (2004) Parallel substrate binding sites in a beta-agarase suggest a novel mode of action on double-helical agarose. Structure 12:623–32. doi: 10.1016/j.str.2004.02.020
- Allouch J, Jam M, Helbert W, Barbeyron T, Kloareg B, Henrissat B, Czjzek M (2003) The three-dimensional structures of two beta-agarases. J Biol Chem 278:47171–80. doi: 10.1074/jbc.M308313200
- Andrykovitch G, Marx I (1988) Isolation of a New Polysaccharide-Digesting Bacterium from a Salt Marsh. Appl Environ Microbiol 54:1061–1063.
- Armstrong E, Rogerson A, Leftley JW (2000) The Abundance of Heterotrophic Protists Associated with Intertidal Seaweeds. Estuarine, Coastla Shelf Sci 50:415–424. doi: 10.1006/ecss.1999.0577
- Barbeyron T, Gerard A, Potin P, Henrissat B, Kloareg B (1998) The Kappa-Carrageenase of the Marine Bacterium *Cytophaga drobachiensis*. Structural and Phylogenetic Relationships within family-16 Glycoside hydrolases. Mol Biol Evol 15:528–537.
- Barbeyron T, Henrissat B, Kloareg B (1994) The gene encoding the kappa-carrageenase of *Alteromonas carrageenovora* is related to β -1,3-1,4-glucanases. Gene 139:105–109.
- Barbeyron T, L'Haridon S, Corre E, Kloareg B, Potin P (2001) *Zobellia galactanovorans* gen. nov., sp. nov., a marine species of *Flavobacteriaceae* isolated from a red alga, and classification of *[Cytophaga] uliginosa* (ZoBell and Upham 1944) Reichenbach 1989 as *Zobellia uliginosa* gen. nov., comb. nov. Int J Syst Evol Microbiol 51:985–97.
- Barbeyron T, L'Haridon S, Michel G, Czjzek M (2008) Mariniflexile fucanivorans sp. nov., a marine member of the *Flavobacteriaceae* that degrades sulphated fucans from brown algae. Int J Syst Evol Microbiol 58:2107–13. doi: 10.1099/ijs.0.65674-0
- Barbeyron T, Lerat Y, Sassi J-F, Le Panse S, Helbert W, Collén PN (2011) *Persicivirga ulvanivorans* sp. nov., a marine member of the family *Flavobacteriaceae* that degrades ulvan from green algae. Int J Syst Evol Microbiol 61:1899–905. doi: 10.1099/ijs.0.024489-0
- Barbeyron T, Michel G, Potin P, Henrissat B, Kloareg B (2000) Iota-Carrageenases constitute a novel family of glycoside hydrolases, unrelated to that of kappa-carrageenases. J Biol Chem 275:35499–505. doi: 10.1074/jbc.M003404200
- Barott KL, Rodriguez-Brito B, Janouškovec J, Marhaver KL, Smith JE, Keeling P, Rohwer FL (2011) Microbial diversity associated with four functional groups of benthic reef algae and the reef-building coral *Montastraea annularis*. Environ Microbiol 13:1192–204. doi: 10.1111/j.1462-2920.2010.02419.x
- Belas R (1989) Sequence analysis of the agrA gene encoding beta-agarase from *Pseudomonas atlantica*. J Bacteriol 171:602–605.
- Bengtsson M, Sjøtun K, Øvreås L (2010) Seasonal dynamics of bacterial biofilms on the kelp *Laminaria hyperborea*. Aquat Microb Ecol 60:71–83. doi: 10.3354/ame01409

- Bengtsson MM, Øvreås L (2010) Planctomycetes dominate biofilms on surfaces of the kelp *Laminaria hyperborea*. BMC Microbiol 10:261. doi: 10.1186/1471-2180-10-261
- Bengtsson MM, Sjoetun K, Lanzen A, Oevreas L (2013) Bacterial diversity in relation to secondary production and succession on surfaces of the kelp *Laminaria hyperborea*. ISME J 6:2188–2198.
- Berteau O, Mulloy B (2003) Sulfated fucans, fresh perspectives: structures, functions, and biological properties of sulfated fucans and an overview of enzymes active toward this class of polysaccharide. Glycobiology 13:29R–40R. doi: 10.1093/glycob/cwg058
- Bhatnagar I, Kim S-K (2010) Immense essence of excellence: marine microbial bioactive compounds. Mar Drugs 8:2673–701. doi: 10.3390/md8102673
- Bhattacharyya S, Liu H, Zhang Z, Jam M, Dudeja PK, Michel G, Linhardt RJ, Tobacman JK (2010) Carrageenan-induced innate immune response is modified by enzymes that hydrolyze distinct galactosidic bonds. J Nutr Biochem 21:906–13. doi: 10.1016/j.jnutbio.2009.07.002
- Bixler HJ, Porse H (2010) A decade of change in the seaweed hydrocolloids industry. J Appl Phycol 23:321–335. doi: 10.1007/s10811-010-9529-3
- Both H, Brown SH, Kelly RM (1993) Characterization of Amylolytic Enzymes, Having Both ox-1, 4 and ao-1, 6 Hydrolytic Activity, from the Thermophilic Archaea *Pyrococcus furiosus* and *Thermococcus litoralis*.
- Burke C, Steinberg P, Rusch D, Kjelleberg S, Thomas T (2011a) Bacterial community assembly based on functional genes rather than species. PNAS 108:14288–14293. doi: 10.1073/pnas.1101591108/-/DCSupplemental.www.pnas.org/cgi/doi/10.1073/pnas.1101591108
- Burke C, Thomas T, Lewis M, Steinberg P, Kjelleberg S (2011b) Composition, uniqueness and variability of the epiphytic bacterial community of the green alga *Ulva australis*. ISME J 5:590–600. doi: 10.1038/ismej.2010.164
- Cantarel BL, Coutinho PM, Rancurel C, Bernard T, Lombard V, Henrissat B (2009) The Carbohydrate-Active EnZymes database (CAZy): an expert resource for Glycogenomics. Nucleic Acids Res 37:D233–8. doi: 10.1093/nar/gkn663
- Chassany O, Bonaz B, Bruley DES Varannes S, Bueno L, Cargill G, Coffin B, Ducrotté P, Grangé V (2007) Acute exacerbation of pain in irritable bowel syndrome: efficacy of phloroglucinol/trimethylphloroglucinol. A randomized, double-blind, placebo-controlled study. Aliment Pharmacol Ther 25:1115–23. doi: 10.1111/j.1365-2036.2007.03296.x
- Colin S, Deniaud E, Jam M, Descamps V, Chevolot Y, Kervarec N, Yvin J-C, Barbeyron T, Michel G, Kloareg B (2006) Cloning and biochemical characterization of the fucanase FcnA: definition of a novel glycoside hydrolase family specific for sulfated fucans. Glycobiology 16:1021–32. doi: 10.1093/glycob/cwl029
- Corre S, Prieur D (1990) Density and Morphology of Epiphytic Bacteria on the Kelp *Laminaria digitata*. Bot Mar 33:515–523.
- Correc G, Hehemann J-H, Czjzek M, Helbert W (2011) Structural analysis of the degradation products of porphyran digested by *Zobellia galactanivorans* β-porphyranase A. Carbohydr Polym 83:277–283. doi: 10.1016/j.carbpol.2010.07.060
- Cosse A, Leblanc C, Potin P (2007) Dynamic Defense of Marine Macroalgae Against Pathogens: From Early Activated to Gene Regulated Responses. Adv Bot Res 46:221–266.
- Cristóbal H a., Schmidt A, Kothe E, Breccia J, Abate CM (2009) Characterization of inducible cold-active β-glucosidases from the psychrotolerant bacterium *Shewanella sp.* G5 isolated from a sub-Antarctic ecosystem. Enzyme Microb Technol 45:498–506. doi: 10.1016/j.enzmictec.2009.06.010
- Cristóbal HA, López MA, Kothe E, Abate CM (2011) Diversity of protease-producing marine bacteria from sub-antarctic environments. J Basic Microbiol 51:590–600. doi: 10.1002/jobm.201000413

- Croft MT, Lawrence AD, Raux-Deery E, Warren MJ, Smith AG (2005) Algae acquire vitamin B12 through a symbiotic relationship with bacteria. Nature 438:90–3. doi: 10.1038/nature04056
- Croft MT, Warren MJ, Smith AG (2006) Algae need their vitamins. Eukaryot Cell 5:1175-83. doi: 10.1128/EC.00097-06
- Cundell AM, Sleeter TD, Mitchell R (1977) Microbial Populations Associated with the Surface of the Brown Alga *Ascophyllum nodosum*. Microb Ecol 4:81–91.
- Descamps V, Colin S, Lahaye M, Jam M, Richard C, Potin P, Barbeyron T, Yvin J-C, Kloareg B (2006) Isolation and culture of a marine bacterium degrading the sulfated fucans from marine brown algae. Mar Biotechnol (NY) 8:27–39. doi: 10.1007/s10126-005-5107-0
- Dimitrieva GY, Crawford RL, Yuksel GU (2006) The nature of plant growth-promoting effects of a pseudoalteromonad associated with the marine algae *Laminaria japonica* and linked to catalase excretion. J Appl Microbiol 100:1159–1169. doi: 10.1111/j.1365-2672.2006.02831.x
- Dobretsov S, Dahms H-U, Qian P-Y (2006) Inhibition of biofouling by marine microorganisms and their metabolites. Biofouling 22:43–54. doi: 10.1080/08927010500504784
- Dong J, Hashikawa S, Konishi T, Tamaru Y, Araki T (2006) Cloning of the novel gene encoding beta-agarase C from a marine bacterium, *Vibrio sp.* strain PO-303, and characterization of the gene product. Appl Environ Microbiol 72:6399–401. doi: 10.1128/AEM.00935-06
- Dong J, Hong Y, Shao Z, Liu Z (2010) Molecular cloning, purification, and characterization of a novel, acidic, pH-stable endoglucanase from *Martelella mediterranea*. J Microbiol 48:393–398.
- Donlan RM (2002) Biofilms: microbial life on surfaces. Emerg Infect Dis 8:881-90. doi: 10.3201/eid0809.020063
- Egan S, Harder T, Burke C, Steinberg P, Kjelleberg S, Thomas T (2013) The seaweed holobiont: understanding seaweed-bacteria interactions. FEMS Microbiol Rev 37:462–76. doi: 10.1111/1574-6976.12011
- Egan S, Thomas T, Kjelleberg S (2008) Unlocking the diversity and biotechnological potential of marine surface associated microbial communities. Curr Opin Microbiol 11:219–25. doi: 10.1016/j.mib.2008.04.001
- Engel S, Jensen PR, Fenical W (2002) Chimical ecology of marine microbial defense. J Chem Ecol 28:1971–1985.
- Enquist-Newman M, Faust AME, Bravo DD, Santos CNS, Raisner RM, Hanel A, Sarvabhowman P, Le C, Regitsky DD, Cooper SR, Peereboom L, Clark A, Martinez Y, Goldsmith J, Cho MY, Donohoue PD, Luo L, Lamberson B, Tamrakar P, Kim EJ, Villari JL, Gill A, Tripathi S a, Karamchedu P, Paredes CJ, Rajgarhia V, Kotlar HK, Bailey RB, Miller DJ, Ohler NL, Swimmer C, Yoshikuni Y (2013) Efficient ethanol production from brown macroalgae sugars by a synthetic yeast platform. Nature. doi: 10.1038/nature12771
- Erwin PM, Olson JB, Thacker RW (2011) Phylogenetic diversity, host-specificity and community profiling of sponge-associated bacteria in the northern Gulf of Mexico. PLoS One 6:1–16. doi: 10.1371/journal.pone.0026806
- Fang Z-M, Li T-L, Chang F, Zhou P, Fang W, Hong Y-Z, Zhang X-C, Peng H, Xiao Y-Z (2012) A new marine bacterial laccase with chloride-enhancing, alkaline-dependent activity and dye decolorization ability. Bioresour Technol 111:36–41. doi: 10.1016/j.biortech.2012.01.172
- Fernandes N, Steinberg P, Rusch D, Kjelleberg S, Thomas T (2012) Community structure and functional gene profile of bacteria on healthy and diseased thalli of the red seaweed *Delisea pulchra*. PLoS One 7:e50854. doi: 10.1371/journal.pone.0050854
- Flament D, Barbeyron T, Jam M, Potin P, Czjzek M, Kloareg B, Michel G (2007) Alpha-agarases define a new family of glycoside hydrolases, distinct from beta-agarase families. Appl Environ Microbiol 73:4691–4. doi: 10.1128/AEM.00496-07
- Fu X, Liu P, Lin L, Hong Y, Huang X, Meng X, Liu Z (2010) A Novel Endoglucanase (Cel9P) From a Marine Bacterium *Paenibacillus sp.* BME-14. Appl Biochem Biotechnol 160:1627–1636.

- Gavrilescu M, Chisti Y (2005) Biotechnology-a sustainable alternative for chemical industry. Biotechnol Adv 23:471–99. doi: 10.1016/j.biotechadv.2005.03.004
- Ge H, Xu P, Xu Y, Fang Z, Xiao Y (2011) Purification, crystallization and preliminary crystallographic analysis of recombinant Lac15 from a marine microbial metagenome. Acta Crystallogr Sect F Struct Biol Cryst Commun 67:956–8. doi: 10.1107/S1744309111024912
- Genilloud O, Pelaez F, Gonzalez I, Diez MT (1994) Diversity of actinomycetes and fungi on seaweeds from the Iberian coasts. Microbiologia 10:413–422.
- Gibbs MD, Saul DJ, Lüthi E, Bergquist PL (1992) The beta-mannanase from "Caldocellum saccharolyticum" is part of a multidomain enzyme. Appl Environ Microbiol 58:3864–7.
- Goecke F, Labes a, Wiese J, Imhoff J (2010) Chemical interactions between marine macroalgae and bacteria. Mar Ecol Prog Ser 409:267–299. doi: 10.3354/meps08607
- Goecke F, Labes A, Wiese J, Imhoff JF (2013) Phylogenetic analysis and antibiotic activity of bacteria isolated from the surface of two co-occurring macroalgae from the Baltic Sea. Eur J Phycol 48:47–60. doi: 10.1080/09670262.2013.767944
- Gontang E a, Fenical W, Jensen PR (2007) Phylogenetic diversity of gram-positive bacteria cultured from marine sediments. Appl Environ Microbiol 73:3272–82. doi: 10.1128/AEM.02811-06
- Gravot A, Dittami S, Rousvoal S, Lugan R, Eggert A, Collén J, Boyen C, Bouchereau A, Tonon T (2010) Diurnal oscillations of metabolite abundances and gene analysis provide new insights into central metabolic processes of the brown alga *Ectocarpus siliculosus*. New Phytol 188:98–110.
- Guibet M, Barbeyron T, Genicot S, Kloareg B, Michel G, Helbert W (2007) Degradation of λ -carrageenan by *Pseudoalteromonas carrageenovora* λ -carrageenase: a new family of glycoside hydrolases unrelated to κ - and ι -carrageenases. Biochem J 114:105–114. doi: 10.1042//BJ20061359
- Ha SC, Lee S, Lee J, Kim HT, Ko H-J, Kim KH, Choi I-G (2011) Crystal structure of a key enzyme in the agarolytic pathway, α-neoagarobiose hydrolase from *Saccharophagus degradans* 2-40. Biochem Biophys Res Commun 412:238–44. doi: 10.1016/j.bbrc.2011.07.073
- Han F, Gong Q-H, Song K, Li J-B, Yu W-G (2004) Cloning, sequence analysis and expression of gene alyVI encoding alginate lyase from marine bacterium *Vibrio sp.* QY101. DNA Seq 15:344–50. doi: 10.1080/10425170400019300
- Harder T, Campbell AH, Egan S, Steinberg PD (2012) Chemical mediation of ternary interactions between marine holobionts and their environment as exemplified by the red alga *Delisea pulchra*. J Chem Ecol 38:442–50. doi: 10.1007/s10886-012-0119-5
- Hatada Y, Mizuno M, Li Z, Ohta Y (2011) Hyper-production and characterization of the t-carrageenase useful for t-carrageenan oligosaccharide production from a deep-sea bacterium, *Microbulbifer thermotolerans* JAMB-A94T, and insight into the unusual catalytic mechanism. Mar Biotechnol (NY) 13:411–22. doi: 10.1007/s10126-010-9312-0
- Hatada Y, Ohta Y, Horikoshi K (2006) Hyperproduction and application of alpha-agarase to enzymatic enhancement of antioxidant activity of porphyran. J Agric Food Chem 54:9895–900. doi: 10.1021/jf0613684
- Hehemann J-H, Correc G, Barbeyron T, Helbert W, Czjzek M, Michel G (2010) Transfer of carbohydrate-active enzymes from marine bacteria to Japanese gut microbiota. Nature 464:908–12. doi: 10.1038/nature08937
- Hehemann J-H, Correc G, Thomas F, Bernard T, Barbeyron T, Jam M, Helbert W, Michel G, Czjzek M (2012a) Biochemical and structural characterization of the complex agarolytic enzyme system from the marine bacterium *Zobellia galactanivorans*. J Biol Chem 287:30571–84. doi: 10.1074/jbc.M112.377184
- Hehemann J-H, Kelly AG, Pudlo N a, Martens EC, Boraston AB (2012b) Bacteria of the human gut microbiome catabolize red seaweed glycans with carbohydrate-active enzyme updates from extrinsic microbes. Proc Natl Acad Sci U S A 109:19786–91. doi: 10.1073/pnas.1211002109

- Hehemann J-H, Smyth L, Yadav A, Vocadlo DJ, Boraston AB (2012c) Analysis of keystone enzyme in Agar hydrolysis provides insight into the degradation (of a polysaccharide from) red seaweeds. J Biol Chem 287:13985–95. doi: 10.1074/jbc.M112.345645
- Hollants J, Leroux O, Leliaert F, Decleyre H, De Clerck O, Willems A (2011) Who is in there? Exploration of endophytic bacteria within the siphonous green seaweed *Bryopsis* (*Bryopsidales*, *Chlorophyta*). PLoS One 6:e26458. doi: 10.1371/journal.pone.0026458
- Ismail-Ben Ali a., El Bour M, Ktari L, Bolhuis H, Ahmed M, Boudabbous a., Stal LJ (2011) *Jania rubens*-associated bacteria: molecular identification and antimicrobial activity. J Appl Phycol 24:525–534. doi: 10.1007/s10811-011-9758-0
- Jam M, Flament D, Allouch J, Potin P, Thion L, Kloareg B, Czjzek M, Helbert W, Michel G, Barbeyron T (2005) The endo-beta-agarases AgaA and AgaB from the marine bacterium *Zobellia galactanivorans*: two paralogue enzymes with different molecular organizations and catalytic behaviours. Biochem J 385:703–13. doi: 10.1042/BJ20041044
- Jiao G, Yu G, Zhang J, Ewart HS (2011) Chemical structures and bioactivities of sulfated polysaccharides from marine algae. Mar Drugs 9:196–223. doi: 10.3390/md9020196
- Jiao G, Yu G, Zhao X, Zhang J, Ewart HS (2012) Natural Polymers with Antioxidant Properties: Poly-/oligosaccharides of Marine Origin. Antioxid. Polym. Synth. Prop. Appl. pp 179–201
- Jung K a, Lim S-R, Kim Y, Park JM (2013) Potentials of macroalgae as feedstocks for biorefinery. Bioresour Technol 135:182–90. doi: 10.1016/j.biortech.2012.10.025
- Kanagasabhapathy M, Sasaki H, Haldar S, Yamasaki S, Nagata S (2006) Antibacterial activities of marine epibiotic bacteria isolated from brown algae of Japan. Ann Microbiol 56:167–173. doi: 10.1007/BF03175000
- Kanagasabhapathy M, Sasaki H, Nagata S (2008) Phylogenetic identification of epibiotic bacteria possessing antimicrobial activities isolated from red algal species of Japan. World J Microbiol Biotechnol 24:2315–2321. doi: 10.1007/s11274-008-9746-y
- Kennedy J, O'Leary ND, Kiran GS, Morrissey JP, O'Gara F, Selvin J, Dobson ADW (2011) Functional metagenomic strategies for the discovery of novel enzymes and biosurfactants with biotechnological applications from marine ecosystems. J Appl Microbiol 111:787–799. doi: 10.1111/j.1365-2672.2011.05106.x
- Kilian SG, Prior BA, Venter JJ, Lategan PM (1985) App Microbiology Biotechnology Production, purification, and properties of beta-glucosidase from *Candida wickerhamii*. Appl Microbiol Biotechnol 21:148–153.
- Kim DE, Lee EY, Kim HS (2009) Cloning and characterization of alginate lyase from a marine bacterium Streptomyces sp. ALG-5. Mar Biotechnol (NY) 11:10–6. doi: 10.1007/s10126-008-9114-9
- Kim HT, Lee S, Kim KH, Choi I-G (2012) The complete enzymatic saccharification of agarose and its application to simultaneous saccharification and fermentation of agarose for ethanol production. Bioresour Technol 107:301–6. doi: 10.1016/j.biortech.2011.11.120
- Kim J, Hong S-K (2012) Isolation and characterization of an agarase-producing bacterial strain, *Alteromonas sp.* GNUM-1, from the West Sea, Korea. J Microbiol Biotechnol 22:1621–8. doi: http://dx.doi.org/10.4014/jmb.1209.08087
- Kobayashi T, Uchimura K, Koide O, Deguchi S, Horikoshi K (2012) Genetic and Biochemical Characterization of the *Pseudoalteromonas tetraodonis* Alkaline κ-Carrageenase. Biosci Biotechnol Biochem 76:506–511. doi: 10.1271/bbb.110809
- Kubanek J, Jensen PR, Keifer P a, Sullards MC, Collins DO, Fenical W (2003) Seaweed resistance to microbial attack: a targeted chemical defense against marine fungi. Proc Natl Acad Sci U S A 100:6916–21. doi: 10.1073/pnas.1131855100
- Labourel A, Jam M, Jeudy A, Hehemann J-H, Czjzek M, Michel G (2014) The β-Glucanase ZgLamA from *Zobellia* galactanivorans evolved a bent active site adapted for efficient degradation of algal laminarin. J Biol Chem 289:2027–2042. doi: 10.1074/jbc.M113.538843

- Lachnit T, Blümel M, Imhoff J, Wahl M (2009) Specific epibacterial communities on macroalgae: phylogeny matters more than habitat. Aquat Biol 5:181–186. doi: 10.3354/ab00149
- Lachnit T, Meske D, Wahl M, Harder T, Schmitz R (2011) Epibacterial community patterns on marine macroalgae are host-specific but temporally variable. Environ Microbiol 13:655–65. doi: 10.1111/j.1462-2920.2010.02371.x
- Lahaye M, Brunel M, Bonnin E (1997) Fine chemical structure analysis of oligosaccharides produced by an ulvan-lyase degradation of the water-soluble cell-wall polysaccharides from *Ulva sp*, (*Ulvales*, *Chlorophyta*). Carbohydr Res 304:325–33.
- Lahaye M, Robic A (2007) Structure and functional properties of ulvan, a polysaccharide from green seaweeds. Biomacromolecules 8:1765–74. doi: 10.1021/bm061185q
- Lemoine M, Nyvall Collén P, Helbert W (2009) Physical state of kappa-carrageenan modulates the mode of action of kappa-carrageenase from *Pseudoalteromonas carrageenovora*. Biochem J 419:545–53. doi: 10.1042/BJ20080619
- Lemos ML, Toranzo AE, Barja JL (1985) Antibiotic activity of epiphytic bacteria isolated from intertidal seaweeds. Microb Ecol 11:149–163. doi: 10.1007/BF02010487
- Li B, Lu F, Wei X, Zhao R (2008) Fucoidan: Structure and Bioactivity. Molecules 13:1671–1695. doi: 10.3390/molecules13081671
- Li J-W, Dong S, Song J, Li C-B, Chen X-L, Xie B-B, Zhang Y-Z (2011) Purification and characterization of a bifunctional alginate lyase from *Pseudoalteromonas sp.* SM0524. Mar Drugs 9:109–23. doi: 10.3390/md9010109
- Liu G-L, Li Y, Chi Z, Chi Z-M (2011) Purification and characterization of κ-carrageenase from the marine bacterium *Pseudoalteromonas porphyrae* for hydrolysis of κ-carrageenan. Process Biochem 46:265–271. doi: 10.1016/j.procbio.2010.08.021
- Liu Y, Lei Y, Zhang X, Gao Y, Xiao Y, Peng H (2012) Identification and phylogenetic characterization of a new subfamily of α -amylase enzymes from marine microorganisms. Mar Biotechnol (NY) 14:253–60. doi: 10.1007/s10126-011-9414-3
- Liu Z, Li G, Mo Z, Mou H (2013) Molecular cloning, characterization, and heterologous expression of a new κ-carrageenase gene from marine bacterium *Zobellia sp.* ZM-2. Appl Microbiol Biotechnol 97:10057–67. doi: 10.1007/s00253-013-5215-0
- Longford S, Tujula N, Crocetti G, Holmes A, Holmström C, Kjelleberg S, Steinberg P, Taylor M (2007) Comparisons of diversity of bacterial communities associated with three sessile marine eukaryotes. Aquat Microb Ecol 48:217–229. doi: 10.3354/ame048217
- Ma C, Lu X, Shi C, Li J, Gu Y, Ma Y, Chu Y, Han F, Gong Q, Yu W (2007) Molecular cloning and characterization of a novel beta-agarase, AgaB, from marine *Pseudoalteromonas sp.* CY24. J Biol Chem 282:3747–54. doi: 10.1074/jbc.M607888200
- Ma S, Duan G, Chai W, Geng C, Tan Y, Wang L, Le Sourd F, Michel G, Yu W, Han F (2013) Purification, cloning, characterization and essential amino acid residues analysis of a new 1-carrageenase from *Cellulophaga sp.* QY3. PLoS One 8:e64666. doi: 10.1371/journal.pone.0064666
- Mai Z, Yang J, Tian X, Li J, Zhang S (2013) Gene cloning and characterization of a novel salt-tolerant and glucose-enhanced β-glucosidase from a marine streptomycete. Appl Biochem Biotechnol 169:1512–1522.
- Malissard M, Duez C, Guinand M, Vacheron M-J, Michel G, Marty N, Joris B, Thamm I, Ghuysen J-M (1993) Sequence of a gene encoding a (poly ManA) alginate lyase acitve on *Pseudomonas aeruginosa* alginate. FEMS Microbiol Lett 110:101–106.
- Mayer AMS, Gustafson KR (2003) Marine pharmacology in 2000: antitumor and cytotoxic compounds. Int J Cancer 105:291–9. doi: 10.1002/ijc.11080

- Mazure HG., Field JG (1980) Density and ecological importance of bacteria on kelp fronds in an upwelling region. J Exp Mar Bio Ecol 43:173–182.
- McHugh DJ (2003) A guide to the seaweed industry. FAO Fish Tech Pap 105.
- Meslet-Cladiere L, Delage L, J.-J. Leroux C, Goulitquer S, Leblanc C, Creis E, Gall E a., Stiger-Pouvreau V, Czjzek M, Potin P (2013) Structure/Function Analysis of a Type III Polyketide Synthase in the Brown Alga *Ectocarpus siliculosus* Reveals a Biochemical Pathway in Phlorotannin Monomer Biosynthesis. Plant Cell 1–16. doi: 10.1105/tpc.113.111336
- Meusnier I, Olsen JL, Stam WT, Destombe C, Valero M (2001) Phylogenetic analyses of *Caulerpa taxifolia* (*Chlorophyta*) and of its associated bacterial microflora provide clues to the origin of the Mediterranean introduction. Mol Ecol 10:931–46.
- Michel G, Barbeyron T, Kloareg B, Czjzek M (2009) The family 6 carbohydrate-binding modules have coevolved with their appended catalytic modules toward similar substrate specificity. Glycobiology 19:615–23. doi: 10.1093/glycob/cwp028
- Michel G, Chantalat L, Duee E, Barbeyron T, Henrissat B, Kloareg B, Dideberg O (2001a) The kappa-carrageenase of *P. carrageenovora* features a tunnel-shaped active site: a novel insight in the evolution of Clan-B glycoside hydrolases. Structure 9:513–25.
- Michel G, Chantalat L, Fanchon E, Henrissat B, Kloareg B, Dideberg O (2001b) The iota-carrageenase of *Alteromonas fortis*. A beta-helix fold-containing enzyme for the degradation of a highly polyanionic polysaccharide. J Biol Chem 276:40202–9. doi: 10.1074/jbc.M100670200
- Michel G, Czjzek M (2013) Polysaccharide-degrading enzymes from marine bacteria. In: Trincone A (ed) Marine enzymes for biocatalysis: sources, biocatalytic characteristic and bioprocesses of marine enzymes, Woodhead Publishing Series in Biomedicine, pp 429-464.
- Michel G, Helbert W, Kahn R, Dideberg O, Kloareg B (2003) The Structural Bases of the Processive Degradation of t-Carrageenan, a Main Cell Wall Polysaccharide of Red Algae. J Mol Biol 334:421–433. doi: 10.1016/j.jmb.2003.09.056
- Michel G, Tonon T, Scornet D, Cock JM, Kloareg B (2010a) The cell wall polysaccharide metabolism of the brown alga *Ectocarpus siliculosus*. Insights into the evolution of extracellular matrix polysaccharides in Eukaryotes. New Phytol 188:82–97. doi: 10.1111/j.1469-8137.2010.03374.x
- Michel G, Tonon T, Scornet D, Cock JM, Kloareg B (2010b) Central and storage carbon metabolism of the brown alga Ectocarpus siliculosus: insights into the origin and evolution of storage carbohydrates in Eukaryotes. New Phytol 188:67–81. doi: 10.1111/j.1469-8137.2010.03345.x
- Mohapatra BR, Bapuji M, A. S (2003) Production of Industrial Enzymes (Amylase, Carboxymethylcellulase and Protease) by Bacteria Isolated from Marine Sedentary Organisms. Acta Biotechnol 23:75–84.
- Morris RM, Rappe MS, Connon SA, Vergin KL, Siebold WA, Carlson CA, Giovannoni SJ (2002) SAR11 clade dominates ocean surface bacterioplankton communities. Nature 420:806–809. doi: 10.1038/nature01281.1.
- Nyvall Collén P, Sassi J-F, Rogniaux H, Marfaing H, Helbert W (2011) Ulvan lyases isolated from the Flavobacteria *Persicivirga ulvanivorans* are the first members of a new polysaccharide lyase family. J Biol Chem 286:42063–71. doi: 10.1074/jbc.M111.271825
- O'Sullivan L, Murphy B, McLoughlin P, Duggan P, Lawlor PG, Hughes H, Gardiner GE (2010) Prebiotics from marine macroalgae for human and animal health applications. Mar Drugs 8:2038–64. doi: 10.3390/md8072038
- Oh C, Nikapitiya C, Lee Y, Whang I, Kim S-J, Kang D-H, Lee J (2010) Cloning, purification and biochemical characterization of beta agarase from the marine bacterium *Pseudoalteromonas sp.* AG4. J Ind Microbiol Biotechnol 37:483–94. doi: 10.1007/s10295-010-0694-9
- Ohta Y, Hatada Y (2006) A novel enzyme, lambda-carrageenase, isolated from a deep-sea bacterium. J Biochem 140:475–481.

- Pasmore M, Costerton JW (2003) Biofilms, bacterial signaling, and their ties to marine biology. J Ind Microbiol Biotechnol 30:407–13. doi: 10.1007/s10295-003-0069-6
- Paul VJ, Puglisi MP (2004) Chemical mediation of interactions among marine organisms. Nat Prod Rep 21:189–209. doi: 10.1039/b302334f
- Penesyan A, Marshall-Jones Z, Holmstrom C, Kjelleberg S, Egan S (2009) Antimicrobial activity observed among cultured marine epiphytic bacteria reflects their potential as a source of new drugs. FEMS Microbiol Ecol 69:113–24. doi: 10.1111/j.1574-6941.2009.00688.x
- Pereira MS, Mulloy B, Mourao PAS (1999) Structure and Anticoagulant Activity of Sulfated Fucans: Comparison between the regular, repetitive, and linear fucans from echinoderms with the more heteroheneous and branched polymers from brown algae. J Biol Chem 274:7656–7667. doi: 10.1074/jbc.274.12.7656
- Pluvinage B, Hehemann J-H, Boraston AB (2013) Substrate recognition and hydrolysis by a family 50 exo-β-agarase, Aga50D, from the marine bacterium *Saccharophagus degradans*. J Biol Chem 288:28078–88. doi: 10.1074/jbc.M113.491068
- Popper Z a, Michel G, Hervé C, Domozych DS, Willats WGT, Tuohy MG, Kloareg B, Stengel DB (2011) Evolution and diversity of plant cell walls: from algae to flowering plants. Annu Rev Plant Biol 62:567–90. doi: 10.1146/annurev-arplant-042110-103809
- Potin P, Bouarab K, Kupper F, Kloareg B (1999) Oligosaccharide recognition signals and defence reactions in marine plant-microbe interactions. Curr Opin Microbiol 2:276–283.
- Potin P, Bouarab K, Salaün J-P, Pohnert G, Kloareg B (2002) Biotic interactions of marine algae. Curr Opin Plant Biol 5:308–317. doi: 10.1016/S1369-5266(02)00273-X
- Potin P, Richard C, Rochas C, Kloareg B (1993) Purification and characterization of the alpha-agarase from *Alteromonas agarlyticus* (Cataldi) comb. nov., strain GJ1B. Eur J Biochem 214:599–607.
- Pushpamali WA, Nikapitiya C, Zoysa M De, Whang I, Kim SJ, Lee J (2008) Isolation and purification of an anticoagulant from fermented red seaweed *Lomentaria catenata*. Carbohydr Polym 73:274–279. doi:10.1016/j.carbpol.2007.11.029
- Rajkumar R, Jayappriyan KR, Rengasamy R (2011) Purification and characterization of a protease produced by *Bacillus megaterium* RRM2: application in detergent and dehairing industries. J Basic Microbiol 51:614–24. doi: 10.1002/jobm.201000517
- Rao TS (2010) Comparative effect of temperature on biofilm formation in natural and modified marine environment. Aquat Ecol 44:463–478. doi: 10.1007/s10452-009-9304-1
- Read SM, Currie G, Bacic a (1996) Analysis of the structural heterogeneity of laminarin by electrospray-ionisation-mass spectrometry. Carbohydr Res 281:187–201.
- Rebuffet E, Barbeyron T, Jeudy A, Jam M, Czjzek M, Michel G (2010) Identification of catalytic residues and mechanistic analysis of family GH82 iota-carrageenases. Biochemistry 49:7590–9. doi: 10.1021/bi1003475
- Rebuffet E, Groisillier A, Thompson A, Jeudy A, Barbeyron T, Czjzek M, Michel G (2011) Discovery and structural characterization of a novel glycosidase family of marine origin. Environ Microbiol 13:1253–70. doi: 10.1111/j.1462-2920.2011.02426.x
- Ren A, Xia Z-X, Yu W, Zhou J (2010) Expression, crystallization and preliminary X-ray analysis of an anomeric inverting agarase from *Pseudoalteromonas sp.* CY24. Acta Crystallogr Sect F Struct Biol Cryst Commun 66:1635–9. doi: 10.1107/S174430911004114X
- Rohwer F, Seguritan V, Azam F, Knowlton N (2002) Diversity and distribution of coral-associated bacteria. Mar Ecol Prog Ser 243:1–10. doi: 10.3354/meps243001
- De Ruiter GA, Rudolph B (1997) Carrageenan biotechnology. Trends Food Sci Technol 8:389–395.

- Salaün S (2009) Interactions entre la macroalgue brune *Laminaria digitata* et ses épibiontes bactériens : Etudes moléculaire et spectroscopiques , et capacité d'adhésion et de formation de biofilm. Thesis, University of Pierre and Marie Curie, Paris, Fr.
- Salaün S, La Barre S, Dos Santos-Goncalvez M, Potin P, Haras D, Bazire A (2012) Influence of exudates of the kelp *Laminaria digitata* on biofilm formation of associated and exogenous bacterial epiphytes. Microb Ecol 64:359–69. doi: 10.1007/s00248-012-0048-4
- Salaün S, Kervarec N, Potin P, Haras D, Piotto M, La Barre S (2010) Whole-cell spectroscopy is a convenient tool to assist molecular identification of cultivatable marine bacteria and to investigate their adaptive metabolism. Talanta 80:1758–70. doi: 10.1016/j.talanta.2009.10.020
- Sawabe T, Takahashi H, Ezura Y, Gacesa P (2001) Cloning, sequence analysis and expression of *Pseudoalteromonas elyakovii* IAM 14594 gene (alyPEEC) encoding the extracellular alginate lyase. Carbohydr Res 335:11–21.
- Schaumann K, Weide G (1995) Efficiency of uronic acid uptake in marine alginate-degrading fungi. Helgoländer Meeresuntersuchungen 49:159–167. doi: 10.1007/BF02368346
- Schroeder DC (2003) Investigation of the role of a (1-4) agarase produced by *Pseudoalteromonas gracilis* B9 in eliciting disease symptoms in the red alga *Gracilaria gracilis*. Microbiology 149:2919–2929. doi: 10.1099/mic.0.26513-0
- Staufenberger T, Thiel V, Wiese J, Imhoff JF (2008) Phylogenetic analysis of bacteria associated with *Laminaria* saccharina. FEMS Microbiol Ecol 64:65–77. doi: 10.1111/j.1574-6941.2008.00445.x
- Steinberg PD, de Nys R (2002) Chemical mediation of colonization of seaweed surfaces. J Phycol 38:621-629.
- Stratil SB, Neulinger SC, Knecht H, Friedrichs AK, Wahl M (2013) Temperature-driven shifts in the epibiotic bacterial community composition of the brown macroalga *Fucus vesiculosus*. Microbiologyopen 2:338–49. doi: 10.1002/mbo3.79
- Sugano Y, Matsumoto T, Kodama H, Noma M (1993) Cloning and Sequencing of agaA, a Unique Agarase 0107 Gene from a Marine Bacterium, *Vibrio sp*. Strain JT0107. Appl Environ Microbiol 59:3750–3756.
- Swanson PE (1999) Dehalogenases applied to industrial-scale biocatalysis. Curr Opin Biotechnol 10:365–369.
- Tait K, Williamson H, Atkinson S, Williams P, Cámara M, Joint I (2009) Turnover of quorum sensing signal molecules modulates cross-kingdom signalling. Environ Microbiol 11:1792–802. doi: 10.1111/j.1462-2920.2009.01904.x
- Takamatsu S, Hodges TW, Rajbhandari I, Gerwick WH, Hamann MT, Nagle DG (2003) Marine natural products as novel antioxidant prototypes. J Nat Prod 66:605–8. doi: 10.1021/np0204038
- Taylor MW, Schupp PJ, Dahllöf I, Kjelleberg S, Steinberg PD (2003) Host specificity in marine sponge-associated bacteria, and potential implications for marine microbial diversity. Environ Microbiol 6:121–130. doi: 10.1046/j.1462-2920.2003.00545.x
- Thomas F, Barbeyron T, Tonon T, Génicot S, Czjzek M, Michel G (2012) Characterization of the first alginolytic operons in a marine bacterium: from their emergence in marine *Flavobacteriia* to their independent transfers to marine Proteobacteria and human gut Bacteroides. Environ Microbiol 14:2379–94. doi: 10.1111/j.1462-2920.2012.02751.x
- Thomas F, Lundqvist LCE, Jam M, Jeudy A, Barbeyron T, Sandström C, Michel G, Czjzek M (2013) Comparative characterization of two marine alginate lyases from *Zobellia galactanivorans* reveals distinct modes of action and exquisite adaptation to their natural substrate. J Biol Chem 288:23021–23037. doi: 10.1074/jbc.M113.467217
- Tsavkelova E a., Klimova SY, Cherdyntseva T a., Netrusov a. I (2006) Microbial producers of plant growth stimulators and their practical use: A review. Appl Biochem Microbiol 42:117–126. doi: 10.1134/S0003683806020013
- Tujula N a, Crocetti GR, Burke C, Thomas T, Holmström C, Kjelleberg S (2010) Variability and abundance of the epiphytic bacterial community associated with a green marine *Ulvacean alga*. ISME J 4:301–11. doi: 10.1038/ismej.2009.107

- Uchida M, Murata M (2004) Isolation of a lactic acid bacterium and yeast consortium from a fermented material of *Ulva spp.* (*Chlorophyta*). J Appl Microbiol 97:1297–310. doi: 10.1111/j.1365-2672.2004.02425.x
- Vera J, Castro J, Gonzalez A, Moenne A (2011) Seaweed polysaccharides and derived oligosaccharides stimulate defense responses and protection against pathogens in plants. Mar Drugs 9:2514–25. doi: 10.3390/md9122514
- Wargacki AJ, Leonard E, Win MN, Regitsky DD, Santos CNS, Kim PB, Cooper SR, Raisner RM, Herman A, Sivitz AB, Lakshmanaswamy A, Kashiyama Y, Baker D, Yoshikuni Y (2012) An engineered microbial platform for direct biofuel production from brown macroalgae. Science 335:308–13. doi: 10.1126/science.1214547
- Wegley L, Edwards R, Rodriguez-Brito B, Liu H, Rohwer F (2007) Metagenomic analysis of the microbial community associated with the coral *Porites astreoides*. Environ Microbiol 9:2707–19. doi: 10.1111/j.1462-2920.2007.01383.x
- Weinberger F, Friedlander M, Gunkel W (1994) A bacterial facultative parasite of *Gracilaria conferta*. Dis Aquat Organ 18:135–141.
- Wiese J, Thiel V, Nagel K, Staufenberger T, Imhoff JF (2009) Diversity of antibiotic-active bacteria associated with the brown alga *Laminaria saccharina* from the Baltic Sea. Mar Biotechnol (NY) 11:287–300. doi: 10.1007/s10126-008-9143-4
- Wilson GS, Raftos D a, Corrigan SL, Nair S V (2010) Diversity and antimicrobial activities of surface-attached marine bacteria from Sydney Harbour, Australia. Microbiol Res 165:300–11. doi: 10.1016/j.micres.2009.05.007
- Wilson GS, Raftos D a, Nair S V (2011) Antimicrobial activity of surface attached marine bacteria in biofilms. Microbiol Res 166:437–48. doi: 10.1016/j.micres.2010.08.003
- Yang J, Li J, Mai Z, Tian X, Zhang S (2013) Purification, characterization, and gene cloning of a cold-adapted thermolysin-like protease from *Halobacillus sp.* SCSIO 20089. J Biosci Bioeng 115:628–32. doi: 10.1016/j.jbiosc.2012.12.013
- Yang J-I, Chen L-C, Shih Y-Y, Hsieh C, Chen C-Y, Chen W-M, Chen C-C (2011) Cloning and characterization of β-agarase AgaYT from *Flammeovirga yaeyamensis* strain YT. J Biosci Bioeng 112:225–32. doi: 10.1016/j.jbiosc.2011.05.016
- Yao Z, Wang F, Gao Z, Jin L, Wu H (2013) Characterization of a κ-Carrageenase from Marine *Cellulophaga lytica* strain N5-2 and Analysis of Its Degradation Products. Int J Mol Sci 14:24592–602. doi: 10.3390/ijms141224592
- Yaphe W (1957) The use of agarase from *Pseudomonas atlantica* in the identification of agar in marine algae (RHODOPHYCEAE). Can J Microbiol 3:987–993.

 Table 1 Most represented bacterial phyla (class and order) on diverse algal species

Algae Phyla	Algae specie	Sample region	Sample season	Most bacterial represented phylum (and/or class)	Source
Phaeophyta	Fucus vesiculosus	Kiel fjord (Western Baltic Sea, Germany)	Winter Summer	Bacteroidetes, Planctomycetes, Proteobacteria (Alpha and Gamma) Bacteroidetes, Cyanobacteria, Proteobacteria (Alpha (Rhodobacterales)), Verrucomicrobia	(Lachnit et al. 2011)
	Fucus vesiculosus	Kiel fjord (Western	Winter	Actinobacteria, Bacteroidetes (Flavobacteria), Firmicutes (Bacilli)	(Goecke et
		Baltic Sea, Germany)	Summer	Bacteroidetes (Flavobacteria), Firmicutes (Bacilli), Proteobacteria (Gamma)	al. 2013)
	Dictyota bartayresiana	Island of Curacao (Netherlands Antilles)	(Not specified)	Bacteroidetes, Cyanobacteria, Proteobacteria	(Barott et al. 2011)
	Laminaria digitata	Roscoff (France)	(Not specified)	Actinobacteria, Bacteroidetes, Proteobacteria (Alpha and Gamma)	(Salaün et al. 2010)
	Laminaria saccharina	Kiel fjord (Western Baltic Sea, Germany)	Summer	Actinobacteria, Bacteroidetes, Firmicutes, Proteobacteria (Alpha, Beta and Gamma)	(Wiese et al. 2009)
	Laminaria hyperborea	Southwest coast of	Summer	Plantomycetes, Proteobacteria (Alpha, Beta and Gamma), Verrucomicrobia	(Bengtsson et
		Norway	Winter	Bacteroidetes, Cyanobacteria, Plantomycetes, Proteobacteria (Alpha, Beta and Gamma), Verrucomicrobia	al. 2010; Bengtsson and Øvreås 2010)
Rodophyta	Gracileria	Kiel fjord (Western	Winter	Bacteroidetes, Proteobacteria (Alpha (Rhodobacterales and Rhizobiales))	(Lachnit et
,	vermiculophylla	Baltic Sea, Germany)	Summer	Bacteroidetes, Proteobacteria (Alpha (Rhodobacterales))	
	Jania rubens	Cap Zebib (northern coast of Tunisia)	Summer	Bacteroidetes, Proteobacteria (Alpha and Gamma)	(Ismail-Ben Ali et al. 2011)
	Delisea pulchra	Bare Island (Sydney, Australia)	Summer	Bacteroidetes, Proteobacteria (Alpha), Planctomycetes	(Fernandes et al. 2012)
	Delesseria sanguinea	Kiel fjord (Western Baltic Sea, Germany)	Winter Summer	Actinobacteria, Bacteroidetes (Flavobacteria), Firmicutes (Bacilli), Proteobacteria (Gamma) Bacteroidetes (Flavobacteria), Firmicutes (Bacilli), Proteobacteria (Gamma)	(Goecke et al. 2013)
Chlorophyta	Ulva intestinalis	Kiel fjord (Western	Winter	Bacteroidetes, Proteobacteria (Alpha (Rhizobiales) and Gamma)	(Lachnit et
		Baltic Sea, Germany)	Summer	Bacteroidetes, Proteobacteria (Alpha (Rhodobacterales and Rhizobiales) and Gamma)	
	Ulva sp.	Wembury Beach, Devon, UK	(Not specified)	Bacteroidetes (Flavobacteria), Proteobacteria (Alpha (Rhodobacterales))	al. 2011) (Tait et al. 2009)
	Ulva australis	Bare Island (Sydney, Australia)	Winter	Bacteroidetes, Planctomycetes, Proteobacteria (Alpha and Gamma)	(Burke et al. 2011b; Tujula et al. 2010)
	Bryopsis hypnoides Bryopsis pennata	Pacific Mexican coast	Winter	Bacteroidetes (Flavobacteria and unclassified), Mollicutes (Mycoplasmataceae), Bacteroidetes (Flavobacteria)	(Hollants et al. 2011)
	Caulerpa taxifolia	Mediterranean Tahiti Philippines Australia	(Not specified)	Cytophaga-Flexibacter-Bacteroides (CFB), Proteobacteria (Alpha and Beta) Proteobacteria (Alpha and Delta) Proteobacteria (Alpha ,Delta and Gamma) Cytophaga-Flexibacter-Bacteroides (CFB), Proteobacteria (Alpha and Beta)	

Table 2 Census of the algal-specific polysaccharidases from seaweed-associated bacteria. Only enzymes characterized at the molecular and biochemical level have been considered. Some bacterial sequences from other environments have been added when they constitute representative enzymatic activities or 3D structure.

Protein	CAZY	Bacterial species	Associated algal species (or isolation habitat) B= browα-1,3-(3,6-Anhydro)-	Genbank	PDB	References
			L-galactosidases			
			n alga, G= green alga, R= Red alga			
Carrageenases						
к-carrageenase PcCgkA	GH16	Pseudoaltermonas carrageenovora ATCC 43555	seawater	CAA50624.1	1DYP	(Barbeyron et al. 1994 Michel et al. 2001a)
к-carrageenase	GH16	Pseudoalteromonas porphyrae LL1	Decayed seaweed	ADD92366.1	-	(Liu et al. 2011)
к-carrageenase ZgCgkA	GH16	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ94309.1	-	(Barbeyron et al. 1998)
к-carrageenase	GH16	Zobellia sp. ZM-2	Decayed seaweed	AGS43006.1	-	(Liu et al. 2013)
ι-carrageenase <i>Af</i> CgiA	GH82	"Alteromonas fortis"ATCC 43554	Seawater	CAC07801.1	1H80 1KTW 3LMW	(Barbeyron et al. 2000 Michel et al. 2001b Michel et al. 2003 Rebuffet et al. 2010)
t-carrageenase ZgCgiA1	GH82	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAC07822.1	-	(Barbeyron et al. 2000)
ι-carrageenase ZgCgiA2	GH82	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ96312.1	-	(Rebuffet et al. 2010)
1-carrageenase ZgCgiA3	GH82	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CBW46642.1	-	(Rebuffet et al. 2010)
1-carrageenase CgiA	GH82	Cellulophaga sp. QY3	Grateloupia livida (R)	AEV89930.1	-	(Ma et al. 2013)
t-carrageenase CgiB	GH82	Cellulophaga sp. QY3	Grateloupia livida (R)	AGN70890.1	-	(Ma et al. 2013)
λ -carrageenase Pc CglA	GHnc	Pseudoaltermonas carrageenovora ATCC 43555	seawater	CAL37005.1	=	(Guibet et al. 2007)
λ-carrageenase CglA	GHnc	Pseudoalteromonas sp. CL19	Deep-sea sediment	BAF35571.1	-	(Ohta et al. 2006)
Agarases						
α-agarase AaAgaA	GH96	"Alteromonas agarlytica" DSM 12513	Seawater	AAF26838.1	-	(Potin et al. 1993) Flament et al. 2007)
α-agarase Aga33	GH96	Thalassomonas agarivorans JAMB A33	Deep-sea sediment	BAF44076.1	-	(Hatada et al. 2006)
β-agarase ZgAgaA	GH16	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ98338.1	104Y 1URX	(Allouch et al. 2003 Allouch et al. 2004 Jam et al. 2005)
β -agarase Zg AgaB	GH16	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	AAF21821.1	1O4Z 4ATF	(Allouch et al. 2003 Jam et al. 2005 Hehemann et al. 2012a)
β-agarase Zg AgaD	GH16	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ98378.1	4ASM	(Hehemann et al. 2012a)
β-agarase AagA	GH16	Pseudoalteromonas gracilis B9	Gracilaria gracilis (R)	AAF03246.1	-	(Schroeder et al. 2003)
β-agarase	GH16	Pseudoalteromonas sp. AG4	Chondrus crispus (R)	ADD60418.1	-	(Oh et al. 2010)
β-agarase AgaYT	GH16	Flammeovirga yaeyamensis YT	Gracilaria tenuistipitata (R)	AEK80424.1	-	(Yang et al. 2011)
β-agarase AgaG1	GH16	Alteromonas sp. GNUM-1	Sargassum serratifolium (B)	AGW43026.1	-	(Kim et al. 2012)
exo-β-agarase AgaD	GH50	Saccharophagus degradans 2-40	Spartina alterniflora	ABD81904.1	4BQ2	(Pluvinage et al. 2013a)

			(salt marsh plant)		4BQ3 4BQ4	
					4BQ5	
β-agarase II (AgrA)	GH86	Pseudoalteromonas atlantica T6c	Palmaria palmata (R)	ABG40858.1	-	(Belas 1986)
β-agarase AgaC	GH118	Vibrio sp. PO-303	seawater	BAF03590.1	-	(Dong et al. 2006)
β-agarase AgaB	GH118	Pseudoalteromonas sp. CY24	seawater	AAQ56237.1	-	(Ma et al. 2007)
Porphyranases						
β -porphyranase Zg PorA	GH16	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CBM41182.1	3ILF 4ATE	(Hehemann et al. 2010)
β-porphyranase ZgPorB	GH16	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ95074.1	3JUU	(Hehemann et al. 2010)
β -porphyranase Bp GH16B	GH16	Bacteroides plebeius DSM 17135	Japanese gut microbiota	EDY95423.1	4AWD	(Hehemann et al. 2013)
β-porphyranase <i>Bp</i> GH86A	GH86	Bacteroides plebeius DSM 17135	Japanese gut microbiota	EDY95427.1	4AW7	(Hehemann et al. 2013)
a-1,3-(3,6-anhydro)-L-galactosida	ises	•				·
ZgAhgA	GH117	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CBM41465.1	3P2N	(Rebuffet et al. 2011)
<i>Sd</i> NABH	GH117	Saccharophagus degradans 2-40	Spartina alterniflora (salt marsh plant)	ABD81917.1	3R4Y 3R4Z	(Ha et al. 2011)
<i>Bp</i> GH117	GH117	Bacteroides plebeius DSM 17135	Japanese gut microbiota	EDY95405.1	4AK5 4AK6 4AK7	(Hehemann et al. 2012b)
Laminarinases						
Algal laminarin-specific β-	GH16	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ96583.1	4BOW	(Labourel et al. 2014)
glucanase ZgLamA		·	,		4BPZ	,
					4BQ1	
Fucoidanases						
Fucoidanase MfFcnA	GH107	Marineflexile fucanivorans SW5	Alginate-extraction factory	CAI47003.1	-	(Colin et al. 2006)
Alginate lyases						
Alginate lyase ZgAlyA4	PL6	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ98265.1	-	(Thomas et al. 2012)
endo-guluronate lyase ZgAlyA1	PL7	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ95239.1	3ZPY	(Thomas et al. 2012) (Thomas et al. 2013)
exo-alginate lyase ZgAlyA5	PL7	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ98266.1	4BE3	(Thomas et al. 2012)
		·				(Thomas et al. 2013)
poly-mannuronate lyase AlxM	PL7	Photobacterium sp. ATCC 43367	Sargassum fluitans (B)	CAA49630.1	-	(Mallisard et al. 1993)
alginate lyase Aly1	PL7	Streptomyces sp. ALG-5	green seaweed (G)	AAP47162.1	-	(Kim et al 2009)
alginate lyase AlyVI	PL7	Vibrio sp. QY101	Laminaria sp. (B)	AAP45155.1	-	(Han et al. 2004)
Alginate lyase ZgAlyA7	PL14	Zobellia galactanivorans Dsij	Delesseria sanguinea (R)	CAZ98462.1	-	(Thomas et al. 2012)
poly-MG alginate lyase alyPEEC	PL18	Pseudoalteromonas elyakovii IAM14594	Laminaria sp. (B)	AAD16034.1	-	(Sawabe et al. 2001)
alginate lyase Aly-SJ02	PL18	Pseudoalteromonas sp. SM0524	Kelp (B)	ACB87607.1	-	(Li et al. 2011)
Ulvan lyases						
ulvan lyase	PLnc	Persicivirga ulvanivorans PLR	Feces of <i>Aplysia punctata</i> (sea hare) feed with <i>Ulva</i> sp. (G)	AEN28574.1	-	(Nyvall-Collen et al. 2011)

Table 3 Percentages and genera of bacterial strains exhibiting antimicrobial activities, isolated on seaweeds.

Algae phyla	Algae species	% of isolated strains with antimicrobial activities (total isolated strains)	Bacteria genera identified with antimicrobial activities	Source
Phaeophyta	Laminaria saccharina	49% (210)	Bacillus, Glaciecola, Kopriimonas, Mesorhizobium, Pseudoalteromonas, , Streptomyces,	(Wiese et al. 2009)
	Pelvetia canaliculata Fucus ceranoides	7 % (55) 13 % (45)	- Alteromonas, Pseudomonas	(Lemos et al. 1985)
	Fucus vesiculosus.	53% (69)	Bacillus, Parracoccus, Pseudomonas, Streptomyces	(Goecke et al. 2013)
	Sargassum serratifolium Sargassum fusiforme Sargassum filicinum Padina arborescens Undaria pinnatifida Petalonia fascia Colpomenia sinuosa Scytosiphon lomentaria Ecklonia cava	20% (116)	Bacillus	(Kanagasabhapathy et al. 2006)
Rodophyta	Delesseria sanguinea	51% (97)	Algoriphagus, Microbacterium, Paenibacillus, Pseudoalteromonas, Streptomyces, Zobellia	(Goecke et al. 2013)
	Jania rubens	36 % (19)	Aquamarina, Bacillus, Paracoccus, Pseudoalteromonas, Pseudomonas	(Ismail-Ben Ali et al. 2011)
	Pachymeniopsis lauceola Plocamium telfairiae Gelidium amansii Chondrus oncellatus Grateloupia filicina 33% (92) Ceramium kondoi Lomentaria catenata Schizymenia dubyi Porphyra yezoensis		Bacillus, Microbacterium, Psychrobacter, Vibrio	(Kanagasabhapathy et al. 2008)
	Delisea pulchra	12% (325)	Micrococcus, Phaeobacter, Pseudoaltermonas, Rhodobacteraceae, Roseobacter, Ruegeri, Schwenalla, Vibrio	(Penesyan et al. 2009)
Chlorophyta	Ulva australis	12% (325)	Bacillus, Flavobacteriaceae, Phaeobacter, Photobacterium, Roseobacter	(Penesyan et al. 2009)
	Enteromorpha intestinalis Enteromorpha compressa Ulva lactuca	34 % (46) 18% (33) 13 % (45)	Alteromonas, Pseudomonas	(Lemos et al. 1985)

Table 4 Industrial applications using algal polysaccharides.

Industrial domain	Industrial use	Algal polysaccharides	Source
Pharmacy	Laxative	Agars	(Bixler and Porse
	Toothpaste	Carrageenans	2010; Jiao et al. 2011;
	Therapeutic peptides	Carrageenans, Fucoidans	Jung et al. 2013; Li et
	Pharamaceutical tablet desintegrant	Alginates	al. 2008; McHugh
	Medical fiber	Alginates	2003)
	Wound dressing	Alginates	
	Controlled release of medical drugs and other chemicals	Alginates	
	Dietary food	Alginates, Carrageenans	
Food industry	Gelling properties	Agars, Carrageenans	(McHugh 2003)
	Stabilizer	Agars, Alginates	
	Thickener	Agars, Alginates	
	Meat substitute	Agars	
	Wine clarification	Agars	
	Prevent pulp precipitation in fruit juices	Alginates	
	Conservation of frozen fish	Alginates	
Laboratory	Bacterial growth	Agar	(McHugh 2003)
Phytopharmacie	Activate signal pathway in plants and enhance their immune system	Alginates, Carrageenans, Fucans,	(Vera et al. 2011)
	Activate signal pathway in plants and emiance their infinitine system	Laminarin, Ulvan	
Paper industry	Smooth paper	Alginates	(McHugh 2003)
Textile industry	thickeners for the paste containing the dye	Alginates	(McHugh 2003)
Biofuel	Diagon highthanal highytanal	Alainata Laminanin Manaital	(Jung et al. 2013;
	Biogas, bioethanol, biobutanol	Alginate, Laminarin, Mannitol	Wargacki et al. 2012)
Other industry	Welding rod	Alginates	(McHugh 2003)
•	Immobilized biocatalyst	Alginates	,

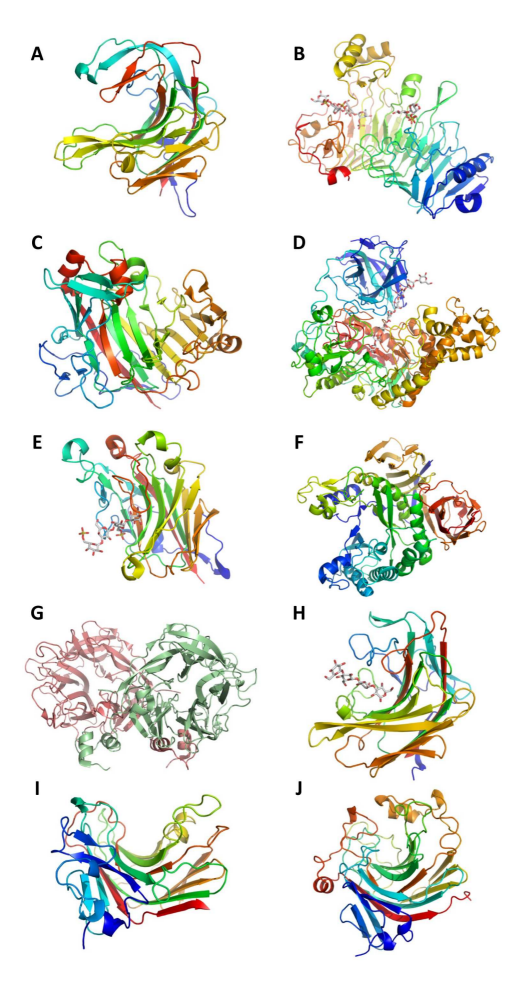


Fig. 1 Representative crystal structures of algal specific-polysaccharidases.

Structure of the GH16 kappa-carrageenase from *Pseudoalteromonas carrageenovora* (A, PDB 1DYP), of the GH82 iota-carrageenase from *Alteromonas fortis* in complex with oligo-iota-carrageenans (B, PDB 1KTW), of the GH16 beta-agarase ZgAgaD from *Zobellia galactanivorans* (C, PDB 4ASM), of the GH50 exo-beta-agarase Aga50D from *Saccharophagus degradans* in complex with an oligo-agar (D, PDB 4BQ5), of the GH16 beta-porphyranase ZgPorA from *Z. galactanivorans* in complex with an oligo-porphyran (E, PDB 3ILF), of the GH86 beta-porphyranase BpGH86A from *Bacteroides plebius* (F, PDB 4AW7), of the GH117 α -1,3-(3,6-anhydro)-L-galactosidase ZgAhgA from *Z. galactanivorans* (G, PDB 3P2N), of the GH16 laminarinase ZgLamA from *Z. galactanivorans* (I, PDB 3ZPY), and the PL7 exo alginate lyase ZgAlyA5 from *Z. galactanivorans* (I, PDB 3ZPY), and the PL7 exo alginate lyase ZgAlyA5 from *Z. galactanivorans* (J, PDB 3ZPY). The β -strands and the α -helices are represented by arrows and ribbons, respectively. The oligosaccharides are displayed with a stick representation. With the exception of ZgAhgA, all the structures are colored with a rainbow spectrum from the N- (blue) to the C-terminus (red). Chains A and B of ZgAhgA are colored in pink and green, respectively. This figure was prepared with the program Pymol.