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Spatial and temporal dynamics of coastal benthic microbial

2 communities along a salinity gradient

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13 Abstract

- 14 The Baltic Sea is a unique brackish water ecosystem studied for decades; however, knowledge about
- 15 the diversity of the benthic communities of bacteria and microbial eukaryotes within this system is
- sparse. Using an amplicon sequencing approach, we evaluated the diversity of shallow water coastal
- microbial sediment communities and their relation to several environmental factors, on both a large
- spatial scale in the Baltic Sea and the adjacent North Sea, as well as on a temporal scale at selected
- sites along the salinity gradient in the Baltic Sea. We found salinity to be among the strongest drivers
- of both bacterial and eukaryote communities' species diversity, and that community network structure
- 21 appeared to change between sites of different salinity. However, for the communities in the poly-to
- 22 mesohaline sections of the study area, diversity seems affected to a higher degree by temperature,
- 23 nutrient, and sediment characteristics.

24 Keywords

- 25 Baltic Sea, benthic bacteria, benthic protists, brackish microbiology, benthic microbial diversity,
- 26 microbial networks

Introduction

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28 Interactions between bacteria and microbial eukaryotes in aquatic environments have been reported 29 in pelagic systems over the last decades (e.g., Miki & Jacquet 2008), and as a result of these interactions the communities of these two taxonomic groups may co-vary. A positive correlation may 30 31 indicate both non-causal relations such as similar preferred conditions, or interactions through commensalism or mutualism. They may however also result in negative correlations reflecting effects 32 33 of parasitism (Anderson & Harvey 2020), grazing or competition for common resources (Chow et al. 34 2014). These interactions between bacteria and eukaryotes may shape diversity for both groups, by 35 applying selection pressure and contribute to the evolution and maintenance of high diversity of 36 microbes (Hiltunen & Becks 2014, Ramanan et al. 2016), and thus shape the community structure of 37 each other besides other ecosystem components e.g., viruses (Chow et al. 2014). 38 Besides biotic interactions, microbial communities are also affected by environmental factors. In 39 marine environments especially salinity is a known driver of changes in diversity (e.g. Herlemann et 40 al. 2011; Campbell & Kirchman 2013). The Baltic Sea is one of the largest brackish water seas in the 41 world, resembling a large estuary with its characteristic mixing of marine and freshwater (Leppäkoski 42 et al. 2009). Due to the physio-chemical properties of the Baltic Sea, its microbial communities are 43 suspected to be uniquely adapted to the brackish water ecosystem (Ininbergs et al. 2015). The salinity 44 gradient in the Baltic Sea has shown to be a strong driver of community structure for both 45 phytoplankton (Olli et al. 2019), planktonic bacteria (Herlemann et al. 2011, Lindh & Pinhassi 2018, 46 Camarena-Gómez et al. 2021) and microbial eukaryotes (Telesh et al. 2011, Hu et al. 2016). Although the first studies of the microbial communities in the Baltic Sea were conducted more than a decade 47 48 ago (Riemann et al. 2008), and despite the few studies examining the benthic microbial communities 49 (Klier et al. 2018, Salonen et al. 2018), little is known of the diversity patterns of the benthic bacteria 50 and eukaryote communities of the littoral zone. Salinity is nevertheless known to affect bacterial diversity in the sublittoral zone (Pavloudi et al. 2016, Klier et al. 2018, Li et al. 2021). Benthic 51 52 bacterial communities have even shown contradictory trends in diversity over salinity gradients, 53 compared to bacterioplankton communities (Campbell & Kirchman 2013, Pavloudi et al. 2016, Klier 54 et al. 2018, Vidal-Durà et al. 2018). However, other physico-chemical driving factors, including 55 seasonal changes in temperature and nutrient availability, could also have a significant influence on the composition of benthic microbial communities, both for bacteria (Vetterli et al. 2015, Lv et al. 56 57 2016), and microbial eukaryotes (Massana et al. 2015, Salonen et al. 2018, Anderson & Harvey 2020).

Microbial communities, including bacteria and eukaryotes, are important components of decomposition and nutrient recycling, and are fundamental for all ecosystems. The use of genetic tools, such as metabarcoding and next generation sequencing for documenting uncultured microbes has widened our knowledge of the composition of these communities particularly in the marine environment. However, the focus of such studies has been primarily on marine pelagic microbial communities, and only within the last decade has widened to include microbial communities in sediments (Edgcomb et al. 2011, Bik et al. 2012, Forster et al. 2016), especially in estuaries and brackish water systems (Chariton et al. 2010, Campbell & Kirchman 2013, Klier et al. 2018).

In this study, we monitored changes in diversity and community structure in shallow coastal benthic bacteria and microbial eukaryote communities along a salinity gradient in the Baltic Sea and adjacent North Sea, covering both a large spatial and local temporal scale. These communities may be affected by the species composition and their interactions, which may also be affected by external factors. By assessing associations between bacteria and microbial eukaryotes along the Baltic Sea, we aim to provide a first insight to the interactions, and potential keystone species in the benthic microbial communities of this brackish water sea. Furthermore, we assessed whether and to what extent selected abiotic factors besides salinity (sediment mean grain size, sorting, sediment water content, porosity, C/N ratio, organic content, and sediment temperature) influenced the observed diversity patterns. We hypothesised that the microbial community composition is strongly driven by salinity, and that the microbial diversity changes between sites along the Baltic Sea salinity gradient. Thus, other environmental factors are of less importance in shaping the communities. Because of the expected decrease in diversity along the Balitc Sea we also expected to see a decrease in complexity of interactions between microbial species in these communities as salinity falls. Testing this hypothesis is realised by presenting networks at three selected sites covering a part of the salinity gradient. We expected larger differences among the microbial communities (sites) than among temporal sampling points within the same site.

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Materials and methods

- 85 Sediment sampling
- 86 Samples for assessing spatial variation in microbial community diversity were collected during
- August 2018, at seven study sites representing a decrease in salinity (List, Saltö, Hersley, Gollwitz,
- 88 Öland, Tvärminne, Pori) in the North Sea and Baltic Sea (Fig. 1). At three of the sampling sites in the

- 89 Baltic Sea with profound differences in salinity (Saltö, Hersley, Öland), samples for assessing
- 90 temporal variation were collected at four time points (August 2018, November 2018, April 2019,
- 91 August 2019). All sites were sampled from the coast at water depth 0-0.80 m, except Tvärminne
- 92 where sampling was performed by SCUBA at 3.8-5.0 m depth. Sediment was collected in three
- 93 replicate cores (5 cm diameter, min. 15 cm depth). After draining, two technical replicates were
- ollected from each core. Samples were taken from the central region of the core (avoiding the core
- 95 liner walls) by scraping off the top 5 mm of sediment (approx. 1.5 g), using sterile razor blades and
- 96 placed in sterile 1.5 ml microcentrifuge tubes. The sediment was preserved in 99% EtOH (Harry et
- 97 al. 2000) and kept frozen at -18°C until DNA extraction.
- 98 Analysis of environmental factors
- 99 Temperature was measured using handheld field thermometer (Frederiksen Scientific), salinity was
- measured using ATACO handheld refractometer (resolution of 0.5 salinity units) and sediment
- 101 characteristics were recorded for each sampling point. Sediment characteristics (including water
- 102 content, porosity, organic content, carbon and nitrogen content, grain size and sorting) were
- determined from three replicate cores per sampling station and analysed following Petersen et al., in
- 104 review.
- 105 DNA extraction and amplification
- 106 In this study three sampling replicates represent each sampling site/time, except for samples
- 107 Tvärminne, Gollwitz, Saltö August 2019, Herslev August 2018, and Öland November 2018, these
- samples are represented by two sampling replicates and one technical replicate.
- 109 DNA from 250 mg of sediment per sample was extracted with DNeasy Powerlyzer PowerSoil Kit
- (Qiagen) according to the manufacturer's protocol. Samples were lysed in Bead Ruptor Elite (OMNI
- International) at 2500 rpm for 45 s with 750 µl of PowerBead solution (Qiagen). DNA concentration
- was quantified using a Qubit 4.0 fluorometer with 1X dsDNA HS Assay Kit (Thermo Fisher,
- 113 Cambridge, UK). Bacterial 16S rRNA gene V1-V2 hypervariable regions were amplified using the
- universal primers 27F (Ludwig et al. 1993) and 338R (Suzuki & Giovannoni 1996). To assess
- diversity of the microbial eukaryotic community V4 hypervariable regions was chosen to amplify the
- eukaryotic 18S rRNA gene. For this we used primer set UNonMetF and UNonMetR (Bower et al.
- 2004), combined with primers E572 and E1009R (Comeau et al. 2011). For primer details see
- 118 Supplementary Table 1. Reactions were performed in 25 μl volumes containing 12.5 μl of iQTM
- 119 SYBR® Green Supermix (2X) (Bio-Rad), 200 nM of each primer, 5.5-9.5 µl of nuclease-free water

- and 3 ng of template DNA or PCR product from UNonMet primer pair amplification for E572 forward
- and E1009R reverse primers. All amplifications were performed in a CFX96 Touch™ Real-Time
- 122 PCR Detection System (Bio-Rad). Thermocycling protocols are shown in Supplementary Table 2.
- 123 Library preparation
- 124 A fusion PCR was performed to add the Ion Torrent PGM sequencing adapters: barcoded M13-tailed
- 125 IonA adapter, target specific P1 adapter and a target specific M13- linker primer (following Mäki et
- al. 2016) (see Supplementary Table 3). Amplification was achieved with the following conditions:
- 127 95°C/5 min, (94°C/45 s, 53°C/1 min, 72°C/1 min)*13, followed by final elongation of 72°C/5 min.
- Reactions (25 μl) consisted of 12.5 μl iQTM SYBR® Green Supermix (2X) (Bio-Rad), 400 nM of
- 129 fusion primers (IonA forward and P1 reverse), 40 nM of target specific M13- linker primer, 8.5 μl of
- 130 nuclease-free water and 1 μl of PCR product from the 16S or 18S rRNA amplification step. Libraries
- were purified with Quanta SparQ PureMag Beads system (Quantabio). Quality and molarity were
- determined with Agilent 2200 TapeStation before pooling in equimolar concentration (22 pM). The
- library pool was sequenced with Ion PGM system using an Ion 318 Chip Kit version 2 (Ion Torrent,
- Life Technologies) at the University of Jyväskylä.
- 135 Analysis of sequencing data
- Single-end raw sequence reads were sorted by barcode prior to exporting from the Ion Torrent SuiteTM
- Software and trimmed to remove sequencing adapters using Cutadapt (version 1.18, Martin 2011).
- Length filtering was performed using Cutadapt as follows: 16S: min. 150 bp, max. 400 bp; 18S: min.
- 139 32 bp max. 400 bp. Quality filtering was performed using FASTX toolkit (Gordon & Hannon 2010),
- with a minimum quality score of 20 for a minimum of 80% of the bases in each read. Dereplication,
- singleton and chimera filtering were performed using VSEARCH (version 2.15.1, Rognes et al.
- 142 2016). All datasets were aligned to the SILVA database (version 138, Yilmaz et al. 2014) using
- Mothur (version 1.44.3, Schloss et al. 2009) before clustering using greedy 97% similarity threshold
- in VSEARCH. Amplicon sequence variants (ASVs) were mapped using VSEARCH, at a similarity
- of 95-97.5%. 16S datasets were classified to the SILVA SEED database (version 138, Yilmaz et al.
- 146 2014) using the Wang method (Wang et al. 2007) with 8 kmer and 80% similarity in Mothur. 18S
- dataset was classified to the PR2 database (version 4.12.0, Guillou et al. 2013; del Campo et al. 2018)
- using the BLAST method with 80% similarity in Mothur.
- We used universal primers, standardised protocols and well curated public databases in order to
- 150 minimize potential technical biases such as PCR error, sequencing error, sequencing depth,

151 bioinformatics analysis, and unassigned classification of the organisms (Guillou et al. 2013, Yilmaz 152 et al. 2014, del Campo et al. 2018). Nevertheless, the 16S spatial dataset generated 7.7% unclassified OTUs, 16S temporal dataset generated 9.0% unclassified OTUs, 18S spatial dataset generated 27.1% 153 154 unclassified OTUs, 18S temporal dataset generated 34.1% unclassified OTUs at phylum/division level. All unclassified OTUs at domain/kingdom were removed prior to further analyses. In addition, 155 156 in the 16S dataset, all OTUs assigned to "Chloroplast" at the order level, and in the 18S dataset, all 157 OTUs classified to phylum Metazoa, Rhodophyta and class Embryophyceae were removed. 158 Statistical analysis 159 All statistical analyses were performed in R version 3.6.3 (2020.02.29). To compensate for large 160 differences in read depth between the samples, all datasets were transformed by rarefaction to an even 161 read depth by repeatable random subsampling (read depth after rarefying in datasets: 16S spatial: 9773, 16S temporal: 15497, 18S spatial: 10957, 18S temporal: 7588), using the package 'phyloseq' 162 (version 1.30.0, McMurdie and Holmes 2013). OTU table plots were made using 'phyloseq' package. 163 Alpha diversity (based on OTU richness and Shannon index) and beta diversity (based on Bray-Curtis 164 165 dissimilarity) measures, dbRDA and ENVFIT analyses based environmental variables were 166 performed in R using 'vegan' package (version 2.5-7, Oksanen et al. 2017). Redundant (i.e., highly 167 correlated) variables (water content and mean grain size) were removed on basis of ENVFIT analysis. All plots were made using 'ggplot2' (version 3.3.0, Wickham 2016). Significant differentially 168 169 diversity matrices were detected using the Kruskal-Wallis test followed by Tukey HSD test for pair 170 wise comparison. Salinity gradient test on community structure was performed by subjecting Bray-171 Curtis dissimilarity matrices in bacterial and microbial eukaryotic communities using the 'adonis' 172 test from the 'vegan' package. 173 Network analysis was performed to show bacterial-eukaryote correlations and network properties 174 were computed using the "igraph" package (Csardi & Tamas Nepusz 2006) as described earlier (Sapkota et al. 2020). We used only temporal datasets with high salinity (all temporal datasets from 175 176 Saltö), medium salinity (all temporal datasets from Herslev) and low salinity (all temporal datasets from Öland). Spatial datasets representing lower number of replicates per salinity level (high, 177 178 medium, and low) were excluded. The bacterial and eukaryote OTU table was trimmed for low 179 abundant OTUs (<50 reads) and normalized as relative abundance counts per million using the 180 "edgeR" package (Robinson et al. 2010). Correlations on all OTU pairs were computed using rcorr 181 function from 'Hmisc' package and only highly significant (p < 0.001) correlated OTUs with 182 spearman's rank correlations > 0.7 for positive correlations and < -0.7 for negative correlations, were

- used for network graphics using a Fruchterman-Reingold layout with 999 permutations. In the
- network, OTUs were set as nodes and the correlation as edges. Nodes (OTUs) with the highest number
- of connections were used as keystone species in the network analysis. For the figures the 10 most
- 186 connected OTUs represent keystone species, and tables show the 5% if OTUs with most connections.

187 **Results**

- 188 Environmental characteristics
- Salinity ranged between 33 and 6 among sites in the spatial study, highest at List and lowest at Pori.
- 190 For the temporal dataset, salinity ranged between 26 and 7, highest at Saltö and lowest at Öland. In
- addition to salinity, most variation among spatial sites was seen in sediment temperature, water
- 192 content, and C/N variables; whereas temperature and C/N variables showed most variation among
- 193 the temporal samples. For details of environmental characteristics for the spatial and temporal
- samplings see Supplementary table 4.
- 195 Bacterial diversity
- 196 In the spatial study, the observed number of OTUs per sample for the bacterial communities (16S)
- ranged between 1300 and 735, highest at Saltö and lowest at List (Fig. 2A), and Shannon index for
- each sampling site ranged between 6.2 and 4.3, highest at Saltö and lowest in Öland (Fig. 2A,
- 199 Supplementary Table 5a). Proteobacteria was the most dominant phylum at all sites, followed by
- 200 Cyanobacteria and Bacteroidota (Fig. 2C) except for the euhaline site List. At List, Patescibacteria
- was the second relative abundant phylum, and for both List and Saltö this group was generally more
- relatively abundant than at the remaining sites (Fig. 2C). The relative abundance of Cyanobacteria
- was also high at all sites, however lowest at Saltö, and it was the second most relative abundant
- 204 phylum at Hersley, Gollwitz and Öland (Fig. 2C). Chloroflexi had highest relative abundance at
- 205 Öland, Tvärminne and Pori (Fig. 2C), compared with other sites. Relative abundance of
- 206 Gemmatimonadota was generally very low at all sites except for Pori (Fig. 2C).
- 207 Considering alpha diversity for the spatial dataset, only Shannon diversity was related to salinity,
- according to both an ANOVA test (Supplementary table 6) and Pearson correlation (Supplementary
- 209 figure 1). Beta diversity was correlated with all environmental factors (Table 1), however when
- 210 represented by dbRDA (Fig. 3A), List and Saltö clustered closest together with a positive relation to
- sorting, salinity and C/N ratio, and negative relation to water content explaining their placement.
- Herslev and Gollwitz formed a discrete cluster, and Öland and one replicate of Tvärminne clustered
- 213 together but not as closely, while the remaining replicates of Tvärminne and Pori were placed within

214 proximity to each other along the same axis. Herslev and Gollwitz were grouped by high organic 215 matter and temperature, and the Öland and Tvärminne cluster was grouped by high water content, well sorted sediment, and low salinity and C/N ratio (Fig. 3A). Pori and the two replicates of 216 217 Tvärminne were characterised by low organic matter and low temperature at the time of sampling (Fig. 3A). Analysis of distance-decay, which tests if communities become more dissimilar with 218 219 increasing distance, for bacterial communities along the extent of the Baltic Sea covered in the spatial 220 part of the study showed a strong relationship (Supplementary figure 2A, B & C). 221 In the temporal study, observed OTUs in bacterial communities (16S) ranged between 2122 and 1058 222 (Fig. 4A). Shannon index ranged between 6.5 and 4.5 and Öland showed significant lower diversity 223 compare to Saltö and Herslev (Fig. 4A, Supplementary table 5B). As seen in the spatial part of the 224 study, Proteobacteria was the most dominant phylum at all sampling sites, followed by 225 Cynanobacteria and Bacteroidota, except for Saltö where Patescibacteria had a higher relative abundance (Fig. 4C). Generally, the distribution and relative abundance of phyla followed the same 226 227 patterns at each sampling time for each sampling site, respectively (Fig. 4C). There were, however, 228 some fluctuations in the relative abundance following the seasonal cycle; Proteobacteria showed a 229 peak in relative abundance in the April 2019 sampling at both Saltö (S3) and Herslev (H3), and 230 Actinobacteria was highest in November 2018 at both Herslev (H2) and Öland (Ö2) (Fig. 4C). 231 Additionally, Actinobacteriota was relatively more abundant at Hersley and Öland compared to Saltö 232 at all sampling times. Desulfobacterota had a lower relative abundance in April 2019 at Saltö (S3) 233 compared to the other sampling dates and had lowest relative abundance at Öland at all times 234 compared to the other sites. 235 Considering alpha diversity of the temporal dataset, ANOVA test showed that only Shannon diversity 236 was related to salinity (Supplementary table 6), however Pearson correlation showed that both 237 observed, and Shannon diversity were correlated to salinity (Supplementary figure 1). In the beta diversity analysis, each site formed its own cluster in the dbRDA plot, but temporal samples from 238 239 Herslev clustered more closely together than did the temporal samples from the other sites (Fig. 3C). 240 Though the beta-diversity of bacterial dataset was correlated to all environmental factors (Table 1), 241 Saltö August 2018 (S1) and 2019 (S4) were best characterised by a positive relation to salinity and 242 sorting, and November 2018 (S2) and April 2019 (S3) were better characterised by higher C/N ratio, 243 water content and lower temperature in the dbRDA (Fig. 3C). Variation between samples at Herslev 244 was related to sediment temperature, water content and C/N ratio, and variation between samples at Öland was related to organic matter (Fig. 3C). Analysis of distance-decay for bacterial communities 245

- 246 along the extent of the Baltic Sea included in the temporal dataset showed a strong relationship 247 (Supplementary figure 2A & B). 248 Analysis of correlation between salinity and bacterial OTUs showed both positive and negative 249 relationships (Supplementary table 7). Especially several OTUs assigned to genus Candidatus Kaiserbacteria, showed positive correlation to the salinity gradient. 250 251 252 Microbial eukaryote diversity 253 In the spatial study of microbial eukaryote diversity (18S), observed number of OTUs per sample 254 ranged between 619 and 146 (Fig. 2B), and Shannon index ranged between 4.6 and 1.9 (Fig. 2B). 255 Ochrophyta had high average relative abundance and was the most abundant phylum/division at all 256 sites except Herslev and Gollwitz (Fig. 2D). Ciliophora was the highest relative abundant phylum/division at Gollwitz, and in Hersley, Dinoflagellata had highest relative abundance (Fig. 2D). 257 Alveolata, Apicomplexa, and Fungi had highest relative abundance in the mesohaline sites Gollwitz, 258 Öland, Tvärminne and Pori, and these sites, together with Saltö, also had high relative abundance of 259 260 Ciliophora (Fig. 2D). Abundance of unclassified eukaryotes was high in the datasets from List and 261 Gollwitz. Neither observed richness nor Shannon diversity of the spatial dataset were related to 262 salinity in both ANOVA test and Pearson correlation (Supplementary table 6, Supplementary figure 263 1 & 3). However, beta diversity was correlated to salinity, together with C/N ratio and water content 264 (Table 1). The dbRDA (Fig. 3B) showed List to cluster together with Öland and one replicate from Saltö and was somewhat close to the Herslev samples also, best described by high C/N ratio, water 265 266 content and organic matter. The remaining samples from Saltö and Gollwitz were placed somewhat 267 close together, characterised by high organic matter, C/N ratio and poor sorting (Fig. 3B). Tvärminne and Pori formed a distinct cluster mainly characterised by low salinity and temperature, and well 268 269 sorted sediment (Fig. 3B). 270 For the temporal study of eukaryotes (18S), between 711 and 136 OTUs were captured (Fig. 4B), and 271 Shannon index ranged between 5.1 and 1.6 (Fig. 4B, Supplementary table 5B). Ochrophyta was the
- Shannon index ranged between 5.1 and 1.6 (Fig. 4B, Supplementary table 5B). Ochrophyta was the most relatively abundant phylum/division in the April 2019 and August 2019 samplings for all sites, except for Öland August 2019, where it was the second most relative abundant, exceeded by Dinoflagellata (Fig. 4D). Ochrophyta was also most relative abundant in August 2018 for both Saltö and Öland (S1 and Ö1) (Fig. 4D). Dinoflagellata was the most relatively abundant phylum/division in the samples from Herslev August 2018 (H1) and November 2018 (H2), as well as Saltö November

277 2018 (S2) (Fig. 4D). The most relatively abundant phylum/division at Öland in November 2018 (Ö2) 278 was Fungi (Fig. 4D). Abundance of unclassified eukaryotes was especially high in the datasets from 279 Saltö. Both alpha diversity measures were significantly correlated to salinity (see Supplementary table 280 6). Community structure was influenced by salinity, as well as temperature, C/N ratio, mean grain size and sediment sorting (Table 1). In the dbRDA plot samples were in general scattered with only 281 little clustering of sites, especially for Herslev and Öland (Fig. 3D), indicating more temporal changes 282 283 than seen in the bacteria communities (Fig. 3C). For most locations, samples from August for two 284 consecutive years were somewhat similar. For example, Saltö August 2018 (S1) and 2019 (S4) 285 samples clustered together characterised by high temperature, organic matter, and poor sorting. 286 Similarly, August 2018 and 2019 for Herslev (H1 and H4) and Öland (Ö1 and Ö4) were placed closely 287 together, somewhat more like the samples from November or April from those stations compared to 288 Saltö (Fig. 3D). For Saltö, samples from November and April formed a distinct cluster described by 289 different environmental factors than those from August samples, specifically high water content and 290 temperature (Fig. 3D). Temperature was an obvious explanatory variable for the observed community 291 patterns, but also salinity, C/N ratio and sediment sorting were important for the differences in 292 eukaryote communities seen in the dbRDA plots. However, analysis of distance-decay for eukaryote 293 communities along the extent of the Baltic Sea covered in this study only showed a weak correlation 294 for the temporal dataset (Supplementary figure 2C).

- Analysis of correlations between salinity and eukaryote OTUs only revealed five positive correlations and one negative correlation (Supplementary table 7). In our eukaryote datasets, a large proportion of OTUs could not be classified to phylum/division level at 80% similarity. This is most likely due to incomplete microbial eukaryote coverage in the database; since the benthic microbes historically have received less scientific attention than the pelagic ones, a large proportion of benthic microscopic eukaryotes are expected to be missing from the databases (Forster et al. 2016, Zhang et al. 2018).
- 301 Microbe-Microbe interactions shown as networks
- To explore bacteria-eukaryote interactions in different salinity environments, we used the temporal dataset and visualised co-occurrence at three different salinity levels as network graphs (Fig. 5). Interestingly, we found distinct co-occurrence patterns in networks from the three different salinity levels. At highest salinity level, co-occurrence networks were strongest, having the more connections compared to networks for the other two salinity levels. Similarly, the lowest salinity level showed the least connections revealing fewer eukaryote-bacteria interactions, and co-occurrence network for the medium salinity level was in between the other two. Several bacterial and eukaryote OTUs were

identified as the keystone species based on the highest number of connections that they showed in the co-occurrence networks (for details see Supplementary table 8A, B & C). In particular, Raphidpennate diatoms (Stramenopiles) appeared several times as keystone eukaryotic species in all three networks. Also, bOTU607 (a member of Rhodobacteraceae) appeared as a keystone species in two networks.

Discussion

Using amplicon sequencing, we surveyed microbial communities of shallow coastal sediments on both a large spatial scale in the Baltic Sea and the adjacent North Sea, and a temporal scale in a subset of these communities. We examined diversity patterns of bacterial communities as well as among microbial eukaryotes, and the correlations between the two kingdoms. Because the two taxonomic groups could respond differently to environmental factors, we evaluated the relationship of diversity patterns to selected environmental factors, particularly salinity, temperature, and sediment characteristics for both groups. These analyses suggested that salinity is a strong driver of diversity of microbial communities, however, salinity is not the only driver, as sediment characteristics, nutrients, and temperature also were important. In addition, our results revealed significant interactions and relationships among several bacterial and eukaryote OTUs. This study provides a broader description of coastal benthic microbial communities and their interactions at shallow sites in the Baltic.

Baltic Sea salinity gradient affects bacterial and microbial eukaryote communities

Though the ANOVA test showed that OTU richness for bacteria was not affected by salinity for either dataset, Shannon diversity of these communities was significantly affected by salinity. Moreover, we did see a significant positive correlation for both diversity measures using Pearson's correlation on the temporal dataset (Supplementary figure 3). This suggests that the missing correlation between salinity and observed OTUs for the spatial dataset could be due to too few data points. Likewise, distance-decay showed beta-diversity of both bacterial datasets were positively correlated with distance between communities, hence when distance increases the communities become more dissimilar along the gradient (Supplementary figure 1A, B & C). While salinity was a significant driver of beta diversity for bacteria, it might not be an equally strong driver of diversity patterns at all sites. As reflected in the dbRDA (Fig. 3A and C), the communities at the sites with highest salinity (List and Saltö) were described to a higher degree by salinity, however the

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communities at lower salinities were not to a high degree described by salinity, except for Öland which was to some degree described by lower salinity. Despite the coarse sampling, we did observe patterns across the salinity gradient: highest diversity was found at the eu- and polyhaline sites (List and Saltö), a diversity minimum at the mesohaline sites (Öland and Tvärminne), followed by a slight increase in diversity at the site with the mesohaline site of lowest salinity (6) (Pori). This suggests that bacterial communities at the sites of highest salinity are more diverse and evenly distributed than communities in the mesohaline. This pattern suggests that estuarine bacterial communities could be adapted to lower salinity. The pattern we observed supported previous findings in the Baltic Sea, where bacterial communities in the mesohaline consisted mainly of bacterial lineages with a broad salinity tolerance (Klier et al. 2018). Although the bacterial taxonomic composition was dominated by the same phyla at all samplings (both spatial and temporal), there were some differences between the salinity extremes. Patescibacteria had highest relative abundance at the eu- and polyhaline sites, List and Saltö, especially genus Candidatus Kaiserbacteria, showed several OTUs positively correlated to salinity, indicating better conditions for this group at higher salinities. However, a few OTUs assigned to Candidatus Kaiserbacteria also showed a negative correlation to salinity, highlighting the complexity in bacterial diversity. Cyanobacteria generally were abundant at all sites except at the eu- and polyhaline ones (List and Saltö), indicating better conditions for these taxa at lower salinities. However, individual OTUs of Cyanobacteria also showed both positive and negative correlation to salinity, except most OTUs of Synechococcales showed a negative correlation, indicating that this group might favour low salinities. The same argument could be made for phylum Chloroflexi, which had highest relative abundance at the low salinity sites (Öland, Tvärminne and Pori), a trend also previously observed for sublittoral bacterial communities in the Baltic Sea (Klier et al. 2018). Several OTUs of Cloroflexi was negatively correlated to salinity, thus favouring low salinity, especially OTUs of the obligate anaerobe genus Anaerolinea, indicating that the low salinity sites are also affected by low oxygen. Gemmatimonadota had low abundance at all sites except the one with lowest salinity (Pori), and three OTUs of family Gemmatimonadacea (member of phylum Gemmatimonadota) were negatively correlated to salinity, indicating a negative affliction to salinity. However, other local factors, such as lack of associated phytoplankton (Mujakić et al. 2021) may also have an influence on this particular phylum.

The spatial patterns we found for microbial eukaryotes were like those for bacterial communities: salinity played an important role, but it was not the only driver of benthic marine microbial eukaryote

community composition and diversity. We observed no clear trends in alpha diversity related to salinity for either eukaryote dataset, but both OTU richness and Shannon diversity of the temporal dataset was significantly affected by salinity (Table 1) in the ANOVA test, although we did not see a correlation for either eukaryote dataset using Pearson's correlation. This suggests the changes were not continuous, but rather in steps. Neither did we see a relation to distance, which could have been expected to follow somewhat the same diversity trend as salinity. We did however observe a weak correlation along the extent of the Baltic Sea when using the temporal samples. There were also some systematic changes in taxonomic composition along the Baltic Sea and its salinity gradient, e.g., while Alveolata, Apicomplexa and Fungi were present at all sites, the three phyla were more abundant in sites with lower salinity. This indicates that lower salinities are a more favourable condition for these taxa, and, for example, supports earlier findings that fungal communities are adversely influenced by salinity (Mohamed & Martiny 2011, Tisthammer et al. 2016). Nevertheless, only one OTU (unclassified Cercozoa) was significantly negatively correlated to salinity. Additionally, only a few OTUs showed a positive correlation to salinity, representing the orders Cercozoa, Bacillariophyta and Apusomonadidae.

- 387 Temporal patterns influence microbial assembly
- Seasonal fluctuations in the bacterial communities' OTU richness and Shannon diversity were observed at all sampling sites, but with no shared patterns between sites. In the summer samples (August 2018 and 2019), communities were more similar in comparison to other times of the year (Fig. 3C), except for Herslev. Changes in the abundance of several phyla during the seasonal cycle were also described, however, no clear trends were observed common to all sites. These cyclic changes support previous assumptions of seasonal effects on both benthic and planktonic bacterial communities in the Baltic Sea (Vetterli et al. 2015, Herlemann et al. 2016). The inconsistency between sites suggests that the seasonal changes are shaped locally, rather than on a larger regional climatic scale, or that local factors modify a possible common seasonal pattern.
- In contrast to the bacterial communities, eukaryote communities seem to have more variation in taxonomic composition on a temporal scale, with each site having their own distinctive pattern, supporting previous observations of microbial eukaryote communities on a smaller geographic scale in the Baltic Sea (Salonen et al. 2018). Additionally, fluctuations in richness were observed at all sites over the season. The eukaryote communities potentially cover more functional groups compared with bacteria (Reynolds 2006), some of which could be favoured at different environmental conditions.

The bacterial community may therefore be more resilient to environmental fluctuations, which could explain the differences in patterns, especially the variation in species composition, between bacteria and microbial eukaryotes observed here. However, changes in diversity could be present on a lower taxonomic level e.g., genus level, which will not be reflected in our overview at phylum level. Furthermore, some of the organisms may be represented by dormant stages, especially among microbial eukaryotes (Marcus & Boero 1998), or inactive or even dead organisms added from surrounding water masses (Stoeck et al. 2007) which could be evident in the two organismal groups differently. Although temporal fluctuations were observed in the benthic microbial eukaryote communities, there were no consistent seasonal trends in the species composition, such as signal of spring bloom of either pelagic diatoms or dinoflagellates as earlier observed in benthic microscopic eukaryote communities in the north-eastern Baltic Sea (Salonen et al. 2018). However, the eukaryote community at Herslev in August 2018 had lower alpha diversity than other temporal samples, and was dominated by Dinoflagellata and Chlorophyta, which was distinctly different from the other samples that were largely dominated by Ochrophyta, suggesting that the August 2018 community may have been influenced by blooming dinoflagellates in Herslev.

418 Inter-kingdom networks contributing to community structure

The diversity of communities is likely not only influenced by environmental factors, but also by competition and predation within the communities. These interactions and the structure of the community, in turn, may be affected by changes in community composition and diversity. We found both positive and negative correlations among bacterial and eukaryote OTUs. The most connected OTUs, putatively regarded as the keystone species, belong to both bacteria and eukaryotes. A few keystone eukaryotes did signify their role in microbe-microbe interactions by having several or only negative correlations. Furthermore, the communities of the low salinity site (Öland) were dominated by a few keystone eukaryotes with only negative correlations, which could indicate a top-down regulated system. The communities of high (Saltö) and medium salinity (Herslev) sites exhibited more complex networks (more OTUs) with higher connectivity; also, keystone species of these communities had more connections, compared to those in the site of low salinity (Öland). This suggest that the communities were functionally affected by either salinity, or by distance from the entrance of the Baltic. This network analyses provides a first insight into the bacteria-eukaryote interactions in benthic sediment communities in the Baltic Sea, however, much remains unknown about these community interactions, not least of which the mechanisms and biological nature of these interactions.

Association with environmental factors other than salinity

Apart from salinity, other environmental factors also affected the diversity of microbial benthic communities. Sediment nutrient composition had a strong influence on the bacterial community diversity, with organic matter being an important driver on the spatial scale, together with temperature (at the mesohaline sites Hersley, Gollwitz, Pori and two replicates of Tvärminne) (Fig. 3A). On the temporal scale, seasonal change in temperature was also an important driver between summer (August) communities and communities from colder winter and spring periods (November and April) (Fig. 3C). Changes in sediment characteristics were also important in explaining temporal changes. C/N ratio and sediment characteristics were mainly related to changes in community at the polyhaline site (Saltö), while changes at the mesohaline site Herslev were better characterised by changes in temperature together with C/N ratio. At the other mesohaline site, Öland, the community was more driven by changes in organic matter and sediment characteristics. For eukaryote communities the roles of environmental variables other than salinity have been described previously (Salonen et al. 2018, Zhang et al. 2018). Similar patterns were seen in the temporal communities in the present study, where composition over the year was driven to a higher degree by change in temperature, C/N ratio, organic matter and sorting, than it was by temporal changes in salinity. On a spatial scale, we found no clear patterns common to both eukaryote datasets for any of the sites.

Conclusion

This study provides an insight into the diversity of coastal sediment bacterial and microbial eukaryote communities in the Baltic Sea, on a broad spatial scale as well as over a seasonal cycle along the natural salinity gradient. We found salinity to be among the strongest drivers of both bacterial and eukaryote communities; however, not an equally strong factor for all communities; beta-diversity of mesohaline communities was better described by temperature, nutrients, and sediment characteristics. Bacteria and eukaryote communities also differed from each other, and thus, their species diversity respond differently to the same environmental conditions. The communities may however not only be influenced by external environmental factors, but also be affected by competition and predation within the communities. Analysis of bacteria-eukaryote networks provide a first insight into the microbial interactions in benthic sediment communities in the Baltic Sea. We found network complexity and connectivity to be higher in the communities of high salinity, but were functionally affected by salinity, showing distance decay. A more in-depth look at the community composition

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465 patterns at genus rather than phylum level could further elucidate bacterial and microbial eukaryote 466 interactions in Baltic Sea sediments. 467 468 Acknowledgements 469 This work was supported by Roskilde University (Department of Science and Environment), 470 University of Jyväskylä (Department of Biological and Environmental Science), Elite Research travel 471 grant, Ministry of Higher Education and Science, Denmark [to HCP], Faxe Fonden [to HCP], H. C. Wegges Mindelegat for Zoologer [to HCP], and Emil Aaltonen Foundation [grant number a6a412 to 472 ALH]. 473 474 We gratefully thank Professor Karsten Reise for help with field sampling and Alfred Wegener Institute Helmholtz Centre for Polar and Marine Research in List, Germany, for use of facilities. 475 476 Assistant professor and research scientist Stefan Forster for field assistance and Rostock University, 477 Germany, for use of facilities. Professor Kerstin Johannesson and Professor Per Jonsson for help and 478 guidance with collecting field samples, and Tjärnö field station, Gothenburg University, Sweden, for use of facilities. Research coordinator Laura Kauppi for assistance in field sampling, and Tvärminne 479 480 field station, University of Helsinki, Finland, for use of facilities. Master student Theophilus Alale for sampling help in Tvärminne and Pori. We thank Laboratory assistant Anne Busk Faarborg for 481 482 assisting with the field sampling and performing the grain size analysis, and Laboratory assistant G. 483 Katrine Bøg for performing CHN analysis. We also thank Master student Aleksi Kolehmainen for 484 assistance with the DNA extraction and amplicon sequencing, and Chief Laboratory Technician Elina Virtanen for assisting with sequencing. We also thank CSC-IT Center for Science, Finland for 485 486 providing computing facilities.

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Figure 1.

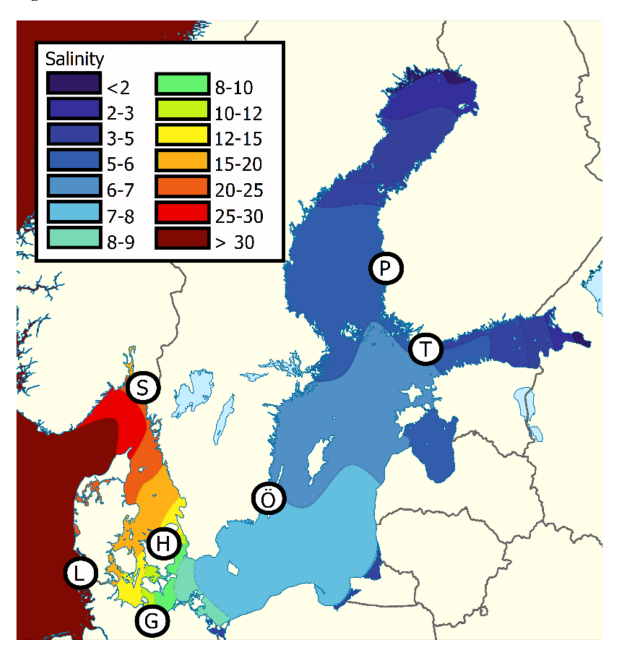


Figure 2.

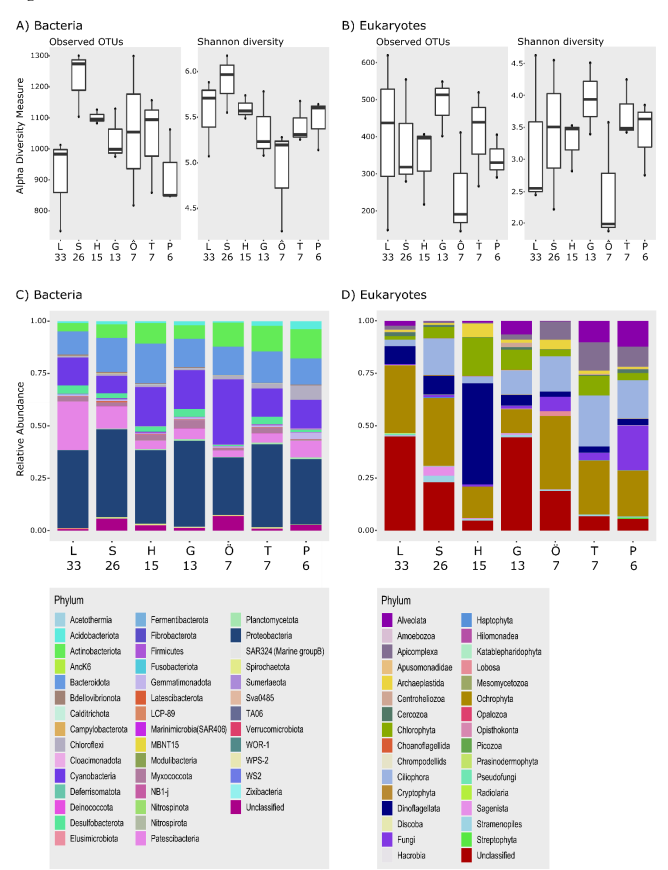


Figure 3.

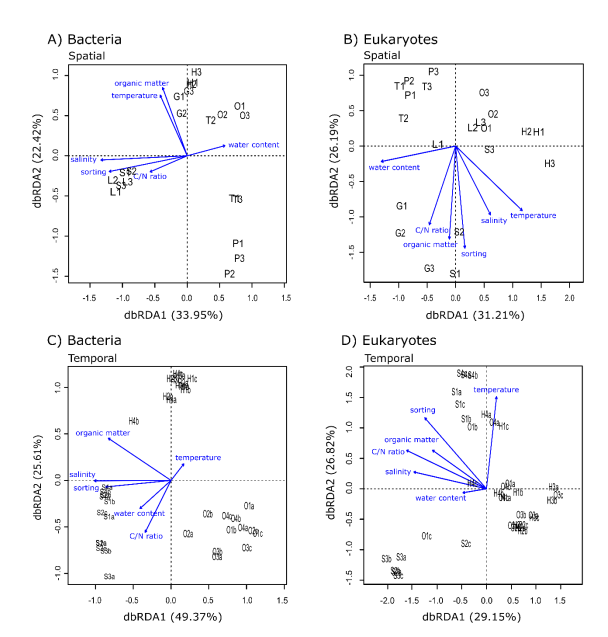


Figure 4.

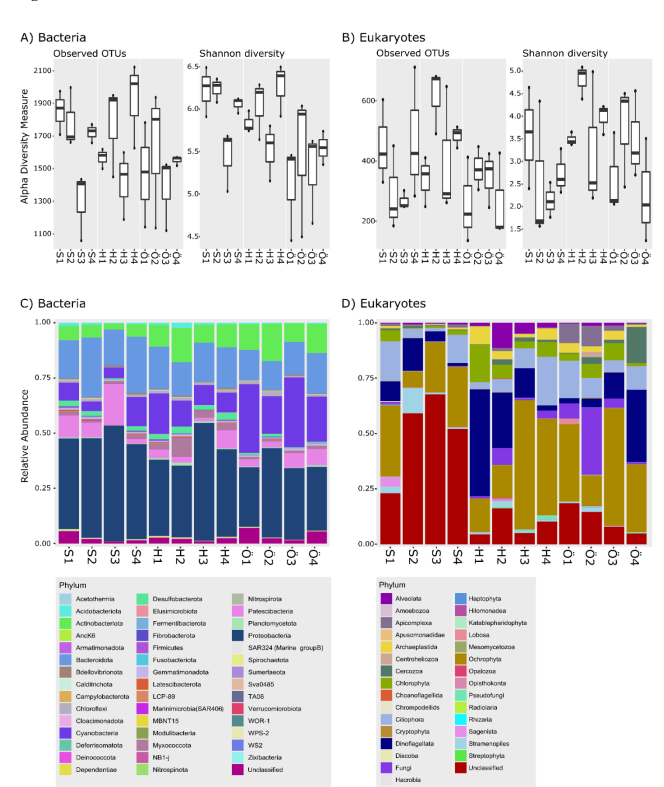


Figure 5a.

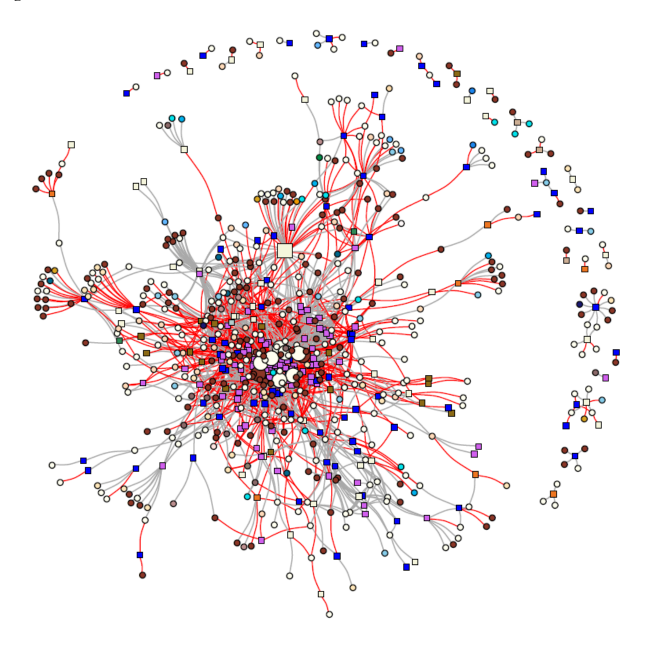


Figure 5b.

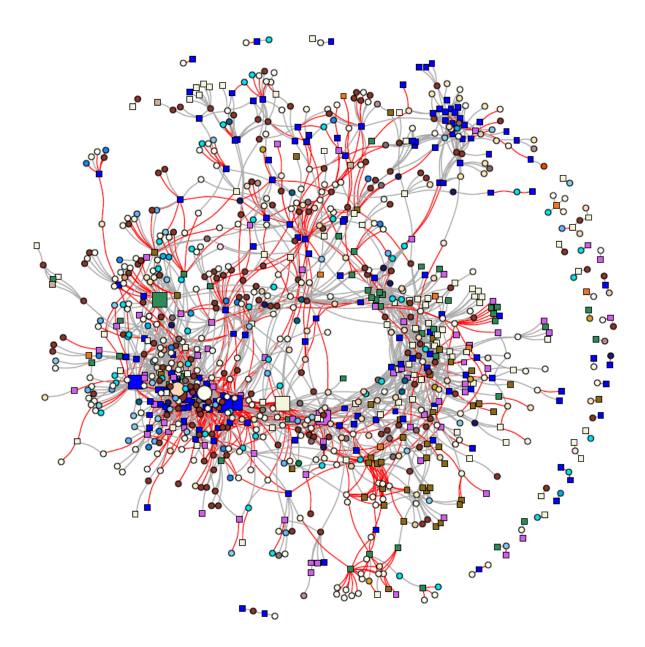
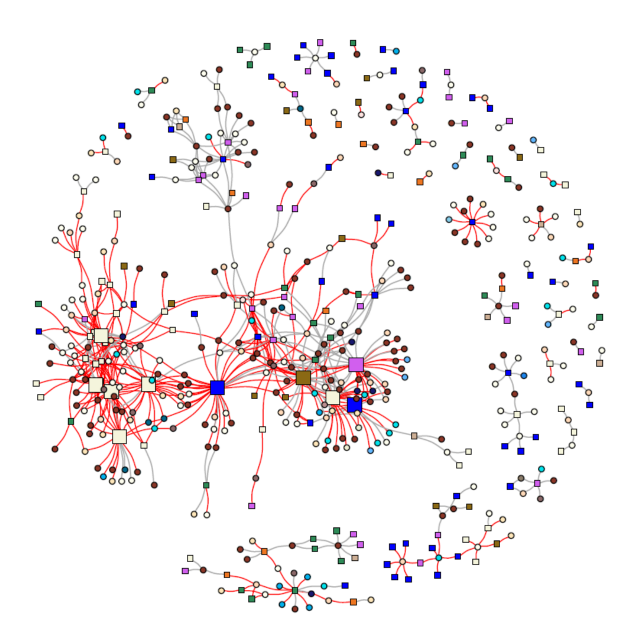


Figure 5c.



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Eukaryotes Bacteria Eukaryota_unclassified(0) Proteobacteria Alveolata Bacteroidota ■Rhizaria Cyanobacteria Latescibacterota ■ Archaeplastida Patescibacteria Stramenopiles ■Opisthokonta Bdellovibrionota ■Hacrobia Actinobacteriota Amoebozoa Spirochaetota ZUnclassified Fibrobacterota Desulfobacterota Gemmatimonadota Acidobacteriota AncK6 Campylobacterota Chloroflexi Planctomycetota •Myxococcota Sva0485 •Marinimicrobia_(SAR406_clade)

Table 1.

Factors	Bacteria	Eukaryotes	Bacteria	Eukaryotes
	spatial	spatial	temporal	temporal
Salinity	0.22***	0.08*	0.29***	0.07***
Temperature	0.09***	ns	0.05**	0.06**
C/N ratio	0.07**	0.08*	0.09***	0.05**
Org. matter	0.12***	ns	0.07***	ns
Water content	0.09***	0.08**	0.05**	ns
Porosity	0.07**	ns	0.07**	ns
Mean grain size	0.10859**	ns	0.09814***	0.05437**
Sediment sorting	0.17089***	ns	0.19869***	0.06962**

665 Figure legends

- 666 Figure 1. Map of the Baltic Sea and the adjacent North Sea, including sampling sites: L (List), S
- 667 (Saltö), H (Herslev), G (Gollwitz), Ö (Öland), T (Tvärminne), P (Pori). Salinity gradient based on
- Snoeijs-Leijonmalm and Andrén (2017).
- 669 Figure 2. Spatial patterns of microbial diversity of A, C) bacteria (16S), and B, D) microbial
- eukaryotes (18S), per site in: L (List), S (Saltö), H (Herslev), G (Gollwitz), Ö (Öland), T (Tvärminne),
- P (Pori). A) observed number of OTUs and alpha diversity index, Shannon Diversity, in bacteria
- 672 (16S) and B) observed number of OTUs and Shannon Diversity, in eukaryotes (18S), note y-axes are
- not equal. C) average relative abundance of phyla/divisions in bacteria (16S) and D) eukaryotes (18S).
- 674 Figure 3. Distance based Redundancy Analysis (dbRDA) based on Bray-Curtis dissimilarity of
- bacteria (16S) on spatial (A) and temporal scale (C) and microbial eukaryotes (18S) on spatial (B)
- and temporal scale (D). A) Sediment characteristics represented by vectors sorting and porosity, B)
- with sediment characteristics represented by vectors water content and sorting, C) and D) with all
- sediment characteristics represented; note axes are not equal.
- 679 Figure 4. Temporal patterns of microbial diversity of A, C) bacteria (16S), and B, D) microbial
- eukaryotes (18S), per site and collection time. Abbreviations of sample site and time: S1 (Saltö
- 681 August 2018), S2 (Saltö November 2018), S3 (Saltö April 2019), S4 (Saltö August 2019), H1
- 682 (Herslev August 2018), H2 (Herslev November 2018), H3 (Herslev April 2019), H4 (Herslev August
- 683 2019), Ö1 (Öland August 2018), Ö2 (Öland November 2018), Ö3 (Öland April 2019), Ö4 (Öland
- August 2019). A) Observed number of OTUs and alpha diversity index, Shannon Diversity, in
- bacteria (16S) and B) observed number of OTUs and Shannon Diversity, in eukaryotes (18S); note
- y-axes are not equal. C) Average relative abundance of phyla/divisions in bacteria (16S) and D)
- eukaryotes (18S).
- 688 Figure 5. Networks of microbial communities estimated from temporal samples; bacterial OTUs
- represented by circles, microbial eukaryote OTUs represented by squares, assignment to specific
- 690 OTUs indicated in legends. Keystone species (OTUs with highest number of connections) indicated
- 691 by larger symbols (see supplementary Tables 8A-C). Negative correlations indicated by red
- 692 connections (lines) and positive correlations indicated by grey connections (lines). Figure 5a
- 693 Communities at the high salinity site Saltö. Figure 5b Communities at the medium salinity site
- 694 Herslev. **Figure 5c** Communities at the low salinity site Öland.

695	Table 1. Correlation of community diversity with measured environmental variables for the spatial
696	and temporal datasets calculated with Adonis on Bray-Curtis distance matrices for bacterial and
697	microbial eukaryote community dissimilarity assessment using 1000 permutations on environmental
698	parameters. Values indicate R2, asterisks indicate significance level, *=p<0.001, **p=0.01,
699	***p=0.001, ns=not significant.
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Supplementary table 1. Table of primers.

Prim	ier pair	Sequences Amplicon	Reference
		size	
16S	27F + 338R	F: 5'-AGA GTT TGA TCM TGG CTC AG-3' 311 bp	Ludwig et
V1-		R: 5'-ATT ACC GCG GCT GCT GG-3'	al. 1993,
V2			Suzuki and
			Giovannoni
			1996
18S	UNonMetF+ UNonMetR	F: 5'-GTG CCA GCA GCC GCG-3' 600 bp	Bower et
V4		R: 5'-TTT AAG TTT CAG CCT TGC G-3'	al. 2004
	E572 + E1009R	F: 5'-CYG CGG TAA TTC CAG CTC-3' 400 bp	Comeau et
		R: 5'-CRA AGA YGA TYA GAT ACC RT-3'	al. 2011

Supplementary table 2. Thermocycling protocols for each target.

	UNonMetF/ UNonMetR	E572/E1009R
PCR	1. 94°C, 2 min	1. 94°C, 2 min
conditions	2. 94°C, 10 s	2. 94°C, 30 s
	3. 51,1 °C, 30 s – 35x	3. 55 °C, 30 s - 30x
	4. 72°C, 1 min _	4. 72°C, 1 min
	5. 72°C, 5 min	5. 72°C, 5 min

Supplementary table 3. Target specific M13-linker primer and fusion primer sequences. Each sample was barcoded with 10-12 bp long unique barcode (marked with N) attached to IonA-forward fusion primer.

Primer	Sequence
M13 _27F	TGTAAAACGACGCCAGTAGAGTTTGATCMTGGCTCAG
M13_E572F	TGTAAAACGACGCCAGTCYGCGGTAATTCCAGCTC
IonP1_338R	CCTCTCTATGGGCAGTCGGTGATTGCTGCCTCCCGTAGGAGT
IonP1_E1009R	CCTCTCTATGGGCAGTCGGTGATCRAAGAYGATYAGATACCRT
IonA _key_bc_ M13	CCATCTCATCCCTGCGTGTCTCCGACTCAGNNNNNNNNNN

Supplementary table 4. Environmental variables measured at all samplings.

Site	Sample time	Salinity	Temp.	C/N ratio	Organic matter	Water- content	Porosity	M. grain	Sorting
List		22	22	9.48	0.940	17.59	0.34		1 27
List	August 2018	33	22	9.48	0.940	17.39	0.34	1.41	1.27
Saltö	August 2018	26	18	10.46	1.16	24.03	0.43	2.24	1.76
	November 2018	25	3	8.86	1.24	27.22	0.50	2.82	0.82
	April 2019	23	11	8.57	1.07	21.20	0.40	2.35	1.53
	August 2019	22	18	7.77	1.34	25.17	0.47	2.16	1.90
Herslev	August 2018	15	23	6.31	1.06	21.56	0.40	1.37	0.70
	November 2018	16	5	4.40	1.16	23.35	0.48	2.15	0.85
	April 2019	15	15	1.94	1.02	22.46	0.40	2.35	0.85
	August 2019	15	17	8.30	1.23	20.37	0.37	2.06	1.05
Gollwitz	August 2018	13	23	9.83	1.42	27.63	0.49	2.78	0.89
Öland	August 2018	7	21	9.89	0.89	22.69	0.43	2.40	0.63
	November 2018	8	1	3.83	0.61	24.15	0.44	2.70	0.41
	April 2019	9	14	7.06	0.55	23.42	0.44	2.50	0.46
	August 2019	8	18	6.90	0.70	20.75	0.37	2.48	0.49
Tvärminne	August 2018	7	13	6.82	1.40	32.41	0.56	2.60	0.67

Pori	August	6	18	8.64	0.24	22.42	0.40	2.49	0.63
	2018								

Supplementary table 5a. Differences in Shannon index between sites in spatial study by TukeyHSD test. Different letters indicate significantly different sites. No significant differences were found by Tukey HSD test for richness of bacteria and eukaryotes.

Spatial Dataset	Shannon_Bacteria
Herslev	ab
List	ab
Pori	ab
Saltö	a
Tvärminne	ab
Öland	ь
Gollwitz	ab

Supplementary table 5b. Differences in Shannon index and richness (observed diversity) between sites in temporal study tested by TukeyHSD test. Different letters for each site indicate significantly different sites.

Temporal				
Dataset	Bac	Bacteria		aryote
Diversity	Shannon	Observed	Shannon	Observed
Saltö	a	a	a	ab
Öland	ь	a	ab	a
Herslev	a	a	ь	ь

Supplementary table 6. ANOVA test of alpha diversity measured by richness (observed) and Shannon against salinity. Note salinity is grouped to three categories as low (<10), medium (>10-20) and high (>20). Values indicate p value, asterisk indicate significance level, *=p<0.001, **p=0.01, ns=not significant.

Dataset	Observed	Shannon
Spatial Bacteria	ns	0.06
Temporal Bacteria	ns	0.00 **
Spatial Eukaryotes	ns	ns

Temporal Eukaryotes	0.04 *	0.03 *	
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- **Supplementary table 7.** See attached excel-sheet. Correlation between OTUs and salinity.
- Supplementary table 8a. Keystone species of high salinity site Saltö, temporal samples. OTU
 identity listed as botu = bacterial OTU, eotu = eukaryote OTU.

OTU	Connections	Phylum	Order	Genus
botu515	60	Gammaproteobacteria	Halieaceae	Proteobacteria
botu607	55	Alphaproteobacteria	Rhodobacteraceae	Proteobacteria
botu331	53	Gammaproteobacteria	BD2-7	Proteobacteria
botu613	52	Gammaproteobacteria	BD7-8_fa	Proteobacteria
botu700	50	Gammaproteobacteria	Saccharospirillaceae	Proteobacteria
botu239	47	Alphaproteobacteria	uncultured	Proteobacteria
eotu228	45	Ochrophyta	Bacillariophyta	Stramenopiles
botu1484	44	Bacteroidia	Crocinitomicaceae	Bacteroidota
botu3889	43	Bacteroidia	Flavobacteriaceae	Bacteroidota
botu183	42	Gammaproteobacteria	Alteromonadaceae	Proteobacteria
eotu520	41	Unclassified	Unclassified Eukaryota	Unclassified Eukaryota
botu2817	41	Bacteroidia	Flavobacteriaceae	Bacteroidota
botu929	41	Alphaproteobacteria	Rhodobacteraceae	Proteobacteria
eotu1227	40	Unclassified	Unclassified Eukaryota	Unclassified Eukaryota
eotu356	40	Ochrophyta	Phaeophyceae	Stramenopiles

- **Supplementary table 8b.** Keystone species of medium salinity site Hersley, temporal samples.
- OTU identity listed as botu = bacterial OTU, eotu = eukaryote OTU.

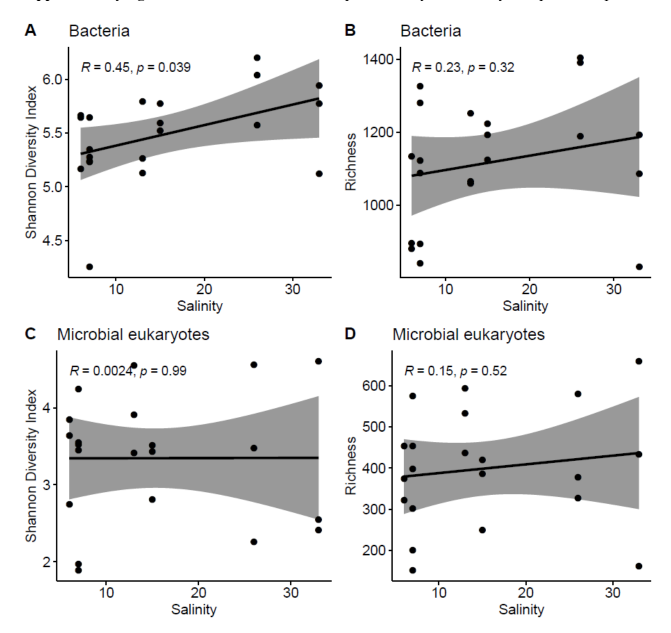
OTU	Connections	Phylum	Order	Genus
botu1673	51	Bacteroidota	Flavobacteriales	Unclassified Flavobacteriaceae
eotu933	45	Alveolata	Dinophyceae	Dinophyceae
eotu3734	40	Alveolata	Dinophyceae	Unclassified Dinophyceae
eotu34	37	Stramenopiles	Bacillariophyta	Unclassified Raphid pennate
eotu1174	36	Opisthokonta	Ascomycota	Unclassified Pezizomycotina
botu607	35	Proteobacteria	Rhodobacterales	Unclassified Rhodobacteraceae
eotu113	30	Unclassified Eukaryota	Unclassified Eukaryota	Unclassified Eukaryota
eotu3124	28	Alveolata	Dinophyceae	Unclassified Dinophyceae
eotu4284	28	Alveolata	Dinophyceae	Unclassified Dinophyceae
botu370	28	Patescibacteria	Candidatus Campbellbacteria	Candidatus Campbellbacteria_ge
eotu363	27	Unclassified Eukaryota	Unclassified Eukaryota	Unclassified Eukaryota
eotu681	27	Stramenopiles	Unclassified Stramenopiles	Unclassified Stramenopiles
botu1222	27	Bacteroidota	Chitinophagales	uncultured
botu290	27	Bacteroidota	Flavobacteriales	Lutibacter

botu4620	27	Bacteroidota	Flavobacteriales	Aquibacter
eotu852	26	Alveolata	Unclassified Alveolata	Unclassified Alveolata

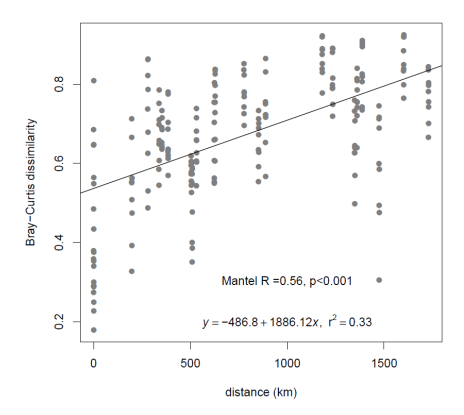
 Supplementary table 8c. Keystone species of low salinity site Öland, temporal samples. OTU identity listed as botu = bacterial OTU, eotu = eukaryote OTU.

OTU	Connections	Phylum	Order	Genus
eotu368	34	Alveolata	Dinophyceae	Unclassified Gymnodiniaceae
eotu338	32	Unclassified Eukaryota	Unclassified Eukaryota	Unclassified Eukaryota
eotu67	30	Archaeplastida	Chlorophyceae	Unclassfied Chlamydomonadales
eotu3341	29	Stramenopiles	Phaeophyceae	Unclassfied Phaeophyceae
eotu231	26	Stramenopiles	Bacillariophyta	Unclassfied Raphid-pennate
eotu135	22	Alveolata	Unclassfied Alveolata	Unclassfied Alveolata
eotu176	22	Stramenopiles	Bacillariophyta	Unclassfied Raphid-pennate
eotu497	21	Stramenopiles	Bacillariophyta	Unclassfied Raphid-pennate
eotu393	20	Stramenopiles	Unclassified Ochrophyta	Unclassified Ochrophyta
eotu863	20	Stramenopiles	Bacillariophyta	Unclassfied Raphid-pennate
eotu730	17	Stramenopiles	Unclassified Sagenista	Unclassified Sagenista

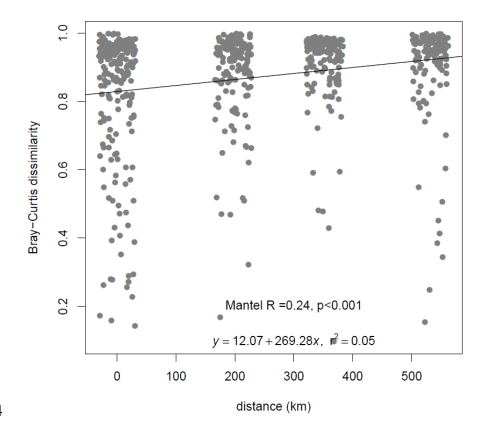
Supplementary figure 1. Pearson correlation of alpha diversity and salinity for spatial samples.



Supplementary figure 2a. Analysis of distance-decay for spatial bacterial dataset.

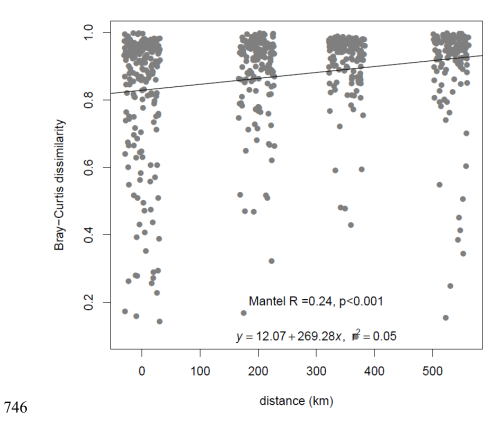


Supplementary figure 2b. Analysis of distance-decay for temporal bacterial dataset.



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Supplementary figure 1c. Analysis of distance-decay for temporal microbial eukaryote dataset.



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Supplementary figure 3. Pearson correlation of alpha diversity and salinity for temporal samples.

