THE MICROBIOME OF COASTAL SEDIMENTS

Graham J. C. Underwood, gjcu@essex.ac.uk ORCID 0000-0001-5605-0697

Alex J. Dumbrell, adumb@essex.ac.uk ORCID 0000-0001-6282-3043

Terry J. McGenity, tjmcgen@essex.ac.uk ORCID 0000-0002-1497-8822

Boyd A.McKew, boyd.mckew@essex.ac.uk . ORCID 0000-0001-5607-6619

Corinne Whitby, cwhitby@essex.ac.uk ORCID 0000-0003-1998-2748

School of Life Sciences, University of Essex, Colchester, Essex. U.K. CO4 3SQ.

ABSTRACT

Coastal zones are amongst the most productive marine environments and many are highly impacted by anthropogenic activity. Coastal zones are key regions for the transformation of land-based inputs of nutrients and pollutants and provide many essential ecosystem services for human society. Periods of tidal exposure and submergence, coupled with seasonal variation in land-based inputs, result in intertidal habitats characterized by highly variable environmental conditions that pose crucial adaptive challenges for organisms. This review focusses on the microbiome of coastal sediments consisting of protists (especially diatoms), bacteria, archaea and fungi. The diversity, distribution, production, adaptations, and interactions between these groups are reviewed. Coastal microbiomes are characterized by high rates of biogeochemical activity. Photoautotrophic diatoms exhibit complex patterns of behavior to cope with a highly variable light climate. Multiple species-species interactions between autotrophs and heterotrophs contribute to the cycling of carbon and nitrogen. In sediments, autotrophic and heterotrophic processes are closely coupled both spatially and temporally. Bacteria and archaea control the nitrogen- and carbon cycles while taxonomic diversity is influenced by gradients of organic matter, nitrogen compounds, sulfide, and oxygen. Fungi are important components of coastal salt marsh sediment microbiomes but their role in unvegetated sediments is less well understood. This review considers the high human impact on coastal sediments and the importance of nutrient gradients and pollution pressures (hydrocarbons) in affecting diversity and species distribution.

1. Introduction

Coastal zones are amongst the most productive marine environments. Located at the interface between marine, freshwater, and terrestrial environments and receiving inputs from both the open ocean and from the land coastal zones consist of a matrix of diverse habitats positioned along various physical, chemical, and biological gradients. Organisms living within transitional coastal zones have to be adapted to major gradients of conditions that can be subject to seasonal variability. In addition to seasonal changes (particularly present at temperate- and polar latitudes), there is a strong influence of the lunar tidal cycle (bi-weekly) resulting in periods of varying length of aerial exposure and saline water submergence in intertidal environments. These exposure cycles can result in important changes in environmental conditions on an hourly basis. Thus, the coastal zone is characterized by highly variable environmental conditions that pose considerable adaptive challenges for organisms living within it. Despite these challenges, coastal habitats support characteristic microbiomes (defined as a characteristic microbial community with distinct physiological and chemical properties and activities resulting in the formation of specific ecological niches, Berg et al. 2020) that underpin the ecological functioning of these habitats. Coastal microbiomes play crucial roles in biogeochemical cycling, food webs, and habitat modification, resulting in the provision of important ecosystem services to human society.

The coastal zone encompasses a wide range of habitats. Rocky shores are extensive worldwide and are generally characterized by steep spatial gradients from land to sea. Rocky shores host abundant communities of macroscopic organisms but their microbiology is less well described (Maggi et al. 2017). The impervious nature of rocky shores means that the influence of microbes present as thin epilithic biofilms are strongly affected by cycles of desiccation, extreme salinity fluctuations, macroalgal spore settlement and germination, and grazing by macroinvertebrates. This chapter does not focus on these environments and readers are referred to Dal Bello et al. (2017) and Notman et al. (2016).

Estuaries and intertidal flats are amongst the most productive of the various coastal ecosystems and provide important ecosystem services such as the provisioning of food resources, water purification, carbon storage, and coastal storm surge tidal defense (Waltham et al. 2020). Estuaries are also some of the most human-modified environments because of historic and current concentration of human populations and industries along their coastlines (Van Niekerk et al. 2013; Henderson et al. 2020). This has resulted in a considerable degradation of estuarine ecosystems, which changed the ecological processes that govern

their health and ecosystem services (Duarte et al. 2020; Van Niekerk et al. 2019; Waltham et al 2020). Coastal habitats (salt marsh, mangrove, seagrass, muddy and sandy intertidal flats) are important zones of nutrient cycling (Nedwell et al. 2016) and valuable sites of organic carbon generation and accumulation ("blue carbon", the carbon stored in the sediment, living and non-living above- and below-ground biomass of salt marsh and seagrass habitats (Alongi 2020; Beaumont et al. 2014; Burdon et al. 2019; Legge et al. 2020; Waltham et al. 2020). Because coastal environments are located in the transition zone between the land and the sea they are particularly susceptible to pollution, including excess nutrients (particularly inorganic N and P), heavy metals, pesticides, pharmaceuticals, numerous industrial persistent organic pollutants and plastics. One of the major types of organic pollutants in coastal ecosystems are petroleum hydrocarbons from crude oil and its many refined products.

Estuarine sedimentary systems are extensively distributed across the globe. All estuaries have their own characteristics influenced by the local geology and catchment features and the local tidal range, from microtidal (<2 m tidal range), mesotidal (2-4 m tidal range), and macrotidal (>4 m tidal range). Tidal range and wind and wave-climate are major factors influencing the geomorphology of a coastal-estuarine system. A typical meso- or macro-tidal estuary is usually characterized by a well-mixed salinity gradient from freshwater to fully marine with fine-grained sediment and mudflats in the sheltered regions of the estuary towards its head and mixed- and sandy sediments towards the mouth where tides and winddriven waves and currents are stronger (Baas et al. 2019; Green and Coco 2014; Zhu et al. 2020). In agricultural and populous catchments most nutrient loading is land derived. Hence, an estuarine gradient reflects co-varying conditions of increasing salinity, decreasing nutrient loading, increasing sediment particle size, and varying levels of tidal exposure (Nedwell et al. 2016). Approximately perpendicular to this linear gradient is the gradient of tidal exposure with upper shores often colonized by vascular macrophytes (salt marshes in temperate regions and mangroves in the tropics, Alongi 2020), and by sand dune habitats on winddominated sandy shores (Galiforni-Silva et al. 2020). Mid-tide level shores tend to be dominated by micro- and macro-algal mats, and the lower shores more physically disturbed, but also colonized by biogenetic reefs of bivalves or polychaete worms. These environmental gradients, the large surface area provided by sediment particles, and accounting for areal and depth dimensions, results in an extensive mosaic of habitats that support productive and diverse microbiomes (Heip et al. 1995; Luna et al. 2013; Underwood and Kromkamp 1999).

2. Coastal autotrophic microbiomes: microphytobenthic biofilms.

On intertidal mud and sand flats and in shallow subtidal systems where sunlight reaches the sediment surface diverse and abundant microbial biofilms occur. Collectively these assemblages are termed microphytobenthos (MPB) or benthic microalgae (BMA) biofilms, terms which emphasize the important role played by the photoautotrophic components of these complex agglomerates of autotrophic and heterotrophic protists, bacteria, archaea, and fungi (An et al. 2020; Chen et al. 2017; Cibic et al. 2019; Pinckney 2018; Sahan et al 2007; Underwood and Kromkamp 1999). The photoautotrophic diatoms (Stramenopiles, Bacillariophyceae) are major components of most MPB or BMA biofilms with net primary production of 29 - 314 g C m⁻² y⁻¹ (Pinckney 2018; Underwood and Krompkamp, 1999). The primary production of MPB provides the main energy resource to biofilm consumers (protozoans and metazoans), and their predators (Christianen et al. 2017; Green et al. 2012; Herman et al. 2000; Hope et al. 2020) while heterotrophic bacteria and archaea are the primary remineralizers of MPB-derived organic material, including volatile compounds, and detrital organic matter present in the sediment (Acuña Alvarez et al. 2009; Bohórquez et al. 2017; Gaubert-Boussarie et al. 2020; Luna et al. 2013; Nedwell et al. 2016).

Two types of microphytobenthic biofilms are recognized: transient microbial biofilms that form and reform over daily and weekly timescales and more permanent microbial mats. Microbial mats are characterized by higher biomass and are usually dominated by cyanobacteria. Microbial mats show long—term temporal persistence (months to years) such that a macroscopic structure is formed, and they are often closed systems with much internal recycling of nutrients (Long et al. 2013; Stal et al. 2019). Stromatolites are a particular type of microbial mat that possess a laminated calcified structure, which is considered to be the outcome of an intense coupling between microbial (cyanobacteria, heterotrophic bacteria, archaea, eukarya) and geochemical processes leading to a remnant geological formation. A specialized type of a coastal microbial mat is the supratidal microbialite. These microbialites have been found in the supratidal zone of rocky shores in South Africa, Australia, and the U.K., where there is a freshwater input (Rishworth et al. 2020). Microbial mats and stromatolite microbiomes have been reviewed by Stal (2016), Stal et al. (2019) and Rishworth et al. (2020) and are not considered further here.

2.1 Diversity of microphytobenthos in coastal sediments

Transient marine benthic biofilms have a high potential species richness of photoautotrophs. Although both photosynthetic and heterotrophic microeukaryotes such as flagellates and ciliates are present (Chen et al. 2017; Gong et al. 2015; Massana et al. 2015),

their ecology and importance in intertidal biofilm is in most cases unresolved. A few genera of cyanobacteria (e.g. Lyngbya, Oscillatoria) and motile euglenophytes (e.g. Euglena deses, E. proxima) are found and are often in high abundance in transient biofilms (Bellinger et al. 2005; Kingston 1999; Perkins et al. 2002; Underwood et al. 2005) (Table 1). However, the dominant group of MPB in terms of biomass and activity are benthic diatoms with well over 1,500 benthic diatom (morpho)species described from different geographical regions (Witkowski et al. 2000). Within a particular environment however, especially on estuarine intertidal mudflats it is more usual to find only a few (20+) species that are numerically dominant within MPB assemblages (Forster et al. 2006; Park et al. 2014; Redzuan and Underwood 2020; Redzuan and Underwood 2021; Ribeiro et al. 2013; Ribeiro et al. 2020; Sahan et al. 2007; Thornton et al. 2002; Underwood 1994; Underwood and Barnett 2006). The majority of the literature on benthic diatom diversity relies on microscopy-based identification and a morphology-based taxonomy, an approach which is time-consuming and requires a high level of expertise (Ribeiro et al. 2020). Where detailed studies have been conducted it is clear that deterministic (niche-based) factors rather than neutral factors determine the community composition of the abundant species (Plante et al. 2016; Plante et al. 2020; Thornton et al. 2002). Taxonomic composition of the dominant components in the microbiome is strongly influenced by sediment particle size (the balance of sands, silts, and clays) selecting for a range of epipelic (mud inhabiting), highly motile biraphid epipsammic (attached to sand grains), less motile mono- or biraphid, or araphid diatom taxa (Table 1) (Hamels et al. 1988; Sabbe 1993; Underwood and Barnett 2006). The distribution of sediment types corresponds to gradients of physical energy, salinity, and water flow, with sands present in the more exposed marine sediments and clays and silts settling in the more sheltered, low-energy, upper reaches of estuaries, often exposed to a greater range of salinity conditions over tidal and seasonal cycles (Baas et al. 2019; Green and Coco 2014). These gradients are major factors that determine both alpha- and beta diversity in MPB microbiomes (Gong et al. 2015; Park et al. 2014; Plante et al. 2016; Plante et al. 2020; Ribeiro et al. 2013; Witkowski et al. 2000).

Salinity within an estuarine gradient and position on the shore, which relates to the degree of tidal exposure and period of subtidal disturbance, are important controls on diatom species distribution (Forster et al. 2006; Oppenheim 1991; Peletier 1996; Ribeiro et al. 2003; Ribeiro et al. 2013; Sahan et al. 2007; Thornton et al. 2002; Underwood 1994; Underwood et al. 1998). These physical factors vary with seasonal changes in irradiance, thermal stress, and

winter mixing and storminess, and alter the species composition of estuarine benthic diatom communities (Oppenheim 1991; Underwood 1994; Underwood 2005). For example, species such as *Fallacia pygmaea* and *Navicula salinarum* are found in cold and warm months, respectively (Admiraal et al. 1984). Inorganic nutrient concentrations are a strong driver of dominant taxa. They correlate with microphytobenthic biomass and species composition on spatial and temporal scales (Thornton et al. 2002; Underwood et al. 1998).

Many patterns of species distribution in estuaries are based on correlative field surveys. Because of covarying gradients, especially of exposure at low tide, sediment type, salinity, and nutrient concentrations, it is not clear how much variability in community composition occurs in the absence of changes in nutrients (or by changes in other variables such as sediment particle size distribution) and over what time scales such variability operates. Experimental manipulations have shown that nutrients are a selective force in determining species composition (Sullivan et al. 1999; Underwood et al., 1998, Underwood and Provot 2000). High concentrations of ammonium and sulfide (often due to sewage inputs or organic enrichment) can be inhibitory or selective for particularly resistant taxa (Admiraal 1984). Significant decrease in Chl a and changes in species composition in the Ems Dollard estuary occurred between 1979 and 1993 after the lowering of organic waste input from local potato starch industries onto adjacent mudflats (Peletier 1996). Small spatial scale (10 – 100 m) patterns in biomass and species composition were documented in the Colne Estuary (Thornton et al. 2002; Underwood et al. 1998), which have been experimentally demonstrated to relate to species-specific preferences (Underwood and Provot 2000) and tolerance to sulfide and anoxia (McKew et al. 2013).

High throughput sequencing (HTS) methodologies have been applied to the coastal benthic eukaryotic microbiome in the last decade. The results of these analyses have in general supported the conclusions of microscope-based approaches about patterns of richness, distribution of species, and correlation with environmental variables. However, HTS report higher operational taxonomic unit (OTU) richness than that recognized by morphological taxonomic approaches. Chen et al (2017) found 6,627 benthic microeukaryotic OTUs (18S rRNA gene, 97% sequence similarity cut-off) at Xiamen Island, China; Plante et al. (2020) identified 4,411 different diatom OTUs (18S rRNA gene, sequence similarity threshold of 98%) in South Carolina sediments, while An et al. (2020) reported 9,582 diatom OTU (*rbcL* gene, 98% similarity cut off) on Korean intertidal mudflats. Chen et al. (2017) found strong deterministic control of the abundant and

conditionally rare taxa (total phosphorus, total nitrogen, salinity, phosphate, and total oxidized nitrogen for the dominant taxa) in benthic microbial eukaryote communities but rare species showed no spatial, environmental, or distance-decay pattern. Comparisons of the diatoms present on mudflat-salt marsh transitions on a number of barrier island sites in South Carolina, USA, revealed that 95% of all OTUs were rare, less than 0.1% of total sequence count and that a few key taxa (e.g. *Navicula* and *Gyrosigma* species) were dominant (Plante et al. 2020). Spatial effects (dispersal limitation) and spatially-structured environmental factors affected these dominant taxa caused the significant differences in beta diversity between island sites. The presence of planktonic or tychoplankton taxa, e.g. *Thalassiosira* was also recorded in benthic samples (Plante et al. 2020). Within single geographical sites factors such as physical disturbance and sediment type (sands, muds) determined the dominance patterns of a few key taxa but with greater neutral or stochastic process elements influencing the composition of the different patches (Plante et al. 2016).

High throughput sequencing provides a much greater resolution of the richness of the eukaryotic members of the sediment microbiome than revealed by morphological microscopic approaches. This higher richness is due in part to a higher intensity of sampling (greater sample sizes) and because of multiple copies of target genes within single cells, which varies between taxa (Gong and Marchetti 2019). But it could also be caused by cryptic diversity within some groups of diatoms. Vanelslander et al. (2009) found that the common estuarine diatom morphospecies Navicula phyllepta was in fact two difference species separated spatially along the salinity gradient of the Westerscheldt estuary. Clonal cultures of Nitzschia inconspicua isolated from across a range of freshwater, brackish, and marine habitats in the River Ebro were paraphyletic with six different genotypes and a range of different reproductive strategies and salinity tolerances (Rovira et al. 2015). Growth of two clonal cultures of Cylindrotheca closterium, isolated from the oligosaline and mesosaline regions of the Colne estuary, showed different growth optima to salinity and nitrogen gradients (Underwood and Provot 2000). Morphological approaches with their much lower sample sizes (usually 200-400 individuals) cannot sample as many rare taxa as HTS and cannot resolve cryptic diversity. Moreover, rare valves require greater identification skills and may just be reported as unidentified by the untrained eye. These factors hamper the understanding of community assembly processes for microphytobenthos. Comparison of morphological species lists from spatially-distant sites, especially when studied by different research teams, is difficult because of the judgement required for many morphological species. HTS data lends itself to larger spatial analyses (Clark et al. 2017) and avoids individuals' decisions on species attribution. However, ecological interpretation of HTS datasets also relies on accurate reference databases and consideration of the issues of numbers of copies of phylogenetically informative genes when translating to relative abundance (Gong and Marchetti 2019). There is a need for better taxonomic alignment in the libraries used to assign HTS DNA sequence data to coastal marine morphological species in order to minimize the risk of misidentification and to ensure that the latest taxonomic phylogenies are reported as is being done for freshwater diatoms (Perez Burillo et al. 2020). This is particular necessary in order to understand the role of the many rare taxa that HTS detects in coastal microbiomes. Are these OTUs representing cryptic diversity within recognized species complexes or are these new, unknown, species or false identifications made by the original sequence depositors? These challenges need addressing to understand the rules for community assembly in these coastal microbiomes.

Intertidal diatom-rich biofilms exhibit a positive relationship between assemblage biodiversity (species richness and Shannon diversity of abundant taxa) and net primary production (Forster et al. 2006). Experimental studies with cultured benthic diatoms show niche complementarity and transgressive over-yielding (increased biovolumes) for mixtures of up to eight species combinations but antagonistic interactions were present between some taxa (Koedooder et al. 2019; Vanelslander et al. 2009). Facilitation, possibly through mixotrophic growth on of organic substrates produced by other diatoms, or by associated hetertrophic bacteria, was shown for Cylindrotheca closterium, which grew strongly in the spent media from Navicula cultures (Vanelslander et al. 2009). The nature of the competitive interactions between species is not well described. Nitzschia c.f. pellucida releases cyanogen bromide immediately after the onset of light. Cyanogen bromide is toxic to other diatom species that thrive in the immediate vicinity and cause their death (Vanelslander et al. 2012). There is generally a negative relationship between biofilm biomass and diatom diversity in mudflat biofilms with lower shore beach sediments exposed to a greater level of disturbance having a higher diversity and a more even species distribution than upper and mid tide sites that support high biomass (Forster et al. 2006; Hill-Spanik et al. 2019; Underwood 1994). At times, conditions can be favorable for the rapid growth of just a few taxa or even a single species may "bloom" resulting in a biofilm with low species diversity and high biomass (Forster et al. 2006; Underwood 1994; Underwood et al. 1998). These studies were conducted in high nutrient status estuaries with traditional microscopic assessments of composition. In lower nutrient environments a positive relationship between biofilm biomass and Shannon diversity (determined by HTS) has been found (Plante et al. 2020). When comparing biomass-diversity relationships between sediment types, epipsammic habitats can have both lower biomass and lower alpha-diversity than mudflat sites (Plante et al. 2016). This illustrates how more detailed studies on the causes and patterns of benthic diatom assemblage composition and functioning are needed.

2.2 Adaptations of photoautotrophs to living in intertidal sediments

The consequence of periods of tidal exposure for photosynthetic microorganisms living on intertidal sediments is that they can experience high incident radiation (including UVB) at varying times over the day and over a year (Laviale et al. 2015; Mouget et al. 2008; Waring et al. 2007). Additionally, sediment disturbance and mobilization by waves and tidal flows mixes cells out of the photic zone of the sediment surface (De Jonge and van Beusekom 1995; Redzuan and Underwood 2020; Savelli et al. 2019). Autotrophic microphytobenthos has adapted to these environmental pressures of high irradiation and sediment disturbance in various ways. Motility is a key adaptation for MPB taxa living in mud and silty sediments allowing them to (re)position at locatons with a favorable light climate within the sediment. Motile MPB is primarily composed of pennate diatoms (motile taxa found on mud are termed epipelon) and euglenophytes, as well as some taxa of filamentous cyanobacteria (Underwood and Kromkamp 1999). Upon tidal exposure the populations of phototrophic organisms undergo mass vertical migration, which brings cells to the surface (Consalvey et al. 2004; Jesus et al. 2009) in order to be able to photosynthesize. This vertical migration has been recognized for over a century (Perkins 1960) and is a macroscale feature visible to the naked eye as a color change of the sediment and is even detectable by remote sensing (Méléder et al. 2020; Savelli et al. 2020).

The motility of MPB exhibits a number of features of ecological relevance. There is an underlying endogenous rhythm of motility, which is maintained for a number of days in the absence of light or tidal stimuli. This rhythmicity is detectable in terms of changing cell density at the sediment surface, intensity of photosynthetic pigments, and rate of photosynthesis and carbohydrate production (Coelho et al. 2011; Haro et al. 2019; Perkins 1960; Round et al. 1966; Serôdio et al. 1997; Smith and Underwood 1998). These patterns of rhythmicity is evidence for a circadian rhythm and circadian rhythm regulator genes, such as *kaiA*, *kaiB*, *kaiC*, and peroxiredoxin (*prx*) are expressed by cyanobacteria and diatoms in microbial mats (Hörnlein et al. 2018). Circadian rhythms of activity have also been found in

the expression patterns of conserved gene transcripts for photosynthesis (PSI and PSII) and CO₂ fixation (RuBisCo) of cyanobacteria and diatoms in intertidal cyanobacterial mats (Hörnlein et al. 2018). Circadian rhythms can be entrained by external stimuli. MPB shows entrainment of rhythmicity by tidal exposure cycles as well as by day-night light cycles (Haro et al. 2019; Hörnlein et al. 2018; Perkins 1960; Serôdio et al 1997). Haro et al. (2019) found that endogenous rhythms of migration and net community primary production were lost after exposure to continuous light for 3 days but were reset by re-imposition of an alternating light regime. Although easily demonstrated and reported quite some time ago (Perkins 1960; Round et al. 1966) the mechanism by which MPB maintains its endogenous vertical migration rhythms in synchrony with the progressive daily movement of the tidal exposure window (Happey-Wood and Jones 1988) is not yet resolved.

Taxon-specific differences are present within the mass movement of the whole community during tidal emersion-immersion cycles. Some taxa only appear at the sediment surface for short periods of the exposure period (e.g. Round and Palmer 1966; Round 1979; Underwood et al. 2005) or have a specific movement in response to light stimuli (*Gyrosigma balticum*, Jönnson et al. (1994)). Barranguet et al. (1998) proposed that cells of different species micro-migrated into and out of the surface photic zone of the biofilm during illumination thus avoiding photo-inhibition while maintaining an overall high assemblage photosynthesis. Single cell imaging of intact biofilms provided evidence for this (Oxborough et al. 2000; Underwood et al. 2005), correlating time and light intensity with the species present at the sediment surface during tidal exposure.

Microphytobenthos shows strong behavioral responses to light intensity with populations migrating down into sediment to avoid high light (Perkins et al. 2002; Perkins et al. 2010; Prins et al. 2020; Underwood 2002). MPB can also detect spectral composition with high and low intensities of blue and red light generating different patterns of surface active biomass and photo-acclimation in diatom-rich mudflat biofilms (Prins et al. 2020). Benthic diatoms can also sense UVB radiation and will move away from the surface even when PAR light intensities are constant (Waring et al. 2007). Benthic diatoms respond to light through a combination of positive and negative phototaxis (directional movement) and photokinesis (changing speed) (Cohn et al. 2016). There are differences in response between diatom species. *Nitzschia perminuta* exhibits positive phototaxis at low levels of blue light and negative phototaxis at high intensities as well as a photokinetic response to red light. However, under the same conditions *Cylindrotheca closterium* only displayed photokinetic

responses to red light and no blue light response (McLachlan et al. 2009). Craticula cuspidata, Stauroneis phoenicenteron, and Pinnularia viridis have different positive and negative photophobic (changing direction of movement) responses to red, green, and blue light (Cohn et al. 2016). Light sensing appears to take place at the apices of the cell valves and is therefore not necessarily directly associated with the chloroplast (McLachlan et al. 2012; Cohn et al. 2016). High intensity blue light at the apex of N. perminuta causes an increase in intracellular calcium concentration along the line of the raphe in the cell wall followed by a reversal of the direction of movement (McLachlan et al. 2012). Diatoms possess genes that code for phytochromes, cryptochromes, aureochromes, and other light receptor proteins for harvesting red/far red and blue light (Blommaert et al. 2020; König et al. 2017; Mann et al. 2020). Benthic diatoms in sediment experience a gradient of the light spectrum with red light most rapidly attenuated and blue light penetrating deepest (Lassen et al. 1992). Differential motility behavior modulated by spectral quality would allow cells to position themselves in a favorable light climate within the narrow photic zone of intertidal sediments.

Benthic diatoms are photo-physiologically flexible and are able to use rapid photochemical and non-photochemical quenching (NPQ), the xanthophyll cycle (XC), and longer term acclimation of Chl *a* and other photopigments in order to maintain high rates of primary production in a rapidly varying light climate over tidal emersion and during the year (Barnett et al. 2015; Juneau et al. 2015; Prins et al. 2020; Underwood et al. 2005; Waring et al. 2010). The ability to dissipate light energy through NPQ and XC is particularly important for non-motile species that cannot migrate away from damaging light conditions. Experimental work has demonstrated differences in the ability of diatom species to induce high levels of NPQ and XC (Barnett et al. 2015; Blommaert et al. 2018). Epipsammic diatoms, which are found attached to sand grains, have high capacity for non-photochemical quenching (NPQ), while epipelic diatoms have lower potential NPQ, and non-motile tychoplankton, which live under low light conditions in frequently-mixed and resuspended sediments, also possess a low capacity for NPQ (Barnett et al. 2015; Blommaert et al. 2018).

2.3 Distribution of MPB biomass in coastal sediments

The biomass of MPB present at any location and time is a consequence of the physical and environmental conditions of the preceding period. MPB shows rapid growth responses and can increase their biomass over a period of a few days when conditions are conducive to growth (Nedwell et al. 2016). Nutrients, light availability, and sediment type are

major controls of biomass (Cibic et al. 2019; Underwood and Kromkamp 1999), while grazing and desiccation (McKew et al. 2011, Savelli et al. 2018), physical disturbance by waves, wind, and tides (de Jonge and van Beusekom 1995; Redzuan and Underwood 2020; Redzuan and Underwood 2021; Savelli et al. 2018), and macrofauna (birds) (Booty et al. 2020) may have local impacts.

Combined, these environmental factors produce general patterns of higher biomass on upper intertidal flats and in sheltered regions of estuaries (Daggers et al. 2020; Underwood and Kromkamp 1999). Seasonal patterns are variable between locations. In temperate zones biomass may peak at any time throughout the year, though summer peaks are common. However, summer declines have also been reported as the result of grazing or temperature stress (Daggers et al. 2020; Nedwell et al. 2016; Park et al. 2014; Savelli et al. 2018; Underwood and Paterson 1993). This seasonal variability reflects in part the spatial patchiness of MPB, which occurs on a range of scales from cm to km (Redzuan and Underwood 2021; Spilmont et al. 2011; Taylor et al. 2013; Weerman et al. 2012). Only a few long-term (>3 years) data sets of MPB biomass exist, and these indicate that sediment type, exposure or tidal position, windiness, and, to a lesser extent, air temperature are the main drivers of biomass. De Jonge et al. (2012) found similar inter-annual patterns of biomass at different stations in the Ems estuary (Netherlands) and a long-term positive relationship between biomass and annual air temperatures with higher Chl a content during the 1990s during the monitoring period from 1976 to 1999. Van der Wal et al. (2010) used remote sensing data to determine MPB biomass on mudflats in the southern North Sea over the period 2001 to 2009 and found broad synchrony in the patterns of occurrence and biomass between estuaries, although stronger relationships were present within regional data sets (e.g. within Dutch estuaries). Weather and summer temperatures strongly influenced MPB biomass in the Loire estuary (France) from 1993 to 1998 and from 2006 to 2010 (Benyoucef et al. 2014).

2.4 Interactions between photoautotrophs and chemoheterotrophs and the turnover of organic carbon in coastal microbiomes

Photosynthetic activity by MPB produces oxygen and a variety of labile carbon compounds. MPB, particularly diatoms, produces extracellular polymeric substances (EPS) as well as low molecular weight labile carbon compounds (Bellinger et al. 2005; 2009; Underwood and Paterson 2003). EPS molecules are important in creating a biofilm matrix that increases sediment stability (Baas et al. 2019; Hope et al. 2020) and provide protection

from desiccation and salinity stress (Steele et al. 2014). The production of these molecules is variable and is moderated by environmental factors (e.g. light and nutrients, Staats et al. 2000; Underwood and Paterson 2003) and rhythms of vertical migration (Hanlon et al. 2006; Perkins et al. 2001; Smith and Underwood 1998). These environmental factors drive a distinct seasonality in the balance of labile and recalcitrant exudates produced over a year (Moerdijk et al. 2018a; 2018b). There is evidence from freshwater studies that EPS production and biofilm formation by diatoms is enhanced by the presence of certain bacterial taxa (Bruckner et al. 2008; Bruckner et al. 2011; Grossart et al. 2005). These interactions appear to be species-specific. Bacterial-diatom interactions have negative effects on estuarine diatom biomass in cultures with a single diatom species but are neutral in co-cultures of different diatom species (Koedooder et al. 2019). Different benthic diatom taxa promoted the growth of assemblages of sediment bacteria. For example, Seminavis robusta cultures supported Alphaproteobacteria Thalassospira the sp., Roseobacter Kordiimonadaceae sp. and the Bacteroidetes Mangrovimonas sp. and Owenweeksia sp., while monocultures of the diatoms Cylindrotheca closterium and Navicula phyllepta had different bacterial assemblage profiles (Koedooder et al. 2019). There is evidence of antagonistic interactions between the bacterial assemblage associated with a certain diatom species with other benthic diatom taxa (Stock et al. 2019). This suggests that different diatom species may have their own associated bacterial microbiome.

Microphytobenthos-derived dissolved organic carbon compounds contribute 30 to 50% of the total organic matter in the sediments (Bellinger et al. 2009) and represent the key source of labile organic carbon (Nedwell et al. 2016). The importance of this carbon source varies in different intertidal habitats (sandy to muddy; temperate to tropical) (Cook et al. 2007; Oakes et al. 2010; 2012). MPB-fixed carbon has a characteristic ^{δ13}C signal that can be tracked through food webs (Christianen et al. 2017). EPS ¹³C-carbon has been tracked into the phospholipid fatty acids (PLFA) and RNA of various bacterial groups (Bellinger et al. 2009; Gihring et al. 2009; Middelburg et al. 2000; Taylor et al. 2013). Major utilizers of diatom EPS in aerobic sediments are Alphaproteobacteria, Gammaproteobacteria and Bacteriodetes, and in anaerobic conditions Deltaproteobacteria (Bohórquez et al. 2017; McKew et al. 2013; Miyatake et al. 2014; Taylor et al. 2013). A subset of Alphaproteobacteria and Gammaproteobacterial taxa was adapted to utilize diatom EPS before it became available to the rest of the bacterial assemblage (Taylor et al. 2013). Different bacterial groups (for Sphingobacteria Tenacibaculum (Bacteroidetes), example and two classes of

Verrucomicrobiae and Opitutae)) grow preferentially on labile and refractory diatom EPS (Bohórquez et al. 2017; Underwood et al. 2019). Turnover rates of these different DOC fractions vary under aerobic and anaerobic conditions. Anaerobic conditions lead to preferential breakdown of refractory compounds and enhance the growth of Firmicutes (Clostridia, Lachnospiraceae, Peptostreptococcaceae, Ruminococcaceae and unclassified Clostridiales) other and sulfate-reducing Deltaproteobacteria (Desulfobacteraceae and Desulfobulbaceae) (McKew et al. 2013). There are close linkages between photoautotrophic and chemoheterotrophic microorganisms present in the coastal sediment microbiome and evidence of antagonistic, synergistic, and mutualistic interactions. Hörnlein et al. (2019) proposed the *Choirmaster-Choir* theory. This theory predicts that the rhythmic release of photosynthate and other metabolites is controlled by the circadian clock of the photoautotrophic members of the microbiome (cyanobacteria, diatoms) and dictates the genetic clocks of other microbes either directly or in association with external Zeitgebers such as light and temperature, which results in a synchronized activity during a 24-h cycle. This idea remains to be further explored.

3. Nitrogen cycling in the marine coastal microbiome

The dominant heterotrophic bacteria found in aerobic coastal sediments are Actinobacteria, Alphaproteobacteria, Gammaproteobacteria, Chloroflexi, Verrumicrobiae, and Bacteriodetes (Bohórquez et al. 2017; McKew et al. 2013; Yao et al. 2019; Yi et al. 2020). The relative abundance of the different groups is strongly influenced by sediment characteristics particularly by sediment grain size, organic content, pH, and nitrogen and phosphorous availability (Yao et al. 2019; Yi et al. 2020). In estuarine environments, salinity and freshwater inputs influence taxonomic composition, with Actinobacteria and Betaproteobacteria more abundant in lower salinity zones or during periods of higher rainfall. For example, Actinobacteria, Chloroflexi, and Verrucomicrobia showed significant differences between rainfall seasons in the Yangtze estuary (Yi et al. 2020). However, overall salinity-related changes in assemblages appear to be more pronounced in estuarine bacterioplankton assemblages (Gołębiewski et al. 2017, Osterholz et al. 2018) than in estuarine benthic assemblages where organic matter and sediment properties are most influential (McKew et al. 2013; Yao et al. 2019; Yi et al. 2020). Marine coastal sediment microbiomes exhibit profound depth profiles of the distribution of taxonomic groups of bacteria reflecting the gradient of electron acceptors (Böer et al. 2009; Webster et al. 2010; Wilms et al. 2006). In muds with high organic carbon content, high rates of bacterial activity,

and limited diffusion of oxygen, the anaerobic zone can be as near as a few millimetres below the surface and sometimes even reaches the surface. In the anaerobic zone the processing of organic carbon is largely driven by sulfate reduction and nitrogen cycling with Delta- and Epsilonproteobacteria, including sulfate-reducing bacteria, e.g. Desulfomonadales, as well as Archaea as prominent players (McKew et al. 2013; Nedwell et al. 2016; Webster et al. 2010).

The nitrogen (N) cycle (Fig. 1) is mediated by metabolically diverse groups of microorganisms. The location of the different processes in the sediment is determined by sediment redox state. Microbial-driven N transformations are especially crucial in coastal systems, which often receive high anthropogenic N inputs (e.g. via fluvial discharges) resulting in organic matter breakdown and oxygen depletion (Nedwell et al. 2016). Characterizing N cycle communities in the environment by traditional microbiological methods has been problematic due to the difficulties in obtaining pure cultures of the responsible microorganisms. However, molecular methods and HTS techniques have enabled to uncover the functional nitrogen-cycle microbiome of these ecosystems.

3.1 Nitrogen cycling in aerobic coastal sediments: nitrification and aerobic ammonia oxidation and commamox.

Autotrophic ammonia oxidation is the rate-limiting step in nitrification and important in the N cycle (Fig. 1). During nitrification, aerobic ammonia-oxidizing bacteria (AOB) and archaea (AOA) oxidize ammonium by ammonia monooxygenase (encoded by *amoA*) (McTavish et al. 1993). With AOB, the second step is the dehydrogenation of hydroxylamine to nitrite by hydroxylamine oxidoreductase (encoded by *hao*) (Arp et al. 2002). However, AOA genome data does not appear to contain *hao* gene homologues, and an alternative mechanism has been proposed (Hallam et al. 2006). Nitrite is oxidized to nitrate by nitrite-oxidizing bacteria (NOB) (e.g. *Nitrospira*). Previously, based on 16S rRNA gene sequencing autotrophic ammonia oxidation was thought to be restricted to two monophyletic lineages of aerobic ammonia-oxidizing bacteria (AOB) (Head et al. 1993). The first lineage belongs to the Betaproteobacteria (Beta-AOB) (e.g. *Nitrosomonas, Nitrosospira*) and the second lineage belongs to the Gammaproteobacteria (Gamma-AOB) (e.g. *Nitrosococcus* sp.) (Head et al. 1993). However, metagenome libraries from seawater (Venter et al. 2004) and soil (Treusch et al. 2005) revealed putative genes involved in ammonia oxidation from uncultured Thaumarchaeota. AOA *amoA* gene sequences form five clusters four with cultured

representatives (*Nitrosocaldus*, *Nitrososphaera*, *Nitrosopumilus*, *Nitrosotalea*) and the fifth is known as '*Nitrososphaera* sister' cluster (Pester et al. 2011).

Aerobic ammonia-oxidizers are found in most environments (Table 1) (Francis et al. 2005; Jiang et al. 2009; Philips et al. 1999; Stehr et al. 1995; Whitby et al. 1999, 2001). In some ecosystems AOA outnumber AOB often by a factor of 10 to 1,000 (Beman et al. 2008; Wuchter et al. 2006). This is the case for example in North Sea coastal sediments (Lipsewers et al. 2014) and coastal waters (Smith et al. 2014a), suggesting a greater contribution of AOA to nitrification in these systems (Jiang et al. 2009). However, in some coastal and estuarine sediments AOB are more abundant than AOA (Caffrey et al. 2007). In a hypernutrified temperate estuary (Colne, U.K.) with gradients of salinity and ammonia concentration, benthic AOB (notably Nitrosomonas spp.) were significantly more abundant (by 100-fold) than AOA, suggesting that AOB were the main contributors to nitrification (Li et al. 2015a). Seasonal differences in nitrification in coastal sediments have been observed with the highest rates often in the summer (Li et al. 2015a). However, in North Sea coastal sediments AOA 16S rRNA gene transcriptional activity was higher in the winter despite the lower abundance of these organisms (Lipsewers et al. 2014; 2017). In contrast, higher AOA abundances were found in the winter in the North Sea water column, which was attributed to ammonia availability and the lack of competition for ammonia with phytoplankton (Pitcher et al. 2011; Wuchter et al. 2006). Differences in spatial distribution between Nitrosospira and Nitrosomonas have also been found. For example, in freshwater lake sediments, N. europaea was present in littoral sediments whilst N. eutropha was found in profundal sediments, whilst members of Nitrosospira were ubiquitous (Whitby et al. 1999; 2001); and in the water column of the Mediterranean Sea, where different members of the beta-proteobacterial ammonia-oxidizers were associated with particulate material and planktonic samples (Phillips et al. 1999).

The oxidation of ammonia via nitrite to nitrate was originally considered to be a two-step process catalyzed by two functionally distinct groups of chemolithoautotrophs (ammonia-oxidizers and nitrite oxidizers). However, a nitrifying bacterium belonging to *Nitrospira* was discovered and sequencing of its genome revealed that it has all the genes necessary for the oxidation of ammonia and nitrite (Daims et al. 2015). The discovery of the complete oxidation of ammonia to nitrate in one organism (comammox) (Fig. 1) has changed the paradigm that this process requires two distinct functional groups of microbes and raises questions about the role of comammox *Nitrospira* in N-cycling.

Comammox organisms belong as far as known to the *Nitrospira* lineage II (Daims et al. 2016; Koch et al. 2019). Based on *amoA* gene sequences from metagenomes comammox bacteria comprise two clades, A and B (Daims et al. 2016; Palomo et al. 2018; Van Kessel et al. 2015). Putative comammox (clade A) *amoA* gene sequences were previously misidentified as "unusual" methanotroph *pmoA* genes relating to *Crenothrix* in the Gammaproteobacteria (Stoecker et al. 2006) or presumed to belong to a methanotroph from Alphaproteobacteria (clade B, *amoA*) (Radajewski et al. 2002). Clade A is further delineated in two groups: clades A.1 and A.2 (Xia et al. 2018). Because comammox bacteria do not form a monophyletic group within *Nitrospira*, lineage II comammox and canonical nitrite-oxidizing *Nitrospira* cannot be distinguished by 16S rRNA-based methods (Pjevac et al. 2017). Comammox bacteria have been found in various habitats (Table 2) with high proportions in estuarine and coastal environments (Xia et al. 2018). In the open ocean, however, comammox *amoA* genes were either rarely detected (Daims et al. 2015) or absent (Xia et al. 2018).

3.2 Environmental factors influencing nitrification and ammonia oxidation

Although AOA and AOB co-exist there is evidence of niche differentiation linked to various environmental factors (e.g. temperature, ammonium concentration, oxygen, pH, salinity, light, macrofaunal activity) (Caffrey et al. 2007; Cao et al. 2011; Dang et al. 2010; Erguder et al. 2009; Scarlett et al. 2020; Stehr et al. 1995). Ammonium concentration and availability are major factors for niche partitioning of AOA versus AOB with lower concentrations generally favoring AOA (Clark et al. 2020; Martens-Habbena et al. 2009). In some coastal sediments low phosphate availability selects for AOA over AOB (Lipsewers et al. 2014). In subsurface sediments *Nitrosomonas* dominated and was linked to nitrite concentration (Cao et al. 2012). In estuarine sediments decreased dissolved oxygen altered AOB *amoA* expression but not AOA (Abell et al. 2010). Phytoplankton may also outcompete nitrifiers for substrates in surface waters (Smith et al., 2014b). In estuarine sediments benthic microalgae have a high demand for ammonium (Thornton et al. 1999) and can outcompete AOB, reducing the rates of nitrification (Risgaard-Petersen 2003).

Differential sensitivity to pollutants between AOB and AOA in coastal environments has also been found with higher Beta-AOB diversity in polluted sites whilst AOA were unaffected (Cao et al. 2011). Agriculturally-impacted estuarine sediments were dominated by AOA and *Nitrosomonas* spp. *amoA* sequences whilst *Nitrosospira* spp. dominated less impacted sites (Wankel et al. 2010). Distinct clusters of *Nitrosomonas* and *Nitrosospira* lineages have been found in eutrophic coastal sediments subjected to inputs from nearby

wastewater treatment plants and polluted rivers (Dang et al. 2010). Silver nanoparticles inhibited AOB-driven nitrification but not AOA in a temperate eutrophic estuary (Beddow et al. 2017). Addition of titanium nanoparticles resulted in increased ammonium fluxes from sediments into overlying water, which could be due to lower rates of ammonia oxidation and nitrification as well as a decrease of net MPB primary productivity (Passarelli et al. 2020). Beta-AOB *N. europaea* and *N. communis* lineages also thrive in heavy metal-polluted environments and in environments with high ammonium concentrations (Dang et al. 2010; Stein et al. 2007). In estuaries *Nitrosospira*-like lineages appear to be better adapted than *Nitrosomonas* (Cao et al. 2011) and Beta-AOB (particularly the *N. oligotropha* lineage) could be used as bioindicators of pollution in coastal systems (Dang et al. 2010).

Comammox bacteria are functionally versatile and adaptative to many environments (Hu and He 2017). Comammox bacteria exhibit niche partitioning influenced by various environmental factors (Shi et al. 2020) and differences in abundance among clades have been found (Xia et al. 2018). Co-occurrences of comammox with canonical ammonia oxidizers indicates a potential functional differentiation between these groups (Bartelme et al. 2017; Palomo et al. 2018; Pjevac et al. 2017) and may depend on whether the main activity of comammox in an environment is ammonia oxidation or nitrite oxidation (Xia et al. 2018). Comammox bacteria may outnumber AOB (Xia et al., 2018) and can functionally outcompete other canonical nitrifiers in highly oligotrophic systems (Hu and He 2017). However, which factors drive niche specialization between comammox and canonical ammonia oxidizers currently remains unknown.

3.3 Nitrogen cycling in anaerobic coastal sediments: anammox, denitrification and dissimilatory reduction of nitrate to ammonium

Anaerobic ammonia oxidation (anammox) involves the conversion of ammonium and nitrite to N_2 in the absence of oxygen (Fig. 1). Some anammox bacteria are facultative chemoorganotrophs that can also metabolize organic compounds notably formate, acetate, and propionate (Kartal et al. 2007; Strous et al. 2006), allowing anammox bacteria to adopt a 'disguised' denitrifying lifestyle (Kartal et al. 2007). Anammox bacteria form a monophyletic order of the Brocadiales within the Planctomycetes (Jetten et al. 2010), and consist of five candidate genera: *Candidatus* Kuenenia (Strous et al. 2006); *Candidatus* Brocadia (Oshiki et al. 2011; Strous et al. 1999); *Candidatus* Anammoxoglobus (Kartal et al. 2007); *Candidatus* Jettenia (Quan et al. 2008), and *Candidatus* Scalindua (Schmid et al. 2003).

Anammox bacteria are found in virtually any anoxic environment that contains fixed N (Table 2). Although anammox is responsible for a large proportion of N₂ production in marine sediments, in eutrophic coastal sediments, and saline tidal marsh sediments, anammox is not important relative to denitrification (Koop-Jakobsen and Giblin 2010; Thamdrup and Dalsgaard 2002). Although anammox bacteria have been found in coastal and estuarine sediments (Li et al. 2011a; Tal et al. 2005; Trimmer et al. 2003) and coastal mangrove wetlands (Cao et al 2011; Li et al. 2011a), greater anammox bacterial diversity occurs in the Oxygen Minimum Zones (OMZs) of oceans (Woebken et al. 2009). Anammox bacteria are abundant and active in oxygenated upper sediments and bioturbated marine coastal sediments in the North Sea (Lipsewers et al. 2014). High anammox bacterial abundances have also been found in surface sediments of hypernutrified estuarine tidal flats (Zhang et al. 2013). In some environments anammox bacteria are scarce like suboxic and anoxic aquatic systems where low anammox bacterial diversity was found and comprised mostly *Scalindua* (Penton et al. 2006).

Denitrification is fundamental in the N cycle releasing nitric oxide (NO), nitrous oxide (N₂O), and dinitrogen gas (N₂) to the atmosphere (Fig. 1). As denitrifying bacteria belong to different phylogenetic groups the 16S rRNA gene is not very useful for analyzing denitrifier communities. Instead, functional genes involved in denitrification have been targeted e.g. *napA*, *narG* (nitrate reductase), *nirS*, *nirK* (nitrite reductases), and *nosZ* (nitrous oxide reductase) (Nogales et al. 2002) (Fig. 1). Denitrifiers are facultative organoheterotrophic anaerobes that constitute a phylogenetically diverse group spanning >50 different genera (Jones and Hallin, 2010; Zumft, 1997). Most denitrifiers belong to the alpha, beta-, gamma-, and epsilon-Proteobacteria (Braker and Conrad, 2011). The most frequently isolated denitrifying bacteria belong to the Pseudomonads (Herbert, 1999). Denitrification has also been found among Firmicutes, Actinomycetes, Bacteroidetes, Aquificaceae and Archaea (Braker and Conrad, 2011). Denitrification is also widespread among Foraminifera, *Gromiida* (Piña-Ochoa et al. 2010; Risgaard-Petersen et al. 2006) and fungi (Braker and Conrad, 2011).

Denitrification is widely distributed in the environment (Table 2). In the ocean however, denitrification is geographically restricted to a few oceanic regions (e.g. OMZs and hemipelagic sediments) (Jayakumur et al. 2009) and distinct *nirS* and *nirK* populations have been found within the oxygen-deficient zone in marine sediments (Liu et al. 2003). In eutrophic estuaries denitrification can mediate the lowering of N load and contribute to

eutrophication control (Nogales et al. 2002). Indeed, in coastal and estuarine sediments denitrification can remove >50% of inorganic N inputs from terrestrial systems (Nedwell et al. 2016; Rivera-Monroy et al. 2010; Seitzinger, 1988). In coastal and estuarine sediments denitrification rates are generally higher than in shallower waters (Herbert, 1999).

In addition to denitrification microbial nitrate reduction may also take place via alternative pathways. Dissimilatory nitrate reduction to ammonium (DNRA) (Fig. 1) is particularly important in organic-rich sediments (King and Nedwell, 1987; Laverman et al. 2006) and tends to retain bioavailable N in aquatic ecosystems. DNRA is common in bacteria (e.g. Proteobacteria, Firmicutes, Verrucomicrobia, Planctomycetes, Acidobacteria, Chloroflexi, Beggiatoa, Thioploca, and Chlorobia) (Papaspyrou et al. 2014; Preisler et al. 2007), and also occurs in eukaryotes (e.g. diatoms, fungi) (Pajares and Ramos, 2019). Anammox bacteria may also perform DNRA in the presence of small organic compounds (Kartal et al. 2007) or ammonium might be released from fermentative reactions (Herbert, 1999; Lam et al. 2009). DNRA is commonly found in environments low in oxygen, such as OMZs (Lam et al. 2009) and sediments with steep oxygen gradients (Kamp et al. 2011). DNRA has also been found in the Namibian inner-shelf bottom waters (Kartal et al. 2007) and deep-sea sediments (Pajares and Ramos, 2019).

3.4 Environmental factors influencing the anaerobic nitrogen cycling biome

Anammox is controlled by several environmental factors including salinity (Sonthiphand et al. 2014), temperature (Qian et al. 2018), organic matter content (Trimmer and Engström 2011), and inorganic N availability (Trimmer et al. 2005). Interactions between AOA, AOB, and anammox bacteria have been shown where nitrifiers supply nitrite to anammox (Lam et al. 2007; 2009). In mangrove sediments, positive correlations occur with AOA diversity and abundance and anammox *hzo* gene abundances (Li et al. 2011a,b; 2013), suggesting that complex interactions exist between anammox bacteria and ammonia oxidizers. Sulfide may also inhibit anammox bacteria (Dalsgaard et al. 2003; Jensen et al. 2008).

Nitrogen removal via denitrification may cause a decrease in N availability, which in coastal environments can severely impact primary producers and levels of eutrophication (Seitzinger, 1988; Herbert 1999). Numerous environmental factors (e.g. N availability and concentration, temperature, oxygen concentration, water depth, organic matter quality and quantity, bioturbation), affect denitrifier distribution and abundance (Braker et al. 2000; Dang

et al. 2009; Liu et al. 2003; Prokopenko et al. 2011; Zhang et al. 2014). Denitrification rates also show distinct seasonal patterns driven largely by temperature, nitrate, and availability of organic carbon (Kaplan et al. 1977). Denitrification rates decrease in the spring (in estuarine sediments) (Jørgensen and Sorensen 1988) and in the summer (in subtropical macrotidal estuarine sediments) where *nirS:nirK* ratios are negatively correlated with temperature (Abell et al. 2010). Nitrate concentration and oxygen have an impact on denitrifying communities (Liu et al. 2003) and nitrate availability drives *nirS* communities whilst *nirK* communities respond to other parameters (Jones and Hallin, 2010). To date, the ecological function of these denitrifying communities and the factors that determine the composition of *nirS / nirK* communities remains unknown (Jones and Hallin, 2010). Sulfide also decreases denitrification rates (Porubsky et al. 2009). Yet, paradoxically in sulfidic sediments some microorganisms use sulfide as an electron donor for denitrification (Bowles et al. 2012). Bioturbated sediments from large burrowing macrofauna also increase coupled nitrification-denitrification (Laverock et al. 2011; Papaspyrou et al. 2014).

Seasonal and spatial differences in DNRA have been found with increased rates in the summer throughout sediment depths compared to other times when activity was restricted to deeper sediments (Jørgensen 1989). In intertidal and sub-tidal environments DNRA may change on a daily basis due to the growth and photosynthetic activity of benthic microalgae. Photosynthetically evolved oxygen diffuses into the surface of the sediment during daylight which inhibits DNRA (Herbert, 1999). MPB photosynthesis can decrease the rate of denitrification of nitrate that diffuses into the sediment from the water column (Dw) but stimulates the rate of coupled nitrification-denitrification (Dong et al. 2000; Risgaard-Petersen 2003). In estuaries high *nrfA* gene abundances (encoding cytochrome c nitrite reductase) have been found and change along gradients of salinity and nitrate (Papaspyrou et al. 2014).

3.5 Nitrogen fixation in coastal sediments

Biological nitrogen fixation involves specialized groups of autotrophic and heterotrophic bacteria and archaea that possess molybdenum (Mo)—Fe protein (dinitrogenase) (encoded by *nifDK*) and Fe protein (dinitrogenase reductase) (encoded by *nifH*) (Fig. 1). Oxygen exposure deactivates nitrogenase and oxygenic phototrophs must separate dinitrogen fixation from oxygenic photosynthesis either spatially (e.g. in heterocysts) or temporally (Berman-Frank et al. 2003).

Nitrogen-fixing organisms (diazotrophs) are a diverse group of bacteria and archaea that include members of the Chromatiaceae, Chlorobiaceae, Chloroflexaceae, Rhodospirillaceae, and chemoautotrophic bacteria and archaea (Bergman et al 1997; Capone, 1988; Raymond et al. 2004). Marine diazotrophs mainly include non-heterocystous, heterocystous, symbiotic, and unicellular cyanobacteria (e.g. *Ca.* Atelocyanobacterium thalassa [UCYN-A]; *Crocosphaera watsonii* [UCYN-B] and *Cyanothece* [UCYN-C]) (Capone, 1988; Martinez-Perez et al. 2016; Pajares and Ramos 2019). Other marine diazotrophs include heterotrophic bacteria (e.g. *Klebsiella*), anoxygenic phototrophic bacteria (e.g. *Chlorobium, Chromatium*), strict anaerobic chemotrophs (e.g. *Clostridium, Desulfovibrio*), methanogenic Euryarchaeota and Planctomycetes (Pajares and Ramos 2019). Nitrogen-fixing eukaryotes are not known and it seems that these organisms solved the problem by entering in symbiosis with nitrogen-fixing bacteria (Kuypers et al. 2018).

The main factors that affect marine diazotroph distribution are oxygen, light, temperature, inorganic N, phosphorus, iron, and organic matter (Pajares and Ramos 2019). In estuaries and coastal regions UCYN-A are highly abundant (Moreira-Coello et al. 2019) along with heterotrophic bacteria (Pajares and Ramos 2019). Several factors influence nitrogen fixation activity in benthic sediments including carbon availability, temperature, light, pH, oxygen, inorganic N, salinity, and trace metal availability (Herbert, 1999). Organic carbon availability is generally the main factor limiting the nitrogen fixation in unvegetated sediments (Herbert, 1999). In unvegetated shallow coastal lagoons and intertidal sediments where light is not limiting dense communities of benthic nitrogen-fixing cyanobacteria may occur (Herbert, 1999; Stal et al. 2016; Stal et al. 2019). In tropical coastal marine lagoons sediment nitrogen fixation contributes 11% of the annual N input (Hanson and Gundersen 1977) and high rates occur in temperate sediments, mudflats, and salt marshes, especially in organically rich sediments (Nedwell and Aziz 1980; Herbert, 1999). Cyanobacterial mats (both temperate and tropical) exhibit high nitrogen fixation rates linked to dark-light cycles and are under the control of circadian clocks (Herbert 1999; Hörnlein et al. 2018; Stal et al. 2016; Stal et al. 2019). High nitrogen fixation rates have been found in salt marsh sediments which has been attributed to organic compounds excreted from plant roots coupled to plant photosynthetic activity (Moriarty and O'Donohue 1993; Whiting et al. 1986) whilst rates in bare marine sediments were low (Herbert, 1999).

4. Archaea in marine sediment microbiomes

4.1 An array of coastal Archaea: Marine Group III (putative Pontarchaea), Asgard Archaea, Marine Benthic Group D, and Woesarchaeota

Archaea are an important component in the surface sediments of intertidal communities with an abundance of one to two orders of magnitude lower than bacteria (Li et al. 2012; McKew et al. 2011; Wang et al. 2020). Deeper in the sediment, e.g. in the sulfate-methane transition zone, they can be in equal abundance (Li et al. 2012; Wang et al. 2020). Until the 1990s the domain Archaea was divided into Euryarchaeota and Crenarchaeota but this view is rapidly changing (see Baker et al. 2020). Methanogens were the only Archaea in coastal environments that were well known. Advances in sequencing technology unveiled the uncultured archaeal diversity in coastal settings. The awareness of the presence of Archaea in non-extreme environments such as the open ocean was thanks to the pioneering work of Norman Pace, Ed DeLong, Jed Fuhrman and colleagues (DeLong 1992; Fuhrman et al. 1992; Pace 1997). Archaea were described from Colne Point salt marsh in Essex, U.K. by Munson et al. (1997) who detected 16S rRNA gene sequences of methanogens, haloarchaea and an archaeal lineage that was distinct from any known taxon. Subsequently, this unknown archaeal taxon was detected elsewhere, e.g. from the deep sea (Fuhrman and Davis 1997) and continental shelf samples (Vetriani et al. 1998). It became known as Marine Group III (MG-III) Euryarchaeota with a proposed phylum-level reassignment to Pontarchaea (Li et al. 2015b). Further surveys, using fosmid clones and metagenome assembled genomes (MAGs), revealed the distribution and putative functions of MG-III. For example, Haro-Moreno et al. (2017) showed that MG-III phylotypes living in the photic zone probably have a photoheterotrophic lifestyle, which they based on the presence of photolyase and rhodopsin genes as well as of genes for peptide and lipid uptake and degradation. It remains to be seen whether the coastal MG-III found by Munson et al. (1997) are similar to epipelagic or bathypelagic phylotypes (Haro-Moreno et al. 2017).

Kim et al. (2005) found MG-III Euryarchaeota in tidal flat sediments from Ganghwa Island, Korea, together with many sequences that were considered to be Crenarchaeota, which had not been detected in the Colne Estuary salt marshes by Munson et al. (1997). This phylogenetic lineage was referred to as Marine Benthic Group B (MBG-B) by Vetriani et al. (1998), a sister group of the Deep-Sea Archaeal Group (DSAG), which have been reclassified as members of the Asgard Archaea. Specifically, MBG-B are now known as Thorarchaeota (Seitz et al. 2016) and DSAG as Lokiarchaeota (Spang et al. 2015). Phylogenomic analysis places Eukaryotes within the archaea most closely related to the Asgard archaea, which

possess a range of eukaryote features. The classification of the Asgard archaea has contributed to redefining the tree of life from three domains into one with two-domains (Williams et al. 2020). Thorarchaeota (MBG-B) have been found in a number of different estuaries (Fig. 2; Zou et al. 2020a), including the Colne, Essex, U.K. (Webster et al. 2015). Thorarchaeota from White Oak River estuary, North Carolina, USA, have the genetic capacity for protein degradation with the formation of acetate as well as for the reduction of elemental sulfur and thiosulfate and therefore this group of organisms may play an important role in carbon and sulfur cycling in estuarine sediments (Seitz et al., 2016). Lokiarchaeaota in Namibian shelf sediments anaerobically consume necromass and extracellular polymeric substances from diatoms and also fixed CO₂ via the H₂-dependent Wood–Ljungdahl pathway (Orsi et al., 2019). This archaeal mixotrophic activity was more rapid than that of bacteria emphasizing that archaea should not be overlooked in sediment biogeochemical processes (Orsi et al. 2019).

Another archaeal group that is commonly found in estuaries is the Marine Benthic Group D or MBG-D (also called DHVE1), which has been variously called Thermoprofundales or Izemarchaea (Baker et al. 2020; Zhou et al. 2019). MBG-D were the second most abundant archaea in the Paerl River Estuary, China, after the Bathyarchaeota (Wang et al. 2020; see also Zou et al. 2020b) and were also detected in sediments along the Colne Estuary, U.K. (Webster et al., 2015). MBG-D seem to be heterotrophic with the capacity for degrading proteins (Lloyd et al. 2013). In addition, the potential for mixotrophic growth was revealed upon the reconstruction of MBG-D genomes from a mangrove sediment (Zhou et al. 2019).

Woesaerchaeota, formerly known as DHVEG-6 (Liu et al., 2018a, b), are globally distributed in many environments including estuaries where they are usually a minor component of the microbial community but occasionally reach a high abundance (Fig. 2; Zou et al. 2020a). In *Zostera marina* seagrass beds and nearby bare sediment in Rongcheng Bay, Yellow Sea, China, Woesearchaeota (42% of Archaea) were the most abundant archaea followed by Bathyarchaeota (21%) and Thaumarchaeota (17%) with specific sub-clades of Woesearchaeota and Bathyarchaeota enriched in the vegetated areas (Zheng et al., 2019).

4.2 Bathyarchaeota (Miscellaneous Crenarchaeota Group) and Thaumarchaeota are generally the most abundant archaea in marine sediments

There are two other even more widely distributed and abundant examples of novel archaeal taxa in estuarine environments. The Marine Group I, which together with species

from terrestrial environments belongs to the Thaumarchaeota, consists predominantly of ammonia-oxidizing Archaea, which are discussed in section 3.1. The Marine Group I are abundant in estuarine sediments (Fig. 2) and dominated in a large-scale study of eastern Chinese marginal seas (Liu et al. 2020). The Miscellaneous Crenarchaeota Group (MCG), now known as Bathyarchaeota, is one of the most abundant phyla on Earth and generally the most abundant archaea in estuarine sediments (Fig. 2; Li et al. 2012). Bathyarchaeota have an anaerobic organoheterotrophic lifestyle (Seyler et al. 2014), probably degrading proteins (Lloyd et al. 2013), carbohydrates (Lazar et al. 2016), aromatic (Dong et al., 2019; Meng et al. 2014), and aliphatic (Dong et al. 2019) compounds, as well as a variety of other organic matter (Seyler et al. 2014). The aforementioned taxonomic groups and the Bathyarchaeota are phyla and hence comprise a variety of different microorganisms with an array of genetic and functional capacities, which will likely reflect their ecological distribution in estuarine sediments. Bathyarchaeota, for example, comprise 25 subclades (Zhou et al. 2018). Lazar et al. (2014) propose that the Bathy-6 in contrast to other lineages prefers suboxic sediment with minimal free sulfide. Bathy-6 also has the genetic capacity to take up and catabolize a wide range of carbohydrates and proteins (Lazar et al. 2016) and may be able to carry out dissimilatory nitrite reduction to ammonium (DNRA) (Lazar et al. 2016). By performing diverse enrichments from estuarine sediments Yu et al. (2018) showed that subclade Bathy-8 grew on lignin as an energy source. Then, by using lipid stable-isotope probing, these authors demonstrated that lignin-degrading cultures used bicarbonate as a carbon source. This organoautotrophic growth on an abundant biopolymer may partially explain its dominance particularly in estuarine sediments that receive input from plant debris (Yu et al. 2018).

Thus, a variety of different archaea contribute to the turnover of organic matter in coastal sediments while some of them may be autotrophic at the same time. There is a need to better understand the contribution of archaea to benthic cycling of carbon, sulfur, and nitrogen as well as their interactions with other organisms. Obtaining enriched or pure cultures, as was done for Lokiarchaeota (Imachi et al., 2020), will be necessary in order to understand the ecophysiology of archaea in coastal sediments.

4.3. Archaea drive the methane cycle in coastal sediments

Strictly anaerobic methane-producing archaea perform the final step in the anaerobic degradation of organic matter. Much is known about methanogens, primarily because many strains from different classes have been isolated and studied in detail. All characterized methanogens belong to the phylum Euryarchaeota and include: Methanobacteriales,

Methanococcales, Methanomicrobiales, Methanosarcinales, Methanopyrales, Methanocellales, and Methanomassiliicoccales (Lyu and Liu, 2018) and the class Methanonatronarchaeia (Sorokin et al. 2017; 2018). The first-described methanogens use the major products of microbial fermentation either hydrogen plus CO₂ (hydrogenotrophic) or acetate (acetoclastic) (Thauer et al., 2008). Fermentation and methanogenesis occur when energetically favorable electron acceptors such as oxygen, nitrate, and sulfate have been depleted such as is the case in deeper coastal sediments (Wilms et al. 2006; 2007). However, methanogenesis also occurs near the surface of coastal sediments, where sulfate reducers outcompete methanogens for hydrogen and acetate. Here, methanogens coexist with sulfatereducing bacteria by using non-competitive methylated substrates, such as methylamine (e.g. Oremland et al., 1982). Methylamines, methylsulfides, and other methylated compounds are common in marine and hypersaline environments as breakdown products of osmolytes (McGenity and Sorokin 2018) and also as components of lipid polar head groups, e.g. choline (Jameson et al. 2018). Methylotrophic methanogenesis could be distinguished into two mechanisms: (1) hydrogen-independent carried out by several representatives of the Methanosarcinales, and (2) hydrogen-dependent carried out by several other groups (Feldewert et al. 2020). Hydrogen-dependent methylotrophic methanogens also appear to compete with H₂-utilizing sulfate-reducing bacteria as long as the partial pressure of hydrogen is low and there is a supply of suitable C₁-compounds owing to their superior affinity for hydrogen (Feldewert et al. 2020).

A summary of the numerous investigations on coastal/estuarine methanogenesis is beyond the scope of this chapter. For a discussion of methanogenesis in the Colne Estuary see Nedwell et al. (2016). The Colne Estuary, U.K., is typical for many global estuaries as methane production occurs along the length of the estuary together with sulfate reduction but at a rate almost two orders of magnitude lower than sulfate reduction (Nedwell et al. 2004). In the Colne Estuary a change from acetoclastic and hydrogenotrophic taxa to methylotrophic (*Methanococcoides*) and versatile (*Methanosarcina*) taxa from the head to the mouth was observed (Webster et al., 2015). The salinity (and, hence, sulfate) gradient that characterizes estuaries, together with proximity to land and sea, are major reasons why these environments have a higher diversity of methanogens than other ecosystems (Wen et al., 2017). For example, *Methanoregula* is typically freshwater while *Methanococcoides* is typically marine but both are common in estuaries (Wen et al., 2017). An investigation on methanogens and methanogenesis in mangrove sediments showed that the dominant taxa were

Methanomicrobiales and Methanosarcinales together with putative hydrogen-dependent methyl-reducing methanogens *Candidatus* Methanofastidiosa and Methanomassiliicoccales, the latter exhibiting the highest activity (Zhang et al., 2020). Thus, even for a well-known archaeal process much has to be learned about the diversity of the responsible microbes.

It came as a surprise that methylotrophic methanogenesis may be a property of non-euryarchaeal candidate phyla within the archaea such as Verstraetearchaeota (Vanwonterghem et al. 2016) and Bathyarchaeota (Evans et al. 2015). This was based on the possession of *mcr* genes coding for methyl coenzyme M reductase (Mcr) complex, which catalyzes the terminal step of methanogenesis. Subsequently, *mcr* in *Candidatus* Syntrophoarchaeum was proposed to code for an enzyme involved in short-chain alkane oxidation and its sequence was similar to the *mcr* sequences from Verstraetearchaeota and Bathyarchaeota (Evans et al. 2019). Thus, it is supposed that these archaea are not methanogens but oxidize short-chain alkanes.

Consumption of short-chain alkanes may not be a common process in estuaries because propane and butane are not present in large amounts. However, anaerobic oxidation of methane produced by methanogens occurs ubiquitously especially in the sulfate methane transition zone (SMTZ) (Boetius et al. 2000; Hoehler et al. 1994), and is a near-quantitative sink for the methane produced (Egger et al. 2018). This process is carried out by polyphyletic groups of uncultured archaea, which are related to methanogenic Euryarchaeota and referred to as ANME (anaerobic methane oxidizers) (Evans et al. 2019; Knittel et al. 2018). The original mechanistic explanation for anaerobic oxidation of methane was that it occurs as a syntrophic process in which the ANME methanotrophs convert methane to hydrogen, which is consumed by associated bacteria most typically in marine sediments by sulfate-reducing bacteria (Boetius et al., 2000). However, the precise mechanisms of anaerobic oxidation of methane is debated in terms of: 1) the main interacting bacterial species and their terminal electron acceptors (e.g. sulfate, Fe III, Mn IV, and nitrate) 2) the internal metabolic processes in the ANME methanotrophs, which genetically resemble methanogens, and 3) the energetics and mode of exchange e.g. metabolite transfer or direct interspecies electron exchange (McGlynn 2017).

In estuaries, which overall are a methane source, more methane reaches the atmosphere as salinity decreases due to a combination of greater methane production and less effective anaerobic removal where sulfate reduction is lower (Dean et al. 2018). However, there remains much to be learned about the sources and sinks of methane in coastal

environments and particularly in estuaries. For example, in a brackish Baltic Sea estuary, anaerobic oxidation of methane was identified as an important process presumably coupled to iron III and manganese IV reduction (Myllykangas et al. 2020).

4.4 Haloarchaea are consistently present and locally abundant in coastal sediments

Extremely halophilic Euryarchaeota belonging to the Halobacteria (more commonly referred to as haloarchaea) dominate in coastal environments where seawater evaporates to create hypersaline conditions such as in sabkhas, hypersaline lagoons, as well as artificial salt pans (McGenity and Oren 2012). However, Munson et al. (1997) reported that haloarchaea were abundant in a temperate salt marsh. This observation led Purdy et al. (2004) to culture haloarchaea from creek or saltmarsh pan sediments, aerobically, over a range of salinities, with either glucose or glycerol as carbon and energy sources, and with antibiotics to inhibit growth of bacteria and eukarya. They isolated three taxa of haloarchaea one of which had strains that grew slowly at seawater salinity and optimally with 10% NaCl, a property that is unusual for haloarchaea (Purdy et al, 2004). Subsequently, haloarchaea have been found in coastal environments across the globe sometimes locally at high abundance. This is particularly true for members of *Haladaptatus*, which are likely to contribute to carbon cycling during periods of desiccation.

5. The coastal fungal microbiome

Fungi are a ubiquitous component of all ecosystems. They support the decomposition of lignocellulosic compounds (Bani et al. 2019; Francioli et al. 2020), provide industry-relevant bioactive products (Overy et al. 2019), and in vegetated habitats (e.g. coastal marshes) produce mycorrhizal networks that facilitate nutrient uptake by plants (Smith and Read, 2008). In addition, through the differential accumulation of fungal pathogens, they can promote plant biodiversity and productivity (Mommer et al. 2018).

Within coastal marshes, mycorrhizal associations sustain specific interactions between plants and fungi that are beneficial to survival and growth. For example, the arbuscular mycorrhizal (AM) fungi (Phylum: Glomeromycota), which are obligate plant-root endosymbionts of most terrestrial plant species, decrease salt stress and increase water uptake in plants growing in coastal marshes (Evelin et al. 2009). They may also decrease the impacts of localized hypoxia experienced by plant roots during tidal inundation. AM fungi play similar roles in plant nutrient acquisition (P and N uptake in exchange for plant-derived C) in coastal marshes as they do in most terrestrial habitats (Fitter 2005). The rewards to a given

plant species of this symbiosis depends both on the identification of the AM fungal species present and the soil/sediment nutrient levels (Hoeksema et al. 2010). Thus, these interactions influence plant competition dynamics at the ecosystem scale (van der Heijden 2002). In salt marshes the interaction between nutrient levels and AM fungi can influence plant zonation via changing the competitive ability of different plant species (Daleo et al. 2008), although this less extensively studied than for terrestrial habitats. For example, in the presence of AM fungi and at low nutrient concentrations Spartina densiflora has a competitive advantage over S. alterniflora but this is reversed by increased nutrient concentrations and/or suppression of AM fungi (Daleo et al. 2008). Other fungal phyla (e.g. Basidiomycota and Ascomycota) comprise species that provide the primary route for the decomposition of vascular plant litter and the remineralization of carbon in salt marshes, although bacteria may supersede fugal decomposers under more saline conditions (Cortes-Tolalpa et al. 2018). Some specificity in fungal decomposer communities appears to be present with different fungal species decomposing different plants and being present at different geographic locations (Calado et al. 2019; Cortes-Tolalpa et al. 2017; Lyons et al. 2010). Decomposition activity shows limited spatial variability on marshes (Buchan et al. 2003) and is largely uninfluenced by changes in salinity (Connolly et al. 2014), although which fungal decomposers are present may be affected by salinity (see below). However, decomposition rates vary seasonally (Buchan et al. 2003) and related microbial activity varies across diel cycles (Kuehn et al. 2004). Invasive (non-native) plant species affect bacteria-to-fungal ratios in marshes (Zhang et al. 2018), potentially altering decomposition rates via the introduction of novel substrates. This link between decomposition and substrate type means that there is also a certain degree of top-down control on fungal saprotrophs; where the modification of the physical structure of aboveground vegetation by animal grazers can alter both recalcitrant autochthonous input as well as the capacity to trap less-recalcitrant allochthonous inputs of plant material (Mueller et al. 2017).

Fungal biodiversity within coastal marshes is regulated by the identity and abundance of the plant species present, tidal inundation and salinity, alongside environmental gradients in physiochemistry that co-vary with salinity (Alzarhani et al. 2019; Mohamed and Martiny 2011). Across three marshes of different salinity (27-33, 15-25, and 0-10 ppt) fungal species richness was predominantly influenced by the presence of plant species and less influenced by salinity and other environmental gradients (Mohamed and Martiny, 2011). This most likely reflects increased heterogeneity in the microhabitats that fungi occupy and differential

patterns of co-occurrence (and or host specificity) across fungi and plant species. In contrast, the community composition of fungi from the same marshes was primarily determined by the underlying salinity gradient and not by the identity of plant species present, reflecting levels of halotolerance across fungal species limiting their occurrences to marshes within their salinity tolerances (Mohamed and Martiny 2011). However, larger-scale studies examining multiple marshes (n = 3) in high and low salinity environments (33-43 and 3.3-5.9 ppt) have shown contrasting results; with abiotic factors primarily determined the patterns of fungal richness while the interaction between abiotic and biotic factors determined community composition (Alzarhani et al. 2019). Moreover, the relationship between abiotic variables and fungal species richness was not generalizable across marshes and the relative influence of abiotic and biotic factors on community composition also varied. Subsequently, statistical models relating fungal biodiversity to the abiotic and/or biotic factors on a particular salt marsh performed poorly at predicting fungal biodiversity on other marshes despite the similarities between these environments (Alzarhani et al., 2019). These context-dependencies can be attributed, among other things, to differences in the functional groups of fungi present in salt marshes where certain abiotic or biotic variables were more strongly related to specific functional groups than over others (Alzarhani et al., 2019).

In coastal marine and aquatic estuarine environments adjacent to salt marshes much less is known about the diversity and functionality of fungi. Typically, estuaries support more diverse fungal communities than coastal marine environments, which in turn are more diverse than oceanic environments (Jeffries et al. 2016). This gradient of fungal diversity reflects the flow of terrestrial matter into estuarine systems and the role of the critical transition zone between freshwater and marine systems that determines benthic biodiversity (Levin et al. 2001). Fungal turnover is highest in estuarine environments when compared to other marine systems because terrestrial, freshwater, and non-halotolerant species give way to halotolerant fungi found in near-shore environments (Burgaud et al. 2013). In coastal environments that experience extreme salt stress (e.g. hyper-saline lagoons) fewer fungal species are present (e.g. Trimmatostroma spp., Emericella spp., and Phaeotheca spp.) and in general there are only a few known halophilic fungal species (Gostinčar et al. 2010). In the absence of terrestrial plant species with which to interact, the major functional role of fungi in these environments is the decomposition of lignocellulosic compounds and recycling of vascular plant litter (Newell 1996), alongside those that are pathogens. This also includes lignin degraders (Bucher et al. 2004), which contribute to the primary decomposition of woody

debris in estuarine environments (Poole and Price 1972; Tsui and Hyde 2004). The primarily saprotophic role fungi play in estuary environments has led to them being dominated by Basidiomycota and Ascomycota (Burgaud et al., 2013; Wang et al., 2019), but with a reduction in terrestrial subsidies in coastal waters, many species of Chytridiomycota become abundant (Sun et al., 2014; Jeffries et al., 2016). Generalizing these patterns of presence of fungal phyla is problematic as the dominance of species from different phyla changes with geographic locations. For example, in the Baltic Sea, low-salinity areas (< 8 ppt) contain fungal communities compositionally similar to those in local freshwaters, but higher-salinity areas (> 8 ppt) contain fungal communities similar to those in marine systems (Rojas-Jimenez et al. 2019). However, in these locations Basidiomycota and Ascomycota dominate in the marine environment and Chytridiomycota in freshwaters (Rojas-Jimenez et al. 2019). It is also worth noting that DNA sequences from Glomeromycota (AM fungi) have been detected during surveys in near-shore fungal communities, which if they originate from spores suggests a previously underexplored dispersal route (Lacerda et al. 2020). Given the important role of fungi in degrading complex organic material close species-species interactions and biogeochemical coupling between the fungal and algal, bacterial, and archaeal constituents are expected in coastal sediment biomes. These questions are currently unanswered.

6. Impacts of oil pollution on coastal microbiomes.

An estimated 1.3 million tons of oil enters the marine environment each year (National Research Council 2003). This includes oil from natural seeps and spills associated with the extraction and transportation of petroleum (e.g. tanker, pipeline, and coastal facility spills). The largest offshore oil spill in history was The Deepwater Horizon spill resulting from the Macondo well blowout that resulted in the release of 134 million gallons of crude oi in to the Gulf of Mexico. Despite the deposition of large quantities of oil in deep water systems, or being biodegraded, chemically dispersed, or burned in situ, large quantities of oil still reached coastal ecosystems and contaminated 2100 km of coastline. This pollution caused serious negative effects on marine life and coastal saltmarsh, seagrass, and reef systems (Beyer et al. 2016). Many coastal environments are particularly vulnerable to oil spills because many oil refineries are situated at the coast or at large estuaries. The majority of the 20 largest oil tanker spills to date also occurred close to the coast when tanker vessels ran aground (ITOPF, 2019). Oil spills have major effects on coastal ecosystems such as mass mortality of invertebrates, birds and mammals. For example, the Prestige oil spill caused a decrease of

66% of total species richness on Spanish Galician beaches (de La Huz et al. 2005) and the Exxon Valdez spill on the Alaskan coast caused major sea otter (Monson et al. 2000) and seabird (Piatt and Ford 1996) mortalities. Oil can also have toxic affects on microorganisms because the accumulation of hydrocarbon molecules in the membrane can result in loss of membrane integrity and impaired cellular homeostasis (Sikkema et al. 1995). This can have major impacts on key microbial ecosystem services such as coastal nitrogen cycling (Horel et al. 2014; Zhao et al. 2020). For example, some ammonia oxidizing bacteria and archaea are respectively 100 and 1000 times more sensitive to hydrocarbon toxicity than model heterotrophs respectively (Urakawa et al. 2019). Whilst large oil spills are thankfully rare coastal ecosystems are threatened continually with chronic oil and hydrocarbon pollution from rivers and land runoff (National Research Council 2003) and intensive industrial and recreational activities around coasts particularly near estuaries and harbors (Duran et al. 2015; McGenity 2014; Nogales and Bosch 2019).

Crude oil contains a complex mix of hydrocarbons that includes saturated aliphatic hydrocarbons such as cycloalkanes, linear *n*-alkanes (ranging from short chains to long chains with over 40 carbon atoms), and branched alkanes such as pristine and phytane (Weisman 1998). There are also many aromatic hydrocarbons such as the monoaromatic BTEX compounds (benzene, toluene, ethylbenzene and xylene) and polyaromatic hydrocarbons that include a wide range of both parent and methylated 2- to 5-ring compounds such as naphthalenes, phenanthrenes, pyrenes and perylenes. In addition to these two main classes of hydrocarbons there is a variety of large and highly recalcitrant asphaltene and resin compounds. Whilst certain processes remove some components of oil from the environment (e.g. evaporation of the lighter fractions, chemical- or photo-oxidation), unless oil is physically removed, the primary loss route will be via natural biodegradation by hydrocarbon-degrading microbes that utilize hydrocarbons as their carbon and energy source (Harayama et al. 1999; Head et al. 2006; McGenity et al. 2012). Because of the complexity of oil its biodegradation requires a diverse consortium of species that can degrade different hydrocarbons. There is niche partitioning between the different species in the consortium in the utilization of different hydrocarbon substrates (Head et al. 2006; McGenity et al. 2012; McKew et al. 2007).

6.1 Diversity of hydrocarbon-degrading microbes in coastal sediments.

High concentrations of hydrocarbons can dramatically alter the composition of coastal microbial communities leading to large decreases in species richness and diversity coupled

with the selection for specialist hydrocarbon-degrading bacteria (Head et al. 2006; McGenity et al. 2012). In estuarine and coastal sediments hydrocarbons are particularly used by Alphaand Gammaproteobacteria (Chronopoulou et al. 2013; Coulon et al. 2012; Greer 2010). The selection for specific bacterial species may be influenced by numerous factors including the concentration or type of oil and/or its degree of weathering (Head et al. 2006) or environmental conditions such as temperature or the concentration of key nutrients such as nitrogen and phosphorus (Coulon et al. 2007). However, many common patterns are observed globally, such as an increase in the relative abundance of obligate hydrocarbonoclastic bacteria (OHCB; Yakimov et al. 2007) in oxygenated oil-contaminated marine sediments. The OHCB include key genera such as Alcanivorax, Thalassolituus, Oleispira, and Oleibacter, which typically degrade alkanes as well as Cycloclasticus, which degrade a wide range of PAHs. The name 'OHCB' is a slightly misleading because these organisms are not truly "obligate". For example, in pure culture Alcanivorax degrades some other compounds (Radwan et al. 2019) and also some polyesters (Zadjelovic et al. However, there is still very limited evidence that the OHCB are competitive for nonhydrocarbon substrates in the environment and their lifestyle is often restricted to the use of hydrocarbons or their fatty acid or alcohol derivatives. The OHCB are often in low abundance in marine environments when hydrocarbons are absent but respond quickly and grow rapidly in response to oil pollution. Their streamlined genomes are specifically geared towards a hydrocarbon-degrading lifestyle (Kube et al. 2013; Schneiker et al. 2006; Yakimov et al. 2007) and their marine distribution is truly global (Yakimov et al. 2007). In muddy and sandy coastal sediments OHCB such a Alcanivorax, Oleibacter, Cycloclasticu, and Marinobacter hydrocarbonaoclasticus tend to increase in abundance and often dominate the bacterial community after the addition of crude oil (Chronopoulou et al. 2013; Coulon et al. 2012; Kostka et al. 2011; Thomas et al. 2020).

Alcanivorax borkumensis (Yakimov et al. 1998), a specialist *n*-alkane and branched alkane degrader, was first isolated from North Sea sediments and was the first OHCB to have its genome sequenced (Schneiker et al. 2006). Since its discovery 14 named species and a large diversity of unclassified *Alcanivorax* have been recorded in the NCBI database, including species isolated from deep sea sediments such as *A. dieselolei* (Liu and Shao 2005), *A. pacificus* (Lai et al. 2011), and *A. mobilis* (Yang et al. 2018), or from intertidal sediments such as *A. jadensis* (Fernandez-Martinez 2003) and *A. gelatiniphagus* (Kyoung Kwon et al. 2015). *Alcanivorax* is often dominant in oil contaminated intertidal sediments globally,

including in the Gulf of Mexico, in the Atlantic Galicean coast and in Mediterranean beaches following the Deepwater Horizon (Newton et al. 2013; Rodriguez-R et al. 2015), Prestige (Acosta-González et al. 2015), and Agia Zoni (Thomas et al. 2020) oil spills, respectively.

Another key alkane degrader in intertidal sediments is *Thalassolituus*, the type strain of which was isolated from harbor seawater/sediment samples in Milazzo, Italy (Yakimov et al., 2004). Thalassolituus is a highly competitive n-alkane degrader in estuarine environments and mudflats (McKew et al. 2007; Sanni et al. 2015) but is also found globally including deep water environments such as the oil plume from the DWH oil spill in the Gulf of Mexico (Camilli et al. 2010). *Oleibacter* related to the type strain *O. marinus* 201 (Teramoto et al. 2011) or a variety of unclassified strains is important in coastal fine-grained (Chronopoulou et al. 2013; Coulon et al. 2012; Sanni et al. 2015) and sandy (Thomas et al. 2020) sediment communities. However, like Alcanivorax and Thalassolituus, Oleibacter species are not specific to oil-degrading communities within coastal sediments as they (along with many other oil-degrading bacteria) are also found in a variety of marine environments, including for example seawater at 10,400 m in the Challenger Deep at the southern end of the Mariana Trench (Liu et al. 2019). This suggests that it is the availability of hydrocarbon that selects for oil degraders rather than the specific environmental conditions themselves. Low temperature often results in the selection for Oleispira. Bacteria closely related to the psychrophilic alkane-degrading Oleispira antarctica (Gregson et al. 2020; Kube et al. 2013; Yakimov et al. 2003) are important in oil-contaminated temperate coastal microbial communities at winter temperatures (4°C) (Coulon et al. 2007), as well as in cold environments such as deep arctic sediments (Dong et al. 2015) and sea-ice (Gerdes et al. 2005).

Cycloclasticus often plays the primary role in PAH degradation in coastal environments (Chronopoulou et al. 2013; Coulon et al. 2012; Duran and Cravo-Laureau 2016; Kasai et al. 2002; McKew et al. 2007; Sanni et al. 2015; Thomas et al. 2020). This genus comprises a wide variety of PAH-degrading species, including *C. pugetii* (Dyksterhouse et al. 1995), C. *oligotrophus* (Wang et al. 1996), C. *spirillensus* (Chung and King 2001), and *C. zancles* (Messina et al. 2016). There is a bivalve- and sponge symbiont lineage that can also degrade short-chain alkanes (Rubin-Blum et al. 2017).

In addition to these specialist hydrocarbon-degrading genera there are many other species from more nutritionally versatile genera that degrade hydrocarbons and many are regularly found in oil-polluted coastal sediments. There are too many to consider here but

they include for example species from genera such as *Marinobacter* (particularly *Marinobacter hydrcarbonoclasticus* (Gauthier et al. 1992)), *Alteromonas, Erythrobacter, Idiomarina, Microbacterium, Psuedomonas, Pseudoalteromonas, Rhodococcus, Roseovarius, Shewanella, Sphingomonas, Vibrio*, and *Xanthomonas* (e.g. see Prince et al. (2018) for a review of all hydrocarbon degraders, Goñi-Urriza & Duran (2018) for the role of bacteria in hydrocarbon degradation in coastal microbial mats, Greer (2010) for a review of bacterial diversity in hydrocarbon-polluted estuaries and sediments, and Supplementary Table 1 in Thomas et al. (2020) for many such observations in sandy coastal sediments).

Whilst aerobic biodegradation of hydrocarbons dominates in sediments, those that become buried in anoxic sediments can remain there for decades (Reddy et al. 2002) as anaerobic biodegradation of hydrocarbons is slow in comparison to aerobic biodegradation. Aerobic hydrocarbon degraders rely on oxygen not only for respiration but oxygen is also required for the primary step of degradation which is catalyzed by oxygenase enzyme systems (Wang and Shao 2013; Wang et al. 2018). Consequently, anaerobic bacteria must employ alternative pathways of biodegradation in the absence of oxygen. Due to the abundance of sulfate in coastal sediments many of the known anaerobic hydrocarbon oxidizers in coastal sediments are sulfate-reducing bacteria related to *Desulfosarcina*, *Desulfococcus*, *Desulfonema*, *Desulfobacula*, *Desulfotomaculum*, *Desulfotignum*, and *Geobacter* (McGenity 2014; Païssé et al. 2008; Rabus et al. 2016). Also, marine sediments are subjected to oscillations in oxygen concentration as the result of the tides, burrowing, and oxygenic phototrophic activity, which can encourage the growth of certain phylotypes of *Alcanivorax* (Terrisse et al. 2017).

The relative abundance of genera with potential hydrocarbon-degrading abilities can be used to estimate hydrocarbon exposure in an environment using the Ecological Index of Hydrocarbon Exposure (Lozada et al. 2014), which is based on microbial composition determined by 16S rRNA gene sequencing, as the numbers of hydrocarbon-degrading bacteria typically correlate with the concentration of hydrocarbons in sediments (Thomas et al. 2020).

6.2 Association of hydrocarbon-degrading bacteria with photoautotrophs

Many hydrocarbon-degrading bacteria in intertidal sediments have close association with photoautotrophs. Hydrocarbons can alter the composition of phototrophic communities considerably for example by inhibiting enzyme activities and photosynthesis (Megharaj et al.

2000). This toxicity may favor hydrocarbon-resistant species. For example, cyanobacteria belonging to Phormidium, Planktotrix, and Oscillatoria have shown varying degrees of tolerance to oil pollution (Van Bleijswijk and Muyzer 2004). In sediment mesocosms with oil polluted fine-sediments that were dominated with the hydrocarbonoclastic bacteria Alcanivorax, Cycloclasticus, and Oleibacter, there was also an increased abundance of MPB, primarily due to a 10-fold increase in the abundance of cyanobacteria (Chronopoulou et al. 2013; Coulon et al. 2012). This increase was attributed to a lower grazing pressure and/or nitrogen depletion, which encouraged the growth of diazotrophic cyanobacteria. Some microalgae co-exist with hydrocarbon-degrading bacteria (Amin et al. 2009; Chernikova et al. 2020; Gutierrez et al. 2013) and diatom-OHCB floating biofilms have been seen in mudflat sediments after an experimental oil spill (Coulon et al. 2012). There are numerous hypothesized ways that phototrophs enhance hydrocarbon degradation including direct degradation (although evidence for this is limited), supplying key resources (e.g. oxygen, N, Fe) to hydrocarbon-degrading bacteria, or assisting in immobilizing hydrocarbon-degrading bacteria within EPS. Phototrophs in turn can benefit from higher CO₂ concentrations from hydrocarbon-degrading bacterial respiration (e.g. see reviews by Abed 2019; Ardelean 2014; McGenity 2014; McGenity et al. 2012).

6.3 Mechanisms of oil biodegradation.

Generally, the bioavailability and rate of degradation of hydrocarbons decreases with increasing carbon number. Saturated hydrocarbons are often degraded at higher rates than light aromatics. The high-molecular-weight aromatics and polar compounds are degraded at low rates (Leahy and Colwell 1990). Most hydrocarbon-degrading bacteria will typically degrade a small range of either aliphatic or aromatic compounds, although some bacteria may possess pathways for catabolism of both aliphatic and PAH compounds such as some *Pseudomonas* (Whyte et al. 1997), *Rhodococcus* (Andreoni et al. 2000) or *Colwellia* (Mason et al. 2014) strains. Uptake via transport systems lowers the substrate concentration around the cell driving diffusive flux of hydrophobic hydrocarbons towards the cell (Harms et al. 2010), whilst the production of extracellular or cell-bound surface-active compounds (e.g. glucolipid biosurfactant produced by *Alcanivorax* (Yakimov et al., 1998)) increases bioavailability by decreasing the interfacial tension between water and oil (Marchant and Banat 2012). Aerobic hydrocarbon degrading bacteria are equipped with a wide array of genes that code for monooxygenase enzyme systems to activate hydrocarbons such as the two integral-membrane non-haem iron alkane monooxygenase systems AlkB1 and AlkB2 and

three haem-containing P450 cytochromes employed by *Alcanivorax* (Gregson et al. 2019; Schneiker et al. 2006; Yakimov et al. 1998) that convert medium-chain *n*-alkanes or branched alkanes to a primary alcohol that can be further degraded by an array of alcohol- and aldehyde dehydrogenases. Long-chain alkanes can be biodegraded with AlmA flavin-binding monooxygenases (Wang and Shao 2014) or similar sub-terminal Baeyer-Villiger monooxygenases in *Thalassolituus oleivorans* (Gregson et al. 2018). Dioxygenase systems are typically employed by aerobic PAH-degraders such as the PhnA1, PhnA2, PhnA3, and PhnA4 proteins (alpha and beta subunits of an iron-sulfur protein, a ferredoxin and a ferredoxin reductase, respectively) that make up a PAH dioxygenase system in *Cycloclaticus* strain A5 (Kasai et al. 2003).

Fine-grained intertidal sediments such as mudflats are typically anoxic below 1-2 mm. This prevents the activation of hydrocarbons using oxygenase enzymes. The exact mechanisms of the anaerobic activation of hydrocarbons are less well understood but may include direct carboxylation, methylation followed by addition to fumarate, or even utilization of nitrite to activate alkanes (McGenity et al. 2012; Meckenstock and Mouttaki 2011; Rabus et al. 2016; Widdel and Musat 2010; Zedelius et al. 2011).

7 Acknowledgements

The authors wish to acknowledge the many Ph.D., postdoctoral researchers, and technical staff who have contributed to the research of the Ecology and Environmental Microbiology group at the University of Essex, and whose work is cited in this chapter. The following funding bodies and awards are gratefully acknowledged: U.K. Natural Environment Research Council (NERC) Coastal Biodiversity and Ecosystem Services programme (NE/J01561X/1); NERC EPStromNet (NE/V00834X/1); NERC SSB programme, module Blue Carbon, (NE/K001914/1); NERC Quantifying a marine ecosystem's response to a catastrophic oil spill. (NE/R016569/1); NERC PRINCE-A new dynamic for Phosphorus in RIverbed Nitrogen Cycling (NE/P011624/1); European Union's Horizon 2020 research and innovation programme, grant agreement No 702217; Defra funding "Ascertaining Predisposing Factors that Affect Oak Health in the U.K. and Advancements towards Management for Resilient Oak Populations"; Eastern ARC Academic Research Consortium; The Royal Society, Understanding microalgal biofilm contributions to sediment "blue carbon" in contrasting salt marsh habitats in the U.S. and Europe (IES\R1\201260).

8 References

Abed RMM (2019) Phototroph-Heterotroph Oil-Degrading Partnerships. In: McGenity T. (eds) Microbial Communities Utilizing Hydrocarbons and Lipids: Members, Metagenomics and Ecophysiology. Handbook of Hydrocarbon and Lipid Microbiology. Springer, Cham. 37-50

Abell GCJ, Revill AT, Smith C et al (2010) Archaeal ammonia oxidizers and nirS-type denitrifiers dominate sediment nitrifying and denitrifying populations in a subtropical macrotidal estuary. ISME J 4:286-300

Acosta-González A, Martirani-von Abercron SM, Rosselló-Móra R et al (2015) The effect of oil spills on the bacterial diversity and catabolic function in coastal sediments: a case study on the Prestige oil spill. Environ Sci Pollut Res Int 22:15200-15214

Acuña Alvarez L, Exton DA, Suggett DJ et al (2009) Characterization of marine isoprene-degrading communities. Environ Microbiol 11:3280–3291.

Admiraal W (1984) The ecology of estuarine sediment inhabiting diatoms. Progr Phyc Res 3:269-322

Admiraal W, Peletier H, Brouwer T (1984) The seasonal succession patterns of diatom species on an intertidal mudflat: an experimental analysis. Oikos 42:30-40

Alongi DM (2020) Carbon balance in salt marsh and mangrove ecosystems: A global synthesis. J Mar Sci Eng 8:767 doi:10.3390/jmse8100767

Alzarhani AK, Clark DR, Underwood GJC et al (2019) Are drivers of root-associated fungal community structure context specific? ISME J 13:1330–1344

Amin SA, Green HD., Hart MC et al (2009) Photolysis of iron-siderophore chelates promotes bacterial-algal mutualism. Proc Nat Acad Sci USA 106:17071–17076

An SM, Choi DH, Noh JH (2020) High-throughput sequencing analysis reveals dynamic seasonal succession of diatom assemblages in a temperate tidal flat. Estuar Coast Shelf Sci 237:106686

Andreoni V, Bernasconi S, Colombo M et al (2000) Detection of genes for alkane and naphthalene catabolism in *Rhodococcus* sp. strain 1BN. Environ Microbiol 2: 572-577

Ardelean II (2014) The involvement of cyanobacteria in petroleum hydrocarbons degradation: fundamentals, applications and perspectives. In: Cyanobacteria: ecological importance, biotechnological uses and risk management, pp 41–60

Arp DJ, Sayavedra-Soto LA, Hommes NG (2002) Molecular biology and biochemistry of ammonia oxidation by *Nitrosomonas europaea*. Arch Microbiol 178:250–255

Baas JH, Baker ML, Malarkey J et al (2019) Integrating field and laboratory approaches for ripple development in mixed sand–clay–EPS. Sedimentol 66:2749-2768

Baker BJ, De Anda V, Seitz KW et al (2020) Diversity, ecology and evolution of Archaea. Nat Microbiol 5:887-900

Bani A, Borruso L, Nicholass KJM et al (2019) Site-specific microbial decomposer communities do not imply faster decomposition: results from a litter transplantation experiment. Microorganisms 7:349

Barnett A, Méléder V, Blommaert L et al (2015) Growth form defines physiological photoprotective capacity in intertidal benthic diatoms. ISME J 9:32–45

Barranguet C, Kromkamp J, Peene J (1998) Factors controlling primary production and photosynthetic characteristics of intertidal microphytobenthos. Mar Ecol Prog Ser 173:117-126

Bartelme RP, McLellan SL, Newton RJ (2017) Freshwater recirculating aquaculture system operations drive biofilter bacterial community shifts around a stable nitrifying consortium of ammonia-oxidizing Archaea and comammox *Nitrospira*. Front Microbiol 8:101

Beaumont NJ, Jones L, Garbutt A et al (2014) The value of carbon sequestration and storage in coastal habitats. Estuar Coast Shelf Sci 137:32–40

Beddow J, Stolpe B, Cole PA et al (2017) Nanosilver inhibits nitrification and reduces ammonia-oxidizing bacterial but not archaeal amoA gene abundance in estuarine sediments. Environ Microbiol 19:500-510

Bellinger BJ, Abdullahi AS, Gretz MR et al (2005) Biofilm polymers: relationship between carbohydrate biopolymers from estuarine mudflats and unialgal cultures of benthic diatoms. Aquat Microb Ecol 38:169–180

Bellinger BJ, Underwood GJC, Ziegler SE et al (2009) Significance of diatom-derived polymers in carbon flow dynamics within estuarine biofilms determined through isotopic enrichment. Aquat Microb Ecol 55:169–187

Beman JM, Francis CA (2006) Diversity of ammonia-oxidizing archaea and bacteria in the sediments of a hypernutrified subtropical estuary: Bahia del Tobari, Mexico. Appl Environ Microbiol 72:7767–7777

Beman JM, Popp BN, Francis CA (2008) Molecular and biogeochemical evidence for ammonia oxidation by marine Crenarchaeota in the Gulf of California. ISME J 2:429–441

Benyoucef I, Blandin E, Lerouxel A et al (2014) Microphytobenthos interannual variations in a north-European estuary (Loire estuary, France) detected by visible-infrared multispectral remote sensing. Estuar Coast Shelf Sci 136:43-52

Berg G, Rybakova D, Fischer D et al (2020) Microbiome definition re-visited: old concepts and new challenges. Microbiome 8:103

Bergman B, Gallon JR, Rai AN et al (1997) N₂ fixation by non-heterocystous cyanobacteria. FEMS Microbiol Rev 19:139-185

Berman-Frank I, Lundgren P, Falkowski P (2003) Nitrogen fixation and photosynthetic oxygen evolution in cyanobacteria. Res Microbiol 154:157–164

Beyer J, Trannum HC, Bakke T et al (2016) Environmental effects of the Deepwater Horizon oil spill: A review. Mar Pollut Bull 15;110(1):28-51

Blommaert L, Lavaud J, Vyverman W et al (2018) Behavioural versus physiological photoprotection in epipelic and epipsammic benthic diatoms, Eur J Phycol 53:146-155

Blommaert L, Vancaester E, Huysman MJJ et al (2020) Light Regulation of LHCX Genes in the Benthic Diatom *Seminavis robusta*. Front Mar Sci 7:192

Böer SI, Hedtkamp SIC, van Beusekom JEE et al (2009) Time- and sediment depth-related variations in bacterial diversity and community structure in subtidal sands. ISME J 3:780–791.

Boetius A, Ravenschlag K, Schubert CJ et al (2000) A marine microbial consortium apparently mediating anaerobic oxidation of methane. Nature 407:623–626

Bohórquez J, McGenity TJ, Papaspyrou S et al (2017) Different Types of Diatom-Derived Extracellular Polymeric Substances Drive Changes in Heterotrophic Bacterial Communities from Intertidal Sediments. Front Microbiol 8:245

Booty JM, Underwood GJC, Parris A et al (2020) Wading birds affect ecosystem functioning on an intertidal mudflat. Front Mar Sci 7:685.

Bowles M, Nigro L, Teske A et al (2012) Denitrification and environmental factors influencing nitrate removal in Guaymas Basin hydrothermally altered sediments. Front Microbiol 3:377

Braker G, Conrad R (2011) Diversity, structure, and size of N₂O-producing microbial communities in soils—what matters for their functioning? Adv Appl Microbiol 75:33-70

Braker G, Zhou JZ, Wu LY et al (2000) Nitrite reductase genes (*nirK* and *nirS*) as functional markers to investigate diversity of denitrifying bacteria in Pacific Northwest marine sediment communities. Appl Environ Microbiol 66:2096–2104

Bruckner CG, Bahulikar R, Rahalkar M et al (2008) Bacteria associated with benthic diatoms from Lake Constance: Phylogeny and influences on diatom growth and secretion of extracellular polymeric substances. Appl Environ Microbiol 74:7740–7749

Bruckner CG, Rehm C, Grossart H-P et al (2011) Growth and release of extracellular organic compounds by benthic diatoms depend on interactions with bacteria. Environ Microbiol 13:1052-1063

Buchan A, Newell SY, Butler M et al (2003) Dynamics of bacterial and fungal communities on decaying salt marsh grass. Appl Environ Microbiol 69:6676–6687

Bucher VVC, Hyde KD, Pointing SB et al (2004) Production of wood decay enzymes, mass loss and lignin solubilization in wood by marine ascomycetes and their anamorphs. Fungal Divers 15:1–14

Burden A, Garbutt A, Evans CD (2019) Effect of restoration on saltmarsh carbon accumulation in Eastern England. Biol Lett 15:20180773

Burgaud G, Woehlke S, Rédou V et al (2013) Deciphering the presence and activity of fungal communities in marine sediments using a model estuarine system. Aquat Microb Ecol 70:45–62

Caffrey JM, Bano N, Kalanetra K et al (2007) Ammonia oxidation and ammonia-oxidizing bacteria and archaea from estuaries with differing histories of hypoxia. ISME J. 1:660–662

Calado ML, Carvalho L, Barata M et al (2019) Potential roles of marine fungi in the decomposition process of standing stems and leaves of *Spartina maritima*. Mycologia 111:371–383

Camilli R, Reddy CM, Yoerger DR et al (2010). Tracking hydrocarbon plume transport and biodegradation at Deepwater Horizon. Science 330: 201–204

Cao H, Hong Y, Li M et al (2012) Community shift of ammonia-oxidizing bacteria along an anthropogenic pollution gradient from the Pearl River Delta to the South China Sea. Appl Microbiol Biotechnol 94:247–259

Cao H, Hong Y, Li M et al. (2011) Diversity and abundance of ammonia-oxidizing prokaryotes in sediments from the coastal Pearl River estuary to the South China Sea. Antonie van Leeuwenhoek 100:545

Capone DG (1988) Benthic nitrogen fixation. In: Nitrogen cycling in coastal marine sediments (Blackburn TH and Sorensen J, Eds.), pp 85-123. John Wiley and Sons, New York.

Chen W, Pan Y, Yu L et al (2017) Patterns and processes in marine microeukaryotic community biogeography from Xiamen coastal waters and intertidal Sediments, Southeast China. Front Microbiol 8:1912

Chernikova TN, Bargiela R, Toshchakov SV et al (2020) Hydrocarbon-degrading bacteria Alcanivorax and Marinobacter associated with microalgae Pavlova lutheri and Nannochloropsis oculata. Front Microbiol 11:2650

Christianen MJA, Middelburg JJ, Holthuijsen SJ, et al (2017) Benthic primary producers are key to sustain the Wadden Sea food web: stable carbon isotope analysis at landscape scale. Ecology 98:1498–1512

Chronoupolou P-M, Fahy A, Coulon F et al (2013) Impact of a simulated oil spill on benthic phototrophs and nitrogen-fixing bacteria. Environ Microbiol 15:241-252

Chung WK, King GM (2001) Isolation, characterization, and polyaromatic hydrocarbon degradation potential of aerobic bacteria from marine macrofaunal burrow sediments and description of *Lutibacterium anuloederans* gen. nov., sp. nov., and *Cycloclasticus spirillensus* sp. nov. Appl Environ Microbio. 67:5585-5592

Cibic T, Fazi S, Nasi F et al (2019) Natural and anthropogenic disturbances shape benthic phototrophic and heterotrophic microbial communities in the Po River Delta system, Estuar Coast Shelf Sci 222:168-182

Clark DR, McKew B, Dong L et al (2020) Mineralization and nitrification: Archaea dominate ammonia-oxidising communities in grassland soils. Soil Biol Biochem 143:107725

Clark DR, Mégane Mathieu M, Mourot L et al (2017) Biogeography at the limits of life: Do extremophilic microbial communities show biogeographic regionalisation? Glob Ecol Biogeogr 26:1435-1446

Coelho H, Vieira S, Serôdio J (2011) Endogenous versus environmental control of vertical migration by intertidal benthic microalgae. Eur J Phycol 46:271-281

Cohn SA, Dunbar S, Ragland R et al (2016) Analysis of light quality and assemblage composition on diatom motility and accumulation rate. Diatom Res 31:173-184

Connolly CT, William V. Sobczak WV, Findlay SEG (2014) Salinity effects on phragmites decomposition dynamics among the Hudson River's freshwater tidal wetlands. Wetlands 34:575–582

Consalvey M, Paterson DM, Underwood GJC (2004) The ups and downs of life in a benthic biofilm: Migration of benthic diatoms. Diatom Res 19:181-202

Cook, PLM, Veuger B, Böer S et al (2007) Effect of nutrient availability on carbon and nitrogen incorporation and flows through benthic algae and bacteria in near-shore sandy sediment. Aquat Microb Ecol 49:165–180

Cortes-Tolalpa L, Norder J, van Elsas JD et al (2018) Halotolerant microbial consortia able to degrade highly recalcitrant plant biomass substrate. Appl Microbiol Biotechnol 102:2913–2927

Coulon F, Chronopoulou PM, Fahy A et al (2012) Central role of dynamic tidal biofilms dominated by aerobic hydrocarbonoclastic bacteria and diatoms in the biodegradation of hydrocarbons in coastal mudflats. Appl Environ Microbiol 78:3638–3648

Coulon F, McKew BA, Osborn AM et al (2007) Effects of temperature and biostimulation on oil-degrading microbial communities in temperate estuarine waters. Environ Microbiol 9:177–186

Daggers TD, Herman PMJ, van der Wal D (2020) Seasonal and spatial variability in patchiness of microphytobenthos on intertidal flats from Sentinel-2 satellite imagery. Front Mar Sci 7:392

Daims H, Lucker S, Wagner M (2016) A new perspective on microbes formerly known as nitrite-oxidizing bacteria. Trends Microbiol 24:699 –712

Dal Bello M, Rindi L, Benedetti-Cecchi L (2017) Legacy effects and memory loss: how contingencies moderate the response of rocky intertidal biofilms to present and past extreme events. Glob Change Biol 23:3259-3268

Daleo P, Alberti J, Canepuccia A et al (2008) Mycorrhizal fungi determine salt-marsh plant zonation depending on nutrient supply. J Ecol 96: 431–437

Dalsgaard T, Canfield DE, Petersen J et al (2003) N₂ production by the anammox reaction in the anoxic water column of Golfo Dulce, Costa Rica. Nature 422:606–608

Dang H, Li J, Chen R et al (2010) Diversity, abundance, and spatial distribution of sediment ammonia-oxidizing betaproteobacteria in response to environmental gradients and coastal eutrophication in Jiaozhou Bay, China. Appl Environ Microbiol 76:4691-4702

Dang H, Wang, C, Li J et al (2009) Diversity and distribution of sediment *nirS*-encoding bacterial assemblages in response to environmental gradients in the eutrophied Jiaozhou Bay, China. Microb Ecol 58:161–169

De Jonge VN, de BoerWF, de Jong DJ et al (2012) Long-term mean annual microphytobenthos chlorophyll *a* variation correlates with air temperature. Mar Ecol Prog Ser 468:43-56

De Jonge VN, van Beusekom JEE (1995) Wind- and tide-induced resuspension of sediment microphytobenthos from the tidal flats in the Ems Estuary. Limnol Oceanogr 40:766-778

de la Huz R, Lastra M, Junoy J et al (2005) Biological impacts of oil pollution and cleaning in the intertidal zone of exposed sandy beaches: Preliminary study of the "Prestige" oil spill. Estuar Coast Shelf Sci 65:19-29

Dean JF, Middelburg JJ, Röckmann T et al. (2018) Methane feedbacks to the global climate system in a warmer world. Rev Geophys 56:207–250

DeLong EF (1992) Archaea in coastal marine environments. Proc Natl Acad Sci USA 89:5685–5689

Dong C, Bai X, Sheng H et al (2015) Distribution of PAHs and the PAH-degrading bacteria in the deep-sea sediments of the high-latitude Arctic Ocean. Biogeosciences 12:2163–2177

Dong LF, Thornton DCO, Nedwell DB et al (2000) Denitrification in the sediments of the River Colne estuary, England. Mar Ecol Prog Ser 203:109-122

Dong X, Greening C, Rattray JE et al. (2019) Metabolic potential of uncultured bacteria and archaea associated with petroleum seepage in deep-sea sediments. Nat Commun 10:1816

Duarte, CM, Agusti S, Barbier E et al (2020) Rebuilding marine life. Nature 580:39–51

Duran R, Cravo-Laureau C (2016) Role of environmental factors and microorganisms in determining the fate of polycyclic aromatic hydrocarbons in the marine environment. FEMS Microbiol Rev 40:814–830

Duran R, Cuny P, Bonin P et al (2015) Microbial ecology of hydrocarbon-polluted coastal sediments. Environ Sci Pollut Res 22:15195–15199.

Dyksterhouse SE, Gray JP, Herwig RP et al (1995) *Cycloclasticus pugetii* gen. nov., sp. nov., an aromatic hydrocarbon-degrading bacterium from marine sediments. Int J Syst Bacteriol. 45:116-23

Egger M, Riedinger N, Mogollo'n JM et al (2018) Global diffusive fluxes of methane in marine sediments. Nat Geosci 11:421–425

Erguder TH, Boon N, Wittebolle L et al (2009) Environmental factors shaping the ecological niches of ammonia-oxidizing archaea. FEMS Microbiol Rev 33:855–869

Evans PN, Boyd JA, Leu AO et al (2019) An evolving view of methane metabolism in the archaea. Nat Rev Microbiol 17:219–232

Evans PN, Parks DH, Chadwick GL et al (2015) Methane metabolism in the archaeal phylum Bathyarchaeota revealed by genome-centric metagenomics. Science 350:434–438

Evelin H, Kapoor R, Giri B (2009) Arbuscular mycorrhizal fungi in alleviation of salt stress: a review. Ann Bot 104:1263–1280

Feldewert C, Lang K, Brune A. 2020. The hydrogen threshold of obligately methyl775 reducing methanogens. FEMS Microbiol Lett 17:fnaa137

Fernandez-Martinez J, Pujalte MJ, Garcia-Martinez J et al (2003) Description of *Alcanivorax venustensis* sp. nov. and reclassification of *Fundibacter jadensis* DSM 12178T (Bruns and Berthe-Corti 1999) as *Alcanivorax jadensis* comb. nov., members of the emended genus *Alcanivorax*. Int J Syst Evol Microbiol 53:331-338

Fitter AH (2005) Darkness visible: reflections on underground ecology. J Ecol 93:231–243

Forster RM, Creach V, Sabbe K et al (2006) Biodiversity-ecosystem function relationship in microphytobenthic diatoms of the Westerschelde estuary. Mar Ecol Prog Ser 311:192-201

Francioli D, van Rijssel SQ, van Ruijven J et al (2020) Plant functional group drives the community structure of saprophytic fungi in a grassland biodiversity experiment. Plant Soil https://doi.org/10.1007/s11104-020-04454-y

Francis CA, Roberts KJ, Beman JM et al (2005) Ubiquity and diversity of ammonia-oxidizing archaea in water columns and sediments of the ocean. Proc Natl Acad Sci USA 102:14683–14688

Fuhrman JA, Davis AA (1997) Widespread archaea and novel bacteria from the deep sea as shown by 16S rRNA gene sequences. Mar Ecol Prog Ser 150:275–285

Fuhrman JA, McCallum K, Davis AA (1992) Novel major archaebacterial group from marine plankton. Nature 356,148–149

Galiforni-Silva F, Wijnberg KM, Hulscher SJMH (2020) On the Relation between Beach-Dune Dynamics and Shoal Attachment Processes: A Case Study in Terschelling (NL) J. Mar. Sci. Eng. 8: 541

Gaubert-Boussarie J, Prado S, Hubas C (2020) An untargeted metabolomic approach for microphytobenthic biofilms in intertidal mudflats. Front Mar Sci 7:250

Gauthier MJ, Lafay B, Christen R et al (1992) *Marinobacter hydrocarbonoclasticus* gen. nov., sp. nov., a new, extremely halotolerant, hydrocarbon-degrading marine bacterium. Int J Syst Bacteriol 42:568–576

Gerdes B, Brinkmeyer R, Dieckmann G et al (2005) Influence of crude oil on changes of bacterial communities in Arctic Sea-ice. FEMS Microbiol Ecol 53:129–139

Gihring TM, Humphrys M, Mills HJ et al (2009) Identification of phytodetritus-degrading microbial communities in sublittoral Gulf of Mexico sands. Limnol Oceanogr 54:1073–1083

Gołębiewski M, Całkiewicz J, Creer S et al (2017) Tideless estuaries in brackish seas as possible freshwater-marine transition zones for bacteria: the case study of the Vistula river estuary. Environ Microbiol Rep 9:129–143

Gong J, Shi F, Ma B et al (2015) Depth shapes α - and β -diversities of microbial eukaryotes in surficial sediments of coastal ecosystems. Environ Microbiol 17:3722–3737

Gong W, Marchetti A (2019) Estimation of 18S gene copy number in marine eukaryotic plankton using a Next-Generation Sequencing approach. Front Mar Sci 6:219

Goñi-Urriza M, Duran R (2018) Impact of Petroleum Contamination on Microbial Mats. In: McGenity T. (eds) Microbial Communities Utilizing Hydrocarbons and Lipids: Members, Metagenomics and Ecophysiology. Handbook of Hydrocarbon and Lipid Microbiology. Springer, Cham. 19-36

Gostinčar C, Grube M, de Hoog GS et al (2010) Extremotolerance in fungi: evolution on the edge. FEMS Microbiol Ecol 71:2–11

Green BC, Smith DJ, Grey J et al (2012) High site fidelity and low site connectivity of temperate salt marsh fish populations: a stable isotope approach. Oecologia 168:245-255

Green M, Coco G (2014) Review of wave-driven sediment resuspension and transport in estuaries. Rev Geophys 52:77–117

Greer CW (2010) Bacterial diversity in hydrocarbon-polluted rivers, estuaries and sediments. In: Timmis KN, McGenity TJ, van der Meer JR, de Lorenzo V (eds) Handbook of Hydrocarbon and Lipid Microbiology. Springer, Berlin, Heidelberg. 2329–2338

Gregson BH, Metodieva G, Metodiev MV et al (2018) Differential protein expression during growth on medium versus long-Chain alkanes in the obligate marine hydrocarbon-degrading bacterium *Thalassolituus oleivorans* MIL-1. Front Microbiol 9:3130

Gregson BH, Metodieva G, Metodiev MV et al (2019) Differential protein expression during growth on linear versus branched alkanes in the obligate marine hydrocarbon-degrading bacterium *Alcanivorax borkumensis* SK2T. Environ Microbiol 21:2347-2359

Gregson BH, Metodieva G, Metodiev MV et al (2020) Protein expression in the obligate hydrocarbon-degrading psychrophile *Oleispira antarctica* RB-8 during alkane degradation and cold tolerance. Environ Microbiol 22:1870-1883

Grossart HP, Levold F, Allgaier M et al (2005) Marine diatom species harbour distinct bacterial communities. Environ Microbiol 7:860–873

Gutierrez T, Green DH, Nichols PD et al (2013) *Polycyclovorans algicola* gen. nov., sp. nov., an aromatic-hydrocarbon-degrading marine bacterium found associated with laboratory cultures of marine phytoplankton. Appl Environ Microbiol 79:205–214

Hallam SJ, Mincer TJ, Schleper C et al (2006) Pathways of carbon assimilation and ammonia oxidation suggested by environmental genomic analyses of marine Crenarchaeota. PLoS Biol 4:e95.

Hamels I, Sabbe K, Muylaert K et al (1998) Organisation of microbenthic communities in intertidal estuarine flats, a case study from the Molenplaat (Westerschelde Estuary, the Netherlands). Eur. J. Protistol. 34:308–320

Hanlon ARM, Bellinger B, Haynes K et al (2006) Dynamics of extracellular polymeric substance (EPS) production and loss in an estuarine, diatom-dominated, microalgal biofilm over a tidal emersion-immersion period. Limnol Oceanogr 51:79–93

Hanson RB, Gundersen K (1977) Relationship between nitrogen fixation (acetylene reduction) and the C:N ratio in a polluted coral reef system, Kaneohe Bay, Hawaii. Est Coast Mar Sci 5:437-444

Happey-Wood CM, Jones P (1988) Rhythms of vertical migration and motility in intertidal diatoms with particular reference to *Pleurosigma angulatum*. Diat Res 3:83-93

Harayama S, Kishira H, Kasai Y et al (1999) Petroleum biodegradation in marine environments. J Mol Microbiol Biotechnol. 1:63-70

Harms H, Smith KEC, Wick LY (2010) Microorganism–hydrophobic compound interactions. In Handbook of Hydrocarbon and Lipid Microbiology. Edited by Timmis KN, McGenity TJ, van der Meer JR, de Lorenzo V. Berlin Heidelberg: Springer:1479-1490.

Haro S, Bohórquez J, Lara M et al (2019). Diel patterns of microphytobenthic primary production in intertidal sediments: circadian photosynthetic rhythm and migration. Sci Rep 9:13376.

Haro-Moreno JM, Rodriguez-Valera F, López-García P et al (2017) New insights into marine group III Euryarchaeota, from dark to light. ISME J 11:1102–1117

Head IM, Hiorns WD, Embley TM et al (1993) The phylogeny of autotrophic ammonia oxidizing bacteria as determined by analysis of 16S ribosomal RNA gene-sequences. J Gen Microbiol 139:1147-1153

Head IM, Jones DM, Röling WFM (2006) Marine microorganisms make a meal of oil. Nat Rev Microbiol 4:173–182

Heip CHR, Goosen NK, Herman PMJ et al (1995) Production and consumption of biological particles in temperate tidal estuaries. Oceanogr Mar Biol Ann Rev. 33:1-149

Henderson CJ, Gilby BL, Schlacher TA et al (2020) Low redundancy and complementarity shape ecosystem functioning in a low-diversity ecosystem. J Anim Ecol 89:784–794

Herbert RA (1975) Heterotrophic nitrogen fixation in shallow estuarine sediments. J Exp Mar Biol Ecol 18:215-225

Herbert RA (1999) Nitrogen cycling in coastal marine ecosystems. FEMS Microbiol Rev 5:563-590

Herman PMJ, Middelburg JJ, Widdows J et al (2000) Stable isotopes as trophic tracers: combining field sampling and manipulative labelling of food resources for macrobenthos. Mar Ecol Prog Ser 204:79–92

Hill-Spanik KM, Smith AS, Plante CJ (2019) Recovery of benthic microalgal biomass and community structure following beach renourishment at Folly Beach, South Carolina. Estuaries Coast 42:157–172

Hoehler TM, Alperin MJ, Albert DB et al (1994) Field and laboratory studies of 534 methane oxidation in an anoxic marine sediment: Evidence for a methanogen-sulfate 535 reducer consortium. Global Biogeochem Cycles 8, 451–463

Hoeksema JD, Chaudhary VB, Gehring CA et al (2010) A meta-analysis of context-dependency in plant response to inoculation with mycorrhizal fungi. Ecol Letts 13:394–407

Hope JA, Paterson DM, Thrush SF (2020) The role of microphytobenthos in soft-sediment ecological networks and their contribution to the delivery of multiple ecosystem services. J Ecol 108: 815–830

Horel A, Bernard RJ, Mortazavi B (2014) Impact of crude oil exposure on nitrogen cycling in a previously impacted *Juncus roemerianus* salt marsh in the northern Gulf of Mexico. Environ Sci Pollut Res Int. 21:6982-93

Hörnlein C, Confurius-Guns V, Stal LJ et al (2018) Daily rhythmicity in coastal microbial mats. NPJ Biofilms Microbiomes 4:11

Hu H, He J (2017) Comammox—a newly discovered nitrification process in the terrestrial nitrogen cycle. J Soils Sediments 17:2709–2717

Imachi H, Nobu MK, Nakahara N et al. (2020) Isolation of an archaeon at the prokaryote-eukaryote interface. Nature 577:519–525

ITOPF (2019) Oil tanker Spill Statistics 2019. Available http://www.itopf.org/

Jameson E, Stephenson J, Jones H et al. (2018) Deltaproteobacteria (*Pelobacter*) and Methanococcoides are responsible for choline-dependent methanogenesis in a coastal saltmarsh sediment. ISME J 13:277–289

Jayakumar A, O'Mullan GD, Naqvi SWA et al (2009) Denitrifying bacterial community composition changes associated with stages of denitrification in oxygen minimum zones. Microb Ecol 58:350–362

Jeffries TC, Curlevski NJ, Brown MV et al (2016) Marine fungal biogeography. Environ Microbiol Rep 8:235–238

Jensen MM, Kuypers MMM, Lavik G et al (2008). Rates and regulation of anaerobic ammonium oxidation and denitrification in the Black Sea. Limnol Oceanogr 53:23–36

Jesus B, Brotas V, Ribeiro L et al (2009) Adaptations of microphytobenthos assemblages to sediment type and tidal position, Cont Shelf Res 29:1624-1634

Jetten MSM, Op den Camp HJM, Kuenen JG et al (2010) Description of the order Brocadiales. Bergey's Manual of Systematic Bacteriology, Vol 4 (Krieg NR, Ludwig W, Whitman WB, Hedlund BP, Paster BJ, Staley JT, Ward N, Brown D & Parte A, eds), pp. 596–603. Springer, Heidelberg.

Jiang H, Dong H, Yu B et al (2009) Diversity and abundance of ammonia-oxidizing archaea and bacteria in Qinghai Lake, northwestern China. Geomicrobiol J 26:199–211

Jones C, Hallin S. (2010). Ecological and evolutionary factors underlying global and local assembly of denitrifier communities. ISME J 4:633–641

Jönnson B, Sundbäck K, Nilsson C (1994) An upright life-form of an epipelic motile diatoms: on the behaviour of *Gyrosigma balticum*. Eur J Phycol 29:11-15

Jørgensen KS (1989) Annual pattern of denitrification and nitrate ammonification in an estuarine sediment Appl Environ Microbiol 55:1841-1847

Jørgensen KS, Sorensen J (1988) Two annual maxima of nitrate reduction and denitrification in estuarine sediment (Norsminde Fjord, Denmark). Mar Ecol Prog Ser 94:267-274

Juneau P, Barnett A, Méléder V et al (2015) Combined effect of high light and high salinity on the regulation of photosynthesis in three diatom species belonging to the main growth forms of intertidal flat inhabiting microphytobenthos. J Exp Mar Biol Ecol 463:95-104

Kamp A, Beer D; Nitsch JL et al (2011) Diatoms respire nitrate to survive dark and anoxic conditions. Proc Nat Acad Sci USA 108:5649–5654

Kaplan WA, Teal JM, Valiela I (1977) Denitrification in saltmarsh sediments: evidence for seasonal temperature selection among populations of denitrifiers. Microb Ecol 3:193-224

Kartal B, Kuypers MM, Lavik G et al (2007) Anammox bacteria disguised as denitrifiers: nitrate reduction to dinitrogen gas via nitrite and ammonium. Environ Microbiol 9:635–642

Kasai Y, Kishira H, Harayama S. (2002) Bacteria belonging to the genus *Cycloclasticus* play a primary role in the degradation of aromatic hydrocarbons released in a marine environment. Appl Environ Microbiol 68:5625–5633

Kasai Y, Shindo K, Harayama S et al (2003) Molecular characterization and substrate preference of a polycyclic aromatic hydrocarbon dioxygenase from *Cycloclasticus* sp. Strain. A5. Appl Environ Microbiol 69:6688-6697

Kim BS, Oh HM, Kang H et al (2005) Archaeal diversity in tidal flat sediment as revealed by 16S rDNA analysis. J Microbiol 43:144-151

King D, Nedwell DB (1987) The adaptation of nitrate reducing bacterial communities in estuarine sediments in response to overlying nitrate load. FEMS Microb Ecol 45:15-21

Kingston MB (1999) Effect of light on vertical migration and photosynthesis of Euglena proxima (Euglenophyta). J Phycol 35:245-253

Knittel K, Wegener G, Boetius A (2018) Anaerobic methane oxidizers. In Microbial Communities Utilizing Hydrocarbons and Lipids: Members, Metagenomics and Ecophysiology, McGenity TJ (Ed.), Springer International Publishing, Cham. 113-132

Koch H, van Kessel MAHJ, Lücker S (2019) Complete nitrification: insights into the ecophysiology of comammox Nitrospira. Appl Microbiol Biotechnol 103:177–189

Koedooder C, Stock W, Willems A et al (2019) Diatom-Bacteria interactions modulate the composition and productivity of benthic diatom biofilms. Front Microbiol 10:1255

König S, Eisenhut M, Bräutigam A et al (2017) The influence of a cryptochrome on the gene expression profile in the diatom *Phaeodactylum tricornutum* under blue light and in darkness. Plant Cell Physiol 58:1914–1923

Koop-Jakobsen K, Giblin AE (2010) The effect of increased nitrate loading on nitrate reduction via denitrification and DNRA in salt marsh sediments. Limnol Oceanog 55:789–802

Kostka JE, Prakash O, Overholt WA et al. (2011) Hydrocarbon-degrading bacteria and the bacterial community response in Gulf of Mexico beach sands impacted by the deepwater horizon oil spill. Appl Environ Microbiol 77:7962–7974.

Kube M, Chernikova T, Al-Ramahi Y et al. (2013) Genome sequence and functional genomic analysis of the oil-degrading bacterium *Oleispira antarctica*. Nat Commun 4:2156

Kuehn KA, Steiner D, Gessner MO (2004) Dielmineralization patterns of standing-dead plant litter: implications for CO2 flux from wetlands. Ecology 85:2504–2518

Kuypers MM, Sliekers AO, Lavik G et al (2003) Anaerobic ammonium oxidation by anammox bacteria in the Black Sea. Nature 422:608–611.

Kuypers MMM, Marchant HK, Kartal B (2018) The microbial nitrogen cycling network. Nat Rev Microbiol 16:263–276

Kwon KK, Oh JH, Yang S-H et al (2015) *Alcanivorax gelatiniphagus* sp. nov., a marine bacterium isolated from tidal flat sediments enriched with crude oil. Int J Syst Evol Microbiol 65:2204-2208

Lacerda ALDF, Proietti MC, Secchi ER et al (2020) Diverse groups of fungi are associated with plastics in the surface waters of the Western South Atlantic and the Antarctic Peninsula. Mol Ecol 29:1903–1918

Lai Q, Wang L, Liu Y, et al (2011) *Alcanivorax pacificus* sp. nov., isolated from a deep-sea pyrene-degrading consortium. Int J Syst Evol Microbiol 61:1370-1374

Lam P, Jensen MM, Lavik G et al (2007) Linking crenarchaeal and bacterial nitrification to anammox in the Black Sea. Proc Nat Acad Sci 104:7104-7109

Lam P, Lavik G, Jensen MM et al (2009) Revising the nitrogen cycle in the Peruvian oxygen minimum zone. Proc Natl Acad Sci USA 106:4752–4757

Lassen C, Ploug H, Jørgensen BB (1992) Microalgal photosynthesis and spectral scalar irradience in coastal marine sediments of Limfjorden, Denmark. Limnol Oceanogr 37:760-772

Laverman AM, Cappellen PV, Van Rotterdam-Los D et al (2006) Potential rates and pathways of microbial nitrate reduction in coastal sediments. FEMS Microbiol Ecol 58:179–192

Laverock B, Gilbert JA, Tait K et al (2011) Bioturbation: impact on the marine nitrogen cycle. Biochem Soc Trans 39:315-20

Laviale M, Barnett A, Ezequiel J et al (2015) Response of intertidal benthic microalgal biofilms to a coupled light–temperature stress: evidence for latitudinal adaptation along the Atlantic coast of Southern Europe. Environ Microbiol 17:3662–3677

Lazar CS, Baker BJ, Seitz K et al (2016) Genomic evidence for distinct carbon substrate preferences and ecological niches of Bathyarchaeota in estuarine sediments. Environ Microbiol 18:1200–1211

Lazar CS, Biddle JF, Meador TB et al (2014). Environmental controls on intragroup diversity of the uncultured benthic archaea of the miscellaneous Crenarchaeotal group lineage naturally enriched in anoxic sediments of the White Oak River estuary (North Carolina, USA). Environ Microbiol 17:2228–2238

Leahy JG, Colwell RR (1990) Microbial degradation of hydrocarbons in the environment. Microbiol Rev 54:305–315

Legge O, Johnson M, Hicks N et al (2020) Carbon on the Northwest European Shelf: Contemporary budget and future influences. Front Mar Sci 7:143

Levin LA, Boesch DF, Covich A et al (2001) The function of marine critical transition zones and the importance of sediment biodiversity. Ecosystems 4:430–451

Li J, Nedwell DB, Beddow J et al (2015a) *amoA* gene abundances and nitrification potential rates suggest that benthic ammonia-oxidizing bacteria (AOB) not archaea (AOA) dominate N cycling in the Colne estuary, UK. Appl Environ Microbiol 81:159-165

Li M, Baker BJ, Anantharaman K et al (2015b). Genomic and transcriptomic evidence for scavenging of diverse organic compounds by widespread deep-sea archaea. Nat Commun 6:8933

Li M, Cao H, Hong Y et al (2011b) Spatial distribution and abundances of ammonia-oxidizing archaea (AOA) and ammonia-oxidizing bacteria (AOB) in mangrove sediments. Appl Microbiol Biotechnol 89:1243–1254

Li M, Gu J (2013) Community structure and transcript responses of anammox bacteria, AOA, and AOB in mangrove sediment microcosms amended with ammonium and nitrite. Appl Microbiol Biotechnol 97:9859–9874

Li M, Hong Y-G, Cao H-L et al (2011a) Mangrove trees affect the community structure and distribution of anammox bacteria at an anthropogenic-polluted mangrove in the Pearl River Delta reflected by 16S rRNA and hydrazine oxidoreductase (HZO) encoding gene analyses. Ecotoxicology 20:1780–1790

Li Q, Wang F, Chen Z et al (2012) Stratified active archaeal communities in the sediments of Jiulong River estuary China. Front Microbiol 3:311-314

Lipsewers YA, Bale NJ, Hopmans EC et al (2014) Seasonality and depth distribution of the abundance and activity of ammonia oxidizing microorganisms in marine coastal sediments (North Sea). Front Microbiol 5:472–483

Lipsewers YA, Vasquez Cardenas D, Seitaj D et al (2017) Impact of seasonal hypoxia on activity and community structure of chemolithoautotrophic bacteria in a coastal sediment. Appl Environ Microbiol 83:e03517-16

Liu C, Shao Z (2005) *Alcanivorax dieselolei* sp. nov., a novel alkane degrading bacterium isolated from sea water and deep-sea sediment. Int J Syst Evol Microbiol 55:1181-1186

Liu J, Zheng Y, Lin H et al (2019) Proliferation of hydrocarbon-degrading microbes at the bottom of the Mariana Trench. Microbiome 7:47.

Liu J, Zhu S, Liu X et al (2020) Spatiotemporal dynamics of the archaeal community in coastal sediments: assembly process and co-occurrence relationship. ISME J 14:1463–1478

Liu X, Li M, Castelle CJ et al. (2018b) Insights into the ecology, evolution, and metabolism of the widespread Woesearchaeotal lineages. Microbiome 6:102

Liu X, Pan J, Liu Y (2018a) Diversity and distribution of Archaea in global estuarine ecosystems. Sci Total Environ 637-638:349–358

Liu XD, Tiquia SM, Holguin G (2003) Molecular diversity of denitrifying genes in continental margin sediments within the oxygen deficient zone off the Pacific coast of Mexico. Appl Environ Microbiol 69:3549–3560

Lloyd K, Schreiber GL, Petersen DG et al (2013) Predominant archaea in marine sediments degrade detrital proteins. Nature 496:215-220

Long RA, Eveillard D, Franco SLM et al (2013) Antagonistic interactions between heterotrophic bacteria as a potential regulator of community structure of hypersaline microbial mats. FEMS Microbiol Ecol 83:74–81

Lozada M, Marcos MS, Commendatore MG et al (2014) The bacterial community structure of hydrocarbon-polluted marine environments as the basis for the definition of an ecological index of hydrocarbon exposure. Microbes Environ 29:269–276

Luna GM, Corinaldesi C, Rastelli E et al (2013) Patterns and drivers of bacterial α- and β-diversity across vertical profiles from surface to subsurface sediments. Environ Microbiol Rep 5:731–739

Lyons JI, Alber M, Hollibaugh JT (2010) Ascomycete fungal communities associated with early decaying leaves of Spartina spp. from central California estuaries. Oecologia 162:435–42

Lyu Z, Liu Y (2018) Diversity and taxonomy of methanogens. In Biogenesis of Hydrocarbons. Handbook of Hydrocarbon and Lipid Microbiology: 2nd Edition. AJM Stams, DZ Sousa (eds.) Springer, Cham. 19-77

Maggi E, Rindi L, Dal Bello M et al (2017) Spatio-temporal variability in Mediterranean rocky shore microphytobenthos. Mar Ecol Prog Ser 575:17-29

Mann M, Serif M, Wrobel T et al (2020) The aureochrome photoreceptor PtAUREO1a is a highly effective blue light switch in diatoms, iScience doi: https://doi.org/10.1016/j.isci.2020.101730

Marchant R, Banat IM (2012) Microbial biosurfactants: challenges and opportunities for future exploitation. Trends Biotechnol. 30:558-565

Martens-Habbena W, Berube PM, Urakawa H et al (2009) Ammonia oxidation kinetics determine niche separation of nitrifying Archaea and Bacteria. Nature 461:976–979

Martinez-Perez C, Mohr W, Löscher CR et al. (2016) The small unicellular diazotrophic symbiont, UCYN-A, is a key player in the marine nitrogen cycle. Nat Microbiol 1:16163

Mason OU, Han J, Woyke T et al (2014) Single-cell genomics reveals features of a *Colwellia* species that was dominant during the Deepwater Horizon oil spill. Front Microbiol. 5:332

Massana R, Gobet A, Audic S et al (2015) Protist diversity in European coastal areas. Environ Microbiol 17:4035-4049

McGenity TJ (2014) Hydrocarbon biodegradation in intertidal wetland sediments. Curr Opin Biotechnol 27:46-54.

McGenity TJ, Folwell BD, McKew BA et al (2012) Marine crude-oil biodegradation: a central role for interspecies interactions. Aquat Biosyst 8:1–19

McGenity TJ, Oren A (2012) Hypersaline environments. In Life at Extremes: Environments, Organisms and Strategies for Survival. EM Bell (ed.) CAB International, UK. pp. 402-437

McGenity TJ, Sorokin D (2018) Methanogens and methanogenesis in hypersaline environments. In Biogenesis of Hydrocarbons. Handbook of Hydrocarbon and Lipid Microbiology: 2nd Edition. AJM Stams, DZ Sousa (eds.) Springer, Cham. 283-309

McGlynn SE (2017) Energy metabolism during anaerobic methane oxidation in ANME archaea. Microbes Environ 32:5–13

McKew BA, Coulon F, Osborn AM et al (2007) Determining the identity and roles of oil-metabolizing marine bacteria from the Thames estuary, UK. Environ Microbiol 9:165–176

McKew BA, Dumbrell A, Taylor JD et al (2013) Differences between aerobic and anaerobic degradation of microphytobenthic biofilm-derived organic matter within intertidal sediments. FEMS Microbiol Ecol 84:495 – 509

McKew BA, Taylor JD, McGenity TJ et al (2011) Resistance and resilience of benthic biofilm communities from a temperate saltmarsh to desiccation and rewetting. ISME J 5:30-41

McLachlan DH, Brownlee C, Taylor AR et al (2009) Light inducted motile responses of the estuarine benthic diatoms *Navicula perminuta* and *Cylindrotheca closterium* (Bacillariophyceae). J. Phycol 45:592–599

McLachlan DH, Underwood GJC, Taylor AR et al (2012) Calcium release from intracellular stores is necessary for the photophobic motility response in the benthic diatom *Navicula perminuta*. J. Phycol 48:675–681

McTavish H, Fuchs JA, Hooper AB (1993) Sequence of the gene coding for ammonia monooxygenase in *Nitrosomonas europaea*. J Bacteriol 175:2436–2444

Meckenstock RU, Mouttaki H (2011) Anaerobic degradation of non-substituted aromatic hydrocarbons. Curr Opin Biotechnol, 22:406-414

Megharaj M, Singleton I, McClure NC et al (2000) Influence of petroleum hydrocarbon contamination on microalgae and microbial activities in a long-term contaminated soil. Arch Environ Contam Toxicol 38:439–445

Méléder V, Savelli R, Barnett A et al (2020) Mapping the intertidal microphytobenthos gross primary production Part I: Coupling multispectral remote sensing and physical modeling. Front Mar Sci 7:520

Meng J, Xu J, Qin D et al (2014) Genetic and functional properties of uncultivated MCG archaea assessed by metagenome and gene expression analyses. ISME J 8:650-659

Messina E, Denaro R, Crisafi F et al (2016) Genome sequence of obligate marine polycyclic aromatic hydrocarbons-degrading bacterium *Cycloclasticus* sp. 78-ME, isolated from petroleum deposits of the sunken tanker Amoco Milford Haven, Mediterranean Sea. Mar. Genomics 25:11-13

Middelburg JJ, Barranguet C, Boschker HTS et al (2000) The fate of intertidal microphytobenthos carbon: An *in situ* ¹³C-labelling study. Limnol Oceanogr 45:1224-1334

Miyatake T, Moerdijk-Poortvliet TCW, Stal LJ et al (2014) Tracing carbon flow from microphytobenthos to major bacterial groups in an intertidal marine sediment by using an *in situ* ¹³C pulse-chase method. Limnol Oceanogr 59:1275–1287

Moerdijk-Poortvliet TCW, Beauchard O, Stal LJ et al (2018b) Production and consumption of extracellular polymeric substances in an intertidal diatom mat. Mar Ecol Prog Ser 592:77–95

Moerdijk-Poortvliet TCW, van Breugel P, Sabbe K et al (2018a) Seasonal changes in the biochemical fate of carbon fixed by benthic diatoms in intertidal sediments. Limnol Oceanogr 63: 550–569

Mohamed D, Martiny J (2011) Patterns of fungal diversity and composition along a salinity gradient. ISME J 5:379–388

Mommer L, Cotton TEA, Raaijmakers JM et al (2018) Lost in diversity: the interactions between soil-borne fungi, biodiversity and plant productivity. New Phytol 218:542–553

Monson DH, Doak DF, Ballachey BE, et al (2000) Long-term impacts of the Exxon Valdez oil spill on sea otters, assessed through age-dependent mortality patterns. PNAS 97(12):6562-6567

Moreira-Coello V, Mouriño-Carballido B, Marañón E et al (2019) Temporal variability of diazotroph community composition in the upwelling region off NW Iberia. Sci Rep 9:3737

Moriarty DJW, O'Donohue MJ (1993) Nitrogen fixation in seagrass communities during summer in the Gulf of Carpentaria, Australia. Aust J Mar Freshw Res 44:117-125

Mouget J-L, Perkins R, Consalvey M et al (2008) Migration or photoacclimation to prevent high irradiance and UV-B damage in marine microphytobenthic communities. Aquat Microb Ecol 52:223-232

Mueller P, Granse D, Nolte S et al (2017) Top-down control of carbon sequestration: grazing affects microbial structure and function in salt marsh soils. Ecol Appl 27:1435-1450

Munson MA, Nedwell DB, Embley TM (1997) Phylogenetic diversity of Archaea in sediment samples from a coastal salt marsh. Appl Environ Microbiol 63:4729–4733

Myllykangas JP, Rissanen AJ, Hietanen S et al (2020) Influence of electron acceptor availability and microbial community structure on sedimentary methane oxidation in a boreal estuary. Biogeochemistry 148:291-309

National Research Council (NRC) (2003) Oil in the sea III: Inputs, fates, and effects. Washington, D.C: National Academy Press.

Nedwell D, Aziz S (1980) Heterotrophic nitrogen fixation in an intertidal salt marsh sediment. Est Coast Mar Sci 10:699-702

Nedwell DB, Embley TM, Purdy KJ (2004) Sulphate reduction, methanogenesis and phylogenetics of the sulphate reducing bacterial communities along an estuarine gradient. Aquat Microb Ecol 37:209–217

Nedwell DB, Underwood GJC, McGenity TJ et al (2016) The Colne Estuary: A Long-Term Microbial Ecology Observatory. Adv Ecol Res 55:227-281

Newell SY (1996) Established and potential impacts of eukaryotic mycelial decomposers in marine/terrestrial ecotones. J Exp Mar Biol Ecol 200:187–206

Newton R, Huse SM, Morrison HG et al (2013). Shifts in the microbial community composition of Gulf Coast beaches following beach oiling. PLoS ONE 8:e74265

Nogales B, Bosch R (2019) Microbial communities in hydrocarbon-polluted harbors and marinas. In: McGenity T. (eds) Microbial Communities Utilizing Hydrocarbons and Lipids:

Members, Metagenomics and Ecophysiology. Handbook of Hydrocarbon and Lipid Microbiology. Springer, Cham. 63-80

Nogales B, Timmis KN, Nedwell DB et al (2002) Detection and diversity of expressed denitrification genes in estuarine sediments after Reverse Transcription-PCR Amplification from mRNA. Appl Environ Microbiol 68:5017-5025

Notman GM, McGill RAR, Hawkins SJ et al (2016) Macroalgae contribute to the diet of *Patella vulgata* from contrasting conditions of latitude and wave exposure in the UK. Mar Ecol Prog Ser 549:113-123

Oakes JM, Eyre BD, Middelburg JJ (2012) Transformation and fate of microphytobenthos carbon in subtropical shallow subtidal sands: A ¹³C-labeling study. Limnol Oceanogr 57:1846-1856

Oakes JM, Eyre BD, Middelburg JJ et al (2010) Composition, production, and loss of carbohydrates in subtropical shallow subtidal sandy sediments: Rapid processing and long-term retention revealed by ¹³C-labeling. Limnol Oceanogr 55:2126–2138

Oppenheim DR (1991) Seasonal changes in epipelic diatoms along an intertidal shore, Berrow flats, Somerset. J Mar Biol Assoc UK 71:579-596

Oremland R, Marsh L, Polcin S. (1982) Methane production and simultaneous sulphate reduction in anoxic, salt marsh sediments. Nature 296:143–145

Orsi WD, Vuillemin A, Rodriguez P et al (2019) Metabolic activity analyses demonstrate that Lokiarchaeon exhibits homoacetogenesis in sulfidic marine sediments. Nat Microbiol 5:248–255

Oshiki M, Shimokawa M, Fujii N et al (2011) Physiological characteristics of the anaerobic ammonium-oxidizing bacterium 'Candidatus Brocadia sinica'. Microbiology 157:1706-1713

Osterholz H, Kirchman DL, Niggemann J et al (2018) Diversity of bacterial communities and dissolved organic matter in a temperate estuary. FEMS Microbiol Ecol 94: fiy119

Overy DP, Rämä T, Oosterhuis R et al (2019) The Neglected Marine Fungi, Sensu stricto, and Their Isolation for Natural Products' Discovery. Mar Drugs 17:42

Oxborough K, Hanlon ARM, Underwood GJC et al (2000) *In vivo* estimation of the photosystem II photochemical efficiency of individual microphytobenthic cells using high-resolution imaging of chlorophyll *a* fluorescence. Limnol Oceanogr 45:1420-1425

Pace NR (1997) A molecular view of microbial diversity and the biosphere. Science 276:734–740

Païssé S, Coulon F, Goñi-Urriza M et al (2008) Structure of bacterial communities along a hydrocarbon contamination gradient in a coastal sediment. FEMS Microbiol Ecol 66:295-305.

Pajares S, Ramos R (2019). Processes and microorganisms involved in the marine nitrogen cycle: Knowledge and gaps. Front Mar Sci 6:739

Palomo A, Pedersen AG, Fowler SJ et al (2018) Comparative genomics sheds light on niche differentiation and the evolutionary history of comammox *Nitrospira*. ISME J 12:1779–1793

Papaspyrou S, Smith CJ, Dong LF et al (2014) Nitrate reduction functional genes and nitrate reduction potentials persist in deeper estuarine sediments Why? PLoS One 9:e94111

Park J, Kwon B-O, Kim M et al (2014) Microphytobenthos of Korean tidal flats: A review and analysis on floral distribution and tidal dynamics Ocean Coast Manag 102:471-482

Passarelli C, Cui X, Valsami-Jones E et al (2020) Environmental context determines the impact of titanium oxide and silver nanoparticles on the functioning of intertidal microalgal biofilms. Environ Sci Nano 7:3020-3035

Peletier H (1996) Long-term changes in intertidal estuarine diatom assemblages related to reduced input of organic waste. Mar Ecol Prog Ser 137:265-271

Penton CR, Devol AH, Tiedje JM (2006) Molecular evidence for the broad distribution of anaerobic ammonium-oxidizing bacteria in freshwater and marine sediments. Appl Environ Microbiol 72:6829–6832

Pérez-Burillo J, Trobajo R, Vasselon Vet al (2020) Evaluation and sensitivity analysis of diatom DNA metabarcoding for WFD bioassessment of Mediterranean rivers. Sci Total Environ 727:138445

Perkins EJ (1960) The diurnal rhythm of the littoral diatoms of the River Eden estuary, Fife. J Ecol 48:725-728

Perkins RG, Lavaud J, Serôdio J et al. (2010) Vertical cell movement is a primary response of intertidal benthic biofilms to increasing light dose. Mar Ecol Prog Ser 416:93–103

Perkins RG, Oxborough K, Hanlon ARM et al (2002) Can chlorophyll fluorescence be used to estimate the rate of photosynthetic electron transport within microphytobenthic biofilms? Mar Ecol Prog Ser 228:47-56

Perkins RG, Underwood GJC, Brotas V et al (2001) Responses of microphytobenthos to light: primary production and carbohydrate allocation over an emersion period, Mar Ecol Prog Ser 223:101-112

Pester M, Rattei T, Flechl S et al (2011) *amoA*-based consensus phylogeny of ammonia-oxidizing archaea and deep sequencing of *amoA* genes from soils of four different geographic regions. Environ Microbiol 14:525-39.

Phillips CJ, Smith Z, Embley TM et al (1999) Phylogenetic differences between particle-associated and planktonic ammonia-oxidizing bacteria of the beta-subdivision of the class Proteobacteria in the northwestern Mediterranean Sea. Appl Environ Microbiol 65:779–786

Piatt JF, Ford, RG (1996) How many seabirds were killed by the Exxon Valdez oil spill? Am Fish Soc Symp 18:712-719

Piña-Ochoa E, Høgslund S, Geslin E et al (2010) Widespread occurrence of nitrate storage and denitrification among Foraminifera and Gromiida. Proc Natl Acad Sci USA 107:1148-1153

Pinckney JL (2018) A mini-review of the contribution of benthic microalgae to the ecology of the continental shelf in the South Atlantic Bight. Estuaries Coast 41:2070-2078

Pitcher A, Wuchter, C, Siedenberg K et al (2011) Crenarchaeol tracks winter blooms of ammonia-oxidizing Thaumarchaeota in the coastal North Sea. Limnol Oceanogr 56:2308–2318

Pjevac P, Schauberger C, Poghosyan L et al (2017) *AmoA*-targeted polymerase chain reaction primers for the specific detection and quantification of comammox Nitrospira in the environment. Front Microbiol 8:11

Plante C, Fleer V, Jones ML (2016) Neutral processes and species sorting in benthic microalgal community assembly: effects of tidal resuspension. J Phycol 52: 827–839

Plante CJ, Hill-Spanik K, Cook M et al (2020) Environmental and Spatial Influences on Biogeography and Community Structure of Saltmarsh Benthic <u>diatoms</u>. Estuaries Coast https://doi.org/10.1007/s12237-020-00779-0

Poole NJ, Price PC (1972) Fungi colonizing wood submerged in the medway estuary, T Brit Mycoll Soc 59:2

Porubsky W P, Weston NB, Joye SB (2009) Benthic metabolism and the fate of dissolved inorganic nitrogen in intertidal sediments. Estuar Coast Shelf Sci 83:392–402

Preisler A, de Beer D, Lichtschlag A et al (2007) Biological and chemical sulfide oxidation in a *Beggiatoa* inhabited marine sediment. ISME J. 1:341–353

Prince RC, Amande TJ, McGenity TJ (2018) Prokaryotic hydrocarbon degraders. In: Taxonomy, Genomics and Ecophysiology of Hydrocarbon-Degrading Microbes, T.J. McGenity, ed. Springer Nature, Cham, pp. 1–41.

Prins A, Deleris P, Hubas C, Jesus B (2020) Effect of light intensity and light quality on diatom behavioral and physiological photoprotection. Front Mar Sci 7:203

Prokopenko MG, Sigman DM, Berelson WM et al (2011) Denitrification in anoxic sediments supported by biological nitrate transport. Geochim Cosmochim Acta 75:7180–7199

Purdy KJ, Cresswell-Maynard TD, Nedwell DB et al (2004) Isolation of haloarchaea that grow at low salinities. Environ Microbiol 6:591-595

Qian G, Wang J, Kan J et al (2018) Diversity and distribution of anammox bacteria in water column and sediments of the eastern Indian Ocean. Int Biodeterior Biodegr 133:52–62

Quan ZX, Rhee SK, Zuo JE et al (2008) Diversity of ammonium-oxidizing bacteria in a granular sludge anaerobic ammonium-oxidizing (anammox) reactor. Environ Microbiol 10:3130-3139

Rabus R, Boll M, Heider J et al (2016) Anaerobic microbial degradation of hydrocarbons: From enzymatic reactions to the environment. J Mol Microbiol Biotechnol; 26:5-28

Radajewski S, Webster G, Reay DS et al (2002) Identification of active methylotroph populations in an acidic forest soil by stable isotope probing. Microbiology 148:2331–2342

Radwan SS, Khanafer MM, Al-Awadhi HA (2019) Ability of the so-called obligate hydrocarbonoclastic bacteria to utilize nonhydrocarbon substrates thus enhancing their activities despite their misleading name. BMC Microbiol 19:41

Raymond J, Siefert JL, Staples CR et al (2004) The natural history of nitrogen fixation, Mol Biol Evol 21:541–554

Reddy CM, Eglinton TI, Hounshell A et al (2002) The West Falmouth oil spill after thirty years: the persistence of petroleum hydrocarbons in marsh sediments. Environ Sci Technol 36: 4754–4760

Redzuan NS, Underwood GJC (2021) The importance of weather and tides on the resuspension and deposition of microphytobenthos (MPB) on intertidal mudflats. Estuar Coastal Shelf Sci (in press)

Redzuan NS, Underwood GJC (2020) Movement of microphytobenthos and sediment between mudflats and salt marsh during spring tides. Front Mar Sci 7:496

Ribeiro L, Brotas V, Hernández-Fariñas T et al (2020) Assessing alternative microscopy-based approaches to species abundance description of intertidal diatom communities. Front Mar Sci 7:36

Ribeiro L, Brotas V, Mascarell G et al (2003) Taxonomic survey of the microphytobenthic communities of two Tagus estuary mudflats. Acta Oecologica 24:S117-S123

Ribeiro L, Brotas V, Rincé Y et al (2013) Structure and diversity of intertidal benthic diatom assemblages in contrasting shores: a case study from the Tagus estuary. J Phycol 49:258 - 270

Risgaard-Petersen N (2003) Coupled nitrification-denitrification in autotrophic and heterotrophic estuarine sediments: On the influence of benthic microalgae. Limnol Oceanogr 48:93-105

Risgaard-Petersen N, Langezaal A, Ingvardsen S et al (2006) Evidence for complete denitrification in a benthic foraminifer. Nature 443:93–96

Rishworth GM, Dodd C, Perissinotto R et al (2020) Modern supratidal microbialites fed by groundwater: functional drivers, value and trajectories. Earth Sci Rev 210:103364.

Rivera-Monroy VH, Lenaker P, Twilley RR et al (2010) Denitrification in coastal Louisiana: A spatial assessment and research needs. J Sea Res 63:157-172

Rodriguez-R LM, Overholt WA, Hagan C et al (2015) Microbial community successional patterns in beach sands impacted by the Deepwater Horizon oil spill. ISME J. 9 1928–1940

Rojas-Jimenez K, Rieck A, Wurzbacher C et al (2019) A salinity threshold separating fungal communities in the Baltic Sea. Front Microbiol. 10:680

Round FE (1979) Occurrence and rhythmic behaviour of *Tropidoneis lepidoptera* in the epipelon of Barnstable Harbor, Massachusetts, USA. Mar Biol 54:215-217

Round FE, Palmer JD (1966) Persistent, vertical-migration rhythms in benthic microflora. II. Field and laboratory studies on diatoms from the banks of the River Avon. J Mar Biol Assoc UK 46:191-214

Rovira L, Trobajoa R, Satob S et al (2015) Genetic and physiological diversity in the diatom *Nitzschia inconspicua*. J Eukaryot Microbiol 62:815–832

Rubin-Blum M, Antony CP, Borowski C et al (2017). Short-chain alkanes fuel mussel and sponge *Cycloclasticus* symbionts from deep-sea gas and oil seeps. *Nat Microbiol* 2:17093

Sabbe K (1993) Short-term fluctuations in benthic diatom numbers on an intertidal sandflat in the Westerschelde Estuary (Zeeland, The Netherlands). Hydrobiol 269/270:275-284

Sahan E, Muyzer G (2008) Diversity and spatio-temporal distribution of ammonia-oxidizing Archaea and Bacteria in sediments of the Westerschelde estuary. FEMS Microbiol Ecol 64:175–186

Sahan E, Sabbe K, Creach V et al (2007) Community structure and seasonal dynamics of diatom biofilms and associated grazers in intertidal mudflats. Aquat Microb Ecol 47:253-266

Sanni GO, Coulon F, McGenity TJ (2015) Dynamics and distribution of bacterial and archaeal communities in oil-contaminated temperate coastal mudflat mesocosms. Environ Sci Pollut Res Int 22:15230-47.

Savelli R, Dupuy C, Barille L et al (2018) On biotic and abiotic drivers of the microphytobenthos seasonal cycle in a temperate intertidal mudflat: a modelling study. Biogeosciences 15:7243-7271

Savelli R, Méléder V, Cugier P et al (2020) Mapping the intertidal microphytobenthos gross primary production, Part II: Merging remote sensing and physical-biological coupled modeling. Front Mar Sci 7:521

Savelli R, Bertin X, Orvain F et al (2019) Impact of chronic and massive resuspension mechanisms on the microphytobenthos dynamics in a temperate intertidal mudflat. J Geophys Res: 124:3752-3777

Scarlett K, Denman S, Clark DR et al (2020) Relationships between nitrogen cycling microbial community abundance and composition reveal the indirect effect of soil pH on oak decline. ISME J https://doi.org/10.1038/s41396-020-00801-0

Schmid M, Walsh K, Webb R et al (2003) *Candidatus "Scalindua brodae*", sp. nov., *Candidatus "Scalindua wagneri*", sp. nov., two new species of anaerobic ammonium oxidizing bacteria. Syst Appl Microbiol 26:529-38

Schneiker S, Dos Santos VAPM, Bartels D et al (2006) Genome sequence of the ubiquitous hydrocarbon-degrading marine bacterium *Alcanivorax borkumensis*. Nat Biotechnol 24: 997–1004

Seitz KW, Lazar CS, Hinrichs K-U et al (2016) Genomic reconstruction of a novel, deeply branched sediment archaeal phylum with pathways for acetogenesis and sulfur reduction. ISME J 10:1696–1705

Seitzinger SP (1988). Denitrification in fresh-water and coastal marine ecosystems-ecological and geochemical significance. Limnol Oceanogr 33:702–724

Serôdio J, Marques da Silva J, Catarino F (1997) Nondestructive tracing of migratory rhythms of intertidal benthic microalgae using in vivo chlorophyll a fluorescence. J Phycol 33:542-553

Seyler LM, McGuinness LM, Kerkhof LJ (2014) Crenarchaeal heterotrophy in salt marsh sediments ISME J 8:1534-1543

Shi Y, Jiang YY, Wang SY et al (2020) Biogeographic distribution of comammox bacteria in diverse terrestrial habitats. Sci Total Environ 717:137257

Sikkema J, de Bont JAM, Poolman B (1995) Mechanisms of membrane toxicity of hydrocarbons. Microbiol Rev 59(2): 201-222

Smith CJ, Nedwell DB, Dong LF et al (2007) Diversity and abundance of nitrate reductase genes (*narG* and *napA*), nitrite reductase genes (*nirS* and *nrfA*), and their transcripts in estuarine sediments. Appl Environ Microbiol 73:3612-3622

Smith DJ, Underwood GJC (1998) Exopolymer production by intertidal epipelic diatoms. Limnol Oceanogr 43:1578-1591

Smith JM, Cascotti KL, Chavez FP et al (2014a) Differential contributions of archaeal ammonia oxidizer ecotypes to nitrification in coastal surface waters. ISME J 8:1704-1714

Smith JM, Chavez FP, Francis CA (2014b) Ammonium uptake by phytoplankton regulates nitrification in the sunlit ocean. PLoS One 9:e108173.

Smith SE, Read DJ (2008) Mycorrhizal symbiosis, 3rd edn. San Diego, CA, USA: Academic Press.

Sonthiphand P, Hall MW, Neufeld JD (2014) Biogeography of anaerobic ammonia-oxidizing (anammox) bacteria. Front Microbiol 5:399

Sorokin DY, Makarova K, Abbas B, et al (2017) Discovery of extremely halophilic methylreducing euryarchaea provides insight into the evolutionary origin of methanogenesis. Nat Microbiol 2:17081

Sorokin DY, Merkel AY, Abbas B et al (2018) *Methanonatronarchaeum thermophilum* gen. nov., sp. nov. and '*Candidatus* Methanohalarchaeum thermophilum', extremely halo(natrono)philic methyl-reducing methanogens from hypersaline lakes comprising a new euryarchaeal class Methanonatronarchaeia classis nov. Int J Syst Evol Microbiol. 68:2199–2208

Spang A, Saw JH, Jørgensen SL et al (2015) Complex Archaea that bridge the gap between prokaryotes and eukaryotes. Nature 521:173–179

Spilmont N, Seuront L, Meziane T et al (2011) There's more to the picture than meets the eye: Sampling microphytobenthos in a heterogeneous environment. Estuar Coast Shelf Sci 95:470-476

Staats N, Stal LJ, Mur LR (2000). Exopolysaccharide production by the epipelic diatom Cylindrotheca closterium: Effects of nutrient conditions. J Exp Mar Biol Ecol 249:3–27

Stal LJ (2016) Coastal sediments: Transition from land to sea. In, LJ Stal and MS Cretoiu (eds.), *The Marine Microbiome*, Springer International Publishing Switzerland, pp 283-304

Stal LJ, Bolhuis H, Cretoiu MS (2019) Phototrophic marine benthic microbiomes: the ecophysiology of these biological entities. Environ Microbiol 21:1529–1551

Steele DJ, Franklin DJ, Underwood GJC. (2014) Protection of cells from salinity stress by extracellular polymeric substances in diatom biofilms. Biofouling 30:987-998

Stehr G, Biittcher B, Dittberner P et al (1995) The ammonia-oxidizing nitrifying population of the River Elbe estuary. FEMS Microbiol Ecol 17:177–186

Stein LY, Arp DJ, Berube PM et al (2007) Whole-genome analysis of the ammonia-oxidizing bacterium, *Nitrosomonas eutropha* C91: implications for niche adaptation. Environ Microbiol 9:2993–3007

Stock W, Blommaert L, De Troch M et al (2019) Host specificity in diatom-bacteria interactions alleviates antagonistic effects, FEMS Microbiol Ecol 95: fiz171

Stoecker K, Bendinger B, Schoning B et al (2006) Cohn's *Crenothrix* is a filamentous methane oxidizer with an unusual methane monooxy- genase. Proc Natl Acad Sci USA 103:2363–2367

Strous M, Fuerst JA, Kramer EH et al (1999) Missing lithotroph identified as new planctomycete. Nature 400:446–449

Strous M, Pelletier E, Mangenot S et al (2006) Deciphering the evolution and metabolism of an anammox bacterium from a community genome. Nature 440:790–794

Sullivan, MJ (1999) Applied diatom studies in estuarine and shallow coastal environments. In The Diatoms: Applications for the Environmental and Earth Sciences (Stoermer, EF and Smol, JP, editors). 334-351. Cambridge University Press, Cambridge

Sun Z, Li G, Wang C et al (2014) Community dynamics of prokaryotic and eukaryotic microbes in an estuary reservoir. Sci Rep 4:6966

Tal Y, Watts JE, Schreier HJ (2005) Anaerobic ammonia-oxidizing bacteria and related activity in Baltimore inner harbor sediment. Appl Environ Microbiol 71:1816–1821

Taylor JD, McKew BA, Kuhl A et al (2013) Microphytobenthic extracellular polymeric substances (EPS) in intertidal sediments fuel both generalist and specialist EPS-degrading bacteria. Limnol Oceanogr 58:1463-1480

Teramoto M, Ohuchi M, Hatmanti A et al (2011) *Oleibacter marinus* gen. nov., sp. nov., a novel bacterium that degrades petroleum aliphatic hydrocarbons in the tropical marine environment. Int J Syst Evol Microbiol 61:375-380

Terrisse F, Cravo-Laureau C, Noël C et al (2017) Variation of oxygenation conditions on a hydrocarbonoclastic microbial community reveals *Alcanivorax* and *Cycloclasticus* ecotypes. Front Microbiol 8:1549

Thamdrup B, Dalsgaard T (2002) Production of N_2 through anaerobic ammonium oxidation coupled to nitrate reduction in marine sediments. Appl Environ Microbiol 68:1312–1318

Thauer RK, Kaster A-K, Seedorf H et al (2008) Methanogenic archaea: ecologically relevant differences in energy conservation. Nat Rev Microbiol 6:579–591

Thomas GE, Cameron TC, Campo P et al (2020) Bacterial community legacy effects following the Agia Zoni II oil-spill, Greece. Front Microbiol 11:1706

Thornton DCO, Dong LF, Underwood GJC et al (2002) Factors affecting microphytobenthic biomass, species composition and production in the Colne estuary (UK). Aquatic Microb Ecol 27:285-300

Thornton DCO, Underwood GJC, Nedwell DB (1999) Effect of illumination and emersion period on the exchange of ammonium across the estuarine sediment-water interface. Mar Ecol Prog Ser 184:11-20

Treusch AH, Leininger S, Kletzin A et al (2005) Novel genes for nitrite reductase and Amorelated proteins indicate a role of uncultivated mesophilic crenarchaeota in nitrogen cycling. Environ Microbiol 7:1985–1995

Trimmer M, Engström, P (2011) Distribution, activity, and ecology of anammox bacteria in aquatic environments, in Nitrification, eds BB Ward, DJ Arp, and MG Klotz, (Washington, D.C: ASM Press), 201–235

Trimmer M, Nicholls J, Deflandre B (2003) Anaerobic ammonium oxidation measured in sediments along the Thames estuary, United Kingdom. Appl Environ Microbiol 69:6447–6454

Trimmer M, Nicholls JC, Morley N et al (2005) Biphasic behavior of anammox regulated by nitrite and nitrate in an estuarine sediment. Appl Environ Microbiol 71:1923–1930

Tsui CKM, Hyde KD (2004) Biodiversity of fungi on submerged wood in a stream and estuary in the Tai Ho Bay, Hong Kong. Fungal Divers 15:205–220

Underwood GJC (1994) Seasonal and spatial variation in epipelic diatom assemblages in the Severn estuary. Diatom Res 9:451-472

Underwood, GJC (2002) Adaptations of tropical marine microphytobenthic assemblages along a gradient of light and nutrient availability in Suva Lagoon, Fiji. Eur J Phycol 37:449–462

Underwood GJC (2005) Microalgal (microphytobenthic) biofilms in shallow coastal waters: how important are species? Proc Calif Acad Sci 56:162-169

Underwood GJC, Barnett M (2006) What determines species composition in microphytobenthic biofilms? Functioning of microphytobenthos in estuaries. Proceedings of the Microphytobenthos symposium, Amsterdam, The Netherlands, August 2003. Ed. J Kromkamp. pp 121-138. Royal Netherlands Academy of Arts and Sciences.

Underwood GJC, Kromkamp J (1999) Primary production by phytoplankton and microphytobenthos in estuaries. Adv Ecol Res 29:93–153

Underwood GJC, Michel C, Meisterhans G et al (2019) Organic matter from Arctic sea ice loss alters bacterial community structure and function. Nat Clim Change 9:170–176

Underwood GJC, Paterson DM (2003) The importance of extracellular carbohydrate production by marine epipelic diatoms. Adv Bot Res 40:184-240

Underwood GJC, Perkins RG, Consalvey M et al (2005) Patterns in microphytobenthic primary productivity: Species-specific variation in migratory rhythms and photosynthetic efficiency in mixed-species biofilms. Limnol Oceanogr 50:755-767

Underwood GJC, Phillips J, Saunders K (1998) Distribution of estuarine benthic diatom species along salinity and nutrient gradients. Eur J Phycol 33:173-183

Underwood, GJC, Provot L (2000) Determining the environmental preferences of four estuarine epipelic diatom taxa – growth across a range of salinity, nitrate and ammonium conditions. Eur J Phycol 35:173–182

Urakawa H, Rajan S, Feeney ME et al (2019) Ecological response of nitrification to oil spills and its impact on the nitrogen cycle. Environ Microbiol 21(1):18–33.

Van Bleijswijk J, Muyzer G (2004) Genetic diversity of oxygenic phototrophs in microbial mats exposed to different levels of oil pollution. Ophelia 58:157–164

van der Heijden MGA. 2002. Arbuscular mycorrhizal fungi as a determinant of plant diversity: in search for underlying mechanisms and general principles. In: MGA van der Heijden, IR Sanders, eds. Mycorrhizal Ecology. Berlin, Germany: Springer-Verlag, 243–266.

van der Wal D, Wielemaker-van den Dool A., Herman, PMJ (2010) Spatial synchrony in intertidal benthic algal biomass in temperate coastal and estuarine ecosystems. Ecosystems, 13:338–351

Van Kessel MAHJ, Speth DR, Albertsen M et al (2015) Complete nitrification by a single microorganism. Nature 528:555–559.

Van Niekerk L, Adams JB, Bate GC et al (2013) Country-wide assessment of estuary health: an approach for integrating pressures and ecosystem response in a data limited environment. Estuar Coast Shelf Sci 130:239-251

Van Niekerk L, Adams JB, Lamberth SJ et al (eds) 2019. South African National Biodiversity Assessment 2018: Technical Report. Volume 3: Estuarine Realm. CSIR report number CSIR/SPLA/EM/EXP/2019/0062/A. South African National Biodiversity Institute, Pretoria. Report Number: http://hdl.handle.net/20.500.12143/6373

Vanelslander B, De Wever A, Van Oostende N et al (2009) Complementarity effects drive positive diversity effects on biomass production in experimental benthic diatom biofilms. J Ecol 97:1075–1082

Vanelslander B, Paul C, Grueneberg J et al (2012) Daily bursts of biogenic cyanogen bromide (BrCN) control biofilm formation around a marine benthic diatom. Proc Nat Acad Sci USA 109:2412-2417

Vanwonterghem I, Evans PN, Parks DH et al (2016) Methylotrophic methanogenesis discovered in the archaeal phylum Verstraetearchaeota. Nat Microbiol 1:16170

Venter JC, Remington K, Heidelberg JF et al (2004) Environmental genome shotgun sequencing of the Sargasso Sea. Science 304:66–74

Vetriani C. Reysenbach A.-L. Dore J. (1998) Recovery and phylogenetic analysis of archaeal rRNA sequences from continental shelf sediments. FEMS Microbiol Lett 161:83–88

Waltham NJ, Elliott M, Lee SY et al (2020) UN decade on ecosystem restoration 2021-2030 – What chance for success in restoring coastal ecosystems? Front Mar Sci doi 10.3389/Fmars.2020.00071

Wang W, Shao Z (2013) Enzymes and genes involved in aerobic alkane degradation. Front Microbiol 4:116

Wang W, Shao Z (2014) The long-chain alkane metabolism network of *Alcanivorax dieselolei*. Nat Commun 5:5755

Wang W, Tao J, Liu H et al (2020). Contrasting bacterial and archaeal distributions reflecting different geochemical processes in a sediment core from the Pearl River Estuary. AMB Expr 10:16

Wang W, Wang L, Shao Z (2018) Polycyclic Aromatic Hydrocarbon (PAH) degradation pathways of the obligate marine PAH degrader *Cycloclasticus* sp. Strain P1. Appl Environ Microbiol 84:e01261-18

Wang Y, Feng Y, Ma X et al (2013) Seasonal dynamics of ammonia/ammonium-oxidizing prokaryotes in oxic and anoxic wetland sediments of subtropical coastal mangrove. Appl Microbiol Biotechnol 97:7919–7934

Wang Y, Lau PC, Button DK. (1996) A marine oligobacterium harboring genes known to be part of aromatic hydrocarbon degradation pathways of soil pseudomonads. Appl Environ Microbiol. 62:2169-73

Wang YQ, Sen K, He YD et al (2019) Impact of environmental gradients on the abundance and diversity of planktonic fungi across coastal habitats of contrasting trophic status. Sci Total Environ 683:822–833

Wankel SD, Mosier AC, Hansel CM et al (2010). Spatial variability in nitrification rates and ammonia-oxidizing microbial communities in the agriculturally impacted Elkhorn Slough estuary, California. Appl Environ Microbiol 77:269-280

Waring J, Baker NR, Underwood GJC (2007) Responses of estuarine intertidal microphytobenthic algal assemblages to enhanced UV-B radiation. Global Change Biol 13:1398–1413

Waring J, Klenell M, Bechtold U et al (2010) Light-induced responses of oxygen photoreduction, reactive oxygen species production and scavenging in two diatom species. J Phycol 46:1206 – 1217

Webster G, O'Sullivan LA, Meng Y et al (2015) Archaeal community diversity and abundance changes along a natural salinity gradient in estuarine sediments. FEMS Microbiol Ecol 91:1-18

Webster G, Rinna J, Roussel EG et al (2010) Prokaryotic functional diversity in different biogeochemical depth zones in tidal sediments of the Severn Estuary, UK, revealed by stable-isotope probing. FEMS Microbiol Ecol 72:179-197

Weerman, EJ, Van Belzen J, Rietkerk M et al (2012) Changes in diatom patch-size distribution and degradation in a spatially self-organized intertidal mudflat ecosystem. Ecology 93:608-618

Weisman WE (1998) Analysis of petroleum hydrocarbons in environmental media. Massachusetts: Amherst Scientific Publishers.

Wen X, Yang S, Horn F et al (2017) Global biogeographic analysis of methanogenic archaea identifies community-shaping environmental factors of natural environments. Front Microbiol 8:1339

Whitby C, Saunders JR, Pickup RW et al (2001) A comparison of ammonia-oxidiser populations in eutrophic and oligotrophic basins of a large freshwater lake. Ant Van Leeuwenhoek 79:179-201

Whitby C, Saunders JR, Pickup RW et al (1999) Phylogenetic differentiation of two closely related *Nitrosomonas* spp. that inhabit different sediment environments in an oligotrophic freshwater lake. Appl Environ Microbiol 65:4855-4862

Whiting, GJ, Gandy EL, Yoch DC (1986) Tight coupling of root-associated nitrogen fixation and plant photosynthesis in the salt marsh grass *Spartina alterniflora* and carbon dioxide enhancement of nitrogenase activity. Appl Environ Microbiol 52:108-113

Whyte LG, Bourbonniere L, Greer CW (1997) Biodegradation of petroleum hydrocarbons by psychrotrophic *Pseudomonas* strains possessing both alkane (alk) and naphthalene (nah) catabolic pathways. Appl Environ Microbiol 63:3719-3723

Widdel F, Musat F (2010) Energetic and other quantitative aspects of microbial hydrocarbon utilization. In Handbook of Hydrocarbon and Lipid Microbiology. Edited by Timmis KN, McGenity TJ, van der Meer JR, de Lorenzo V. Berlin Heidelberg: Springer: 731-763

Williams TA, Cox CJ, Foster PG et al (2020) Phylogenomics provides robust support for a two-domains tree of life. Nature Ecol Evol 4:138–147

Wilms R, Sass H, Köpke B et al (2006) Specific Bacterial, Archaeal, and Eukaryotic communities in tidal-flat sediments along a vertical profile of several meters. Appl Environ Microbiol 72:2756-2764

Wilms R, Sass H, Köpke B et al (2007) Methane and sulfate profiles within the sub-surface of a tidal flat are reflected by the distribution of sulfate-reducing bacteria and methanogenic archaea. FEMS Microbiol Ecol 59:611–621

Witkowski A, Lange-Bertalot H, Metzeltin D (2000) Diatom flora of marine coasts. Iconographia diatomologica v. 7. Ganter, Ruggell. Koeltz Scientific Books. Königstein, Germany

Woebken D, Lam P, Kuypers MM et al (2009) A microdiversity study of anammox bacteria reveals a novel *Candidatus Scalindua* phylotype in marine oxygen minimum zones. Environ Microbiol 10:3106–3119

Wuchter C, Abbas B, Coolen MJ et al (2006) Archaeal nitrification in the ocean. Proc Natl Acad Sci USA 103:12317–12322

Xia F, Wang JG, Zhu T et al (2018) Ubiquity and diversity of complete ammonia oxidizers (Comammox). Appl Environ Microbiol 84:e01390-18

Yakimov MM, Giuliano L, Denaro R et al (2004) *Thalassolituus oleivorans* gen. nov., sp. nov., a novel marine bacterium that obligately utilizes hydrocarbons. Int J Syst Evol Microbiol. 54:141-148

Yakimov MM, Giuliano L, Gentile G et al (2003) *Oleispira antarctica* gen. nov., sp. nov., a novel hydrocarbonoclastic marine bacterium isolated from Antarctic coastal sea water. Int J Syst Evol Microbiol 53:779–785

Yakimov MM, Golyshin PN, Lang S et al (1998) *Alcanivorax borkumensis* gen. nov., sp. nov., a new, hydrocarbon-degrading and surfactant producing marine bacterium. Int J Syst Bacteriol 48:339-348

Yakimov MM, Timmis KN, Golyshin PN (2007) Obligate oil-degrading marine bacteria. Curr Opin Biotechnol. (3):257-66

Yang S, Li M, Lai Q et al (2018) *Alcanivorax mobilis* sp. nov., a new hydrocarbon-degrading bacterium isolated from deep-sea sediment. Int J Syst Evol Microbiol 68:1639-1643

Yao Z, Du S, Liang C et al (2019) Bacterial community assembly in a typical estuarine marsh with multiple environmental gradients. Appl Environ Microbiol 85:e02602-18

Yi J, Lo LSH, Cheng J (2020) Dynamics of microbial community structure and ecological functions in estuarine intertidal sediments. Front. Mar. Sci. 7:585970.

Yu T, Wu W, Liang W et al (2018) Growth of sedimentary Bathyarchaeota on lignin as an energy source. Proc Natl Acad Sci USA 115:6022–6027

Zadjelovic V, Chhun A, Quareshy M et al (2020) Beyond oil degradation: enzymatic potential of *Alcanivorax* to degrade natural and synthetic polyesters. Environ Microbiol 22: 1356–1369

Zedelius J, Rabus R, Grundmann O et al (2011) Alkane degradation under anoxic conditions by a nitrate reducing bacterium with possible involvement of the electron acceptor in substrate activation. Environ Microbiol Rep 3:125-135

Zhang C-J, Pan J, Liu Y et al (2020) Genomic and transcriptomic insights into methanogenesis potential of novel methanogens from mangrove sediments. Microbiome 8:94

Zhang P, Neher DA, Li B et al (2018) The impacts of above- and belowground plant input on soil microbiota: invasive *Spartina alterniflora* versus native *Phragmites australis*. Ecosystems 21:469–481

Zhang X, Agogué H, Dupuy C et al (2013) Relative abundance of ammonia oxidizers, denitrifiers, and Anammox bacteria in sediments of hypernutrified estuarine tidal flats and in relation to environmental conditions. Clean-Soil Air Water 42:815-823

Zhang Y, Xie X, Jiao N et al (2014) Diversity and distribution of *amoA*-type nitrifying and *nirS*-type denitrifying microbial communities in the Yangtze River estuary. Biogeoscience 11:2131–2145

Zhao Y, Chen W, Wen D (2020) The effects of crude oil on microbial nitrogen cycling in coastal sediments. Environment International. 139:105724

Zheng P, Wang C, Zhang X et al (2019) Community structure and abundance of archaea in a *Zostera marina* meadow: a comparison between seagrass-colonized and bare sediment sites. Archaea 2019:5108012

Zhou Z, Liu Y, Lloyd KG et al (2019) Genomic and transcriptomic insights into the ecology and metabolism of benthic archaeal cosmopolitan, Thermoprofundales (MBG-D archaea). ISME J 13:885–901

Zhou Z, Pan J, Wang F et al (2018) Bathyarchaeota: globally distributed metabolic generalists in anoxic environments. FEMS Microbiol Rev 42:639–655

Zhu Z, van Belzen J, Zhu Q et al (2020) Vegetation recovery on neighboring tidal flats forms an Achilles' heel of saltmarsh resilience to sea level rise. Limnol Oceanogr 65:51-62

Zou D, Liu H, Li M (2020a) Community, distribution, and ecological roles of estuarine Archaea. Front Microbiol 11:2060

Zou D, Pan J, Liu Z et al (2020b) The distribution of Bathyarchaeota in surface sediments of the Pearl River Estuary along salinity gradient. Front Microbiol 11:285.

Zumft WG (1997) Cell biology and molecular basis of cell biology and molecular basis of denitrification. Microbiol Mol Biol Rev 61:533–610

NO N2H4 OXIC NO2 NO2 NO2 NO2 NO2 OXIC NO2 OXIC NO2 OXIC NO2 OXIC NO3

Figure 1. The nitrogen cycle in coastal sediments indicating transformations (purple), functional genes (red), and oxic / anoxic zones (green).

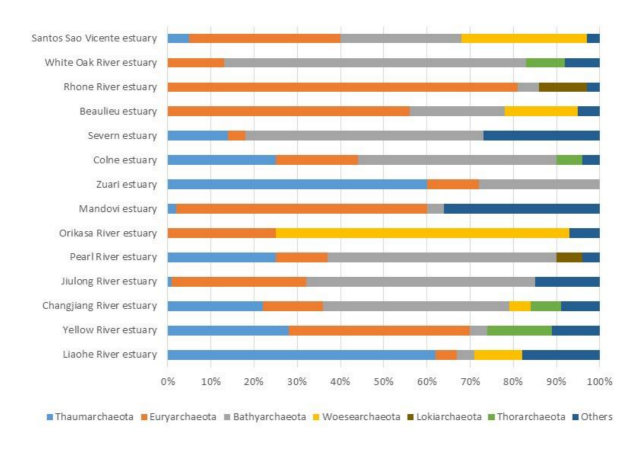


Figure 2. Relative abundance of archaeal taxa in estuarine sediments, based on 16S rRNA gene clone libraries. This figure is produced from data collated by Zou et al. (2020a). Refer to Zou et al. (2020a) for details of the source papers and the primers pairs used.

Table 1. Representative taxa of microphytobenthos found in European coastal sediment microbiomes, characterised by life form and habitat. Note that sediment type represents a continuum of sediment grain size and properties, and individual taxa may occur across this gradient.

Epipelon	Epipsammon	Tychoplankton
(Clays and muds) (Silts and silty sand)	(sand)	(resuspended)
Diatoms (Stramenopiles, Bacillariophyceae)		
Navicula phyllepta, N. gregaria, N. perminuta, N. flanatica,	Planothidium delicatulum	Rhaphoneis minutissima,
N. spartinetensis, N. salinarum	Biremis lucens, Achnanthes sp. Nitzschia frustulum	R. amphiceros
N. peregrina, N. digitoradiata, , N. arenaria		
Gyrosigma limosum, G. fasciola, G. accuminatum	Amphora ovalis. A, salina. A. c.f. tenuissima	Cymatosira belgica
Gyrosigma balticum		Staurosira construens
Pleurosigma angulatum, Scolioneis tumida	Opephora guenter-grassi	Thalassiosira sp., Actinoptychus
Hantzschia virgata, Tropidoneis vitrea		senarius, Odontella aurita
Nitschia c.f. panduriformis, N. sigma, Tryblionella apiculata	Dimeregramma minor	Opephora sp.
Cylindrotheca gracilis. C. signata, C. closterium	Catenula adhaerens	
Euglenids (Stramenopiles, Euglenophyceae)		
Euglena deses, E. proxima		
Cyanobacteria		
Microcoleus chthonoplastes, Lyngbya aestuarii, Spirulina sp.	Merismopedia glauca	
Oscillatoria limosa, O. princeps		

Taken from: Sabbe 1993; Underwood 1994; Underwood et al. 1998; Hamels et al. 1998; Bellinger et al. 2005; Forster et al. 2006; Ribeiro et al. 2013; Redzuan & Underwood 2020; 2021.

Table 2 Representative N-cycle microorganisms found in various coastal marine and estuarine environments

N-cycle microorganisms	Environment	Reference
Aerobic Ammonia Oxidising Bacteria (AOB)		
N. europaea, N. communis, N. oligotropha (Cluster 6a), N. marina (Cluster 6b), Nitrosospira spp.(Clusters 13-15)	Coastal environments (Jiaozhou Bay)	Dang et al. 2010
Nitrosomonas spp.	Hypernutrified estuary (Bahía del Tóbari)	Beman and Francis 2006
Nitrosomonas spp.	Estuarine (Brackish) (Westerschelde estuary)	Sahan and Muyzer 2008.
Nitrosospira spp.	Estuarine (marine) (Westerschelde estuary)	
Estuarine/marine Nitrosospira-like cluster and Nitrosomonas-like cluster	Estuarine sediments (Elkhorn Slough)	Wankel et al 2010
Nitrosospira spp., Nitrosomonas spp. (N. marina, N. oligotropha, N. ureae, N. eutropha)	Wetland sediments of subtropical coastal mangroves	Wang et al. 2013
Nitrosospira sp. and Nitrosolobus multiformis	Coastal sediments (North Sea)	Lipsewers et al. 2014
Aerobic Ammonia Oxidising Archaea (AOA)		
Nitrosopumilus maritimus, Nitrososphaera gargensis	Wetland sediments of subtropical coastal mangroves	Wang et al. 2013
Nitrosopumilus maritimus	Mangrove sediments, South China Sea sediments	Li et al. 2011b, Cao et al. 2012
Nitrosopumilus subclusters 12 and 16 (stable marine cluster), Nitrosopumilus subcluster 4.1 (estuarine cluster)	Marine coastal sediments (North Sea)	Lipsewers et al. 2014
Nitrosopumilus maritimus, Nitrosphaera gargensis	Subtropical macrotidal estuarine sediments	Abell et al 2010
Comammox		
Clade A.1 (e.g. <i>Cand. Nitrospira nitrificans</i> , <i>Cand. Nitrospira nitrosa</i> , Clade A.2, Clade B	Various sediments (tidal flat, saltmarsh, coastal), coastal waters	Xia et al. 2018
Anammox		
Scalindua spp., Kuenenia spp.	Wetland sediments of subtropical coastal mangroves	Wang et al. 2013

Uncultured Planctomycetes, Cand. Scalindua spp., Cand. Brocadia spp., Cand. Kuenenia spp.	Anoxic basin on Black Sea	Kuypers et al. 2003
Denitrification		
Anaeormyxobacter delahogens 2CP-C, Thermus thermophilus strain HB8, Geobacter metallireducans, Rhodoferax ferrireducans, Halomonas halodenitrificans, Rhodobacter sphaeroides, Cupriavidus necator, Hahella chejuensis, Shewanella spp., Vibrio spp., Saccharophagus degradans, Rhodopseudomonas palustris, Magnetospirillum magneticum AMB1, Helicobacter hepaticus ATCC51449, Pseudomonas spp.	Hypernutrified estuarine sediments	Smith et al. 2007
Alcaligenes spp. (A. faecalis, A. xylosoxidans) Pseudomonas spp. (P. stutzeri), Bradyrhizobium japonicum, Blastobacter denitrificans	Sediments within the Oxygen-Deficient Zone, Pacific Coast	Liu et al. 2003
Dissimilatory nitrate reduction to ammonium (DNRA)		
Shewanella frigidimarina, Chlorobium phaeobacteroides	Hypernutrified estuarine sediments	Smith et al. 2007
Nitrogen Fixation		
Azotobacter spp., Azospirillum spp., Campylobacter spp., Beggiatoa spp., Enterobacter spp., Klebsiella spp., Vibrio spp., Desulfobacter spp., Desulfovibrio spp., Clostridium spp. Cyanobacteria including unicellular and non-heterocystous species, (Chromatiaceae, Chlorobiaceae, Chloroflexaceae, Rhodospirillaceae), Archaea (e.g. Methanococcus spp., Methanosarcina spp.)	Marine/ seagrass sediment, estuarine sediment, salt marsh sediment, <i>Spartina</i> roots, <i>Zostera</i> roots, Beach sediment, intertidal sediments, seawater	Herbert 1999