



Benthic protists: the undercharted majority

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1 1 Benthic protists: the undercharted majority

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1
2 **52 Abstract**

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4 53 Marine protist diversity inventories have largely focused on planktonic
5 environments, while benthic protists have received relatively little attention. We
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7 54 therefore hypothesize that current diversity surveys have only skimmed the surface
8 of protist diversity in marine sediments, which may harbour greater diversity than
9
10 55 planktonic environments. We tested this by analyzing sequences of the hypervariable
11 56 V4 18S rRNA from benthic and planktonic protist communities sampled in European
12 57 coastal regions. Despite a similar number of OTUs in both realms, richness
13 58 estimations indicated that we recovered at least 70% of the diversity in planktonic
14 59 protist communities, but only 33% in benthic communities. There was also little
15 60 overlap of OTUs between planktonic and benthic communities, as well as between
16 61 separate benthic communities. We argue that these patterns reflect the heterogeneity
17 62 and diversity of benthic habitats. A comparison of all OTUs against the Protist
18 63 Ribosomal Reference database showed that a higher proportion of benthic than
19 64 planktonic protist diversity is missing from public databases; similar results were
20 65 obtained by comparing all OTUs against environmental references from NCBI's Short
21 66 Read Archive. We suggest that the benthic realm may therefore be the world's
22 67 largest reservoir of marine protist diversity, with most taxa at present undescribed.
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2 70 **Introduction**

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5 71 Molecular studies of marine plankton have uncovered an enormous diversity
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7 72 of protists, many of which could not be assigned to any accession in taxonomic
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9 73 reference databases (e.g., de Vargas *et al.* 2015). Planktonic studies have thus
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11 74 found a very large pool of unknown marine protist species. However, initial molecular
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13 75 studies unveiled highly diverse protist communities from marine benthic habitats
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15 76 which exhibit key ecosystem functions and whose diversity may even exceed that of
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17 77 planktonic protists. Most of these benthic studies were conducted in hydrothermal
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19 78 vent systems (Edgcomb *et al.* 2002; López-García *et al.* 2003; López-García,
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22 79 Vereshchaka and Moreira 2007) and anoxic sediments (Dawson and Pace 2002;
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24 80 Stoeck and Epstein 2003; Takishita *et al.* 2005), both of which environments
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26 81 emerged as hotspots of protist biodiversity. More recently, similarly high diversity was
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28 82 reported from deep-sea and coastal sediments, with many genetic signatures only
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30 83 distantly related to taxonomically described protists (Scheckenbach *et al.* 2010;
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32 84 Pawłowski *et al.* 2011; Bik *et al.* 2012; Gong *et al.* 2015). At least for the three major
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34 85 protist lineages of ciliates, diatoms and foraminifera, the existing species inventories
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36 86 of morphologically delineated species support a higher and distinctively different
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38 87 diversity of benthic compared to planktonic species (Patterson, Larsen and Corliss
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40 88 1989; Mann and Evans 2007; Pawłowski, Holzmann and Tyszka 2013).

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42 89 Marine benthic studies, though, remain scarce and limited compared to the
43
44 90 considerably larger amount of planktonic studies. Based on the sparse and locally
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46 91 restricted amount of data, no consensus has been reached on the extent of microbial
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48 92 eukaryotic diversity in marine sediments (Epstein and López-García 2007; Fierer
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50 93 2008; Bik *et al.* 2012). Methodological difficulties are a partial explanation for this
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52 94 undersampling. The clean extraction of nucleic acids from environmental marine
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54 95 sediment samples is challenging (Hurt *et al.* 2001) and extensive amounts of

1
2 96 extracellular DNA may severely bias the environmental sequencing studies
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4 97 (Dell'Anno and Danovaro 2005). Furthermore, specific techniques and expensive
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6 98 equipment are required to access these physically remote environments (Orcutt *et al.*
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8 99 2011). Published benthic studies of protists are thus not only restricted in geographic
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10 100 scope, but also in their comparisons to the overlying planktonic protists. Despite our
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12 101 limited knowledge of benthic protists, there is strong agreement among microbial
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14 102 ecologists that marine coastal sediments play a pivotal role for the diversity and
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16 103 dynamics of overlying plankton communities by acting as seedbanks (Marcus and
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18 104 Boreo 1998). The benthic species reservoir consists of both truly active benthic
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20 105 species and resting stages of dormant planktonic species. Most of the latter occur in
21
22 106 low abundances, but blooms can be initiated in response to environmental changes
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24 107 (Marcus and Boreo 1998). Furthermore, studies on microfossil protists have
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26 108 suggested that several planktonic lineages have evolved from benthic ancestors,
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28 109 which have colonized the pelagic realm on different occasions (Leckie 2009).

34 110 In this study, we used previously published data from Massana *et al.* (2015)
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36 111 that surveyed protist diversity in European coastal waters and sediments from
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38 112 Norway to Bulgaria using the V4 region of the 18S rRNA. Massana *et al.* (2015)
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40 113 provided a general taxonomic overview of planktonic and benthic protists, with
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42 114 particular attention on examining differences in size-fractionated planktonic protist
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44 115 communities and comparing results obtained from DNA and RNA templates. Building
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46 116 upon this initial study, here we focus on a more detailed comparison of benthic and
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48 117 planktonic protist diversity, with a special emphasis on the richness and phylogenetic
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50 118 novelty contained in marine benthic protist assemblages, two topics that were not
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52 119 covered by Massana *et al.* (2015). Our results not only support previous notions of
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54 120 more diverse benthic than planktonic protist communities, but also present clear

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2 121 indications of a higher degree of novelty in genetic signatures within benthic
3 122 communities.
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11 125 **Methods**
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14 126 *Sampling, pyrosequencing and data processing*
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16 127 In the framework of the BioMarKs project (Dunthorn *et al.* 2014a; Logares *et*
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18 128 *al.* 2014; Massana *et al.* 2015), water samples from the surface and deep chlorophyll
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20 129 maximum (DCM) layer, and sediment samples were collected from six different
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22 130 European coastal sites including the Skagerrak, the English Channel, the Atlantic
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24 131 Ocean, the Mediterranean Sea and the Black Sea (Table 1). For details on sampling
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26 132 protocols, nucleic acid extraction, 454-pyrosequencing of the hypervariable V4 18S
27
28 133 rDNA region and data processing see Massana *et al.* (2015). Briefly, RNA from
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30 134 benthic samples was extracted from 2.5 g of surficial sediment using the Power Soil
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32 135 RNA kit (MoBio, Carlsbad, United States). RNA from planktonic samples was
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34 136 extracted with the NucleoSpin RNA kit (Macherey-Nagel, Düren, Germany) from
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36 137 filters of DCM and surface water samples collected with Niskin bottles. Both
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38 138 extraction kits are specifically optimized to gain high RNA yields from the respective
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40 139 samples and are routinely used in environmental high-throughput sequencing (HTS)
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42 140 studies. Because of the difficulties in RNA extractions from sediment samples (Hurt
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44 141 *et al.* 2001), using an optimized kit for the recovery of nucleic acids from benthic
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46 142 samples was especially important. By targeting environmental (e)RNA rather than
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48 143 environmental (e)DNA we minimized potential biases induced by accumulation and
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50 144 preservation of extracellular eDNA in sediments (Dell'Anno and Danovaro 2005;
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52 145 Stoeck *et al.* 2007). The effect of extracellular eDNA is less critical when only water
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54 146 samples are analyzed (Logares *et al.* 2014), but relevant for a comparison of water

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2 147 with sediment samples. Extracted RNA was then transcribed to cDNA for further
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4 148 processing.
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7 149 Targeted amplification of the hypervariable V4 18S rDNA region was
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9 150 performed with the eukaryotic primer pair TAReuk454FWD1 and TAReukREV3
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11 151 (Stoeck *et al.* 2010). This primer pair, though, does not cover Foraminifera and
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13 152 several excavate lineages that are important contributors to benthic protist
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15 153 communities but whose taxonomy is not very well delineated by the V4 region
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17 154 (Pawlowski *et al.* 2011; Lax and Simpson 2013). **In contrast to other barcode genes,**
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19 155 **the V4 region does allow for better comparisons with published full-length 18S rRNA**
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21 156 **Sanger sequencing studies and has a better database coverage for taxonomic**
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23 157 **assignment** (Stoeck *et al.* 2010; Dunthorn *et al.* 2012). Pyrosequencing of the
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25 158 amplified PCR product was conducted on a 454 GS FLX Titanium system (454 Life
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27 Sciences, USA). Resulting 454 reads were subject of a strict quality filtering,
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29 159 including two steps of chimera checking in UCHIME (Edgar *et al.* 2011) and
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31 160 ChimeraSlayer (Haas *et al.* 2011). All quality sequences were then clustered into
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33 161 OTUs using USEARCH (Edgar 2010) on a 97% sequence similarity value. In a
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35 162 second step of quality filtering, one representative of each OTU was subject to a
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37 163 BLASTn analysis against NCBI's nucleotide database release 183.0. All OTUs
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39 164 assigned to Bacteria, Archaea, Metazoa, Embryophyta and OTUs with less than 80%
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41 165 similarity to database entries were removed from the dataset. The final dataset
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43 166 included 430 894 V4 18S RNA sequences, which clustered into 12 438 distinct
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45 167 OTUs. The complete BioMarkS sequencing dataset is available at the European
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47 168 Nucleotide Archive under the study accession number PRJEB9133.
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53 171 **Taxonomic assignment of OTUs**
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2 172 To search for the best hit of each OTU to a described organism, we conducted
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4 173 a local BLASTn analysis (using default settings) against the protist reference
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6 174 database PR² (Guillou *et al.* 2012). Additionally, we compared our environmental
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8 175 data against reference data of previous environmental HTS diversity surveys of
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10 176 protists using a similar BLASTn analysis. To build this environmental reference
11 177 database, we manually screened NCBI's Short Read Archive (SRA) for studies that
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13 178 at least partially included protist data of the eukaryotic 18S gene (according to the
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15 179 experiment's descriptions in the SRA). After downloading the respective data, we
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17 180 removed all references shorter than 100 bp. The final customized SRA reference
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19 181 database consisted of 11 708 385 references from 167 datasets (Supplemental
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21 182 Table S1).

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183
184 *Statistical analyses and diversity measures*

185 All statistical and diversity analyses were performed in R Studio (version
186 2.15.1, <http://r-project.org>). Following recommendations of previous studies
187 (Dunthorn *et al.* 2014b) we relied on incidence-based rather than abundance-based
188 data to avoid biases induced by uneven gene copy numbers among different protist
189 taxa. Species richness was estimated with the incidence-based coverage estimator
190 (ICE) as implemented in the 'fossil' package (Vavrek 2011). ICE appropriately
191 estimates asymptotic species richness from datasets containing many rare species
192 (Colwell *et al.* 2012), which we expect in benthic protist communities. Additional
193 species richness estimations in CatchAll (Bunge *et al.* 2012) are provided as
194 supplemental material (Supplemental Fig. S2). Non-metric multidimensional scaling
195 (NMDS) using (binary-) Jaccard distances as a measure of β-diversity was performed
196 with the 'vegan' package (Oksanen *et al.* 2015). Non-parametrical, two-sided
197 Kolmogorov-Smirnov tests (KS-tests) using 1000 bootstrap replicates were used to

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3 198 assess the significance of sequence divergence distribution from planktonic and
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5 199 benthic datasets in the package 'Matching' (Sekhon 2011).
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7 200
8
9 201 **Results**
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11 202 *Comparison of planktonic and benthic protist diversity*
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13 203 The number of quality filtered sequences among the three habitats differed:
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15 204 206 602 from the surface, 184 192 from the DCM, and 40 100 from the sediment
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18 205 (Table 1). Despite these differences, the sequences clustered into similar numbers of
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20 206 OTUs: 5747 in the surface, 5685 in the DCM, and 5616 in the sediment. However,
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22 207 ICE-based richness estimates predicted a difference in the number of total OTUs
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24 208 between the plankton and the sediment: 7763 from the surface, 8140 from the DCM,
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26 209 and 16 652 from the sediment (Fig. 1A). The proportion of OTUs thus not detected
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28 210 through our sequencing effort was 26% in surface, 30.2% in the DCM, and 66.3% in
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30 211 the sediment. Richness estimations based on abundance data revealed similar
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32 212 trends but predicted even more undetected OTUs (Supplemental Fig. S2). The
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34 213 observed trends on OTU richness were further congruent with rarefaction results of
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36 214 sequencing data, which were closer to saturation for water column than for sediment
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38 215 samples (Supplemental Fig. S3). Differences in community composition between
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40 216 plankton and benthos were first shown by the number of shared and exclusive OTUs
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42 217 (Fig. 1B). Of 7729 non-singleton OTUs observed in total, only 708 (9.2%) were
43
44 218 shared among all three habitats. This low number is, in part, explained by the low
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46 219 number of co-occurring OTUs in plankton and benthos: the plankton had 4368 non-
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48 220 singleton OTUs not found in the benthos, of which 701 were exclusively found in the
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50 221 surface and 762 were exclusively found in the DCM; the benthos had 2364 non-
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52 222 singleton OTUs not found in the plankton.
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3 223 NMDS analysis further demonstrated the partitioning of the observed diversity
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5 224 patterns in plankton and benthos (Fig. 2). Since differentiation in size fractions was
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7 225 only possible for planktonic samples and beyond the scope of this work, we pooled
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9 226 planktonic data of different size fractions from the same sampling events. In this
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11 227 analysis, surface and DCM samples clustered closely together, indicating a higher
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13 228 similarity in community composition among plankton samples than among benthos
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15 229 samples. Sediment samples were separated from the water communities in the
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17 230 analysis. In contrast to the narrow cluster of surface and DCM samples, the sediment
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19 231 samples were more widely distributed, indicating high dissimilarity in community
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21 232 composition between individual samples. This also applied to sediment samples from
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23 233 the same sampling site taken in consecutive years, as shown by the large distances
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25 234 between both samples from Naples and both samples from Oslo.
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34 236 *Taxonomic affiliation of plankton and benthos OTUs*
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37 237 Most OTUs in each habitat were assigned to the groups of Alveolata,
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39 238 Stramenopiles and Rhizaria (Fig. 3). While the Alveolata dominated the planktonic
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41 239 communities (3281 OTUs in surface, 3638 OTUs in DCM samples), rhizarian OTUs
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43 240 of the phylum Cercozoa dominated the benthic communities (1566 of 1622 benthic
44
45 241 rhizarian OTUs). Approximately the same proportion of OTUs was assigned to
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47 242 Stramenopiles in all three habitats. The number of OTUs from the Amoeboza,
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49 243 Apusozoa, Opisthokonta (predominantly fungi), and Picozoa were notably higher in
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51 244 the benthos than in the plankton. In contrast, Archaeplastida and Hacrobia were
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53 245 more diverse in planktonic (238 and 309 OTUs in surface samples; 175 and 264
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55 246 OTUs in DCM samples) than in benthic communities (50 and 179 OTUs).
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58 247 On a lower taxonomic level, we observed a higher OTU richness in benthic
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60 248 samples for 10 of the 19 most abundant groups (Fig. 4). In seven of these groups,

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3 249 the OTUs detected in the benthos accounted for more than 70% of the OTUs. These
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5 groups comprised the Apicomplexa (Alveolata), Apusomonadidae (Incetae Sedis),
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7 Centrohelida (Hacrobia), Discosea and Tubulinea (both Amoebozoa), as well as the
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9 previously mentioned Cercozoa and Fungi. Further groups mainly detected in the
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11 benthos included Ciliophora (Alveolata), Bacilliarophyceae and Labyrinthulomycetes
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13 (both Stramenopiles). Dinoflagellates, however, as the taxonomic group with highest
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15 OTU richness in total, were predominantly detected in planktonic samples.
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18 256 Acantharea (Rhizaria), MAST (Stramenopiles) and MALV (Alveolata) are three other
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20 taxonomic groups with heterotrophic members that were distinctively more often
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22 detected in planktonic than in benthic samples. Similar observations were made
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24 among the predominantly phototrophic groups Chlorophyta (Archaeplastida),
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26 Chrysophyceae and Dictyochophyceae (both Stramenopiles), and Haptophyta
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28 (Hacrobia). Supplemental figures S4 and S5 provide a closer look on the occurrence
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30 and distribution of phototrophic taxa and show that Cryptophyta and Rhodophyta
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32 were also mostly found in planktonic samples, while more diatoms were found in the
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34 benthos (677 OTUs) than in the plankton (445 OTUs).
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265
266 *Database coverage and genetic divergence of plankton and benthos OTUs*

267 To assess how well the observed diversity of OTUs matched previously
268 collected data, we calculated their degree of genetic similarity to reference
269 sequences of the taxonomically curated PR² database and to environmental
270 reference sequences of earlier protist diversity inventories deposited in NCBI's SRA
271 (Fig. 5). In both BLAST analyses, the novelty profile was much higher for benthic
272 OTUs, while surface and DCM OTUs exhibited an almost identical profile. Moreover,
273 the genetic similarity of BioMarKs OTUs to PR² references (Fig. 5A) was significantly
274 lower ($p < 0.001$) than to environmental references of the SRA database (Fig. 5B).

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2 275 Considering that 71.8% of the planktonic OTUs in our study can be assigned on a
3 276 97%-sequence similarity value to references in PR², this database misses 28.2% of
4 277 the potentially detectable plankton species (conservative estimate, see discussion).
5
6 278 Among benthic protist communities 74.4% of the OTUs did not retrieve a
7 279 taxonomically assigned hit in PR² at the same threshold of 97% similarity. Regarding
8 280 the SRA BLAST results, 78.6% of the planktonic OTUs in our study (n=8988 OTUs)
9 281 were at least 97% similar to previously deposited environmental references,
10 282 compared to only 42.7% of the OTUs from benthic samples (n=2400 OTUs). These
11 283 numbers illustrate that the vast majority of planktonic OTUs had already been
12 284 detected in previous environmental diversity surveys. This, however, was not the
13 285 case for benthic OTUs. Thus, BioMarKs benthos analyses contributed a high
14 286 proportion of novel OTUs to the environmental reference database.

15
16 287 On closer examination of the PR² BLAST results, the mean sequence
17 288 divergence of OTUs detected in surface and DCM samples from references of the
18 289 taxonomic reference database was 2.6% (Fig. 6). By contrast, the mean sequence
19 290 divergence of OTUs from sediment samples to PR² accessions was 6.6%, indicating
20 291 that the benthic compartment contains protist species more distantly related to
21 292 previously deposited taxa than the planktonic compartment. Specifically among the
22 293 Alveolata, Hacrobia, Opisthokonta, Rhizaria and Stramenopiles, the difference in
23 294 divergence between plankton and benthos was statistically significant ($p<0.001$). For
24 295 example, while the mean sequence divergence of OTUs assigned to Rhizaria from
25 296 PR² references was 3.5% in both the surface and DCM, the averaged divergence
26 297 was 8.2% in the sediment.

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28 298
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30 300 **Discussion**

1
2 301 *High α- and β-diversity shapes coastal benthic protist communities*
3 302 The α- and β-diversity patterns (Figs. 1 and 2) of protists along the European
4 303 coastline support previous notions of highly diverse protist communities from different
5 304 benthic habitats in the world's oceans (Dawson and Pace 2002; Edgcomb *et al.*
6 305 2002; López-García *et al.* 2003; Stoeck and Epstein 2003; Takishita *et al.* 2005;
7 306 López-García, Vereshchaka and Moreira 2007; Scheckenbach *et al.* 2010;
8 307 Pawlowski *et al.* 2011; Bik *et al.* 2012; Gong *et al.* 2015). A direct comparison of OTU
9 308 numbers to many of these initial studies is difficult, since most relied on clone library
10 309 approaches (Dawson and Pace 2002; Edgcomb *et al.* 2002; López-García *et al.*
11 310 2003; Stoeck and Epstein 2003; Takishita *et al.* 2005; López-García, Vereshchaka
12 311 and Moreira 2007; Scheckenbach *et al.* 2010) that produced fewer genetic signatures
13 312 than the current pyrosequencing approach. Consequently, Scheckenbach *et al.*
14 313 (2010) estimated a mean OTU richness of 489 in benthic biodiversity hotspots, such
15 314 as hydrothermal vents, and 1240 in abyssal sediments. These numbers would be
16 315 well below the mean estimated species richness of 2776 OTUs in BioMarKs
17 316 sediment samples. Other studies relying on 454 pyrosequencing detected similarly
18 317 high benthic diversity (Pawlowski *et al.* 2011; Bik *et al.* 2012; Gong *et al.* 2015). One
19 318 of these surveys observed between 393 and 1049 protist OTUs and estimated
20 319 between 421 and 1051 OTUs in coastal sediment sites of the Yellow Sea (Gong *et*
21 320 *al.* 2015). OTU richness in European coastal sediment samples was higher, with
22 321 observed protist OTU numbers ranging from 493 to 2499 and estimated OTU
23 322 numbers ranging from 721 to 3573. The BioMarKs numbers are closer to those
24 323 observed in Arctic and Southern Ocean deep-sea samples (between 942 and 1756
25 324 observed OTUs) (Pawlowski *et al.* 2011).

26 325 In contrast to most previous studies, our data allowed us to analyze benthic
27 326 communities in the context of planktonic communities from overlying water masses of

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2 327 the euphotic zone. Such data are scarcely available for protists and usually limited to
3 328 specific lineages. One example is a Sanger sequencing study that focused on ciliates
4 329 (Doherty *et al.* 2010), which reported little overlap between genetic signatures of
5 330 benthic and planktonic communities in the Gulf of Maine and Long Island Sound.
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7 331 This finding corroborates our observations on whole protist communities in European
8 332 coastal habitats (Fig. 1B). More data exists, however, for benthic-planktonic
9 333 community comparisons of marine bacteria. Zinger *et al.* (2011) showed that
10 334 bacterial OTU richness and β -diversity was much higher in coastal sediments than in
11 335 coastal surface waters. Similar results were obtained in an arctic fjord HTS survey
12 336 (Teske *et al.* 2011). These patterns nicely reflect those obtained in our study on
13 337 protists.

14 338 Building upon the Massana *et al.* (2015) study, we targeted the question of the
15 339 magnitude of benthic compared to planktonic diversity by contrasting the degree of
16 340 observed richness with the degree of estimated richness (Fig. 1A). In the same
17 341 context, we highlighted how much of this observed diversity likely represented novel
18 342 diversity (Figs. 5 and 6), a previously unexamined topic. Furthermore, we contrasted
19 343 the β -diversity among benthic and planktonic communities (Fig. 2), illustrating small
20 344 overlap between benthic and planktonic diversity, but also among benthic
21 345 communities in particular. Given that biodiversity is considered to be higher in coastal
22 346 than in open ocean habitats (Gray 1997; Zinger *et al.* 2011), the BioMarKs data
23 347 suggest that marine coastal sediments may be the world's largest reservoir of protist
24 348 diversity, much of which is still undetected and most of which is still undescribed in
25 349 public databases.

26 350 One major factor that might promote high diversity in benthic protist
27 351 communities is a large number of distinct benthic habitats due to horizontal and
28 352 vertical gradients in both physical and chemical characteristics. Even at microscale,

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2 353 habitat heterogeneity in marine sediments reflects gradients in grain-sizes, oxygen
3 354 concentration or organic matter content (Pedersen, Smets and Dechesne 2015). The
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5 355 diversity of physico-chemical microhabitats likely promotes the existence of highly
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7 356 specialized organisms and is probably an important driver for species-richness
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9 357 patterns (Hortal *et al.* 2009). We thus argue that our findings of high protist richness
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11 358 and heterogeneity at comparably small geographical scales represents a general
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13 359 trend in benthic diversity, which is well supported by previous findings, e.g. in the
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15 360 Yellow Sea, where high α - and β -diversity patterns in coastal marine sediments were
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17 361 unveiled (Gong *et al.* 2015). To further investigate the effect of niche partitioning,
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19 362 species-area relationships and distance-decay relationships on small spatial and
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21 363 temporal scales could be analyzed for planktonic and benthic protists at the same
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23 364 locales (Franzén, Schweiger and Betzholtz 2012; Zinger, Boetius and Ramette
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25 365 2014). In addition to habitat heterogeneity, geological structures at the seafloor may
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27 366 act as biogeographical barriers. Even locally, this results in a spatial separation of
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29 367 protist communities (Scheckenbach *et al.* 2010). Both factors, niche partitioning and
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31 368 allopatric speciation processes may work *in concerto* to generate and maintain a high
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33 369 diversity of protists in sediments. In direct comparison, environmental heterogeneity
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35 370 is surely much more pronounced in the benthos (Orcutt *et al.* 2011) than in the
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37 371 plankton, although the pelagic realm may create patchy distributions of protists as
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39 372 well (Menden-Deuer and Fredrickson 2010; Dolan and Stoeck 2011).

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41 373 Additional reasons for diverging protist communities between individual
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43 374 benthic sites (Fig. 2) may also be of technical nature: we found that all sediment
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45 375 samples were severely undersampled as a result of their high diversity
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47 376 (Supplemental Fig. S3). Because of this undersampling, community divergence
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49 377 among different sediment samples may be artificially inflated. It is, however,
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51 378 reasonable to assume that with increasing sampling effort the proportion of OTUs

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2 379 that are shared between two samples and the proportion of OTUs that are unique to
3 380 each of these samples would remain similar. We therefore argue that the observed
4 381 high Jaccard-distance between the sediment protist communities is more due to true
5 382 (biological) heterogeneity in species memberships rather than to (technical)
6 383 undersampling.
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17 385 *Different protist taxon groups thrive in plankton and benthos*
18 386 Benthic protist diversity uncovered by environmental HTS is comprised of i)
19 387 resident species of truly benthic origin; ii) transient species, which spend at least part
20 388 of their life cycle living actively in or on sediments; iii) non-resident species of
21 389 planktonic origin present as inactive resting stages, or as recently settled cells. OTUs
22 390 of resident species clearly dominated the benthic protist communities in our analyses
23 391 (Fig. 1B). Though transient species are commonly found among different taxonomic
24 392 groups (Garstecki *et al.* 2000), we found only little overlap between benthic and
25 393 planktonic OTUs (Figs. 1B and 2). A notable fraction of this overlap was related to
26 394 diatoms, Chrysophyceae and Chlorophyta (Supplemental Fig. S4). Since all of these
27 395 groups include species of planktonic origin which are able to form benthic resting
28 396 stages (McQuoid and Hobson 1996; Duff, Zeeb and Smol 2013), we cannot rule out
29 397 that at least some of the phototrophic OTUs in sediments could correspond to
30 398 phytoplankton cysts or cells that had sunk to the sea floor shortly before sampling.
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385 386 Benthic protist diversity uncovered by environmental HTS is comprised of i)
386 387 resident species of truly benthic origin; ii) transient species, which spend at least part
387 388 of their life cycle living actively in or on sediments; iii) non-resident species of
388 389 planktonic origin present as inactive resting stages, or as recently settled cells. OTUs
389 390 of resident species clearly dominated the benthic protist communities in our analyses
390 391 (Fig. 1B). Though transient species are commonly found among different taxonomic
391 392 groups (Garstecki *et al.* 2000), we found only little overlap between benthic and
392 393 planktonic OTUs (Figs. 1B and 2). A notable fraction of this overlap was related to
393 394 diatoms, Chrysophyceae and Chlorophyta (Supplemental Fig. S4). Since all of these
394 395 groups include species of planktonic origin which are able to form benthic resting
395 396 stages (McQuoid and Hobson 1996; Duff, Zeeb and Smol 2013), we cannot rule out
396 397 that at least some of the phototrophic OTUs in sediments could correspond to
397 398 phytoplankton cysts or cells that had sunk to the sea floor shortly before sampling.
398 399 Genetic signatures of the planktonic diatom family Leptocylindraceae in sediment
399 400 samples represent such a peculiar case (Nanjappa *et al.* 2014). However, particularly
400 401 among phototrophic protists, diatoms were more often detected in the benthos than
401 402 in the plankton. Indeed this group is known to harbour a larger diversity of benthic
402 403 than planktonic species, especially in shallow coastal waters (Mann and Evans
403 404 2007). Regarding the small amount of shared OTUs between benthos and plankton
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2 405 in general, we argue that genetic signatures of dead or sinking organisms or
3 406 dissolved RNA were considerably limited.
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7 407 A striking difference in the community composition between planktonic and
8 408 benthic protists at a higher taxonomic level is the dominance of numerous previously
9 409 undescribed Rhizaria in coastal sediments. More than 95% of these rhizarian OTUs
10 410 were assigned to the phylum Cercozoa (Supplemental Fig. S6), which have emerged
11 411 as an abundant and diverse lineage in several other benthic protist diversity studies
12 412 (summarized by Epstein and López-Garcia, 2008). This phylum comprises a large
13 413 number of gliding zooflagellates, filose and often large reticulose amoebae, which are
14 414 well adapted to a psammophilic lifestyle (Bass *et al.* 2009; Howe *et al.* 2011), but
15 415 also occur as parasites of invertebrates, algae and stramenopiles with benthos-
16 416 associated stages of their lifecycles (Hartikainen *et al.* 2014). Recent studies on
17 417 cercozoans could link many previously uncultured environmental sequences to novel
18 418 benthic cercozoans (Chantangsi and Leander 2010; Howe *et al.* 2011; Berney *et al.*
19 419 2013). In addition, genetically divergent benthic cercozoans, both free-living and
20 420 parasitic, are common and diverse but rarely detected in eukaryote-wide
21 421 environmental surveys, e.g. the amoeboid-flagellate *Reticulamoeba* (Bass *et al.* 2012)
22 422 or the parasitic *Ascetosporea* (Hartikainen *et al.* 2014). These examples further
23 423 emphasize the importance of this phylum in the marine benthos.
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26 424 Among the most abundant taxonomic groups we observed a trend of
27 425 distinctively higher OTU richness in benthic than in planktonic communities (Fig. 4).
28 426 Beside Cercozoa, this trend was especially prominent for Discosea and Tubulinea,
29 427 two groups of rhizopod Amoebozoa, which are common inhabitants of coastal
30 428 benthic ecosystems (Garstecki and Arndt 2000). Likewise, the detection of a high
31 429 proportion of saprotrophic fungi that contribute to detritus processing in marine
32 430 sediments is not surprising (Richards *et al.* 2012). Higher OTU richness in benthic
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2 431 samples, though less pronounced, was also observed for ciliates. This result is
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4 432 supported by previous morphological and molecular studies that reported higher
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6 433 benthic than planktonic ciliate diversity (Patterson, Larsen and Corliss 1989; Doherty
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8 434 *et al.* 2010). As discussed above, the situation was similar for diatoms.
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11 435 We detected an inverse trend in diversity among the Acantharea,
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13 436 Dinoflagellata and MALV, all of which are commonly observed in planktonic
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15 437 communities (Guillou *et al.* 2008; Jeong *et al.* 2010; Massana 2011; Decelle *et al.*
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17 438 2013). Acantharea and dinoflagellates are mostly planktonic organisms and both
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19 439 groups comprise mixotrophs or members with phototrophic symbionts (Gilg *et al.*
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21 440 2010; Hansen 2010). MALV, on the other hand, can be found as parasites of ciliates,
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23 441 dinoflagellates, radiolarians and fish eggs (Massana 2011) and are known to form
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25 442 planktonic lifecycles (Guillou *et al.* 2008). Unicellular Archaeplastida (Chlorophyta)
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27 443 and Hacrobia (Haptophyta) were also largely missing from benthos samples but
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29 444 occurred in plankton samples. These groups mainly consist of autotrophic organisms
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31 445 performing carbon fixation (Vaulot *et al.* 2008; Not *et al.* 2012) and are major
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33 446 contributors to the pico- and nanoplankton diversity and biomass across the world's
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35 447 oceans (Bittner *et al.* 2013; Egge *et al.* 2015).
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449 *High degree of genetic novelty among benthic protist assemblages*
450 450 The genetic divergence of benthic OTUs from reference sequences described
451 451 the marine coastal benthic realm as a habitat having a vast majority of protist
452 452 diversity still uncharted. This applied to both BLAST analyses, against the curated
453 453 PR² reference database and the environmental genetic signatures of the SRA
454 454 database. Thereby, the mean sequence similarity to taxonomic references (93.4%,
455 455 Fig. 5A) and to environmental references (95.4%, Fig. 5B) was higher than reported
456 456 from abyssal sediment communities of protists (87% similarity to taxonomic

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2 457 references, 89% to environmental references) (Scheckenbach *et al.* 2010). Recent
3 458 advances in sequencing technologies enable the detection and assemblage of a
4 459 broader diversity of genetic signatures in environmental microbial surveys than ever,
5 460 but there still remains a clear discrepancy between what can be detected and what
6 461 can be taxonomically assigned. This discrepancy is especially pronounced among
7 462 benthic organisms, which display a much higher novelty in genetic diversity surveys
8 463 than planktonic organisms (Figs. 5 and 6). A promising approach towards exploring
9 464 this novel diversity is the combination of multiple SRA datasets in network analyses
10 465 to screen for groups of sequences that do not have closely described relatives
11 466 (Forster *et al.* 2015). Such highly divergent groups detected in independent
12 467 environmental samples have a high potential to represent genuine undescribed
13 468 organisms. The design of novel probes and primer-sets specifically for these groups
14 469 will in turn enable the targeted recovery and identification of the respective organisms
15 470 from life samples by molecular methods (Gimmler and Stoeck 2015). As our picture
16 471 of protist diversity heavily depends on the coverage of available public databases
17 472 (Pawlowski *et al.* 2012), increasing the efforts to isolate, cultivate and describe
18 473 benthic protist species will also help to link genetic signatures obtained in
19 474 environmental sequencing studies to a real biological entity with a phylogenetic
20 475 context. Although the isolation of individual specimen remains a challenging task,
21 476 methods exist to successfully address organisms in benthos samples. For example,
22 477 even small flagellates can be isolated by quantitative centrifugation (Starink *et al.*
23 478 1994). More recently, a serial dilution method enabled the recovery of diatom spores
24 479 from coastal sediment samples (Montresor *et al.* 2013). First studies on highly
25 480 divergent BioMarkS sediment OTUs already led to the discovery of novel cercozoan
26 481 vampirellids (Berney *et al.* 2013) and opisthokont Fonticulida (del Campo *et al.*
27 482 2015), two groups that were mostly known from soil or freshwater samples.

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2 483 Regarding the divergence of benthic OTUs from publicly available reference
3 sequences, we suppose that we are just scratching the surface of protist diversity in
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5 coastal sediments. Thus, we understand the BioMarKs data as a starting point that
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7 may guide the discovery of more novel benthic protist diversity by further taxon-
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9 specific screening.
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18 490 **Conclusions**
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20 491 Over the last 60 years, our perception of marine benthic environments has
21 changed from being biologically inert deserts towards being highly heterogeneous
22 habitats teeming with a multitude of microbial organisms. Even though new
23 technologies allow for addressing this vast diversity, the results of the BioMarKs
24 project strongly imply that the most part of benthic protist diversity remains a black
25 box. While sampling the deep-seafloor surely imposes many challenges and
26 restrictions that have lagged the exploration of benthic diversity, our data show that it
27 is not mandatory to focus on such remote environments when looking for highly
28 diverse protist communities. From a quantitative (*i.e.* OTU richness) and qualitative
29 (*i.e.* degree of genetic divergence) point of view, coastal sediments inhabit
30 intriguingly rich protist assemblages on local and regional scales. Increasing the
31 efforts to explore these assemblages will be beneficial to learn more about the
32 dispersal patterns of benthic protists, their roles in ecosystem functioning and to
33 complement current species inventories by identifying many still unknown organisms.
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1
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33 521 **Conflict of interest**
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36 522 The authors declare that they have no conflict of interest.
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691 **Table 1 Overview of BioMarKs samplings along the European coastline.**

Sampling Event	Latitude/Longitude	Oceanic region	Quality sequences per sampled habitat		
			Surface water	DCM	Sediment
Blanes	41°40'N/2°48'E	Mediterranean Sea	16 568	n.d.	3576
Gijon	43°40'N/5°35'W	Atlantic Ocean	20 966	n.d.	n.d.
Naples 2009	40°48'N/14°15'E	Mediterranean Sea	62 250	52 114	15 739
Naples 2010	40°48'N/14°15'E	Mediterranean Sea	22 132	31 268	3159
Oslo 2009	59°16'N/10°43'E	Skagerrak	33 133	36 988	12 835
Oslo 2010	59°16'N/10°43'E	Skagerrak	19 316	24 819	3071
Roscoff	48°46'E/3°57'W	English Channel	9298	n.d.	1720
Varna	43°10'N/28°50'E	Black Sea	22 939	39 003	n.d.

692 DCM, deep-chlorophyll maximum; n.d., no data available.

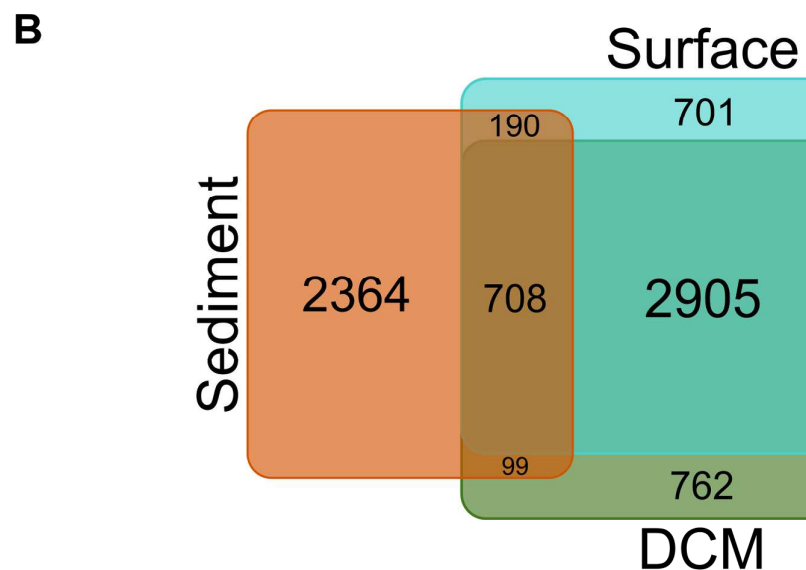
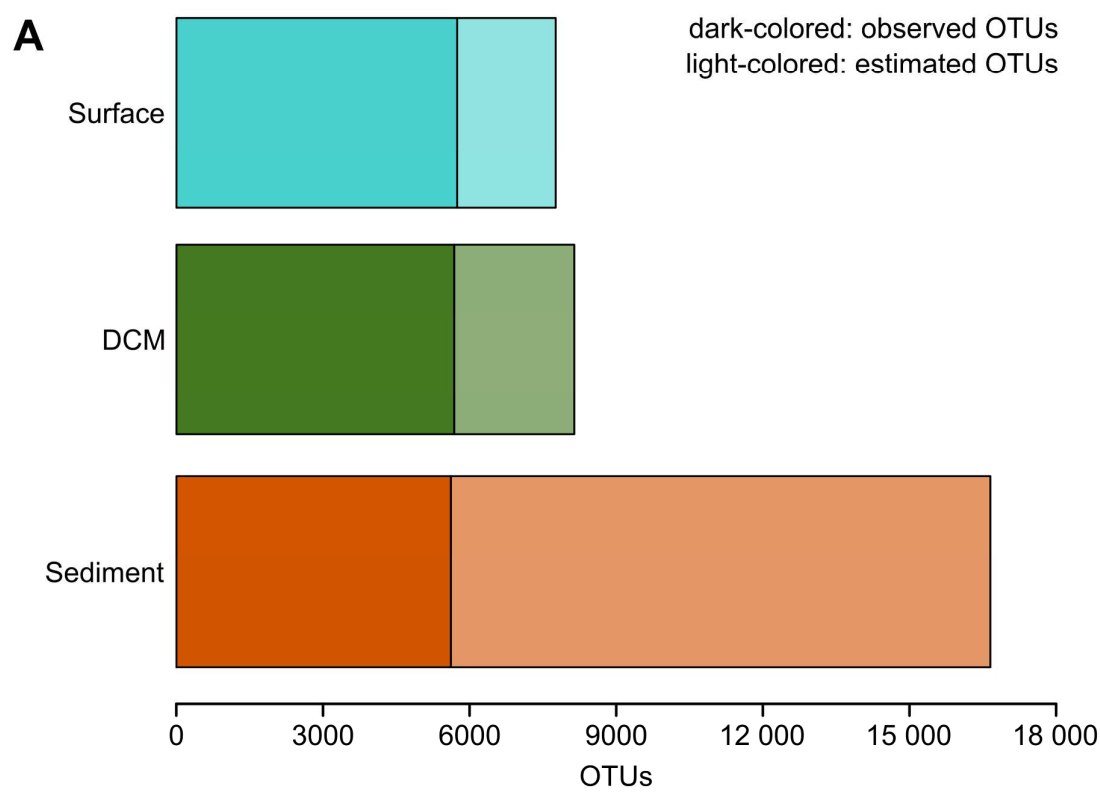
693 Total number of surface water quality sequences: 206 602.

694 Total number of DCM quality sequences: 184 192.

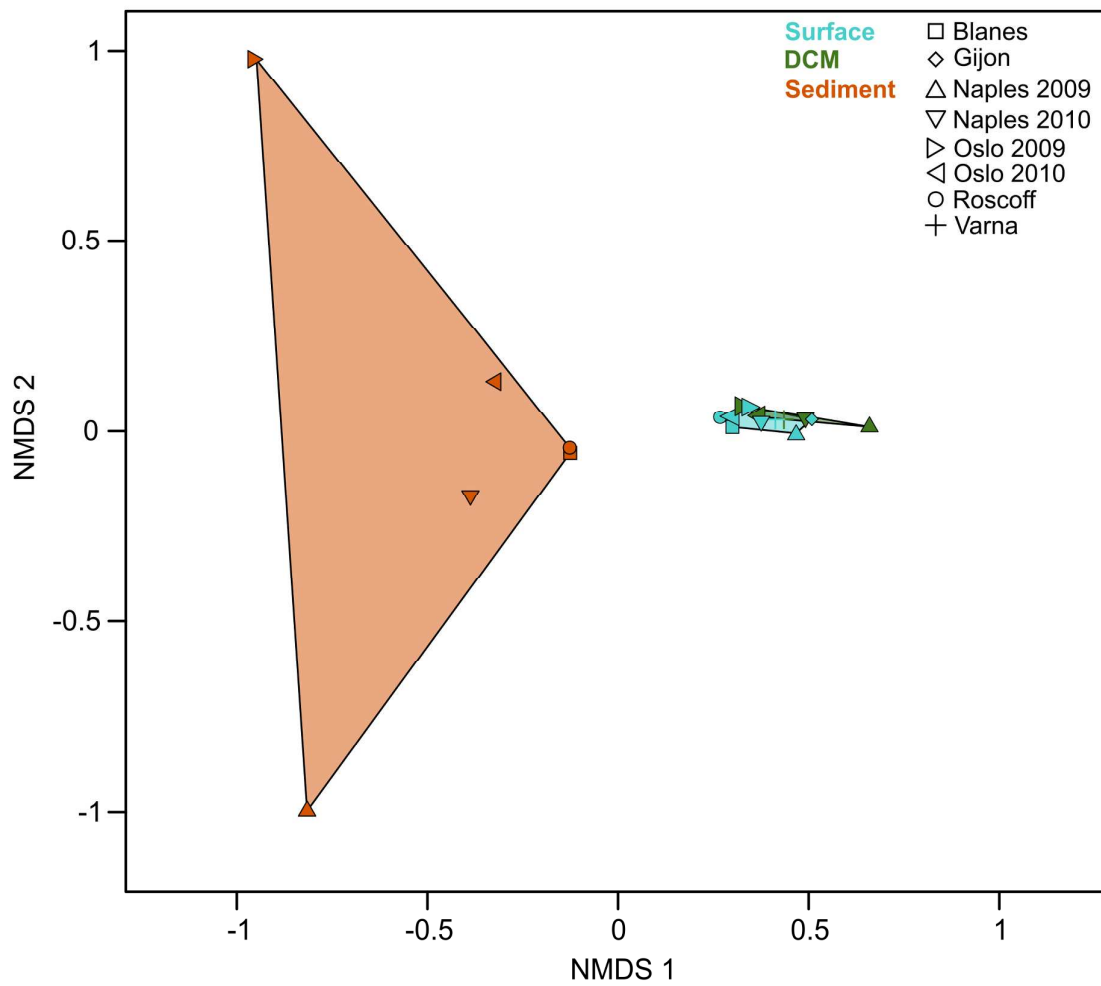
695 Total number of sediment quality sequences: 40 100.

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2 697 **Fig. 1 OTU richness and shared OTUs of planktonic and benthic protist**
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4 698 **communities.** (A) For each habitat the bars show the number of observed (left, dark
5 699 colored part) and estimated OTUs (right, lighter colored part). Estimations are based
6
7 700 on the incidence-based coverage estimator (ICE). (B) The Venn-Diagram highlights
8
9 701 how many non-singleton OTUs were shared between all habitats, shared between
10 702 any two of the sampled habitats or exclusively detected in a single habitat. Numbers
11 703 indicate the amount of OTUs which fall into the respective category. The area of each
12 704 category was proportionally approximated to the number of OTUs from the respective
13 705 habitat.
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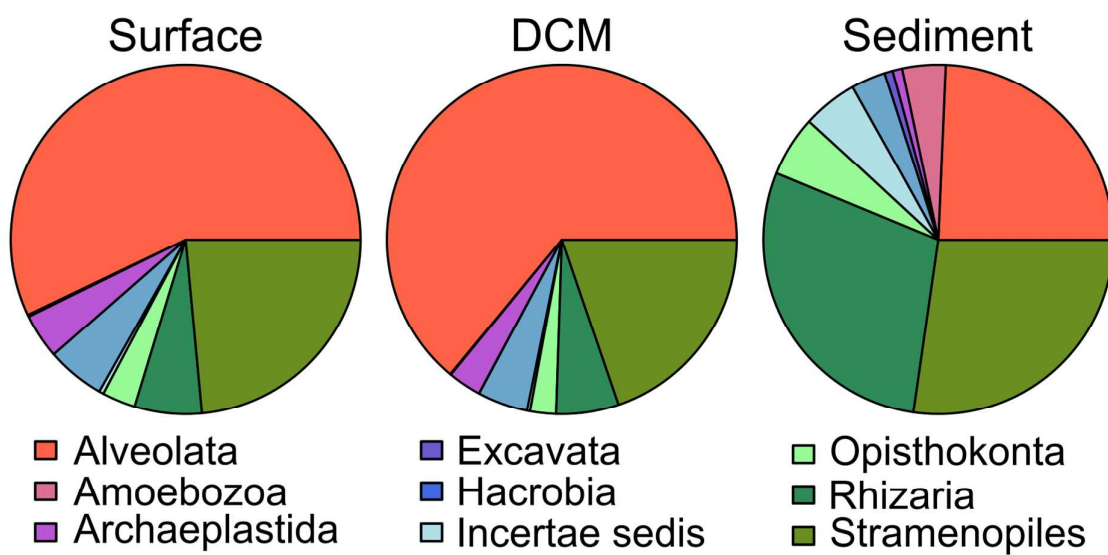
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2 709 **Fig. 2 Beta diversity analysis of single sampling events.** The NMDS is based on
3 710 binary-Jaccard distances between protist communities of single samples (NMDS
4 711 stress level = 0.2). Colors of the data points were chosen with respect to their habitat
5 712 affiliation. Data points of the same affiliation were linked and the area between the
6 713 linked points was colored with respect to their habitat affiliation.
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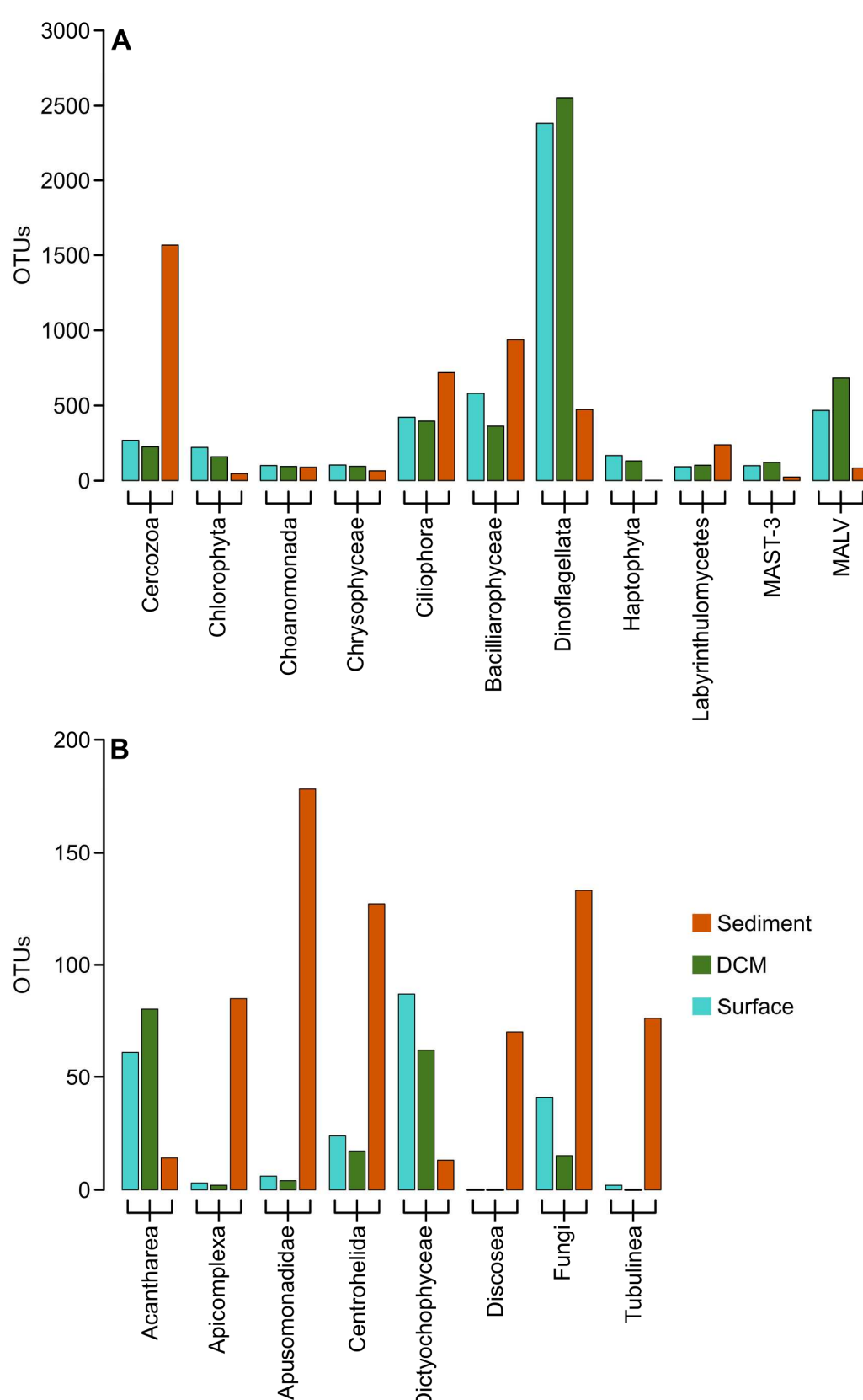
Fig. 3 Taxonomic community compositions as revealed by the PR² database.

The pie charts reflect the distribution of OTUs within major protist groups, by pooling the data of all samples from each habitat. The surface dataset included 5747 distinct OTUs, the DCM dataset 5685 distinct OTUs and the sediment dataset 5616 distinct OTUs.



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2 725 **Fig. 4 Number of OTUs within taxonomic groups in planktonic and benthic**
3
4 726 **samples.** Only groups contributing to at least 1% of the OTU number in either
5 727 surface, DCM or sediment communities were considered. Note the different scale
6 728 among the two panels, to display taxonomic groups with more than 200 OTUs in total
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8 729 (A) or with less than 200 OTUs in total (B).
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3 **Fig. 5 Rank abundance curves displaying the sequence similarity distribution**
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5 **of OTUs against reference databases.** Results are shown for two different BLAST
6
7 queries, against the PR² database (A) and against a custom-built subset of NCBI's
8
9 Short Read Archive (B). The curves display the number of OTUs with a given
10
11 similarity with the closest match from the respective reference database. Sequence
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13 similarities are binned in 0.5 % intervals.
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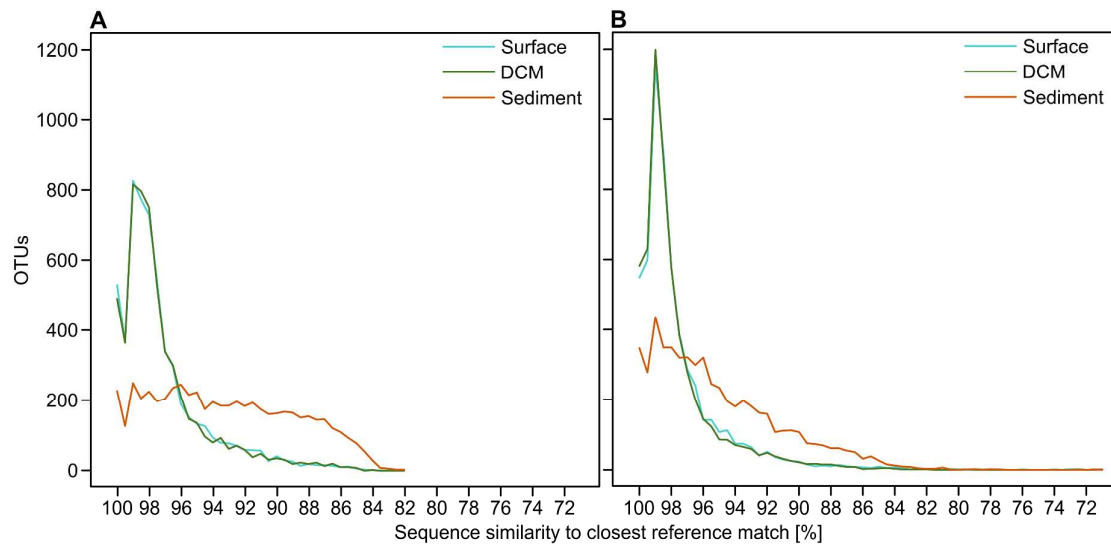
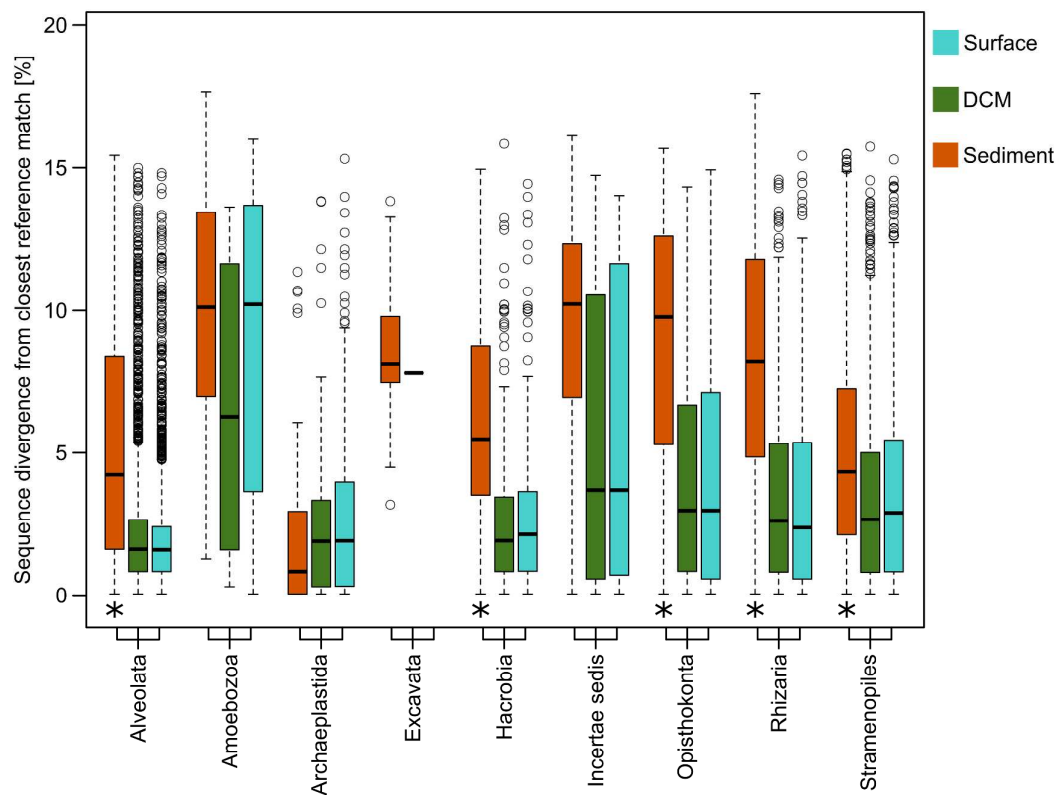


Fig. 6 Sequence divergence of OTUs to their closest taxonomic hit. Each boxplot comprises the sequence divergence values of all OTUs within a given taxonomical group from the same habitat to their closest reference in the PR² database. The taxonomic assignment is shown at the supergroup level. Missing boxplots indicate that the taxonomical group was not detected in the respective habitat. For each taxonomic group, differences between the distribution of sequence dissimilarity values of sediment OTUs to planktonic OTUs was tested for significance ($p<0.001$, indicated by stars). The distribution of sequence dissimilarity values was never significantly different between DCM and surface OTUs. Black bars represent the median of each boxplot, the part of the box above the median represents the upper quartile; accordingly, the part of the box below the median represents the lower quartile. Circles show the outliers of each group of data.



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2 **Table 1 Overview of BioMarKs samplings along the European coastline.**
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Sampling Event	Latitude/Longitude	Oceanic region	Quality sequences per sampled habitat		
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26 Total number of surface water quality sequences: 206 602.
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28 Total number of DCM quality sequences: 184 192.
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30 Total number of sediment quality sequences: 40 100.
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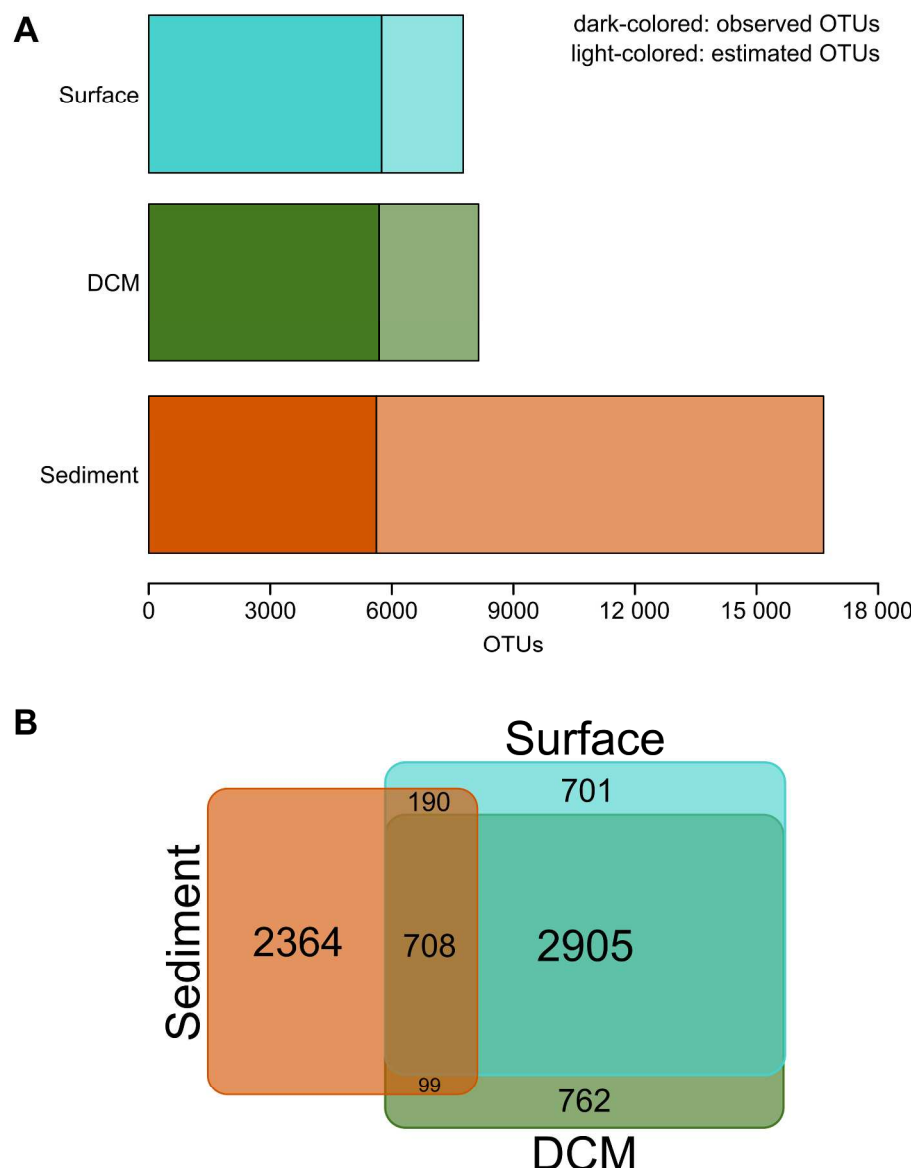


Fig. 1 OTU richness and shared OTUs of planktonic and benthic protist communities. (A) For each habitat the bars show the number of observed (left, dark colored part) and estimated OTUs (right, lighter colored part). Estimations are based on the incidence-based coverage estimator (ICE). (B) The Venn-Diagram highlights how many non-singleton OTUs were shared between all habitats, shared between any two of the sampled habitats or exclusively detected in a single habitat. Numbers indicate the amount of OTUs which fall into the respective category. The area of each category was proportional approximated to the number of OTUs from the respective habitat.

228x290mm (300 x 300 DPI)

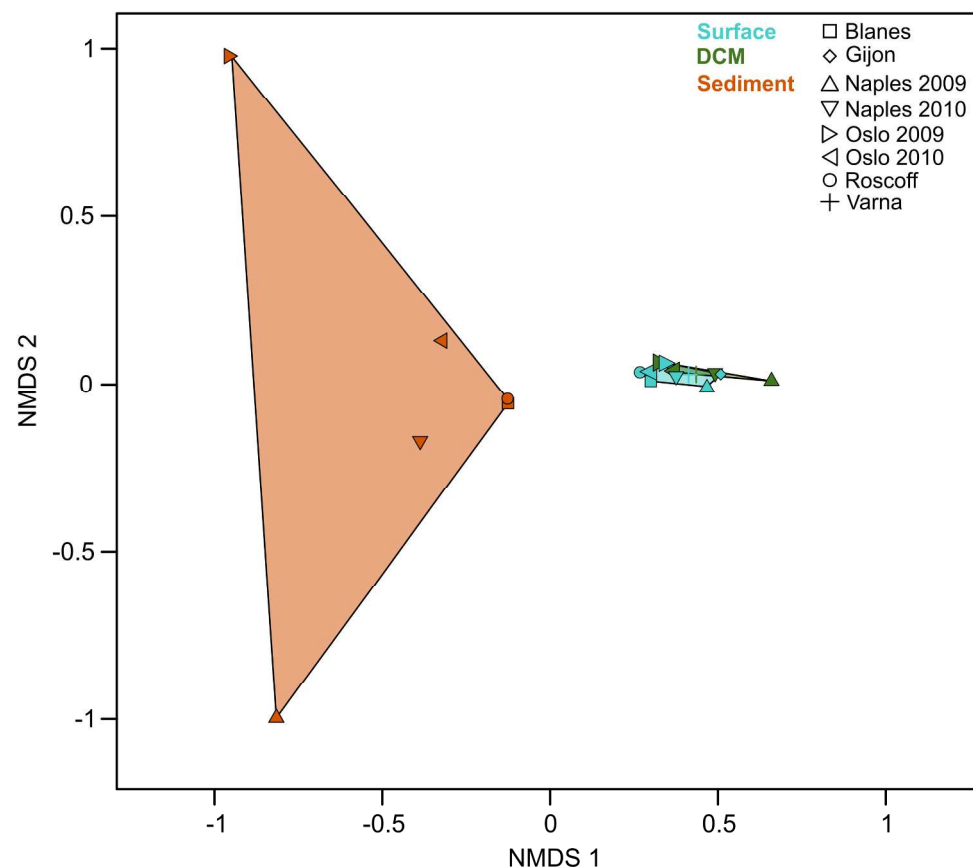


Fig. 2 Beta diversity analysis of single sampling events. The NMDS is based on binary-Jaccard distances between protist communities of single samples (NMDS stress level = 0.2). Colors of the data points were chosen with respect to their habitat affiliation. Data points of the same affiliation were linked and the area between the linked points was colored with respect to their habitat affiliation.

228x200mm (300 x 300 DPI)



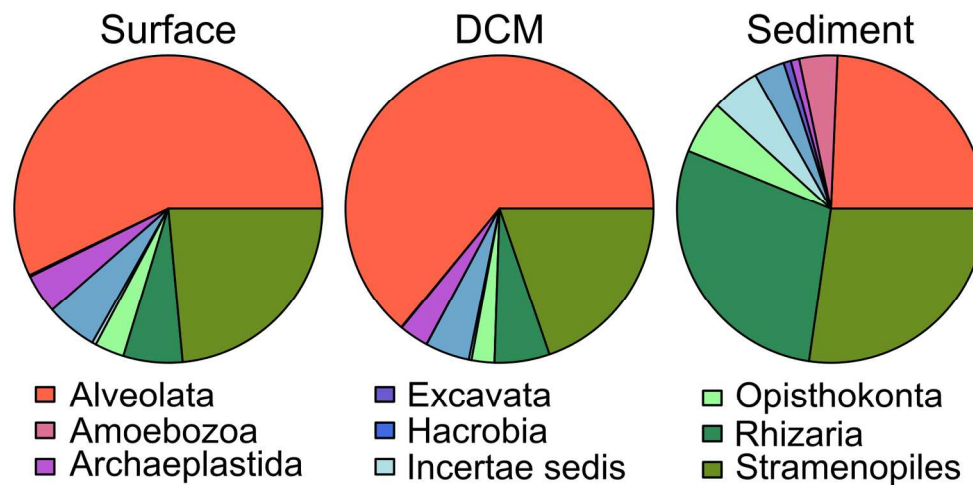


Fig. 3 Taxonomic community compositions as revealed by the PR2 database. The pie charts reflect the distribution of OTUs within major protist groups, by pooling the data of all samples from each habitat. The surface dataset included 5747 distinct OTUs, the DCM dataset 5685 distinct OTUs and the sediment dataset 5616 distinct OTUs.

160x79mm (300 x 300 DPI)

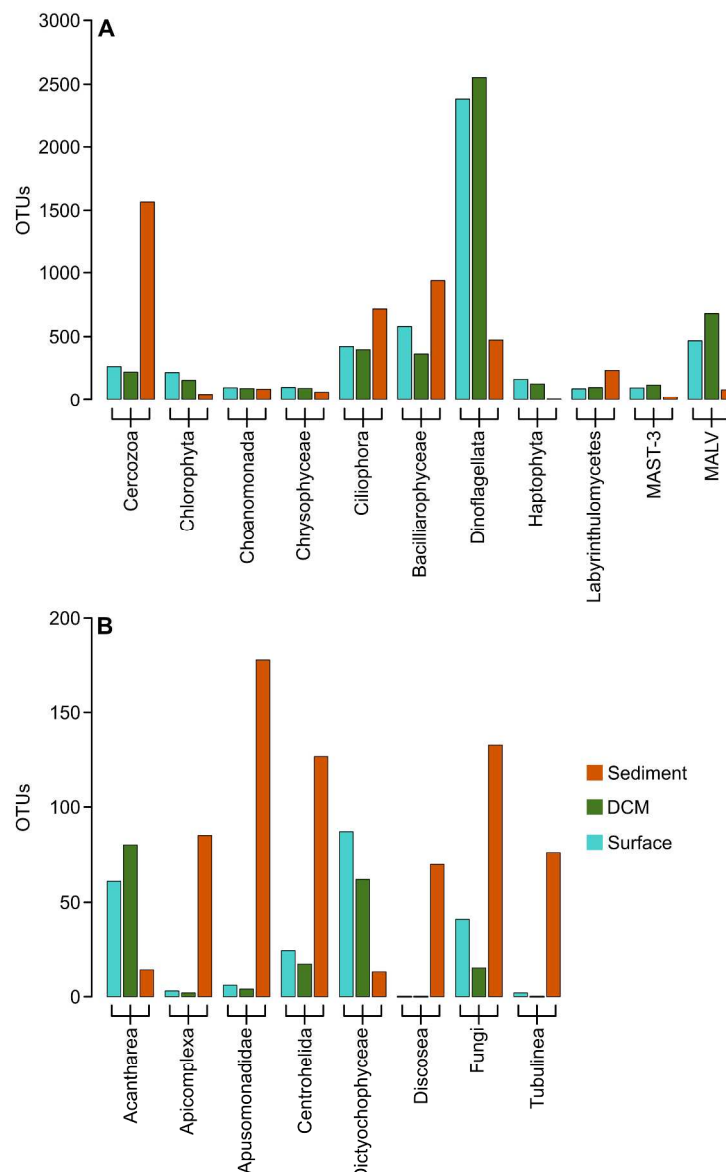


Fig. 4 Number of OTUs within taxonomic groups in planktonic and benthic samples. Only groups contributing to at least 1% of the OTU number in either surface, DCM or sediment communities were considered. Note the different scale among the two panels, to display taxonomic groups with more than 200 OTUs in total (A) or with less than 200 OTUs in total (B).

228x360mm (300 x 300 DPI)

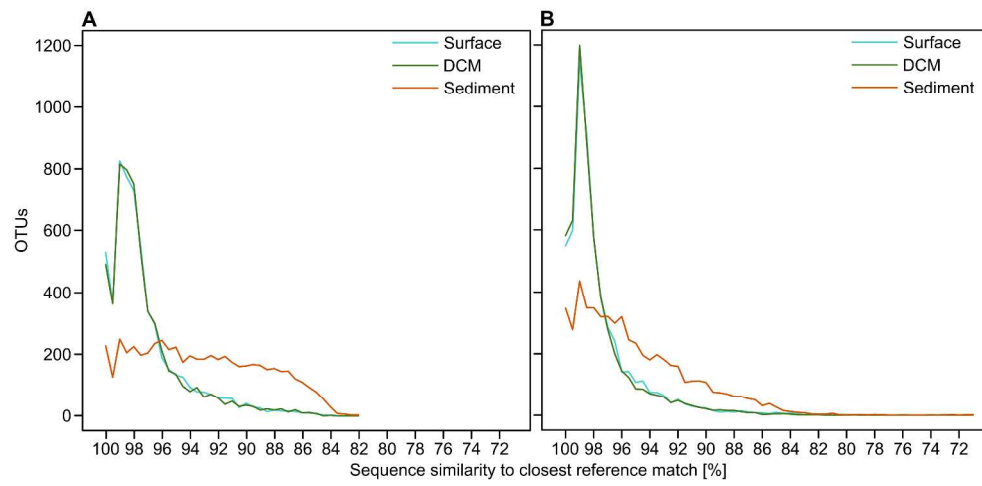


Fig. 5 Rank abundance curves displaying the sequence similarity distribution of OTUs against reference databases. Results are shown for two different BLAST queries, against the PR2 database (A) and against a custom-built subset of NCBI's Short Read Archive (B). The curves display the number of OTUs with a given similarity with the closest match from the respective reference database. Sequence similarities are binned in 0.5 % intervals.

336x169mm (300 x 300 DPI)

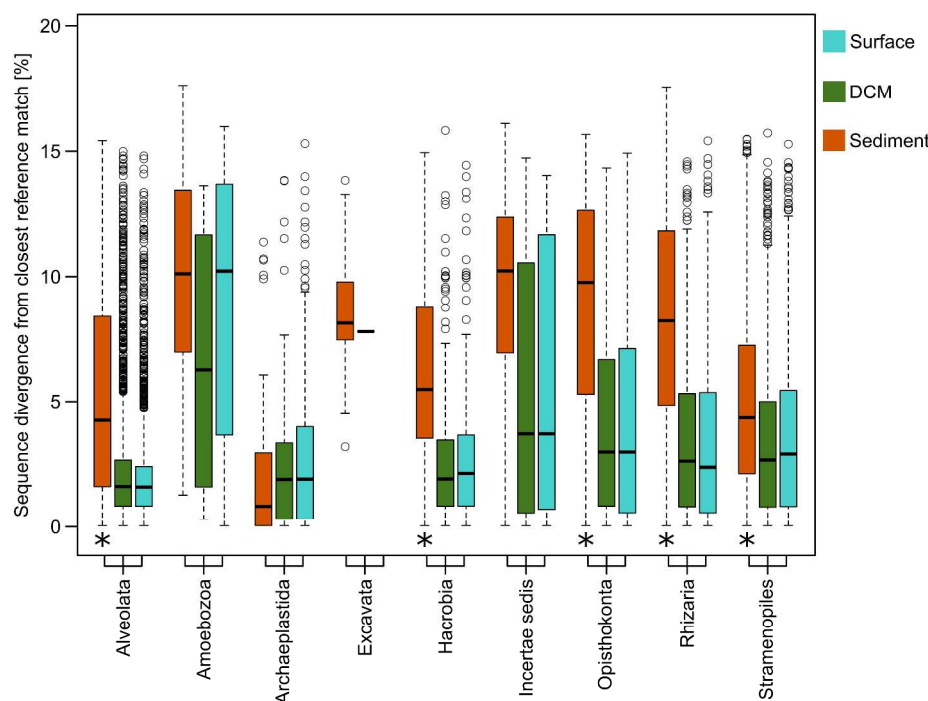


Fig. 6 Sequence divergence of OTUs to their closest taxonomic hit. Each boxplot comprises the sequence divergence values of all OTUs within a given taxonomical group from the same habitat to their closest reference in the PR2 database. The taxonomic assignment is shown at the supergroup level. Missing boxplots indicate that the taxonomical group was not detected in the respective habitat. For each taxonomic group, differences between the distribution of sequence dissimilarity values of sediment OTUs to planktonic OTUs was tested for significance ($p < 0.001$, indicated by stars). The distribution of sequence dissimilarity values was never significantly different between DCM and surface OTUs. Black bars represent the median of each boxplot, the part of the box above the median represents the upper quartile; accordingly, the part of the box below the median represents the lower quartile. Circles show the outliers of each group of data.

336x250mm (300 x 300 DPI)

1 **Benthic protists: the undercharted majority**

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4 Dominik Forster, Micah Dunthorn, Frédéric Mahé, John R. Dolan, Stéphane Audic,
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6 David Bass, Lucie Bittner, Christophe Boutte, Richard Christen, Jean-Michel
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8 Claverie, Johan Decelle, Bente Edvardsen, Elianne Egge, Wenche Eikrem,
9
10 Angélique Gobet, Wiebe H.C.F. Kooistra, Ramiro Logares, Ramon Massana, Marina
11
12 Montresor, Fabrice Not, Hiroyuki Ogata, Jan Pawłowski, Massimo C. Pernice, Sarah
13
14 Romac, Kamran Shalchian-Tabrizi, Nathalie Simon, Thomas Richards, Sébastien
15
16 Santini, Diana Sarno, Raffaele Siano, Daniel Vaulot, Patrick Wincker, Adriana
17
18 Zingone, Colomban de Vargas, Thorsten Stoeck

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2 **Supplemental Table S1 Additional information of the SRA database.** The table
3 includes all 167 distinct datasets used to build our custom subset of the SRA. We
4 screened the complete archive for protist datasets of the eukaryotic 18S gene. After
5 downloading the respective data, we removed all sequences shorter than 100 bp.
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7 The final customized SRA reference database consisted of 11 708 385 sequences.
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9 Information in the table was downloaded directly from NCBI's SRA platform along
10 with the respective sequences.
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20 **Supplemental Fig. 2 Observed and estimated OTU richness calculated by**
21 **abundance-based analyses.** Additionally to incidence-based ICE analyses (Fig.1)
22 we estimated OTU richness with the abundance-based CatchAll tool. In direct
23 comparison ICE and CatchAll results show the same trends, though the estimated
24 richness of the abundance-based analyses always exceeded the one of the
25 incidence-based analyses. For each habitat the plot shows the amount of observed
26 and estimated OTUs. The left, dark colored part of each bar shows the actually
27 observed number of OTUs, the right, light colored part of each bar shows how many
28 more OTUs were estimated to be in each dataset by extrapolation of the data.
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30 **Supplemental Fig. S3 Normalized rarefaction curves of sampled habitats.**
31 Sequence data of all single sampling events were pooled with regard to the
32 respective habitat. Each habitat rarefaction curve was normalized to 40 100
33 sequences, which equals the smallest total amount of sequences found in any of the
34 three habitats. Depicted is the amount of SSU V4 rDNA sequences in relation to the
35 amount of resulting OTUs into which the sequences were clustered.
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2 **37 Supplemental Fig. S4 Community composition among phototrophic protist**
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5 **38 taxon groups.** Represented are all phototrophic groups detected in BioMarkS. The
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7 **39 upper bar shows the phototrophic OTUs exclusively found in the plankton, the middle**
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9 **40 bar the phototrophic OTUs present in both plankton and benthos and the lower bar**
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11 **41 the phototrophic OTUs exclusively found in the benthos. Dinoflagellates were not**
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13 **42 considered in the analyses.**

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18 **43 Supplemental Fig. S5 Occurrence of each phototrophic protist taxon group in**
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20 **44 the sampled habitats.** For each phototrophic taxon group, the bars indicate the
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22 fraction of OTUs which was found exclusively in the plankton, in both plankton and
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24 benthos and exclusively in the benthos. Dinoflagellates were not considered in these
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26 analyses.

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32 **50 Supplemental Fig. S6 Community composition across different taxonomic**
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34 **51 levels.** The pie charts reflect the proportion of taxonomic groups of the total microbial
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36 eukaryote community in each habitat. The inner ring is equivalent to Fig. 4 and
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38 represents the taxonomical assignment into major protist groups (corresponding to
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40 the left part of the taxonomic legend). The outer ring represents the taxonomical
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42 assignment at the phylum level (corresponding to the right part of the taxonomic
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44 legend). Only phyla which contributed with at least 5% to the total community where
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46 considered with a specific color in the outer ring. Exact proportions are given in
47
48 square brackets behind the taxonomical group names.

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2 Experiment Accession
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For Peer Review

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2 Experiment Title
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13 Gene expression during the initiation phase of growth in the dinoflagellate *Alexandrium catenella* (Dinophyceae)
14 community profiling of Mirs Bay seawater by 18S rRNA V4 amplicon 454 sequencing

15 community profiling of Tolo Harbor seawater by 18S rRNA V4 amplicon 454 sequencing

16 Pooled bacterial amplicons from 10 Arctic marine surface water samples from 2003-2009 plus 2010

17 Pooled archaeal amplicons from 10 Arctic marine surface water samples from 2003-2009 plus 2010

18 Pooled eukarya amplicons from 10 Arctic marine surface water samples from 2003-2009 plus 2010

19 Marine microbial eukaryote community analysis in Korea

20 Marine microbial eukaryote community analysis in Korea

21 GS_FLX sequencing for environmental diversity study using the SSU rRNA as taxonomic marker

22 GS_FLX_Titanium sequencing for environmental diversity study using the SSU rRNA as taxonomic marker

23 RN_RepliG-Sequencing

24 Pooled sample of 12 Arctic (Amundsen Gulf and Franklin Bay) sea ice samples from 2008

25 microbial diversity of ornamental fish aquarium water sample D1

26 microbial diversity of ornamental fish aquarium water sample E1

27 microbial diversity of ornamental fish aquarium water sample D2

28 microbial diversity of ornamental fish aquarium water sample A1

29 microbial diversity of ornamental fish aquarium water sample A2

30 microbial diversity of ornamental fish aquarium water sample E2

31 microbial diversity of ornamental fish aquarium water sample E1.V4

32 microbial diversity of ornamental fish aquarium water sample D1.V4

33 microbial diversity of ornamental fish aquarium water sample A1.V4

34 microbial diversity of ornamental fish aquarium water sample A2.V4

35 microbial diversity of ornamental fish aquarium water sample D2.V4

36 microbial diversity of ornamental fish aquarium water sample E2.V4

37 microbial diversity of ornamental fish aquarium water sample B2.V4

38 Pooled samples of protist communities at 8 depth from Lake A in May 2008

39 Pooled samples of protist communities at 4 depths from Lake A in August 2008

40 Pooled samples of protist communities at 5 depths from Lake A in July 2009

41 Environmental Samples from ANT-XXVI/3-NS

42 Partial ribosomal gene sequences Ciliates in DHABs in the Mediterranean Sea

43 V4 18S rRNA amplicons from cDNA prepared from different microbialite sediment samples from Highborne Cay, B

44 V4 amplicons (18S rRNA) of eukaryotes obtained from cDNA prepared from various microbialite samples from Sha

45 ANT XXVI/3 - WE

46 a test of 16-18s universal primers using sponge samples

47 Comparative Microbial Analysis to Understand Taxonomic Fingerprints of Healthy Coral Holobiont Communities at

48 Protistan communities of the upper Arctic Ocean

49 Comparative Microbial Analysis to Understand Taxonomic Fingerprints of Healthy Coral Holobiont Communities at

1 Comparative Microbial Analysis to Understand Taxonomic Fingerprints of Healthy Coral Holobiont Communities at
2 Comparative Microbial Analysis to Understand Taxonomic Fingerprints of Healthy Coral Holobiont Communities at
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18 Comparative Microbial Analysis to Understand Taxonomic Fingerprints of Healthy Coral Holobiont Communities at
19 Comparative Microbial Analysis to Understand Taxonomic Fingerprints of Healthy Coral Holobiont Communities at
20 Exp1.Ward Hunt Lake Dilution Experiment Eukaryotes RNA
21 Characterisation of uncultured marine eukaryotes in the Canada Basin
22 Chukchi Sea < 3 um Stn BS1
23 Chukchi Sea < 3 um Stn AN1
24 Chukchi Sea < 3 um Stn KS6
25 Chukchi Sea < 3 um Stn EC5
26 Chukchi Sea < 3 um Stn HLY
27 Chukchi Sea < 3 um Stn CC18
28 Chukchi Sea < 3 um Stn CN3
29 Chukchi Sea < 3 um Stn CN14
30 Chukchi Sea < 3 um Stn BCH7
31 Chukchi Sea < 3 um Stn IE
32 Chukchi Sea < 3 um Stn HSN5
33 Chukchi Sea > 3 um Stn BS1
34 Chukchi Sea > 3 um Stn AN1
35 Chukchi Sea > 3 um Stn KS6
36 Chukchi Sea > 3 um Stn CHA1
37 Chukchi Sea > 3 um Stn EC5
38 Chukchi Sea > 3 um Stn HLY
39 Chukchi Sea > 3 um Stn CC18
40 Chukchi Sea > 3 um Stn CN3
41 Chukchi Sea > 3 um Stn CN14
42 Chukchi Sea > 3 um Stn BCH7
43 Chukchi Sea > 3 um Stn IE
44 Chukchi Sea > 3 um Stn HSN5

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51 Sequences from three sediments
52

53 Design and validation of four primers for next-generation sequencing to target the 18S rRNA gene of gastrointestinal
54 Design and validation of four primers for next-generation sequencing to target the 18S rRNA gene of gastrointestinal
55 Design and validation of four primers for next-generation sequencing to target the 18S rRNA gene of gastrointestinal
56 454 pyrosequencing of plankton community collected from Hamilton harbour
57 454 pyrosequencing of plankton community collected from Nanaimo harbour

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2 Investigating marine microbial plankton in meso-eutrophic ecosystem (Eastern English Channel)
3 JB121SF_DNA
4 JB121SF_RNA GoMA July 2010
5 JB121SCM_DNA GoMA July 2010
6 JB121SCM_RNA GoMA July 2010
7 JB700SF_DNA GoMA July 2010
8 JB700SF_RNA GoMA July 2010
9 JB700SCM_DNA GoMA July 2010
10 JB700SCM_RNA GoMA July 2010
11 JB601SF_DNA GoMA July 2010
12 JB601SF_RNA GoMA July 2010
13 JB601SCM_DNA GoMA July 2010
14 JB601SCM_RNA GoMA July 2010
15 NEC211SF_DNA
16 NEC211SF_RNA GoMA July 2010
17 NEC211SCM_DNA GoMA July 2010
18 NEC211SCM_RNA GoMA July 2010
19 NEF43SF_DNA GoMA July 2010
20 NEF43SF_RNA GoMA July 2010
21 NEF43SCM_DNA GoMA July 2010
22 NEF43SCM_RNA GoMA July 2010
23 NEF512SF_DNA GoMA July 2010
24 NEF512SF_RNA GoMA July 2010
25 NEF512SCM_DNA GoMA July 2010
26 NEF512SCM_RNA GoMA July 2010
27 Eukaryota Arctic Ocean Survey
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20 Eukaryota Arctic Ocean Survey
21 Marine microbial community structure assessed from combined metagenomic analysis and ribosomal tag pyroseq
22 Protists in Havre-aux-Maisons Lagoon
23 Protists in Havre-aux-Maisons Lagoon
24 Protists in Havre-aux-Maisons Lagoon
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30 Protists in Havre-aux-Maisons Lagoon
31 Protists in Havre-aux-Maisons Lagoon
32 Protists in Havre-aux-Maisons Lagoon
33 Protists in Havre-aux-Maisons Lagoon
34 Eukaryotic Plankton Assemblages Inhabiting Saline Lakes in the Qaidam Basin
35 Marine picoplankton
36 Deep Hypersaline Anoxic Basin sediment microbial eukaryotes
37 Three years diversity in the Eastern English Channel
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2	Organism Name	Instrument
3	marine metagenome	454 GS FLX
4	marine metagenome	454 GS FLX
5	marine metagenome	454 GS FLX
6	marine metagenome	454 GS FLX
7	marine metagenome	454 GS FLX
8	aquatic metagenome	454 GS FLX Titanium
9	aquatic metagenome	454 GS FLX Titanium
10	aquatic metagenome	454 GS FLX Titanium
11	aquatic metagenome	454 GS FLX Titanium
12	aquatic metagenome	454 GS FLX Titanium
13	Alexandrium catenella	454 GS FLX
14	marine metagenome	454 GS FLX
15	marine metagenome	454 GS FLX
16	marine metagenome	454 GS FLX Titanium
17	marine metagenome	454 GS FLX Titanium
18	marine metagenome	454 GS FLX Titanium
19	marine metagenome	454 GS FLX Titanium
20	uncultured marine eukaryote	454 GS FLX Titanium
21	uncultured marine eukaryote	454 GS FLX Titanium
22	metagenomes	454 GS FLX
23	metagenomes	454 GS FLX Titanium
24	Rotylenchulus reniformis	454 GS FLX Titanium
25	marine metagenome	454 GS FLX Titanium
26	metagenomes	454 GS FLX
27	metagenomes	454 GS FLX
28	metagenomes	454 GS FLX
29	metagenomes	454 GS FLX
30	metagenomes	454 GS FLX
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39	metagenomes	454 GS FLX
40	metagenomes	454 GS FLX
41	metagenomes	454 GS FLX
42	metagenomes	454 GS FLX
43	metagenomes	454 GS FLX
44	freshwater metagenome	454 GS FLX Titanium
45	freshwater metagenome	454 GS FLX Titanium
46	freshwater metagenome	454 GS FLX Titanium
47	uncultured marine eukaryote	454 GS FLX Titanium
48	marine metagenome	454 GS FLX Titanium
49	eukaryote XCL-2011	454 GS FLX Titanium
50	eukaryote XCL-2011	454 GS FLX Titanium
51	marine metagenome	454 GS FLX
52	Suberites ficus	454 GS FLX Titanium
53	Isopora palifera	454 GS FLX Titanium
54	marine metagenome	454 GS FLX Titanium
55	Seriatopora hystrix	454 GS FLX Titanium
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1	Montipora digitata	454 GS FLX Titanium
2	Porites cylindrica	454 GS FLX Titanium
3	Isopora palifera	454 GS FLX Titanium
4	Isopora palifera	454 GS FLX Titanium
5	Isopora palifera	454 GS FLX Titanium
6	Montipora digitata	454 GS FLX Titanium
7	Montipora digitata	454 GS FLX Titanium
8	Montipora digitata	454 GS FLX Titanium
9	Seriatopora hystrix	454 GS FLX Titanium
10	Seriatopora hystrix	454 GS FLX Titanium
11	Seriatopora hystrix	454 GS FLX Titanium
12	Porites cylindrica	454 GS FLX Titanium
13	Porites cylindrica	454 GS FLX Titanium
14	Porites cylindrica	454 GS FLX Titanium
15	aquatic metagenome	454 GS FLX Titanium
16	marine metagenome	454 GS FLX Titanium
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18	marine metagenome	454 GS FLX Titanium
19	marine metagenome	454 GS FLX Titanium
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47	marine metagenome	454 GS FLX Titanium
48	marine metagenome	454 GS FLX Titanium
49	marine metagenome	454 GS FLX Titanium
50	gut metagenome	454 GS FLX
51	marine sediment metagenome	454 GS FLX+
52	gut metagenome	454 GS FLX Titanium
53	gut metagenome	454 GS FLX Titanium
54	gut metagenome	454 GS FLX Titanium
55	freshwater metagenome	454 GS FLX Titanium
56	marine metagenome	454 GS FLX Titanium

1 marine metagenome
2 uncultured eukaryote
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21 marine metagenome 454 GS FLX+
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36 marine sediment metagenome 454 GS FLX Titanium
37 marine metagenome 454 GS FLX Titanium
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2	Submitter	Study Accession
3	ESE	ERP001773
4	ESE	ERP001773
5	ESE	ERP001773
6	ESE	ERP001773
7	ESE	ERP001773
8	MPIMM	ERP002513
9	MPIMM	ERP002513
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12	CNRS/UNIVERSITE MONTPELLIER 2	SRP000647
13	The Chinese University of Hong Kong	SRP000936
14	The Chinese University of Hong Kong	SRP000936
15	Universite Laval	SRP005272
16	Universite Laval	SRP005272
17	Universite Laval	SRP005272
18	Inha univ.	SRP006126
19	Inha univ.	SRP006126
20	University of Kaiserslautern	SRP003169
21	University of Kaiserslautern	SRP003169
22	Alabama A&M University	SRP008476
23	Universite Laval	SRP013740
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44	Université Laval	SRP014823
45	Université Laval	SRP014823
46	Université Laval	SRP014823
47	Alfred Wegener Institute for Polar and Marine Rese	SRP016049
48	University of Kaiserslautern	SRP017187
49	Woods Hole Oceanographic Institution	SRP017332
50	Woods Hole Oceanographic Institution	SRP017334
51	Alfred Wegener Institute for Polar and Marine Rese	SRP016049
52	hkust-kaust	SRP021898
53	University of Sydney	SRP022083
54	Laval University	SRP022176
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19	Universit* Laval	SRP027540
20	Universite Laval	SRP029291
21	Université Laval	SRP029300
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50	AWI	SRP032538
51	University of Bergen	SRP033298
52	University of Vermont	SRP034591
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55	University of Vermont	SRP034591
56	RCEES, Chinese Academy of Sciences	SRP038053
57	RCEES, Chinese Academy of Sciences	SRP039005

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2	Universite du Littoral Cote d'Opale	SRP039908
3	Université Laval	SRP040423
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19	Université Laval	SRP040734
20	Université Laval	SRP040734
21	Universite du Littoral Cote d'Opale	SRP039908
22	Universié Laval	SRP043016
23	Universié Laval	SRP043016
24	Universié Laval	SRP043016
25	Universié Laval	SRP043016
26	Universié Laval	SRP043016
27	Universié Laval	SRP043016
28	Universié Laval	SRP043016
29	Universié Laval	SRP043016
30	Universié Laval	SRP043016
31	Universié Laval	SRP043016
32	Universié Laval	SRP043016
33	Universié Laval	SRP043016
34	Institute of Shandong River Wetlands	SRP045389
35	Alfred Wegener Instiute	SRP048617
36	Woods Hole Oceanographic Institution	SRP049010
37	Universite du Littoral Cote d'Opale	SRP039908
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2 Study Title
3 Accuracy of protist diversity assessments: morphology compared to cloning and direct pyrosequencing of 18S rRN
4 Accuracy of protist diversity assessments: morphology compared to cloning and direct pyrosequencing of 18S rRN
5 Accuracy of protist diversity assessments: morphology compared to cloning and direct pyrosequencing of 18S rRN
6 Accuracy of protist diversity assessments: morphology compared to cloning and direct pyrosequencing of 18S rRN
7 Accuracy of protist diversity assessments: morphology compared to cloning and direct pyrosequencing of 18S rRN
8 Microbial community response during the iron fertilization experiment LOHAFEX
9 Microbial community response during the iron fertilization experiment LOHAFEX
10 Microbial community response during the iron fertilization experiment LOHAFEX
11 Microbial community response during the iron fertilization experiment LOHAFEX
12 Microbial community response during the iron fertilization experiment LOHAFEX
13 Alexandrium catenella
14 Composition and genetic diversity of picoeukaryotes in subtropical coastal waters as revealed by 454 sequencing-l
15 Composition and genetic diversity of picoeukaryotes in subtropical coastal waters as revealed by 454 sequencing-l
16 Microbial Community Diversity Response to a Changing Arctic Ocean
17 Microbial Community Diversity Response to a Changing Arctic Ocean
18 Microbial Community Diversity Response to a Changing Arctic Ocean
19 Microbial Community Diversity Response to a Changing Arctic Ocean
20 Marine microbial eukaryote community analysis in Korea
21 Marine microbial eukaryote community analysis in Korea
22 Error rate of 454 GS FLX sequenced V4 and V9 PCR amplicons of various protists
23 Error rate of 454 GS FLX sequenced V4 and V9 PCR amplicons of various protists
24 Rotylenchulus reniformisGenome sequencing
25 Arctic sea ice Targeted Locus (Loci)
26 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
27 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
28 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
29 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
30 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
31 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
32 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
33 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
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41 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
42 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
43 Microbial Diversity and Potential Pathogens in Ornamental Fish Aquarium Water
44 Lake A protist communities
45 Lake A protist communities
46 Lake A protist communities
47 Lake A protist communities
48 Marine eukaryotic protists Metagenome
49 Partial ribosomal RNA gene of DHAB Ciliates
50 Highborne Bay eukaryotic small subunit ribosomal RNA diversity
51 Shark Bay eukaryotic small subunit ribosomal RNA diversity
52 Marine eukaryotic protists Metagenome
53 Sponge metagenome
54 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
55 Protistan communities of the upper Arctic Ocean (18S SSU-rRNA Targeted Locus)
56 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
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1 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
2 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
3 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
4 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
5 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
6 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
7 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
8 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
9 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
10 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
11 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
12 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
13 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
14 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
15 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
16 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
17 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
18 Reef-building coral species harbour Chromera velia and apicomplexan “type-N” (cf. *Gemmocystis cylindrus*) on the
19 Ward Hunt Lake Dilution Experiment Eukarya
20 Uncultured marine eukaryotes Targeted Locus (Loci)
21 Surface ocean Targeted Locus (Loci)
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47 Surface ocean Targeted Locus (Loci)
48 Surface ocean Targeted Locus (Loci)
49 Surface ocean Targeted Locus (Loci)
50 Gut content of salps Targeted Locus (Loci)
51 Environmental eukaryotes Metagenome
52 Design and validation of three primer sets to target the 18S rRNA gene of rumen protozoa.
53 Design and validation of three primer sets to target the 18S rRNA gene of rumen protozoa.
54 Design and validation of three primer sets to target the 18S rRNA gene of rumen protozoa.
55 454 pyrosequencing of plankton sample collected from Hamilton Harbour
56 Plankton sample collected from Nanaimo Harbour Targeted Locus (Loci)

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2 Marine metagenome Targeted Locus (Loci)
3 Uncultured eukaryotes Targeted Locus (Loci)
4 Uncultured eukaryotes Targeted Locus (Loci)
5 Uncultured eukaryotes Targeted Locus (Loci)
6 Uncultured eukaryotes Targeted Locus (Loci)
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32 Uncultured eukaryotes Targeted Locus (Loci)
33 Eukaryota Arctic Ocean Survey Targeted Locus (Loci)
34 Eukaryota Arctic Ocean Survey Targeted Locus (Loci)
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1 Eukaryota Arctic Ocean Survey Targeted Locus (Loci)
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19 Eukaryota Arctic Ocean Survey Targeted Locus (Loci)
20 Eukaryota Arctic Ocean Survey Targeted Locus (Loci)
21 Marine metagenome Targeted Locus (Loci)
22 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
23 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
24 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
25 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
26 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
27 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
28 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
29 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
30 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
31 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
32 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
33 Protists in Havre-aux-Maisons Lagoon, Magdalen Islands, Quebec, Canada
34 aquatic metagenome Genome sequencing
35 Marine picoplankton Targeted Locus (Loci)
36 Environmental pyrotags for eukaryotes Targeted Locus (Loci)
37 Marine metagenome Targeted Locus (Loci)

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	Sample Accession	Sample Title	Total Size, Mb	Total RUNs	Total Spots	Total Bases
1	ERS169722	marine metagenome	23.79	1	21827	9839469
2	ERS169723	marine metagenome	21.43	1	23993	8082496
3	ERS169724	marine metagenome	18.76	1	17244	7794090
4	ERS169725	marine metagenome	17.47	1	19798	6666525
5	ERS249848		85.86	1	72440	38229935
6	ERS249849		102.95	1	81845	47054441
7	ERS249850		118.91	1	98290	53957382
8	ERS249851		100.9	1	88895	43788914
9	SRS002344		35.51	1	71986	18253824
10	SRS004112		61.34	1	100911	27881207
11	SRS004113		53.61	1	87392	24193478
12	SRS152378		75.57	2	74742	32439603
13	SRS152378		105.05	2	93195	46778615
14	SRS152378		190.11	2	174077	86125205
15	SRS182654		3.27	1	3025	1487037
16	SRS182654		4.98	1	4575	2292151
17	SRS086207		22.19	1	53155	10757013
18	SRS086207		14.35	1	15221	5812974
19	SRS265308		1523.59	1	1206043	732000297
20	SRS345723		115.48	1	97702	48782681
21	SRS346704		12.67	1	11368	6229595
22	SRS346705		15.56	1	14233	7736979
23	SRS346706		9.52	1	8332	4512789
24	SRS346707		11	1	9993	5429341
25	SRS346708		14.43	1	13015	7108579
26	SRS346709		17.94	1	16171	8854309
27	SRS346705		3.35	1	8248	1578966
28	SRS346704		2.99	1	7443	1424093
29	SRS346707		3.3	1	8032	1538044
30	SRS346708		3.96	1	9691	1858955
31	SRS346706		3.07	1	7788	1489084
32	SRS346709		3.29	1	8133	1555000
33	SRS348706		3.9	1	9538	1827278
34	SRS355910		86.28	1	72724	36293480
35	SRS355910		42.46	1	35716	17867669
36	SRS355910		76.54	2	66130	34180949
37	SRS367508		378.84	1	300582	180399874
38	SRS375573		540.47	1	513347	236974344
39	SRS377421		277.64	1	260367	120224628
40	SRS377419		293.83	1	273186	128190762
41	SRS415390		351.24	1	278116	166721838
42	SRS417793		4.62	1	11325	6671280
43	SRS418650		0.91	1	7981	1692614
44	SRS419165		322.41	1	273168	144033297
45	SRS419338		0.41	1	3871	806497

1	SRS419339	0.81	1	7392	1559199
2	SRS419340	1.04	1	11758	2445426
3	SRS418650	0.68	1	3132	976006
4	SRS418650	1.64	1	7869	2535594
5	SRS418650	5.82	1	21910	8789284
6	SRS419339	0.82	1	3714	1214580
7	SRS419339	1.77	1	8746	2761817
8	SRS419339	1.93	1	7460	2842999
9	SRS419338	4.94	1	18201	7310788
10	SRS419338	1.04	1	3904	1497539
11	SRS419338	0.82	1	3682	1202397
12	SRS419340	0.99	1	4352	1465489
13	SRS419340	0.91	1	4454	1311120
14	SRS419340	0.89	1	3886	1329835
15	SRS463199	196.76	2	173394	87300958
16	SRS474263	162.05	1	144296	73326694
17	SRS474328	8.68	1	7316	3910652
18	SRS474346	18.1	1	15303	8161494
19	SRS474349	15.9	1	13493	7128393
20	SRS474350	13.83	1	11664	6215030
21	SRS474351	16.15	1	13734	7262782
22	SRS474352	18.29	1	15517	8240291
23	SRS474353	11.44	1	9655	5121533
24	SRS474354	17.62	1	14747	7924648
25	SRS474356	16.06	1	13611	7207989
26	SRS474357	17.41	1	14939	7808117
27	SRS474358	9.22	1	7748	4147573
28	SRS474359	13.97	1	12148	6399423
29	SRS474360	18.17	1	15778	8326048
30	SRS474361	13.18	1	11435	5969057
31	SRS474362	8.88	1	7728	4058666
32	SRS474363	13.63	1	11864	6249808
33	SRS474364	12.67	1	11010	5797710
34	SRS474365	12.04	1	10497	5506280
35	SRS474366	14.59	1	12686	6679449
36	SRS474367	14.41	1	12487	6529205
37	SRS474368	14.11	1	12243	6426116
38	SRS474369	14.04	1	12182	6373006
39	SRS474371	9.96	1	8600	4515861
40	SRS498636	297.51	1	337737	130753424
41	SRS507106	219.12	1	860390	354289445
42	SRS514888	5.06	1	22051	10260945
43	SRS514890	2.82	1	11405	5460929
44	SRS514891	1.69	1	9223	3633683
45	SRS559023	209.76	4	1714374	329548979
46	SRS563175	169.38	2	789405	273242265

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2	SRS569198	521.92	1	498945	236436763
3	SRS579457	11.73	1	10432	5170818
4	SRS579488	10.13	1	8796	4472640
5	SRS579489	10.83	1	9743	4774971
6	SRS579508	10.88	1	9487	4807091
7	SRS579509	10.08	1	8872	4460573
8	SRS579510	8.73	1	7592	3840296
9	SRS579511	11.39	1	10188	5037956
10	SRS579512	8.88	1	7740	3918258
11	SRS579513	11.34	1	9971	5060281
12	SRS579514	9.92	1	8615	4382786
13	SRS579515	12.26	1	10827	5479712
14	SRS579516	10.81	1	9410	4779994
15	SRS579517	11.2	1	9851	4988947
16	SRS579518	10.54	1	9167	4636301
17	SRS579519	10.48	1	9209	4650741
18	SRS579520	10.89	1	9398	4783119
19	SRS579521	11.97	1	10539	5349735
20	SRS579522	11.62	1	10111	5108976
21	SRS579523	10.54	1	9432	4725730
22	SRS579524	9.49	1	8300	4179958
23	SRS579525	11.33	1	10010	5047486
24	SRS579526	11.09	1	9847	4830421
25	SRS579527	10.63	1	9333	4732068
26	SRS579528	8.71	1	7578	3833004
27	SRS584239	9.26	1	9027	4434127
28	SRS584240	8.98	1	8782	4296411
29	SRS584241	12.45	1	12226	5973192
30	SRS584242	12.22	1	12079	5870832
31	SRS584243	9.91	1	9660	4741315
32	SRS584244	11.71	1	11525	5643505
33	SRS584245	12.91	1	12567	6203763
34	SRS584246	12.43	1	12097	5938216
35	SRS584248	18.89	1	18592	9110667
36	SRS584249	13.26	1	13090	6382758
37	SRS584251	12.93	1	12647	6195428
38	SRS584252	12.38	1	12162	5972069
39	SRS584257	10.22	1	9454	4727038
40	SRS584258	9.5	1	8796	4397118
41	SRS584324	11.74	1	10844	5394881
42	SRS584326	11.33	1	10463	5210335
43	SRS584327	11.8	1	10936	5474639
44	SRS584345	13.6	1	12616	6222179
45	SRS584346	12.56	1	11734	5854858
46	SRS584357	11.93	1	11028	5503305
47	SRS584358	13.44	1	12395	6210778
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1	SRS584359	14.03	1	12942	6456278
2	SRS584360	8.6	1	7907	3955921
3	SRS584361	14.47	1	13365	6677072
4	SRS584362	12.43	1	11770	5895814
5	SRS584363	12.81	1	12106	6060868
6	SRS584364	13.81	1	13145	6534390
7	SRS584365	13.88	1	13345	6585414
8	SRS584366	7.25	1	6960	3450835
9	SRS584367	15.2	1	14340	7186457
10	SRS584368	15.66	1	14810	7397424
11	SRS584369	14.13	1	13423	6633286
12	SRS584370	14.5	1	13560	6756945
13	SRS584372	13.97	1	13192	6577063
14	SRS584373	14.7	1	13435	6832284
15	SRS584374	13.14	1	12472	6190537
16	SRS569198	2726.45	1	1135587	1033738707
17	SRS629197	12.05	1	10283	5062010
18	SRS629198	12.16	1	10368	5109882
19	SRS629199	8.25	1	6992	3454326
20	SRS629200	13.76	1	11779	5796806
21	SRS629201	12.28	1	10494	5171193
22	SRS629202	12.39	1	10538	5211005
23	SRS629203	13.89	1	11819	5847093
24	SRS629204	13.02	1	11077	5470294
25	SRS629205	15.87	1	13484	6676694
26	SRS629206	11.71	1	9963	4932541
27	SRS677471	193.28	1	155510	90495203
28	SRS715739	0.72	4	34093	1251455
29	SRS722944	270.9	1	229068	122310596
30	SRS752744	1809.99	1	1566895	865799314
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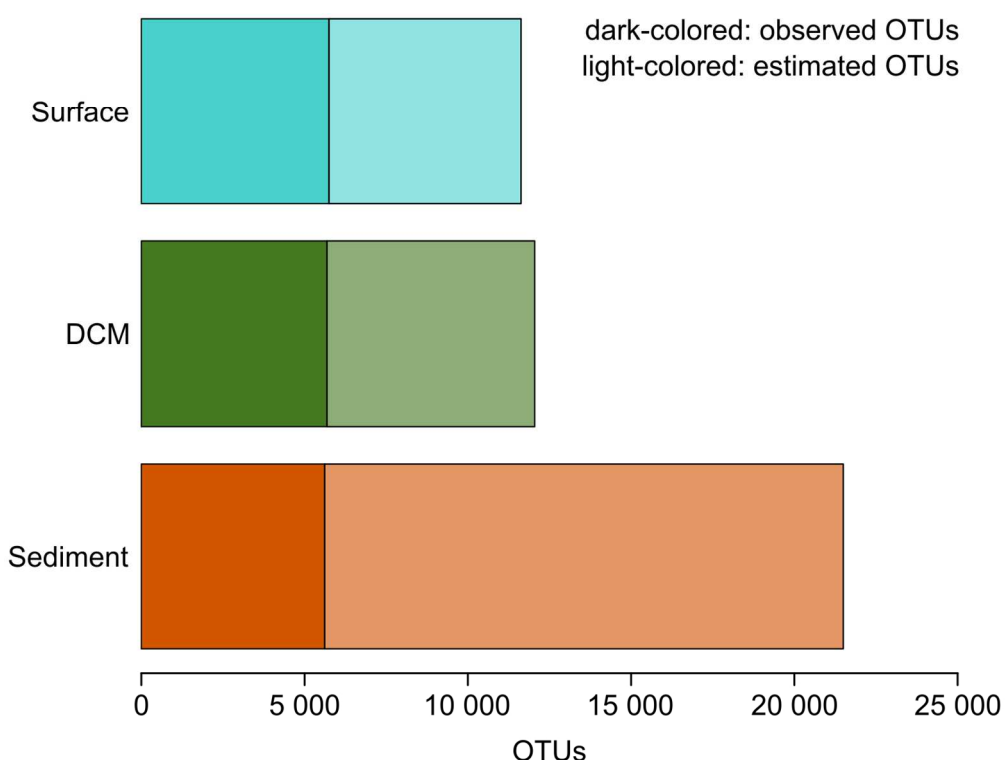
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2	TintinnidITSlib	AMPLICON	GENOMIC	PCR
3	Tintinnid_SSU_lib	AMPLICON	GENOMIC	PCR
4	TintinnidITSlib	AMPLICON	GENOMIC	PCR
5	LOHAFEX library of sample day_-1	AMPLICON	METAGENOMIC	PCR
6	LOHAFEX library of sample day_9	AMPLICON	METAGENOMIC	PCR
7	LOHAFEX library of sample day_18	AMPLICON	METAGENOMIC	PCR
8	LOHAFEX library of sample day_36	AMPLICON	METAGENOMIC	PCR
9	SUBINI	EST	TRANSCRIPTOMIC	RANDOM
10	MB	AMPLICON	GENOMIC	PCR
11	TH	AMPLICON	GENOMIC	PCR
12	AMCEpooledBACT	AMPLICON	OTHER	PCR
13	AMCEpooledARCH	AMPLICON	OTHER	PCR
14	AMCEpooledEUK	AMPLICON	OTHER	PCR
15	Ciliate-specific	AMPLICON	METAGENOMIC	PCR
16	Eukaryote-universal	AMPLICON	METAGENOMIC	PCR
17	GS_FLX	AMPLICON	METAGENOMIC	PCR
18	GS_FLX_Titanium	AMPLICON	METAGENOMIC	PCR
19	Lib_RepliGRN	WGS	GENOMIC	unspecified
20	ASID	AMPLICON	METAGENOMIC	PCR
21	VS-454-MBL:assay:D1.3_Bacteria	AMPLICON	METAGENOMIC	PCR
22	VS-454-MBL:assay:E1.5_Bacteria	AMPLICON	METAGENOMIC	PCR
23	VS-454-MBL:assay:D2.4_Bacteria	AMPLICON	METAGENOMIC	PCR
24	VS-454-MBL:assay:A1.1_Bacteria	AMPLICON	METAGENOMIC	PCR
25	VS-454-MBL:assay:A2.2_Bacteria	AMPLICON	METAGENOMIC	PCR
26	VS-454-MBL:assay:E2.6_Bacteria	AMPLICON	METAGENOMIC	PCR
27	VS-454-MBL:assay:E1.V4.11_Bacteria	AMPLICON	METAGENOMIC	PCR
28	VS-454-MBL:assay:D1.V4.9_Bacteria	AMPLICON	METAGENOMIC	PCR
29	VS-454-MBL:assay:A1.V4.7_Bacteria	AMPLICON	METAGENOMIC	PCR
30	VS-454-MBL:assay:A2.V4.8_Bacteria	AMPLICON	METAGENOMIC	PCR
31	VS-454-MBL:assay:D2.V4.10_Bacteria	AMPLICON	METAGENOMIC	PCR
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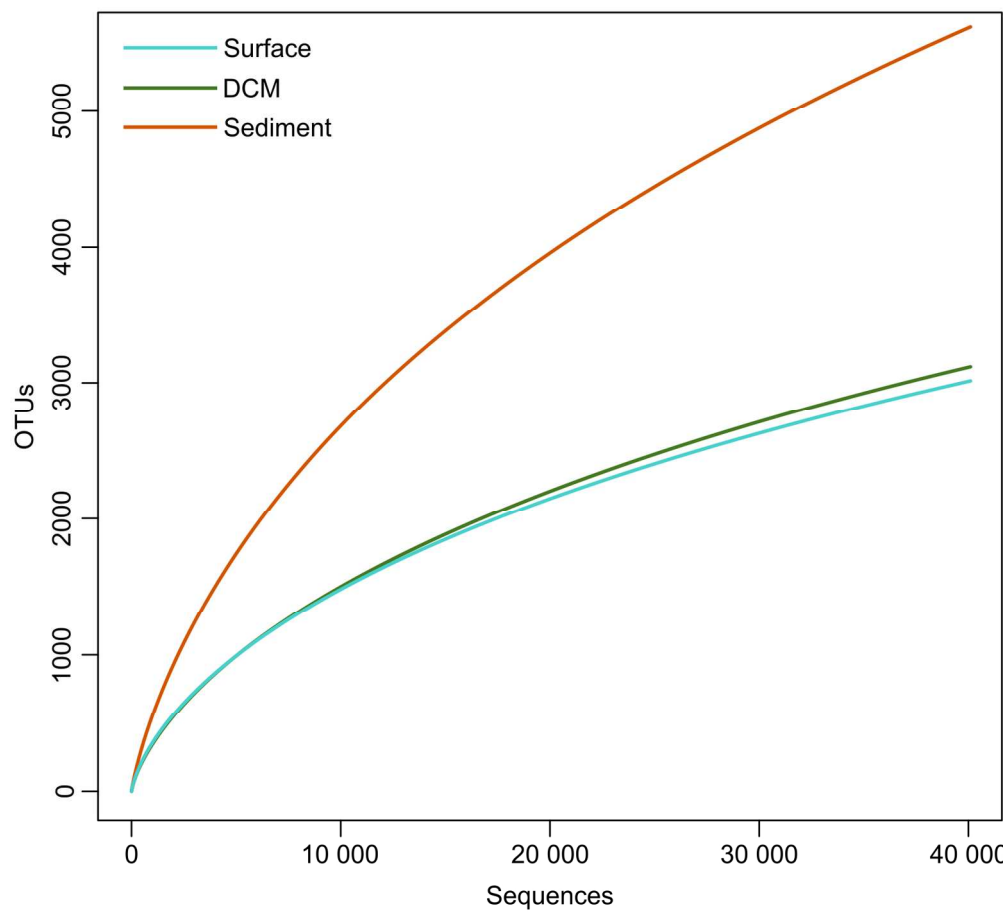
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52	ThreeSeds	AMPLICON	METAGENOMIC	PCR
53		AMPLICON	GENOMIC	PCR
54		AMPLICON	GENOMIC	PCR
55		AMPLICON	GENOMIC	PCR
56		AMPLICON	GENOMIC	PCR
57	Hamilton-18S_Replicate	AMPLICON	METAGENOMIC	PCR
58	Nanaimo plankton	AMPLICON	GENOMIC	PCR
59				
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2	EEC_2013	AMPLICON	GENOMIC	PCR
3		AMPLICON	METAGENOMIC	PCR
4	JB121SF_RNA	AMPLICON	METAGENOMIC	cDNA
5	JB121SCM_DNA	AMPLICON	METAGENOMIC	PCR
6	JB121SCM_RNA	AMPLICON	METAGENOMIC	cDNA
7	JB700SF_DNA	AMPLICON	METAGENOMIC	PCR
8	JB700SF_RNA	AMPLICON	METAGENOMIC	cDNA
9	JB700SCM_DNA	AMPLICON	METAGENOMIC	PCR
10	JB700SCM_RNA	AMPLICON	METAGENOMIC	cDNA
11	JB601SF_DNA	AMPLICON	METAGENOMIC	PCR
12	JB601SF_RNA	AMPLICON	METAGENOMIC	cDNA
13	JB601SCM_DNA	AMPLICON	METAGENOMIC	PCR
14	JB601SCM_RNA	AMPLICON	METAGENOMIC	cDNA
15	NEC211SF_DNA	AMPLICON	METAGENOMIC	PCR
16	NEC211SF_RNA	AMPLICON	METAGENOMIC	cDNA
17	NEC211SCM_DNA	AMPLICON	METAGENOMIC	PCR
18	NEC211SCM_RNA	AMPLICON	METAGENOMIC	cDNA
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21	NEF43SF_RNA	AMPLICON	METAGENOMIC	cDNA
22	NEF43SCM_DNA	AMPLICON	METAGENOMIC	PCR
23	NEF43SCM_RNA	AMPLICON	METAGENOMIC	cDNA
24	NEF512SF_DNA	AMPLICON	METAGENOMIC	PCR
25	NEF512SF_RNA	AMPLICON	METAGENOMIC	cDNA
26	NEF512SCM_DNA	AMPLICON	METAGENOMIC	PCR
27	NEF512SCM_RNA	AMPLICON	METAGENOMIC	cDNA
28				
29	HB_2010_710_10m_euk	AMPLICON	GENOMIC	PCR
30	HB_2010_710_34m_euk	AMPLICON	GENOMIC	PCR
31	HB_2010_699_6m_euk	AMPLICON	GENOMIC	PCR
32	HB_2010_699_15m_euk	AMPLICON	GENOMIC	PCR
33	HB_2010_709_9m_euk	AMPLICON	GENOMIC	PCR
34	HB_2010_709_30m_euk	AMPLICON	GENOMIC	PCR
35	HB_2010_745_10m_euk	AMPLICON	GENOMIC	PCR
36	HB_2010_745_47m_euk	AMPLICON	GENOMIC	PCR
37	HB_2010_706_10m_euk	AMPLICON	GENOMIC	PCR
38	HB_2010_706_30m_euk	AMPLICON	GENOMIC	PCR
39	HB_2010_706_40m_euk	AMPLICON	GENOMIC	PCR
40	LS_2006_301_25m_euk	AMPLICON	GENOMIC	PCR
41	LS_2008_303_16m_euk	AMPLICON	GENOMIC	PCR
42	LS_2008_303_2m_euk	AMPLICON	GENOMIC	PCR
43	CB_2011_314_18m_euk	AMPLICON	GENOMIC	PCR
44	CB_2011_314_2m_euk	AMPLICON	GENOMIC	PCR
45	BB_2005_131_40m_euk	AMPLICON	GENOMIC	PCR
46	BB_2005_131_5m_euk	AMPLICON	GENOMIC	PCR
47	BB_2005_100_22m_euk	AMPLICON	GENOMIC	PCR
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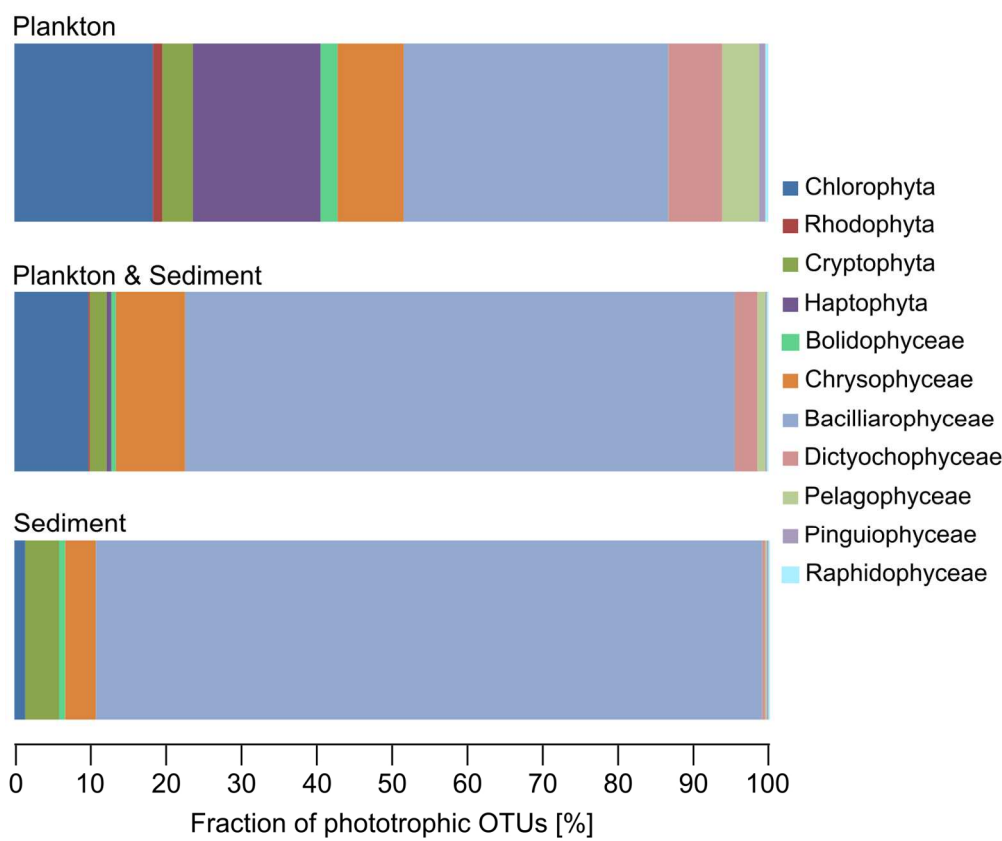
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7	BB_2006_123_45m_euk	AMPLICON	GENOMIC	PCR
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9	Lv_2007_VB0603_5m_euk	AMPLICON	GENOMIC	PCR
10	Lv_2007_VB1007_7m_euk	AMPLICON	GENOMIC	PCR
11	Lv_2007_VB1607_8m_euk	AMPLICON	GENOMIC	PCR
12	Lv_2007_VB2107_7m_euk	AMPLICON	GENOMIC	PCR
13	Lv_2007_VB3807_6m_euk	AMPLICON	GENOMIC	PCR
14	Lv_2007_VB3807_100m_euk	AMPLICON	GENOMIC	PCR
15	Lv_2007_VB7207_5m_euk	AMPLICON	GENOMIC	PCR
16	EEC_2013_De novo	WGS	METAGENOMIC	RANDOM
17		AMPLICON	GENOMIC	PCR
18		AMPLICON	GENOMIC	PCR
19		AMPLICON	GENOMIC	PCR
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31		AMPLICON	GENOMIC	PCR
32		AMPLICON	GENOMIC	PCR
33		AMPLICON	GENOMIC	PCR
34		CLONE	GENOMIC	unspecified
35		AMPLICON	GENOMIC	PCR
36	031814MPtarEuk	AMPLICON	METATRANSCRIPTOMIC	RT-PCR
37		AMPLICON	METAGENOMIC	PCR
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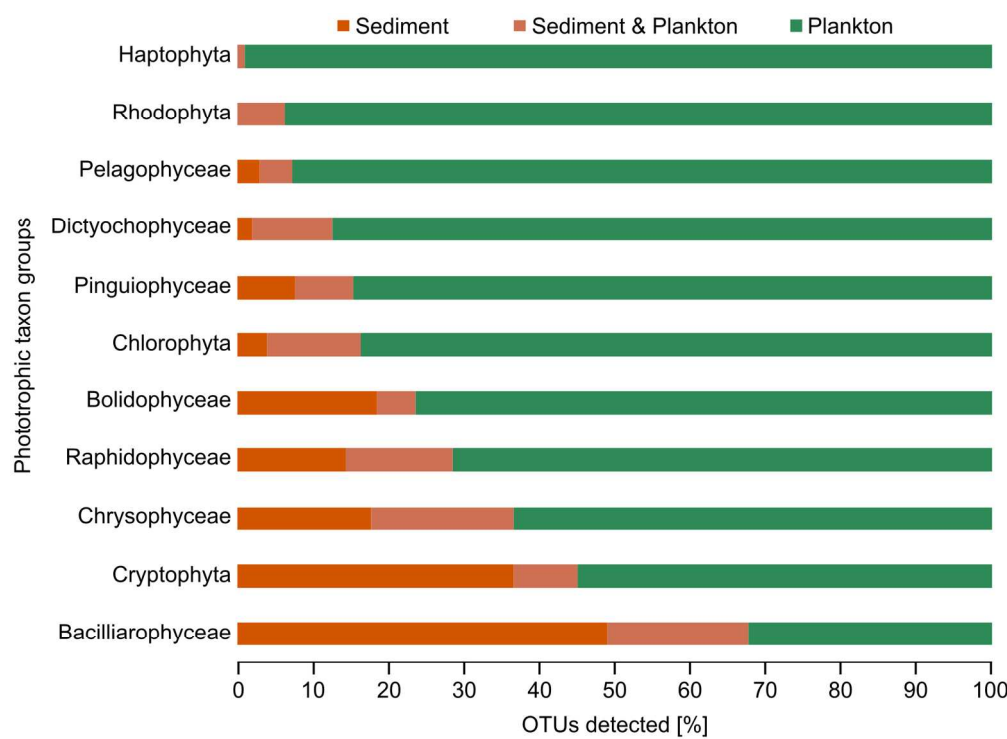


136x102mm (300 x 300 DPI)



152x138mm (300 x 300 DPI)





160x116mm (300 x 300 DPI)

