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Prokaryotic diversity and activity in contrasting productivity regimes in late summer in the  
Kerguelen region (Southern Ocean)

Alejandra Elisa Hernandez-Magana<sup>1,2\*</sup>, Yan Liu<sup>1, 3\*</sup>, Pavla Debeljak<sup>1,4†</sup>, Olivier Crispi<sup>1</sup>,

5 Barbara Marie<sup>1</sup>, Coco Koedooder<sup>1‡</sup>, Ingrid Obernosterer<sup>1#</sup>

<sup>1</sup> Sorbonne Université, CNRS, Laboratoire d'Océanographie Microbienne, LOMIC, F-66650,  
Banyuls/mer, France.

<sup>2</sup> Nordcee, Department of Biology, University of Southern Denmark, Odense M, Denmark

10 <sup>3</sup> School of Life Sciences, Ludong University, Yantai, China

<sup>4</sup> University of Vienna, Department of Functional and Evolutionary Ecology, A-1090 Vienna,  
Austria

present affiliation

15 <sup>†</sup> Institut de Systématique, Évolution, Biodiversité (ISYEB), Muséum National d'Histoire  
Naturelle, CNRS, Sorbonne Université

<sup>‡</sup>The Fredy and Nadine Herrmann Institute of Earth Sciences, Hebrew University of  
Jerusalem, Jerusalem, Israel

\*These authors contributed equally to this study.

20 <sup>#</sup> Corresponding author: Ingrid Obernosterer, Microbial Oceanography Laboratory (LOMIC),  
CNRS-Sorbonne University, 1 Avenue Pierre Fabre, 66650 Banyuls sur mer, France.

Tel: (+33) 04 68 88 73 53

e-mail: [ingrid.obernosterer@obs-banyuls.fr](mailto:ingrid.obernosterer@obs-banyuls.fr)

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25 **Abstract**

Natural iron (Fe) fertilization sustains phytoplankton blooms above the Kerguelen plateau (Indian sector of the Southern Ocean) within otherwise low productive off-plateau waters. In early spring and summer, these diatom-dominated blooms are associated with distinct heterotrophic prokaryotic communities, but whether a structuring effect extends to the post-bloom period has thus far not been investigated. To address this question, we carried out a detailed study of the prokaryotic community composition in the region of Kerguelen Island during late Austral summer (18 February to 27 March 2018; MOBYDICK project).

Concentrations of chlorophyll *a* were seasonally low above the plateau (0.27-0.58  $\mu\text{g Chl } a \text{ L}^{-1}$ ) and in a similar range to those at the 3 off-plateau sites investigated (0.14-0.34  $\mu\text{g Chl } a \text{ L}^{-1}$ ), but we observed an accumulation of dissolved organic carbon and the build-up of heterotrophic prokaryotic biomass in Kerguelen plateau waters. Illumina sequencing of the 16S rRNA gene revealed that the total (DNA-based) and potentially active (RNA-based) prokaryotic communities were structured according to on- and off-plateau sites in the wind-mixed surface layer, in both the free-living (< 0.8  $\mu\text{m}$  size fraction) and particle-attached (> 0.8  $\mu\text{m}$  size fraction) size fractions. The Amplicon Sequence Variants (ASV) with significantly higher relative abundances in on-plateau surface waters as compared to off-plateau waters belonged to *Haliaceae* OM60 group, several *Flavobacteriaceae*, such as the NS5 marine group, *Aurantivirga* and *Ulvibacter*, *Rhodobacteraceae* *Loktanella*, *Saprospiraceae*, and the *Cryomorphaceae* NS10 marine group. ASVs with higher relative abundances in off-plateau waters belonged to the *Flavobacteriaceae* *Formosa*, the *Rhodobacteraceae* *Planktomarina* and the SAR11 clade. We discuss the potential abiotic and biotic drivers of community composition in late Austral summer and the ecological roles of abundant prokaryotic taxa in Kerguelen plateau waters.

## 50 **1. Introduction**

Heterotrophic prokaryotes contribute to the cycling of all elements in the ocean. Within the marine carbon cycle, they process roughly half of recent primary production with most of the organic carbon being respired (Ducklow *et al.*, 2007). This process mediates large fluxes of carbon and energy between autotrophic and heterotrophic microbial communities and has  
55 important consequences for the potential storage of carbon in the ocean interior by the biological pump. Phytoplankton blooms are seasonally re-occurring events that stimulate prokaryotic and higher trophic level growth and metabolic activity. Diverse prokaryotes carry out the transformation of phytoplankton-derived organic matter (reviewed in Buchan *et al.*, (2014)) and the response is generally most pronounced a few weeks after the peak in  
60 phytoplankton biomass (Bunse and Pinhassi, 2017). The observed succession in prokaryotic communities over the course of phytoplankton blooms (Teeling *et al.*, 2012) illustrates the dynamic interplay between the composition of organic matter released by phototrophs, the metabolic capabilities of individual taxa, and the potential species-specific phytoplankton-prokaryote associations.

65 Phytoplankton blooms induced by natural iron (Fe) fertilization provide excellent opportunities to investigate these processes in the otherwise low-productive Southern Ocean. The region east of Kerguelen Island harbors the largest phytoplankton blooms induced by natural Fe fertilization (Blain *et al.*, 2007). Prokaryotes contribute substantially to the transformation of phytoplankton-derived organic matter during early spring (November),  
70 summer (January-February) and late summer (February-March) corresponding to the onset, peak and early decline, and the post-bloom phases in these perennially cold waters (Christaki *et al.*, 2020). These diatom-dominated blooms (Armand *et al.*, 2008; Lasbleiz *et al.*, 2016; Blain *et al.*, 2020) are associated with prokaryotic heterotrophic communities during early spring (Landa *et al.*, 2016) and in summer (West *et al.*, 2008; Obernosterer *et al.*, 2011) that

75 are distinct to those in surrounding low-productive waters. Onboard experiments have highlighted the role of diatom-derived organic matter in shaping prokaryotic community composition in spring (Landa *et al.*, 2016), and potential associations between diatom species and prokaryotic taxa has been identified as mechanism for the temporal structuring of microbial communities (Liu *et al.*, 2020). Observations from the post-bloom period when  
80 phytoplankton biomass has similar levels in on and off-plateau waters are thus far lacking, preventing to address the question on the effects Fe-fertilization on prokaryotic community composition at the seasonal scale in the Southern Ocean.

Numerous studies have described prokaryotic communities in different size-fractions, considered as free-living and particle-attached prokaryotes. The diversity and specific  
85 metabolic activity was shown to be higher in communities attached to particles as compared to free-living ones (Ghiglione *et al.*, 2007, Zhang *et al.*, 2007; Ortega-Retuerta *et al.*, 2013; Rieck *et al.*, 2015; Zhang *et al.*, 2020) or unchanged between size-fractions (Bachmann *et al.*, 2018). Observed differences have been associated to the respective lifestyle and metabolic capabilities. The composition, origin and quality of the particles was shown to have a strong  
90 influence on the associated prokaryotes (Zhang *et al.*, 2007; Ortega-Retuerta *et al.*, 2013; Rieck *et al.*, 2015). Therefore, exploring the free-living and particle-attached community structure can provide a better understanding of the processes influencing the prokaryotic diversity.

The present study was carried out as part of the project MOBYDICK (Marine Ecosystem  
95 Biodiversity and Dynamics of Carbon around Kerguelen: an integrated view) that aimed to provide a detailed picture of the diversity and role in the carbon cycling of biological communities of end-to-end food webs in contrasting nutrient regimes in the Southern Ocean. Our specific objective was to provide a description of the prokaryotic community composition. We considered the total (DNA-based) and potentially active (RNA-based) free-

100 living ( $<0.8 \mu\text{m}$ ) and particle-attached ( $>0.8 \mu\text{m}$ ) prokaryotic communities in the upper 300 m  
water column. Our observations from late Austral summer, corresponding to roughly 2  
months after the peak of the phytoplankton bloom, reveal distinct prokaryotic communities in  
Kerguelen plateau surface waters as compared to off-plateau waters. We further identified the  
prokaryotic taxa that explained the differences between on- and off- plateau waters, providing  
105 insights to microbial processes occurring during post-bloom conditions in this fertilized  
region of the Southern Ocean.

## 2. Material & Methods

### 2.1. Study area

110 Samples were collected in the region of Kerguelen Island in the Indian Sector of the Southern  
Ocean in late Austral summer (MOBYDICK cruise; February 18 to March 27, 2018) (Fig. 1).  
Four stations, identified as MOBYDICK-1 (M1), M2, M3 and M4 were sampled (Table 1).  
Station M2 is located on the central Kerguelen plateau (overall depth 527 m), southeast of  
Kerguelen Island, in naturally Fe-fertilized waters (Blain *et al.*, 2007). The other three stations  
115 are located in off-plateau waters, east (M1) and southwest of Kerguelen Island (M3 and M4).  
Station M3 is located in the polar frontal zone and stations M1, M2 and M4 are in Antarctic  
waters, with station M1 being strongly influenced by the polar front (Pauthenet *et al.*, 2018).  
The sampling period during the MOBYDICK cruise corresponded to roughly two months  
after the seasonal maximum in chlorophyll *a* (Chl *a*) concentrations in on- and off-plateau  
120 waters (Fig. S1). Station M2 was sampled three times (M2-1, M2-2 and M2-3) in an eight-day  
interval, stations M3 (M3-1 and M3-3) and M4 (M4-1 and M4-2) were each sampled twice  
with a 14-day interval and station M1 was visited once (Table 1).

## 2.2. Sample collection

125 Seawater samples were collected with 12 L Niskin bottles mounted on a rosette equipped with  
a conductivity, temperature, depth sensor (Seabird SBE-911 plus CTD unit). Discrete samples  
for concentrations of Chl *a* and dissolved organic carbon (DOC), and prokaryotic abundances  
were taken at 10 to 12 depths in the upper 500m. The sampling depths for prokaryotic  
community composition were chosen according to the CTD profiles with the aim to collect  
130 seawater at the top and at the base of the surface mixed layer ( $Z_{ML}$ ), in the transition layer,  
and in deeper waters (Fig. S2). We used this strategy to obtain 2 samples from the  $Z_{ML}$ , focus  
of our study, and to allow for a comparison of the surface water communities with those from  
deeper layers. Seawater samples were collected at 10m, 60m, 125m and 300m at all sites and  
visits, except for M2-1, where the following depths were sampled 10m, 50m, 100m and  
135 300m. For clarity, the 50 m and 100m sampling depths at station M2-1 will be referred to 60  
m and 125 m, respectively, in the description of the results and figures. The samples collected  
at the two uppermost depths were within the  $Z_{ML}$  at all sites and visits, except for Station M1  
where only the uppermost depth was within the shallow  $Z_{ML}$  (Table 1).

## 2.3. Concentration of Chl *a*

140 For the analysis of Chl *a* concentrations, 2.32 L of seawater were filtered through Whatman  
GF/F filters and the filters were stored in liquid nitrogen until analysis in the laboratory.  
Filters were extracted in 100% methanol, disrupted by sonication and clarified by GF/F  
filtration after 2h. Samples were analysed within 24h using High Performance Liquid  
Chromatography on an Agilent Technologies HPLC 1200 system equipped with a diode array  
145 detector following (Ras *et al.*, 2008).

#### 2.4. Dissolved organic carbon

The concentration of dissolved organic carbon (DOC) was determined in samples filtered through two combusted (450°C, 4h) GF/F filters. Subsamples of 10 mL (in triplicate) were transferred to pre-combusted glass ampoules and acidified with H<sub>3</sub>PO<sub>4</sub> (final pH = 2). The sealed glass ampoules were stored in the dark at room temperature until analysis. DOC measurements were performed on a Shimadzu TOC-V-CSH (Benner and Strom, 1993). Prior to injection, DOC samples were sparged with CO<sub>2</sub>-free air for 6 min to remove inorganic carbon. Hundred µL of each of the sample replicates were injected in triplicate and the analytical precision was 2%. Standards were prepared with acetanilide. Consensus reference materials provided in sealed glass ampoules (<http://www.rsmas.miami.edu/groups/biogeochem/CRM.html>) was injected every 12 to 17 samples to insure stable operating conditions.

#### 2.5. Enumeration of heterotrophic prokaryotes

For the enumeration of heterotrophic prokaryotes by flow cytometry, subsamples (1.44 mL) were fixed with glutaraldehyde grade I 25% (1% final concentration), and incubated for 30 min at 4 °C, then quick-frozen in liquid nitrogen and stored at -80 °C until analysis. Samples were collected from unfiltered seawater, considered as the bulk prokaryotic abundance, and from the < 0.8 µm size fraction, considered as the abundance of free-living prokaryotes. Samples were thawed at room temperature. Counts were performed on a FACSCanto II flow cytometer (Becton Dickinson) equipped with 3 air-cooled lasers: blue (argon 488 nm), red (633 nm) and violet (407 nm). For the enumeration of non-autofluorescent cells, mainly heterotrophic prokaryotes, cells were stained with SYBR Green I (Invitrogen – Molecular Probes) at 0.025% (vol / vol) final concentration for 15 min at room temperature in the dark. Stained prokaryotic cells were discriminated and enumerated according to their right-angle

light scatter (SSC) and green fluorescence at 530/30 nm. In a plot of green versus red fluorescence, non-autofluorescent cells were distinguished from autofluorescent cells. Fluorescent beads (1.002  $\mu\text{m}$ ; Polysciences Europe) were systematically added to each analyzed sample as internal standard. The cell abundance was determined from the flow rate, which was calculated with TruCount beads (BD biosciences).

## 2.6. Prokaryotic community composition

For the analysis of prokaryotic community composition, seawater was passed through a 60  $\mu\text{m}$  nylon mesh and 6 L of seawater was filtered through 0.8  $\mu\text{m}$  pore-size polycarbonate (PC) membranes (47 mm diameter, Nuclepore, Whatman, Sigma Aldrich, St Louis, MO) to retrieve the particle-attached ( $>0.8 \mu\text{m}$ ) fraction. Prokaryotic cells in the  $< 0.8 \mu\text{m}$  fraction were concentrated on 0.22  $\mu\text{m}$  cartridges (Sterivex<sup>TM</sup> Millipore, EMD, Billerica, MA) and considered as free-living fraction. The filters were stored in sterile Eppendorf tubes (2 ml) and the cartridges were sealed at both ends using parafilm. The filters and cartridges were stored at  $-80^\circ \text{C}$  until DNA and RNA extraction.

DNA and RNA extractions were performed simultaneously from the PC membranes ( $> 0.8 \mu\text{m}$  fraction) and the Sterivex<sup>TM</sup> cartridges ( $> 0.2 \mu\text{m}$  fraction) respectively, using the AllPrep Kit (Qiagen, Hilden, Germany) as described in (Liu *et al.*, 2019) with minor modifications. The filter units were thawed and closed with a sterile pipette tip end at the outflow, 425  $\mu\text{l}$  lysis buffer were added per sample (40mM EDTA, 50mM Tris and 0.75 M sucrose) and three freeze-thaw cycles were performed with liquid nitrogen and a water bath at  $65^\circ\text{C}$ . Subsequently, 25 $\mu\text{l}$  of freshly prepared lysozyme solution were added (2 mg  $\text{ml}^{-1}$  final concentration), the filter units were placed in a rotary mixer and incubated at  $37^\circ\text{C}$  during 45 minutes, and then 8 $\mu\text{l}$  of proteinase K solution (0.2 mg  $\text{ml}^{-1}$  final concentration) and sodium

dodecyl sulphate (SDS) (1%) were added and maintained at 55°C with gentle agitation every 10 min for 2 hrs.

To protect the RNA, 10 µl of β-mercaptoethanol were added to 1 ml of RLT plus buffer provided by the kit, 1,550 µl of this solution were added to each filter unit and mixed by inversion. The lysate was recovered by using a sterile 5 ml syringe and loaded in three additions onto the DNA columns by centrifuging at 10 000g for 30 s. DNA and RNA purifications were performed following the manufacturer's guidelines (Qiagen, Germany). The Invitrogen™ SuperScript™ VILO™ cDNA Synthesis Kit (Thermo Fisher Scientific Inc. Carlsbad, CA USA) was utilized to generate cDNA from the RNA extracts. Prior to the reverse transcription, absence of DNA in the RNA extracts was verified by PCR test with general primer sets 341F (5'-CCTACGGGNGGCWGCAG) and 805R (5'-GACTACHVGGGTATCTAATCC) (Herlemann *et al.*, 2011) for the prokaryotic 16S rRNA gene, followed by the examination of amplification products on 1% agarose gel electrophoresis. If a specific band of 1 015 bp was observed the RNA extract was treated with DNase.

The final PCR amplification of DNA and cDNA extracts was performed using the primers 515F-Y (5'-GTGYCAGCMGCCGCGG TAA) and 926R (5'-CCGYCAATTYMTTTRAGTTT) that encompasses the V4 and V5 hypervariable regions of the 16S rDNA (Parada *et al.*, 2016). Triplicate 10 µl reaction mixtures contained 2 µg DNA, 5 µl KAPA2G Fast HotStart ReadyMix, 0.2 µM forward primer and 0.2 µM reverse primer. PCR amplification was performed under the following conditions: an initial denaturation step of 95°C for 3 min, followed by 30 cycles of denaturation at 95°C for 45 s, annealing at 50°C for 45 s, and extension at 68°C for 90 s, and a final elongation step at 68°C for 5 minutes.

The presence of amplification products was confirmed by 1% agarose electrophoresis and  
220 PCR triplicates were pooled and purified by gel-filtration through-Sephadex G-50 Super Fine  
resin (Amersham Biosciences, Uppsala, Sweden). Then the purified samples were recovered  
for sequencing. The 16S rRNA gene amplicons were sequenced via next generation  
sequencing (Illumina MiSeq 2 × 250 bp chemistry on one flow-cell) at the platform GeT-  
PlaGe Genotoul (Toulouse, France). Mock community DNA (LGC standards, UK) was used  
225 as a standard for subsequent analyses and considered as a DNA sample for all treatments. In  
total, 128 samples (32 DNA free-living, 32 DNA particle-attached, 32 cDNA free-living and  
32 DNA particle-attached) were sequenced. After the sequencing one sample (cDNA free-  
living at M2\_1-60m) was discarded, due to low quality in the sequencing.

## 2.7. Sequence analysis

230 The samples obtained in the sequencing run were demultiplexed at the platform GeT-PlaGe  
Genotoul (Toulouse, France). A total of 5,847,892 sequences were obtained. Processing  
sequences was conducted with the DADA2 package for R version 1.10.1 (Callahan *et al.*,  
2016a), following the pipeline by (Callahan *et al.*, 2016b). Amplicon sequence variants  
(ASVs) were inferred through the high resolution DADA2 method (Callahan *et al.*, 2016a;  
235 Callahan *et al.*, 2017). Primers were trimmed and the sequences were filtered based on their  
quality using DADA2 (maxEE=2, truncQ=2). Forward reads with a length of 245 bp, reverse  
reads with a length of 210 pb, and in total 4,253,969 reads were kept after quality filtering.  
Error rates were estimated from the data, and inference of the sequence variants was made  
from the pooled sequences from all the samples. The 1,059,953 unique forward sequences  
240 and 1,619,348 reverse unique sequences were pooled to determine the sequence variants, then  
the forward and reverse sequences were merged and chimeras were removed. We obtained  
31, 227 unique ASVs. Taxonomy was assigned based on the SILVA database release 132 at  
the highest taxonomic level possible (Quast *et al.*, 2013).

From the 5,847,892 reads we obtained a total of 14,204 ASVs for the 127 samples after  
245 singletons removal. The number of reads per sample varied between 2,803 and 48,536. The  
dataset was randomly subsampled to the lowest number of reads (2,803) per sample with  
rarefy\_even\_depth( ) by Phyloseq R package version 1.26.1 (McMurdie and Holmes, 2013).  
Ordinations were performed with both the complete and subsampled datasets, and the trends  
observed were consistent for both datasets. To enable the comparison between samples with  
250 different numbers of reads, we chose to carry out all analysis using the subsampled dataset.  
After subsampling, 12,010 ASVs remained in the dataset.

From a total of 127 samples, the prokaryotic community composition was determined in the  
free-living ( $< 0.8 \mu\text{m}$ ) and in the particle-attached ( $> 0.8 \mu\text{m}$ ) fractions. DNA-based samples  
represent the total prokaryotic community and RNA-based samples were used to identify the  
255 potentially active members of the community (Blazewicz *et al.*, 2013). Our sequencing data  
provided access to 12,010 ASVs, of which 8,871 ASVs were obtained for the free-living  
fraction and 8, 838 ASVs for the particle-attached fraction, and 5,699 ASVs were shared  
between these fractions.

In order to identify the prokaryotic taxa, the ASVs were compared against the SILVA  
260 database release 132 at the highest taxonomic level possible (Quast *et al.*, 2013). The ASVs  
with the same taxonomy at the order level were pooled for illustration of the relative  
abundance in the samples. Sequences alignment was performed using MAFFT online service  
for multiple sequence alignment (Kato *et al.*, 2019) and the phylogenetic tree was built using  
PhyML 3.0 online program, based on the maximum likelihood method, with 100 bootstraps  
265 and the HKY85 substitution model (Guindon *et al.*, 2010). The phylogenetic tree was  
visualized with SeaView version 4.7 and saved as a rooted tree (Gouy *et al.*, 2010). The tree  
was imported in R with the ape package function read.tree( ).

To estimate the absolute cell numbers, the ASVs were corrected for copy numbers of the 16S rRNA gene per cell per specific taxa obtained from the ribosomal RNA database (Stoddard *et al.*, 2015). Total cell numbers per bacterial group and per Liter were then calculated with 16S rRNA gene relative proportions per group and total cell abundance from flow cytometry.

## 2.8. Data analysis

The difference between total (DNA-based) and potentially active communities (RNA-based) was tested by analysis of similarity (ANOSIM), resulting significant differences ( $p=0.001$ ,  $r=0.23$ , permutations=999). Likewise free-living and particle-attached fractions for both, total (DNA) and potentially active (RNA) communities revealed significant differences ( $p=0.001$ ,  $r=0.57$ , for DNA-based set and  $p=0.001$ ,  $r=0.41$  for RNA-based dataset, permutations=999).

The analyses performed thereafter considers therefore separately the four subsets, free-living and particle attached fractions for both, total (DNA) and potentially active (RNA) community.

To explore the distribution patterns of the prokaryotic communities we applied Non-Metric Multidimensional Scaling (NMDS), based on the dissimilarity matrix of the community structure. The statistical analyses were performed in R 3.5.3 version (R Core Team, 2019), Bray–Curtis dissimilarity matrices were generated via `vegdist()` function using the relative abundance of ASVs in each sample. Subsequently, the matrix was used to build the NMDS ordinations using `metaMDS()` function in the package `Vegan` (Oksanen *et al.*, 2019).

Diversity indices were calculated with `phyloseq` command `estimate_richness()`. ANOSIM was performed to test significant differences in microbial communities between sampling depths and among sites in the surface mixed layer.

The contribution of individual species (ASVs) to the average Bray-Curtis dissimilarity between the on- and off-plateau groups was obtained using similarity percentage analysis (SIMPER) (Clarke 1993). Station M3 was chosen as the representative off-plateau site and compared to the on-plateau station M2. The analysis was applied to the prokaryotic communities within the wind mixed surface layer. The ASVs with a relative abundance lower than 1% in at least one sample were discarded prior to the SIMPER analysis. Once the ASVs with significant contribution to the dissimilarity between groups were obtained (p-value < 0.005), the ASVs with relative abundance higher than 5% in at least one sample were selected for plotting them in a heatmap using the package pheatmap version 1.0.12 for R (Kolde, 2019).

Partial Mantel tests for the free-living and particle attached fractions in both total (DNA-based) and active (RNA-based) prokaryotic community composition, diatom community composition and environmental parameters (temperature, salinity, concentrations of dissolved oxygen, ammonium, nitrite, nitrate, phosphate, silicic acid and DOC), were performed in Vegan using mantel.partial() based on the Pearson correlation method. Prior to correlation analysis, environmental variables were z-score transformed. The amount of variance in prokaryotic community composition explained by diatoms and environmental parameters was estimated as the square of the correlation coefficient ( $Rho^2$ ) based on partial Mantel test. The Partial Mantel test was applied to samples in the upper 125m water column, for which diatom community composition was available.

Redundancy analysis (RDA) (Legendre & Legendre, 2012) was applied to evaluate the linkages between the prokaryotic community composition and the environmental parameters in Vegan using rda(). Prior to analysis the environmental variables were z-score transformed

315 and Hellinger transformation was applied to the community matrix (Legendre & Gallagher,  
2001). Permutational multivariate analysis of variance in Vegan with `adonis()` was used to  
select the significant environmental variables ( $p < 0.05$ ). We used the variance inflation factor  
(VIF) to determine the linear dependency among variables. We first evaluated all the  
variables and removed, one at a time, those with a VIF  $> 10$  from the dataset. Only the  
320 selected variables (with a VIF  $< 10$ ) were used to perform the final RDA analysis. As for the  
Partial Mantel test, samples in the upper 125m were considered.

### 3. Results

#### 325 3.1. Environmental conditions

During late Austral summer the water column was well-stratified, with the wind-mixed  
surface layer ( $Z_{ML}$ ) ranging between 49 m and 62 m during the first visits of the off-plateau  
station M4 (M4-1) and of the on-plateau station M2 (M2-1 and M2-2), respectively (Table 1  
and Fig. S2). The occurrence of a storm on March 10<sup>th</sup> led to a deepening of the  $Z_{ML}$  that was  
330 most pronounced at station M4 (87 m). A shallow  $Z_{ML}$  (27 m) was determined during our visit  
at station M1. Temperature in the  $Z_{ML}$  was lowest at the southernmost station M4 (4.5°C) and  
varied between 4.9°C (station M1) and 5.6°C (station M3-1) at the other sites (Table 1, Fig.  
S2). Seasonally low concentrations of Chl *a* were determined at all sites (Fig. S1), and those  
in the  $Z_{ML}$  of the on-plateau station M2 (0.27-0.58  $\mu\text{g L}^{-1}$ ) were 2-3-fold higher as compared  
335 to those at the off-plateau stations M3 (0.14-0.19  $\mu\text{g L}^{-1}$ ) and M4 (0.18-0.21  $\mu\text{g L}^{-1}$ ) (Fig. 2),  
and similar to those at station M1 (0.34  $\mu\text{g L}^{-1}$ ). (Chl *a*, DOC and prokaryotic abundance for  
station M1 are illustrated in Fig. S3). Chl *a* concentrations doubled in the  $Z_{ML}$  between the 2<sup>nd</sup>  
and 3<sup>rd</sup> visit at the on-plateau site M2.

Concentrations of dissolved organic carbon (DOC) were  $54.1 \pm 0.6 \mu\text{M}$  in the  $Z_{\text{ML}}$  of station  
340 M2 (all visits pooled), as compared to  $50.5 \pm 0.7 \mu\text{M}$  at station M3 and  $50.3 \pm 0.8 \mu\text{M}$  at station  
M4 (Fig. 2). This accumulation of DOC is likely a consequence of the enhanced seasonal  
phytoplankton activity in Kerguelen plateau waters. Heterotrophic prokaryotic abundance was  
as high as  $1.17 \times 10^6 \text{ cells mL}^{-1}$  in the  $Z_{\text{ML}}$  during our first visit of the on-plateau station M2,  
and cell abundances decreased to  $0.81 \times 10^5 \text{ cells mL}^{-1}$  and  $0.68 \times 10^5 \text{ cells mL}^{-1}$  during the  
345 following one and two weeks, respectively. Moderate (from  $6.96$  to  $4.45 \times 10^5 \text{ cells mL}^{-1}$  at  
M3) and minor decreases (from  $5.52$  to  $5.01 \times 10^5 \text{ cells mL}^{-1}$  at M4) were observed at the off-  
plateau stations over the 2 weeks that separated the visits (Fig. 2). The prokaryotic cell  
abundance in the  $< 0.8 \mu\text{m}$  fraction accounted for  $89 \pm 11 \%$  ( $n=10$ ) (Table S1) of the  
abundance in unfiltered seawater in surface waters at the different sites.

350

### 3.2. Vertical and spatial structuring of prokaryotic communities

When considering the entire data set, NMDS ordination revealed a clustering by depth layer,  
for the total and active prokaryotic communities in both size fractions. Samples collected for  
the free-living fraction at 10 m and 60 m, corresponding to the  $Z_{\text{ML}}$  except for station M1  
355 (Table 1), clustered and were distinct from those of 125 m and 300 m (Anosim,  $r=0.69$  for  
DNA and  $r=0.61$  for RNA,  $p=0.001$ ) (Fig. 3a for DNA and Fig. S4a for RNA). In the case of  
the particle-attached fraction, the clustering by depth was less pronounced, but still significant  
(Anosim,  $r=0.58$  for DNA and  $r=0.67$  for RNA,  $p=0.001$ ) (Fig. 3b for DNA and Fig. S4b for  
RNA). We determined multiple diversity indices that all revealed an increase with depth in  
360 both size-fractions (Table S2) as illustrated for the Shannon index on Table 2.

To identify possible differences in the prokaryotic community composition among sites, we  
then focused on samples collected in the  $Z_{\text{ML}}$ . For the free-living fraction, samples grouped

according to station (ANOSIM,  $r=0.77$  for DNA and  $r=0.51$  for RNA,  $p=0.001$ )(Fig. 3a and Fig. 4a). This pattern was again less pronounced for the particle-attached fraction (ANOSIM,  $r=0.62$ ,  $p=0.001$  for DNA and  $r=0.41$   $p=0.006$  for RNA)(Fig. 3b and Fig. S4b). Diversity of the total free-living communities was significantly lower in the  $Z_{ML}$  of the on-plateau station M2 (Shannon:  $4.34\pm 0.19$  for DNA,  $n=6$ ) as compared to that in off plateau waters at stations M1, M3 and M4 (Shannon:  $4.86\pm 0.15$  for DNA,  $n=9$ )(one-way ANOVA  $p<0.001$ ). This trend was, however, not observed for the potentially active free-living communities (Shannon:  $5.28\pm 0.71$  at station M2 and  $5.53\pm 0.77$  in the off-plateau stations,  $n=9$ ) and the particle-attached communities in the DNA and RNA data sets (Table 2 and Table S2).

### 3.3. Prokaryotic community composition in surface waters

We further explored in more detail the prokaryotic community composition in surface waters (10 m). The total free-living and particle-attached prokaryotic communities were dominated by *Flavobacteriales* (relative contributions of 53% and 64% in on-plateau waters, 35% and 36% in off-plateau waters, respectively) and *Rhodobacterales* (20% and 10% on-plateau, 17% and 22% off-plateau, respectively). *Cellvibrionales* represented 13% in both size fractions at the on-plateau station (M2), while at the off-plateau stations this order was substantially more abundant in the particle-attached (17%) than in the free-living fraction (3%) (Fig. 4; RNA data are illustrated in Fig. S5). Distinct features of the free-living communities in off-plateau waters were the comparably higher contributions of the SAR11 clade, belonging to *Pelagibacterales* (SAR11, 17-25%), the SAR86 clade (5-8%), *Thiomicrospirales* (2-3%) and *Puniceispirillales* (2-4%). *Verrucomicrobiales* were present only in the total particle-attached communities and their relative abundances were higher in the off-plateau sites (4 - 8%) than in on-plateau waters (1.3%). *Chitinophagales* were detectable only in the particle-attached

fraction at station M2 in both the total (2.2%) and potentially active communities (3.2%), while *Sphingomonadales* (2.5%), *Lactobacillales* (1.4%) and *Rickettsiales* (1.3%) appeared to be specific to the total particle-attached communities at station M1.

390 The high resolution allowed us to identify the relative contribution of the members of the prokaryotic community at the level of ASVs. Interesting information derived at this level of resolution is that in the case of some of the abundant taxonomic groups composed by multiple ASVs, only a few of them were highly abundant. For example, in the total free-living fraction (Fig. S6), *Flavobacteriales* were composed of 24 ASVs, however, only some of them were  
395 more abundant in on-plateau than in off-plateau waters. This was the case for ASV6 *Ulvibacter*, ASV12 (NS2b marine group), ASV23 (*Flavobacteriaceae* ASV15 *Aurantivirga* and ASV36 *Polaribacter*). By contrast, other *Flavobacteriales* ASVs were relatively more abundant in off-plateau waters, such as ASV3 *Formosa*, ASV38 and ASV88 (both NS7 marine group). Another interesting observation is that the most abundant ASV from our  
400 dataset, ASV1 OM60(NOR5), belonging to the gammaproteobacterial clade, was only highly abundant in the free-living fraction of the on-plateau station M2, which was not the case for the off-plateau free-living community (Fig. S6). By contrast, the same ASV was highly abundant at all sites in the particle-attached total community, and less abundant in the potentially active community (Fig. S7). Given these patterns obtained from our high  
405 resolution results of the community composition, we decided to explore which prokaryotes explained the difference among sites.

### 3.4. Identification of site-specific prokaryotes

To address the question of which prokaryotic taxa explained the difference in the community  
410 composition between on- and off-plateau sites (Fig. 3 and Fig. S4), we performed a Simper

analysis (permutations = 999) and selected the ASVs contributing significantly ( $p < 0.05$ ) to the differences between stations at 10 m and 60 m. Because seasonal Chl *a* concentrations were lower at M3 as compared to M4 (Fig. S1) we chose station M3 as a representative off-plateau site for the further comparison with station M2. In order to simplify the visualization of the information, we considered only highly abundant ASVs (relative abundance > 5% in at least one sample).

A total of 9 and 8 highly abundant ASVs, respectively, contributed significantly to the differences in the total (DNA) or potentially active (RNA) free-living communities between stations M2 and M3 at 10 m (Fig. 5; results for 60 m are shown in Fig. S8); among those, 5 ASVs were shared between the DNA and RNA datasets. The 9 and 8 ASVs explained each 24% and 8% of the differences among sites in the total and potentially active free-living communities, respectively. For the particle-attached communities 6 and 3 ASVs contributed significantly to the respective differences and these ASVs explained 28% and 4% of the observed variability. Two ASVs (ASV5 and ASV15) significantly contributed to the differences in both size fractions of the DNA and RNA dataset.

ASVs that had higher relative abundances at station M2 as compared to station M3 belonged to the OM60 group (NOR5 clade, *Haliaceae* ASV1), the *Flavobacteriaceae* NS5 marine group (ASV11), the genera *Aurantivirga* (ASV15) and *Ulvibacter* (ASV6), and ASV5 and ASV23 (not identified on the genus level), the *Saprospiraceae* (ASV33) and the *Cryomorphaceae* NS10 marine group (ASV52). The *Pirellulaceae* genus *Blastopirellula* (ASV8) had low relative abundances in the free-living fraction at station M2 (<0.1%) and this ASV was absent from station M3. ASVs with higher relative abundances at station M3 as compared to station M2 were the *Flavobacteriaceae* genus *Formosa* (ASV3), and the *Rhodobacteraceae* genera *Planktomarina* (ASV16) and *Loktanella* (ASV18). Contrasting patterns were observed for the SAR11 clade (ASV9) that had higher relative abundances in

the total free-living communities at station M3, but a higher contribution to the potentially active free-living communities at station M2.

### 3.5. Changes in the absolute abundance of major taxa

To explore the question of whether the pronounced decrease in bulk prokaryotic abundance  
440 between repeated visits at station M2 (Fig. 2) were due to changes in specific taxa, we  
estimated the absolute abundances of ASVs, grouped at the order level, using previously  
published rRNA operon copy numbers (Stoddard *et al.*, 2015). Our results reveal an increase  
in the abundance of SAR11 from the first to the third visit (by up to 150%) at station M2 at 10  
m and a decrease in the abundance of all other considered taxa by on average  $48\pm 12\%$  and  
445  $65\pm 10\%$  after one and two weeks, respectively (Fig. 6; RNA results are shown in Fig. S9).

### 3.6. Linking prokaryotic community composition to biotic and abiotic factors

The combined set of abiotic environmental parameters and diatom community composition,  
as determined by microscopic observations (Lafond *et al.*, 2020), could each explain changes  
450 in the total and potentially active prokaryotic communities in both size fractions and all depth  
layers ( $p < 0.05$ ; partial Mantel test; Table S3). Environmental parameters could explain 12%  
and 16% of the spatio-temporal changes in the total and potentially active free-living  
communities, respectively, and 9% and 10% of the respective particle-attached communities.  
Diatoms could explain 36% and 28% of the changes in the total and active free-living  
455 communities, respectively, and 20% and 14% of the total and active particle-attached  
communities. An additional RDA-analysis revealed a clear distinction in on and off-plateau  
prokaryotic communities in the  $Z_{ML}$  while samples from 125m of all sites grouped (Fig. S10).  
Among the environmental parameters tested (Table S4), significant predictors of the

community composition were temperature, salinity, dissolved oxygen, ammonium, nitrite and  
460 DOC. In the  $Z_{ML}$ , DOC was a significant predictor for the free-living ( $p=0.039$ ) and particle-  
attached fraction ( $p=0.01$ ), and this was the case for salinity and the communities at 125m  
( $p=0.001$  and  $p=0.007$ , respectively).

#### 4. Discussion

465 We observed distinct prokaryotic communities in surface waters above the central Kerguelen  
plateau as compared to those in off-plateau waters during the post-bloom period in late  
Austral summer. These findings add to observations from early spring (Landa *et al.*, 2016)  
and summer (West *et al.*, 2008; Obernosterer *et al.*, 2011) corresponding to the onset, peak  
and early decline of the phytoplankton bloom. A similar spatial structuring was also reported  
470 for diatoms (Armand *et al.*, 2008; Lasbleiz *et al.*, 2016; Lafond *et al.*, 2020), non-diatom  
phytoplankton and protists (Georges *et al.*, 2014; Irion *et al.*, 2020; Sassenhagen *et al.*, 2020)  
during the different bloom phases, and indicates a seasonally persistent influence of natural  
Fe fertilization on unicellular plankton communities. We discuss in the following the potential  
role of abiotic and biotic factors to better understand this pattern and the role of specific taxa  
475 in the microbial cycling of elements.

##### 4.1. Post-bloom conditions and bulk prokaryotic features

A distinct feature of the surface waters above the Kerguelen plateau in late summer is the  
accumulation of DOC. Both, the Partial Mantel test and RDA analysis indicated that DOC  
concentrations had significant influence on the prokaryotic community composition in the  
480 wind mixed surface layer (Table S3, Fig. S10). The higher DOC concentrations likely result  
from the seasonally enhanced autotrophic activity, the excretion of phytoplankton-derived  
organic matter as cells become less active when entering a senescent stage (Myklestad *et al.*,

1989; Barofsky *et al.*, 2009), and the release of DOC due to grazing (i.e. ‘sloppy-feeding’) (Steinberg and Landry, 2017), leading to a relief of organic carbon limitation (Obernosterer *et al.*, 2015). Besides the quantity, the composition of organic matter could differ among on- and off-plateau sites as a consequence of the varying productivity regimes and thereby contribute to shaping the community composition. The high similarity among prokaryotes at the off-plateau sites, despite their location in different water masses supports this idea.

Seasonal observations in Kerguelen plateau surface waters could provide some insights on organic matter availability and its influence on community composition. Prokaryotic growth rates determined during the post-bloom period (0.04 to 0.15 d<sup>-1</sup>) (Christaki *et al.*, 2020) were in the same range as those in early spring (0.025 to 0.12 d<sup>-1</sup>) (Christaki *et al.*, 2014), and lower than those determined in summer (0.22 to 0.47 d<sup>-1</sup>) (Christaki *et al.* 2008), suggesting an increase in organic matter availability during the peak and early decline of the bloom. The seasonal modifications in organic matter availability were paralleled by changes in prokaryotic diversity. In late summer, the Shannon index in Kerguelen plateau surface waters was in the same range to those determined in early spring at the same site (about 4.5), while a drop to 2.5 was observed following the peak of the spring bloom (Liu *et al.*, 2020). This latter observation points to the dominance of a few fast-growing opportunists taking advantage of the labile organic matter during the peak and just after the bloom (Liu *et al.*, 2020), while the utilization of the post-bloom organic matter pool appears to be attributable to a large number of taxa with well-defined ecological niches.

#### 4.2. Abundant prokaryotic taxa in Kerguelen plateau waters in late summer

*Flavobacteriaceae* were present at all sites, but accounted for up to 53% and 63% of relative abundance of the free-living and particle-attached total community in Kerguelen plateau surface waters. Members of this family contain a diverse repertoire of enzymatic capabilities for the degradation of complex compounds, including those of phytoplankton origin

(Kappelmann *et al.*, 2019). Substrate preferences vary among taxa (Xing *et al.*, 2015; Krüger *et al.*, 2019) leading to the niche partitioning among diverse members of this family, including the late summer community in Kerguelen plateau waters (Sun *et al.*, in press). *Aurantivirga* explained differences between on- and off-plateau waters in the total and potentially active communities in both fractions. *Aurantivirga* was shown to account for about 10% throughout the summer period in Kerguelen plateau waters (Liu *et al.*, 2020). The key role of *Aurantivirga* could be explained by the high number of polysaccharide uptake loci (PULs) and the diverse substrate spectra for glycan degradation reported for members of this genus (Krüger *et al.*, 2019). On the contrary, the *Flavobacteriaceae* genus *Formosa* had higher abundances in off plateau waters, and thus appears to be adapted to a more oligotrophic-type lifestyle.

One ASV belonging to the OM60 group (*Gammprotoebacteria*, NOR5 clade, *Halielaceae*) was among the most abundant taxa of our dataset and this ASV contributed significantly to the differences in communities between on- and off plateau surface waters. This contrasts observations from November through February when the relative abundance of this group remained low (< 2%) in plateau waters (Liu *et al.*, 2020). This clade comprises aerobic anoxygenic phototrophs and members associated to aggregates where they thrive under sub-oxic conditions (Fuchs *et al.*, 2007). In the present study, this ASV was highly abundant in the total particle-attached community (DNA-based) at all sites (Fig. S8). Another interesting feature is the potential implication of members of this clade in the degradation of the sulphur compound dimethylsulfoniopropionate (DMSP) (Nowinski *et al.*, 2019; Steiner *et al.*, 2019). Kerguelen surface waters were not enriched in particulate DMSP in summer during the peak of the diatom-dominated bloom (Belviso *et al.*, 2008). However, the shift to non-diatom phytoplankton, including DMSP producers such as Haptophytes (Schoemann *et al.*, 2005),

dominated by *Phaeocystis Antarctica* in the study region (Irion *et al.*, 2020) could suggest a different scenario in late summer. The changes in the late summer light regime due to the low  
535 phytoplankton biomass in the well-stratified shallow surface mixed layer and changes in organic carbon sources provided by non-diatom phytoplankton could have led to the success of the OM60 group. One ASV belonging to *Saprospiraceae* (*Chitinophagales*) was highly abundant in the particle-attached community, but accounted for <1% on a seasonal scale (Liu *et al.*, 2020). *Saprospiraceae* were shown to respond in terms of relative abundance in  
540 incubation experiments enriched in alginate particles (Mitulla *et al.*, 2016). The associated glycolytic abilities could be advantageous for members of this family in the occupation of a specific substrate niche, provided for example by aggregates of detrital and non-living cells that contributed to up to 65% to total particulate organic carbon in late summer in Kerguelen plateau surface waters (Lafond *et al.*, 2020).

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#### 4.3. Potential role of biotic interactions in shaping community composition

Diatom assemblages explained a substantial part of the changes in prokaryotic community composition, despite their lower biomass as compared to other bloom phases (Irion *et al.*, 2020; Lafond *et al.*, 2020). Diatoms were shown to be drivers of the prokaryotic community  
550 composition in the Southern Ocean on spatial (Liu *et al.*, 2019) and seasonal scales (Liu *et al.*, 2020), with diatom-derived DOM playing an important role in spring (Landa *et al.*, 2016). In late Austral summer, *Corethron inerme* accounted for 60-80% of total diatom biomass and it was among the most actively silicifying species in on-plateau waters (Lafond *et al.*, 2020). On a seasonal scale, *C. inerme* had a large number of positive correlations with free-living and  
555 attached prokaryotes in Kerguelen plateau waters (Liu *et al.*, 2020). These included taxa identified in the present study as being more abundant in on-plateau waters, in particular those belonging to *Flavobacteriaceae* and *Cryomorphaceae*. Another diatom abundant in Austral

summer in the study region is *Rhizosolenia* spp. (Armand *et al.*, 2008; Blain *et al.*, 2020; Lafond *et al.*, 2020). This diatom revealed strong positive correlations with the *Roseobacter* genus *Loktanella* (Liu *et al.*, 2020), a taxon that was highly abundant in off-plateau waters in the particle-attached fraction in the present study. *Corethron* spp. and *Rhizosolenia* spp. are both large diatoms (about 20  $\mu\text{m}$  equivalent spherical radius, ESR), and could thus present habitats for taxa with a particle-attached lifestyle.

The marked decrease in bulk prokaryotic abundances during the consecutive visits at the on-plateau station was predominantly due to grazing by heterotrophic nanoflagellates (Christaki *et al.*, 2020). This raises the question of whether selective ingestion could have influenced prokaryotic community composition. Our observations of a narrow range in the decrease of taxon-specific cells over time suggests grazing to be non-selective, but rather a function of the cell abundance and thus the encounter rate of a given taxon. In addition, prokaryotic community composition did not change substantially between visits at station M2, despite the high grazing activity. While a number of experimental studies carried out with freshwater communities demonstrate the influence of grazers on prokaryotic community composition (see review by Hahn and Höfle, 2001), experimental studies have revealed positive (Teira *et al.*, 2019) or no marked effects (Yokokawa and Nagata, 2005; Landa *et al.*, 2014; Baltar *et al.*, 2016) of the presence of grazers on marine prokaryotic diversity and composition. Our observations appear to be in line with these latter reports and suggest a minor influence of grazers on the temporal changes of the prokaryotic community composition in Kerguelen plateau waters.

#### 4.5. Conclusions

Our observations from late Austral summer add another piece to the pictures obtained during early spring and summer and thereby extend our previous conclusions on the pronounced

effect of natural Fe fertilization on prokaryotic community composition to the post-bloom period. The accumulation of DOC, as a consequence of the high seasonal productivity in Kerguelen plateau waters, together with the potential prokaryote-diatom interactions  
585 contribute to the build-up of prokaryote bulk biomass and to their taxonomic composition. The capabilities in the access to different forms of Fe, key for the processing of organic matter, vary among prokaryotic groups (Debeljak *et al.*, 2019) and likely play an additional role in shaping microbial communities. Our results suggest that the most abundant prokaryotic taxa identified as specific to the productive Kerguelen plateau surface waters in  
590 late summer contribute to the transfer of organic matter to heterotrophic nanoflagellates (Christaki *et al.*, 2020) and potentially higher trophic levels.

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## Figure Legends

**Figure 1.** Map of the MOBYDICK station locations. Surface chlorophyll concentrations are from Global Ocean Satellite Observations (Copernicus-GlobColour). Reprocessed data are  
795 provided by Copernicus Marine Service. Chlorophyll concentrations are the monthly mean for March 2018 at a resolution of 4 km. The black lines denote 1000 m bathymetry.

**Figure 2.** Vertical profiles of Chl *a*, DOC and prokaryotic abundance at the on-plateau station M2 and the off-plateau stations M3 and M4. Chl*a* and prokaryotic abundances are shown for each visit, DOC concentration is given as the mean of the repeated visits. Data for station M1  
800 are shown in Fig. S3.

**Figure 3.** Non-Metric Multidimensional Scaling (NMDS) of total (DNA) prokaryotic communities in the (a) free-living and (b) particle-attached fraction from all depth layers based on Bray-Curtis Dissimilarity. Sample depths are indicated by color, sampling sites by symbol and the number of visit is indicated with a number next to the respective symbol (see  
805 Table 1 for more information about sampling scheme.)

**Figure 4.** Relative abundance of total (DNA) free-living (FL) and particle-attached (PA) taxa grouped at order level in surface waters (10 m). For stations M2, M3 and M4 the relative abundances of the first visit are shown.

**Figure 5.** Relative abundance of ASVs that contribute significantly ( $p < 0.05$ ) to the  
810 dissimilarity between stations M2 and M3 (SIMPER analysis) at 10m. Asterisk highlight significant differences for a given ASV between sites, in either the free-living or particle-attached fractions, for the total (DNA) or potentially active (RNA) communities. Only the ASVs with relative abundance  $> 5\%$  in at least one of the samples are shown. Note that the

ASVs that contribute to the difference among sites are not always the same for the total and  
815 the active communities

**Figure 6.** Changes in the absolute abundance of dominant taxa (DNA, free-living fraction)  
between the first and the second (M2\_3) and the first and the third (M2\_3) visit at station M2  
at 10 m. Deviations are given as percent; positive and negative values indicate an increase and  
a decrease, respectively, in abundance over the repeated visits at station M2. The deviation  
820 was calculated based on a formula published in Agogué *et al.* (2011).

**Sequencing data.** Demultiplexed sequence files are available in NCBI under accession  
number PRJNA679029.

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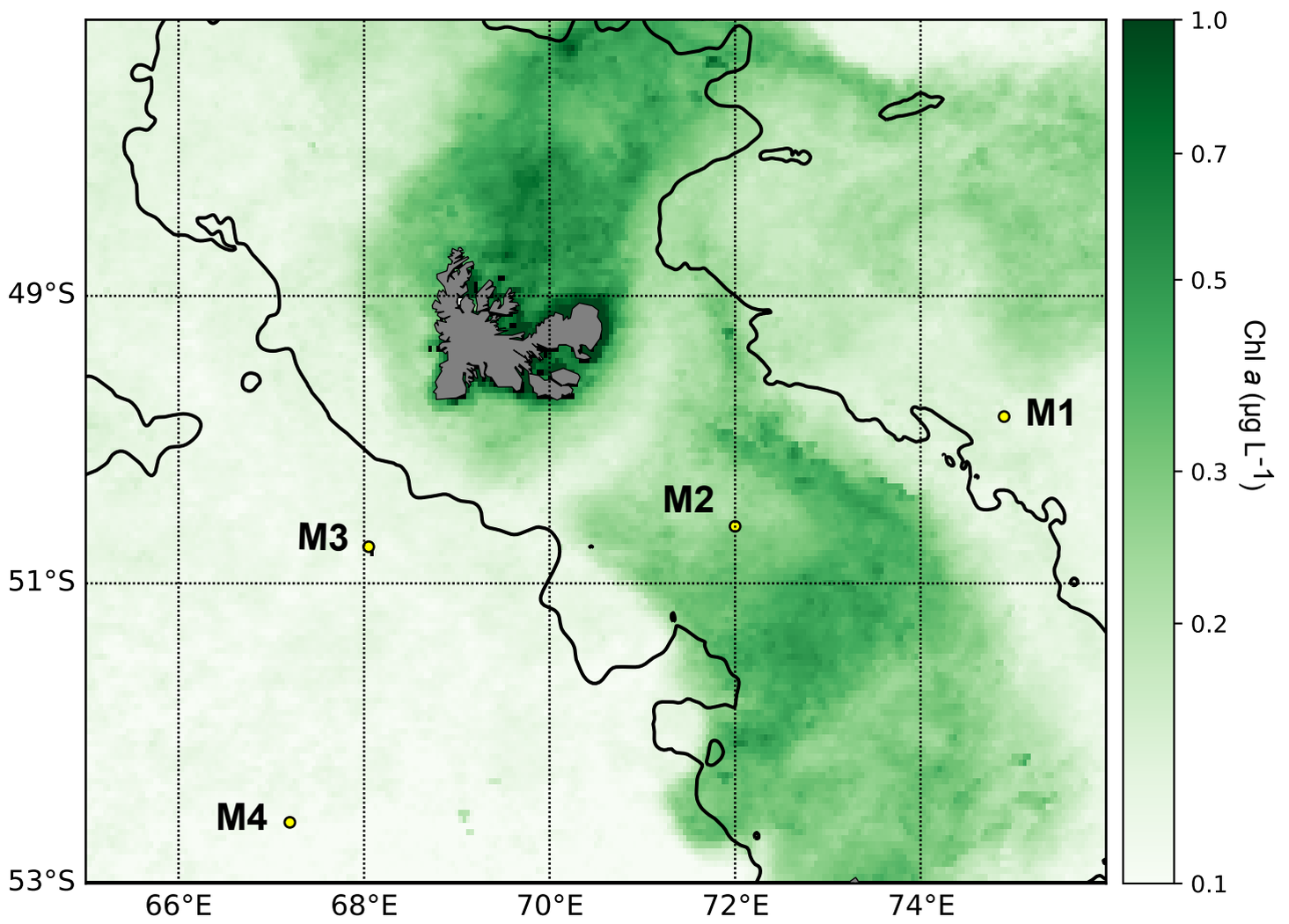


Fig. 1

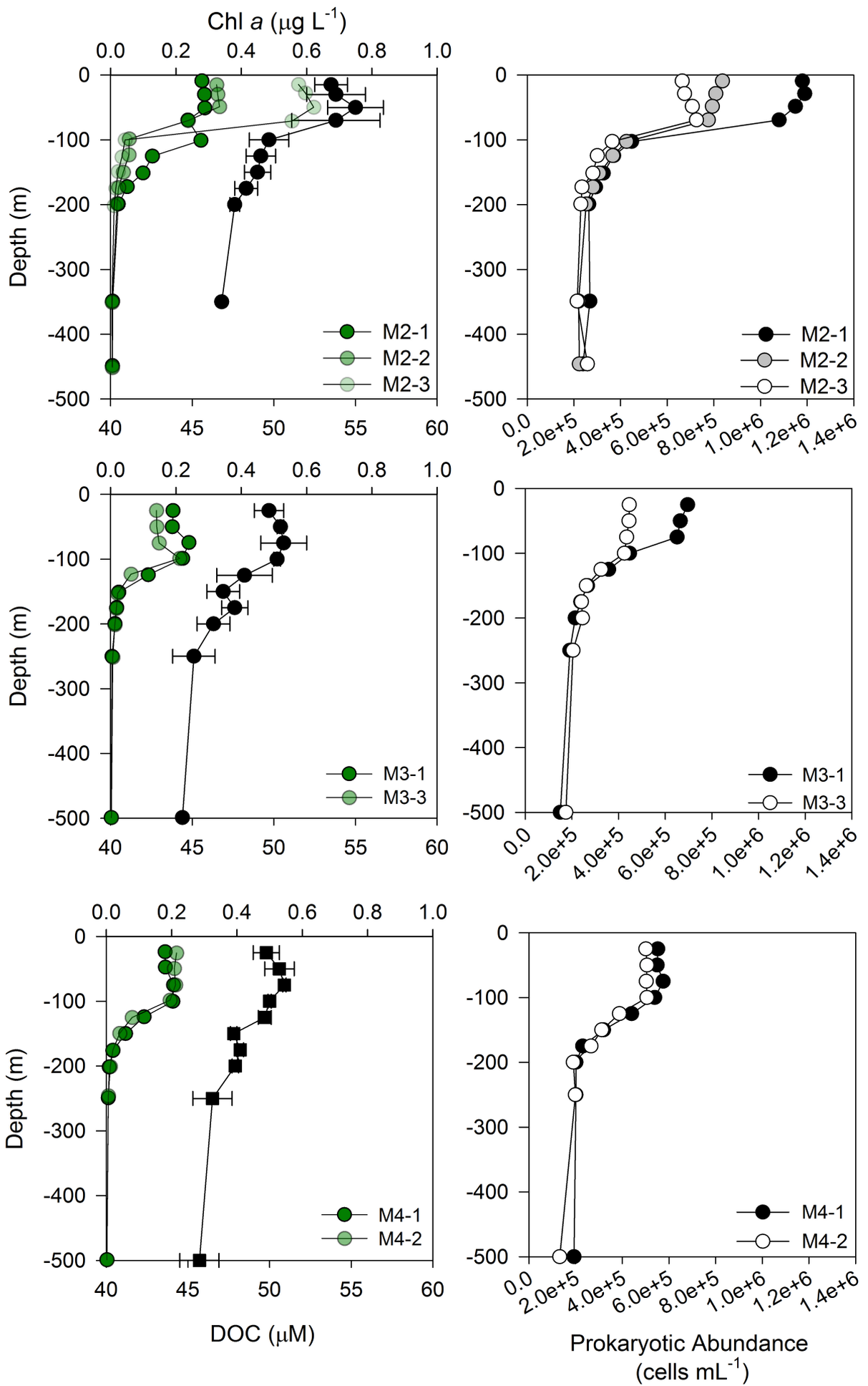


Fig. 2

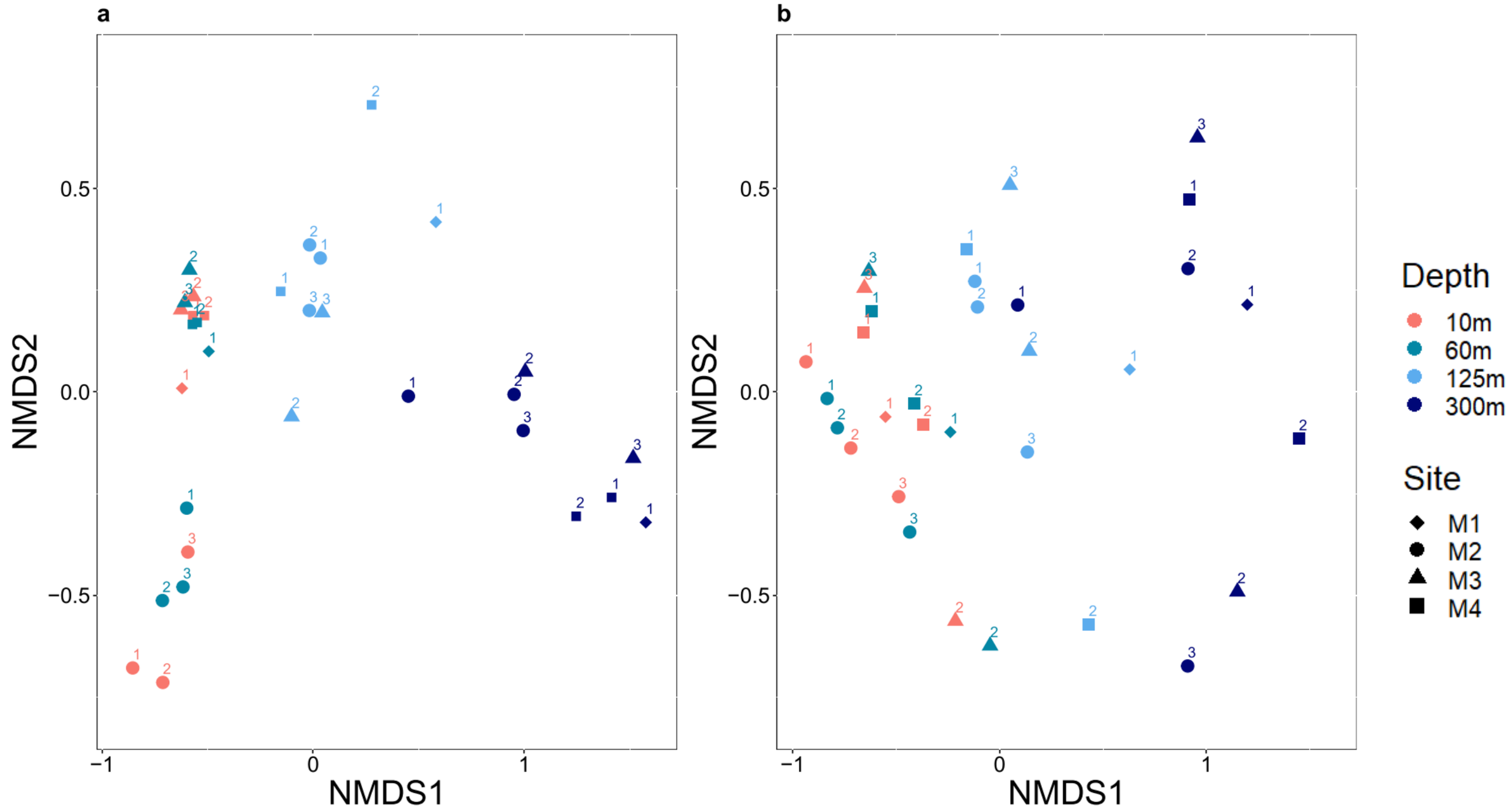


Fig. 3

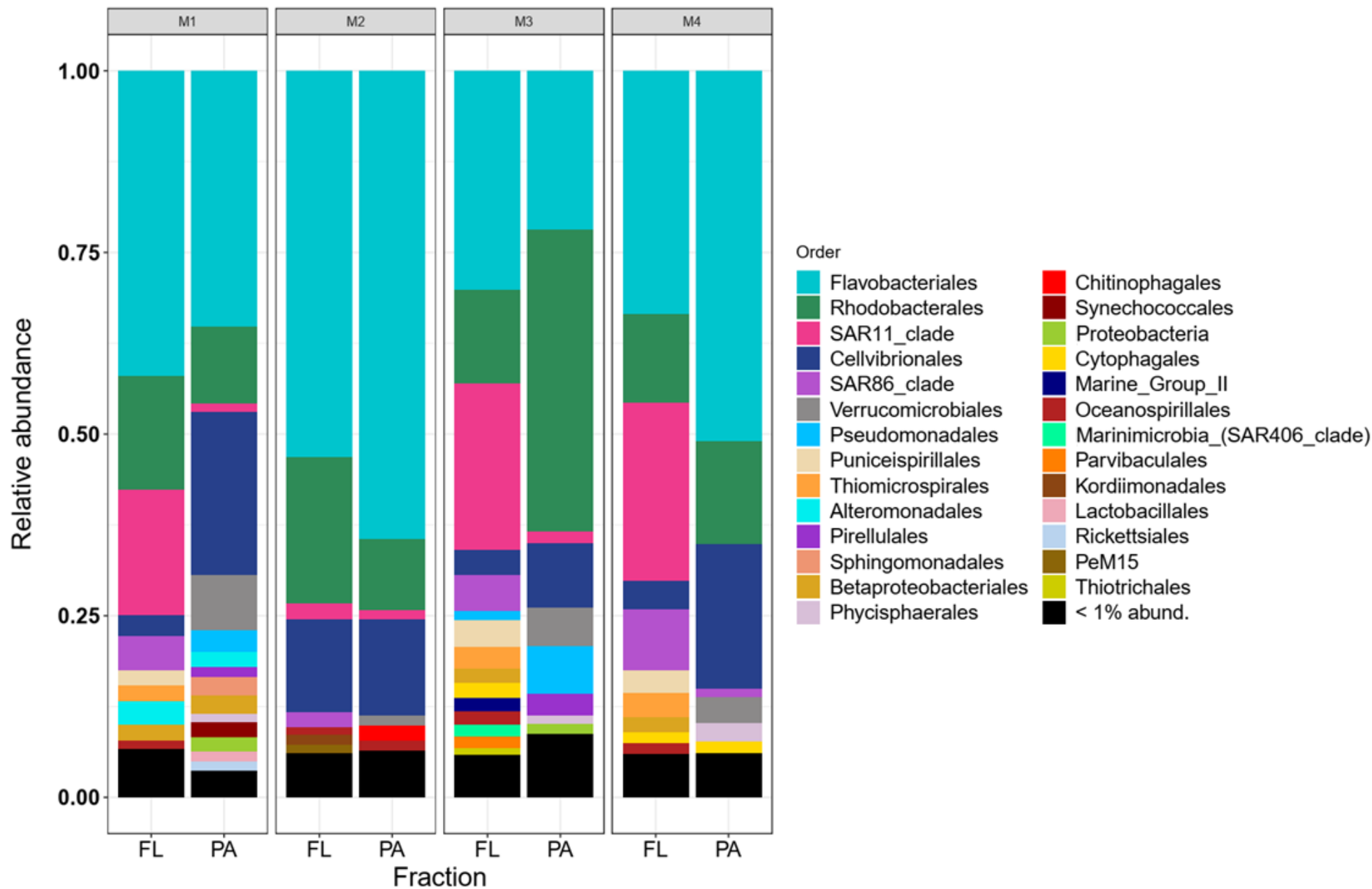


Fig. 4

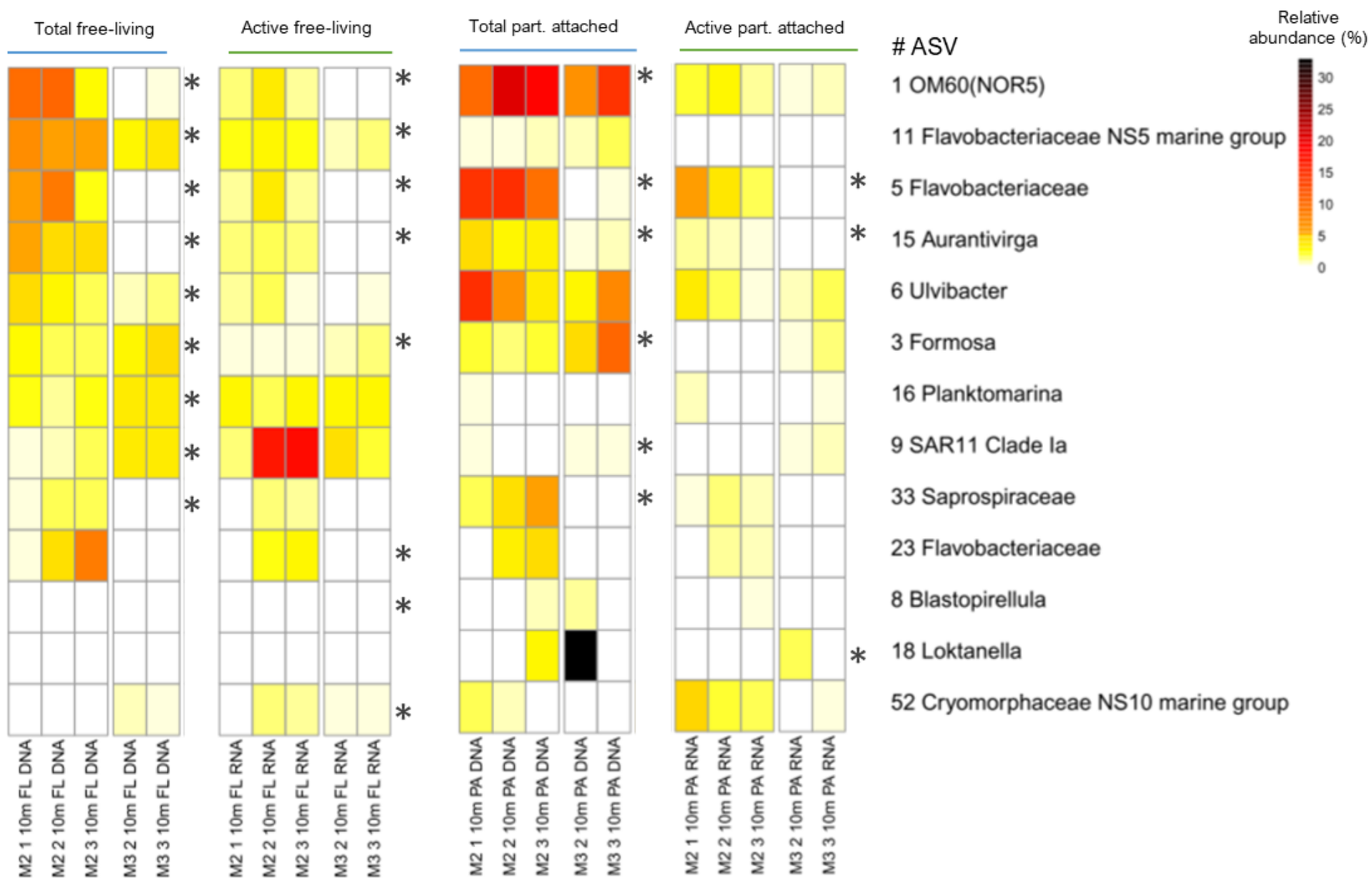


Fig. 5

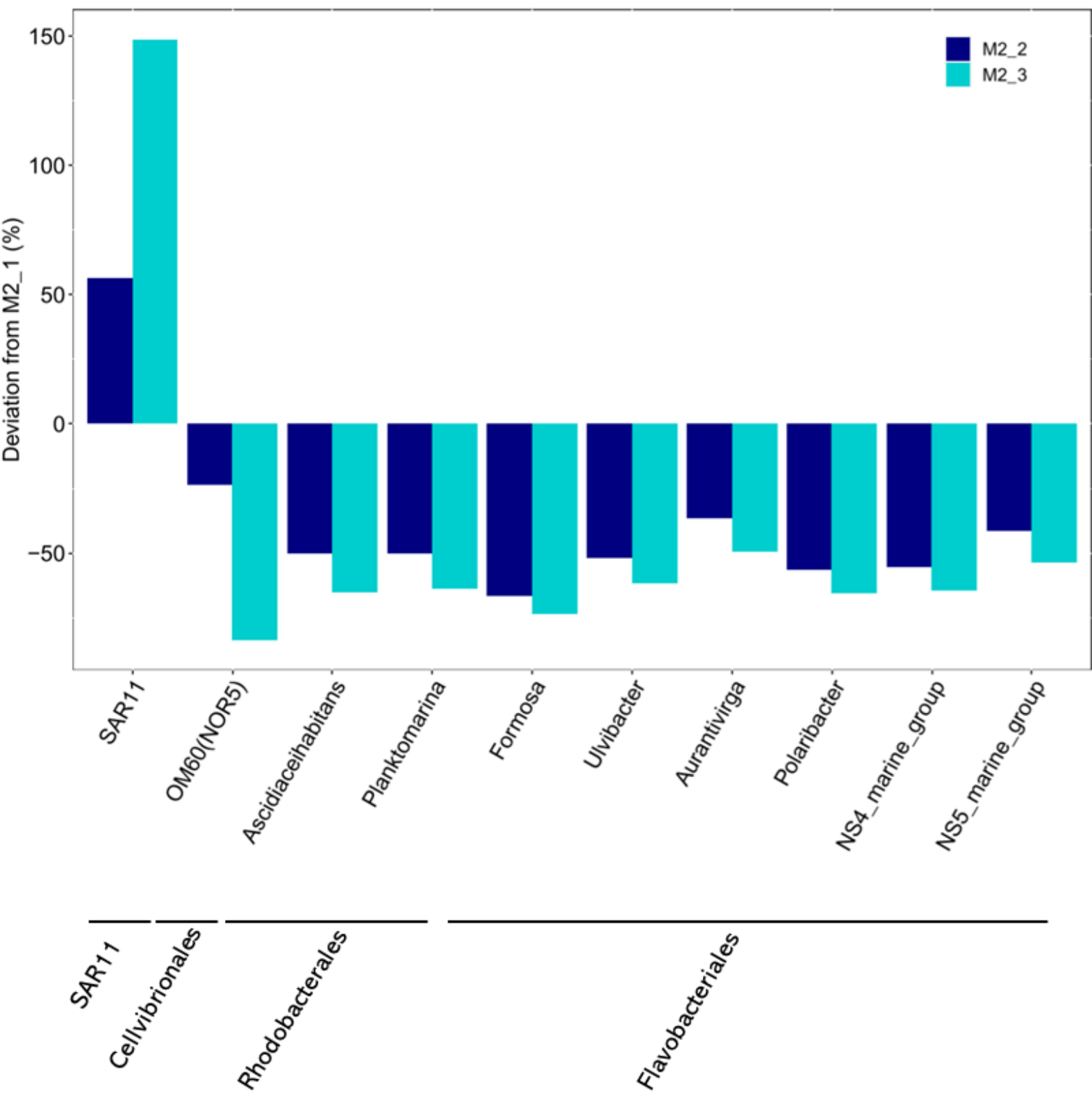


Fig. 6

Table 1. Brief description of the study sites. The wind-mixed surface layer ( $Z_{ML}$ ) is based on a difference in sigma of 0.02 to the surface value. Mean $\pm$ SD of the  $Z_{ML}$  are given

Station	Lat/Long	Depth (m)	Date	$Z_{ML}$ (m)	Temp $Z_{ML}$ ( $^{\circ}$ C)	Chl $a$ $Z_{ML}$ ( $\mu$ g L $^{-1}$ )	DOC $Z_{ML}$ ( $\mu$ M)
On plateau							
M2-1	50.61 $^{\circ}$ S 72.00 $^{\circ}$ E	527	26 Feb	62	5.1 $\pm$ 0.06	0.27 $\pm$ 0.02	52.8 $\pm$ 0.5
M2-2			06 Mar	61	5.2 $\pm$ 0.00	0.30 $\pm$ 0.04	55.7 $\pm$ 1.1
M2-3			16 Mar	68	5.1 $\pm$ 0.07	0.58 $\pm$ 0.02	53.8 $\pm$ 1.1
Off plateau							
M1	49.84 $^{\circ}$ S 74.90 $^{\circ}$ E	2723	09 March	27	4.9 $\pm$ 0.10	0.35 $\pm$ 0.04	50.3*
M3-1	50.68 $^{\circ}$ S	1700	04 Mar	65	5.6 $\pm$ 0.00	0.20 $\pm$ 0.02	50.3 $\pm$ 0.1
M3-3	68.05 $^{\circ}$ E		19 Mar	79	5.3 $\pm$ 0.02	0.14 $\pm$ 0.00	50.4 $\pm$ 1.3
M4-1	52.60 $^{\circ}$ S	4300	01 Mar	49	4.5 $\pm$ 0.06	0.18 $\pm$ 0.01	49.6 $\pm$ 0.5
M4-2	67.19 $^{\circ}$ E		12 Mar	87	4.5 $\pm$ 0.00	0.21 $\pm$ 0.00	50.8 $\pm$ 0.4

\*only one data point in the  $Z_{ML}$  available

Station M3-2 was not sampled for the parameters considered in the present study

Table 2. Prokaryotic diversity as illustrated by the Shannon Index. Results for the first visit to the off-plateau stations are shown. A full description of the diversity indices is provided in Table S1. NA – Not Available

Depth (m)	Total (DNA)		Active (RNA)	
	Free-living	Particle-attached	Free-living	Particle-attached
On Plateau				
Station M2-1				
10	4.05	3.67	6.04	5.79
50	4.52	3.59	NA	5.95
100	4.97	3.96	6.37	6.19
300	5.32	4.61	5.97	6.45
Station M2-2				
10	4.20	3.69	4.59	5.89
60	4.32	3.41	6.04	4.51
125	4.92	4.12	5.21	5.03
300	5.39	5.69	5.99	6.34
Station M2-3				
10	4.56	3.65	4.79	6.37
60	4.38	3.54	4.94	5.85
125	5.04	3.80	6.56	6.22
300	5.37	3.99	6.66	4.90
Off plateau				
Station M1				
10	4.63	4.01	4.39	6.22
60	4.91	3.91	6.26	6.05
125	5.02	5.07	6.55	6.66
300	5.17	6.01	5.64	6.42
Station M3-1				
10	5.01	3.52	6.15	6.04
60	5.12	3.29	6.44	5.91
125	5.00	3.66	4.86	6.20
300	5.57	4.77	6.37	5.76
Station M4-1				
10	4.78	3.72	5.65	6.04
60	4.76	3.88	5.94	6.21
125	5.10	4.40	6.52	6.56
300	5.45	5.65	6.01	6.29

## Supplementary Material

for the manuscript

Prokaryotic diversity and activity in contrasting productivity regimes in late summer in the  
Kerguelen region (Southern Ocean)

Alejandra Elisa Hernandez-Magana<sup>1,2\*</sup>, Yan Liu<sup>1,3\*</sup>, Pavla Debeljak<sup>1,4†</sup>, Olivier Crispi<sup>1</sup>,  
Barbara Marie<sup>1</sup>, Coco Koedooder<sup>1‡</sup>, Ingrid Obernosterer<sup>1#</sup>

This file contains

10 Suppl. Figures

4 Suppl. Tables

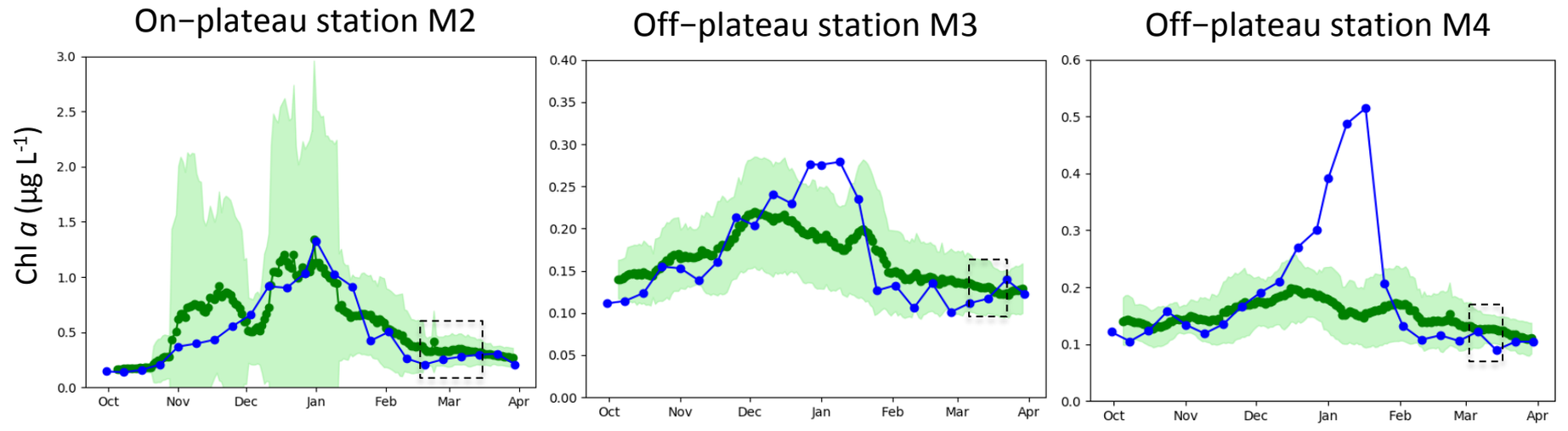
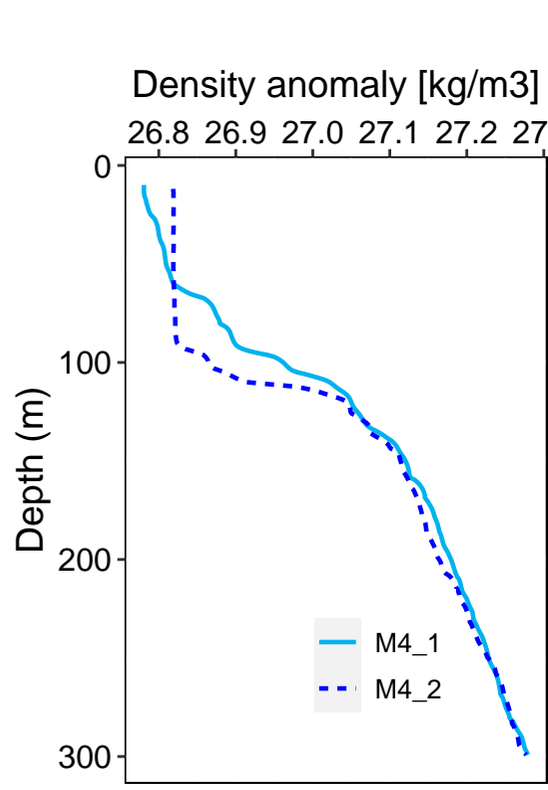
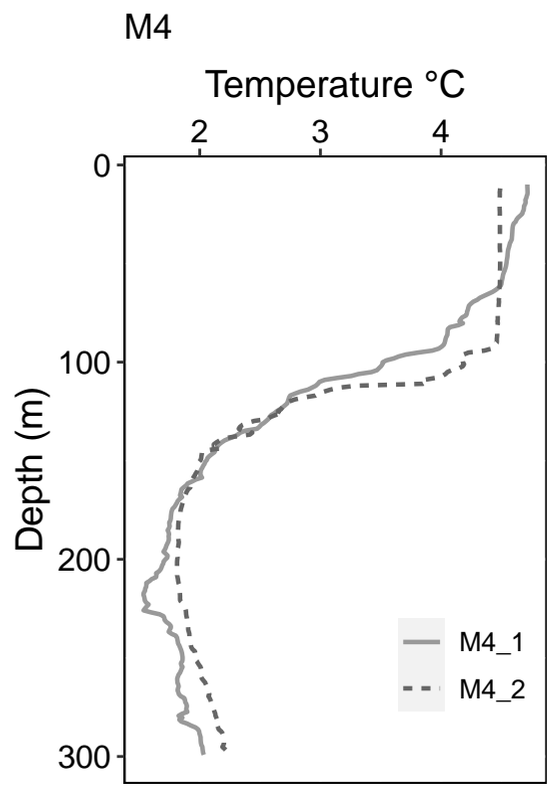
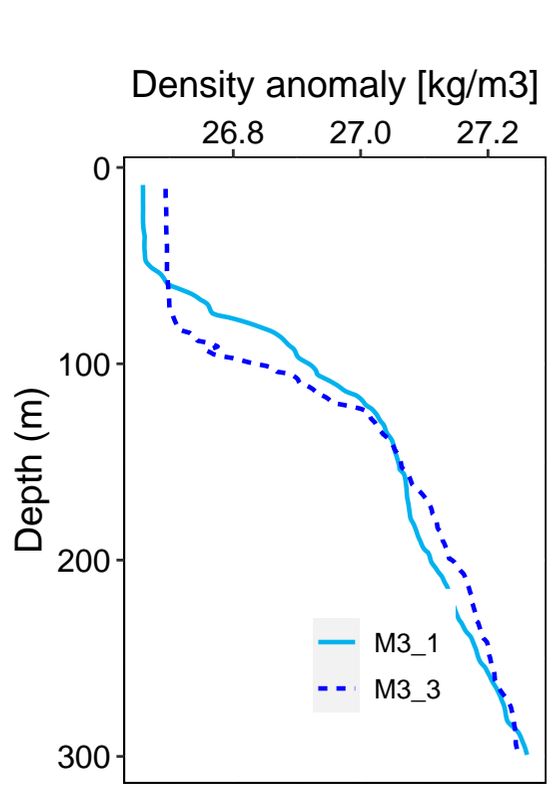
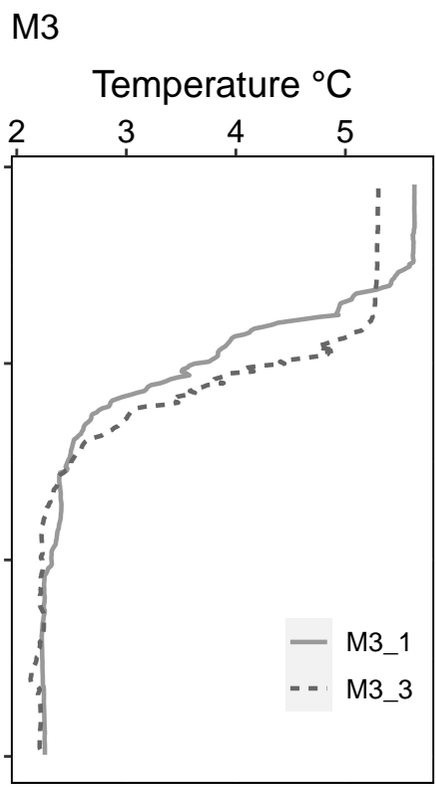
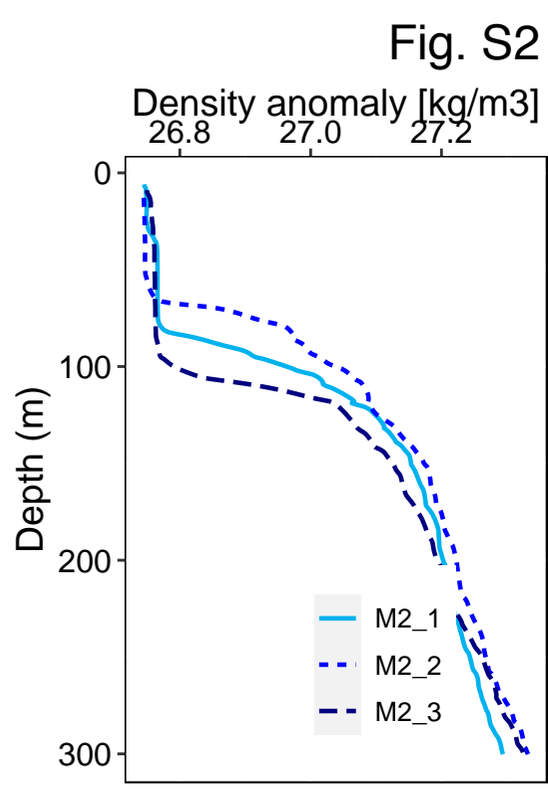
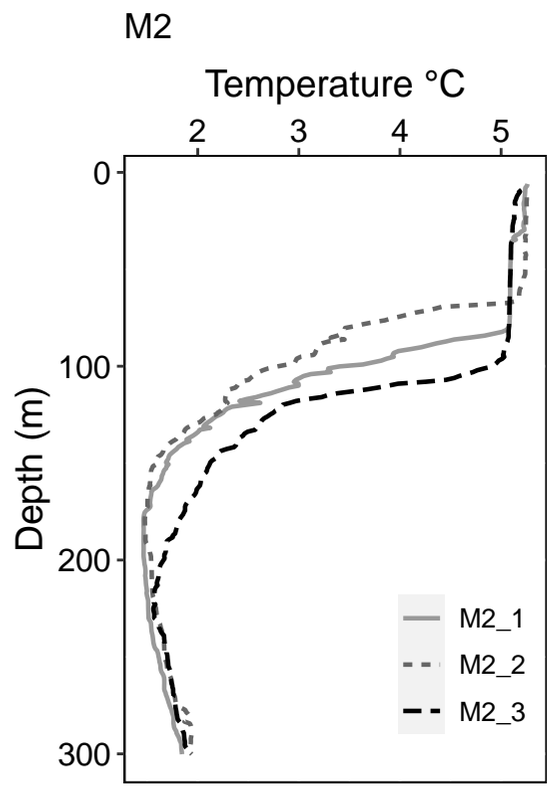
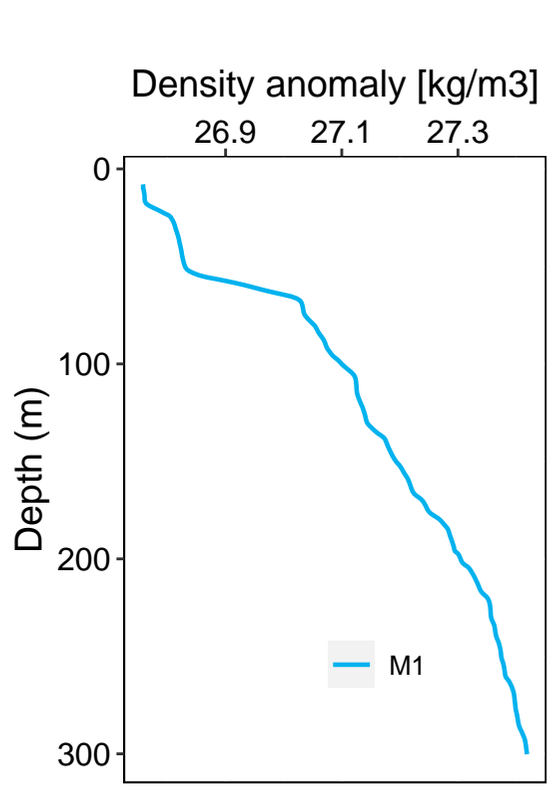


Fig. S1



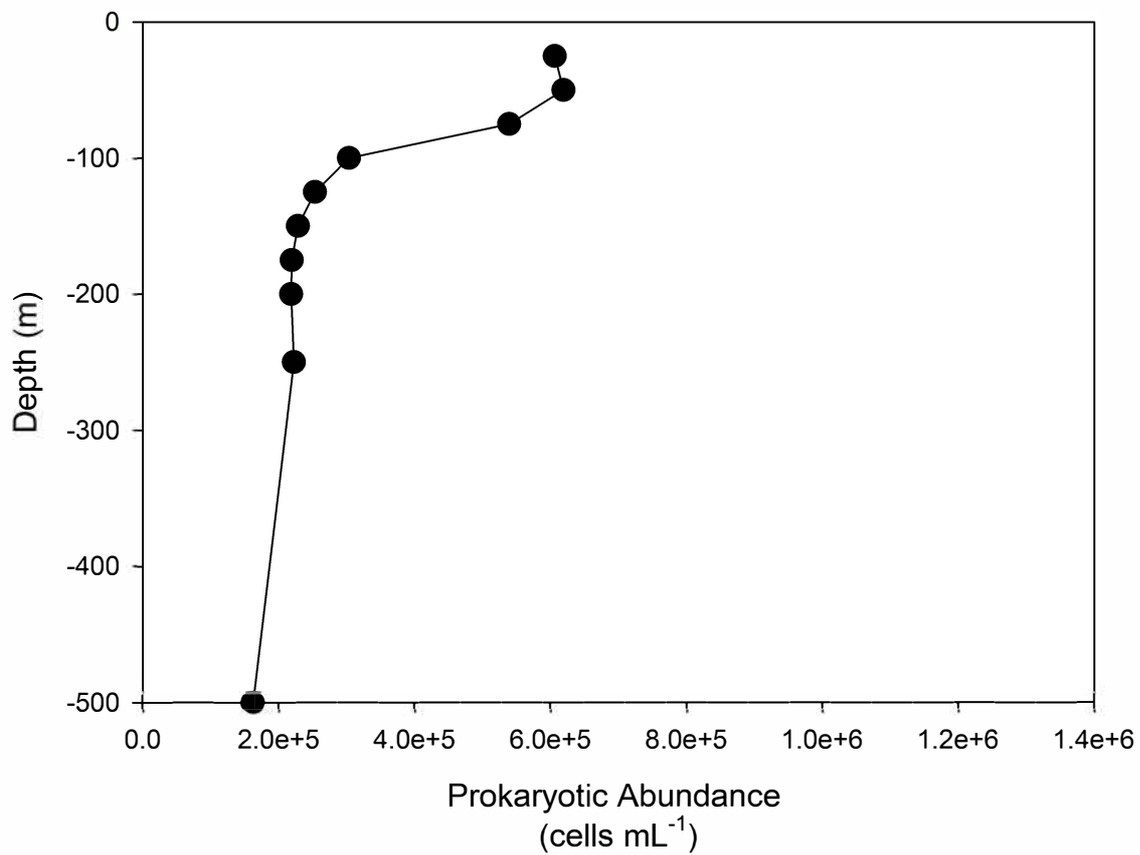
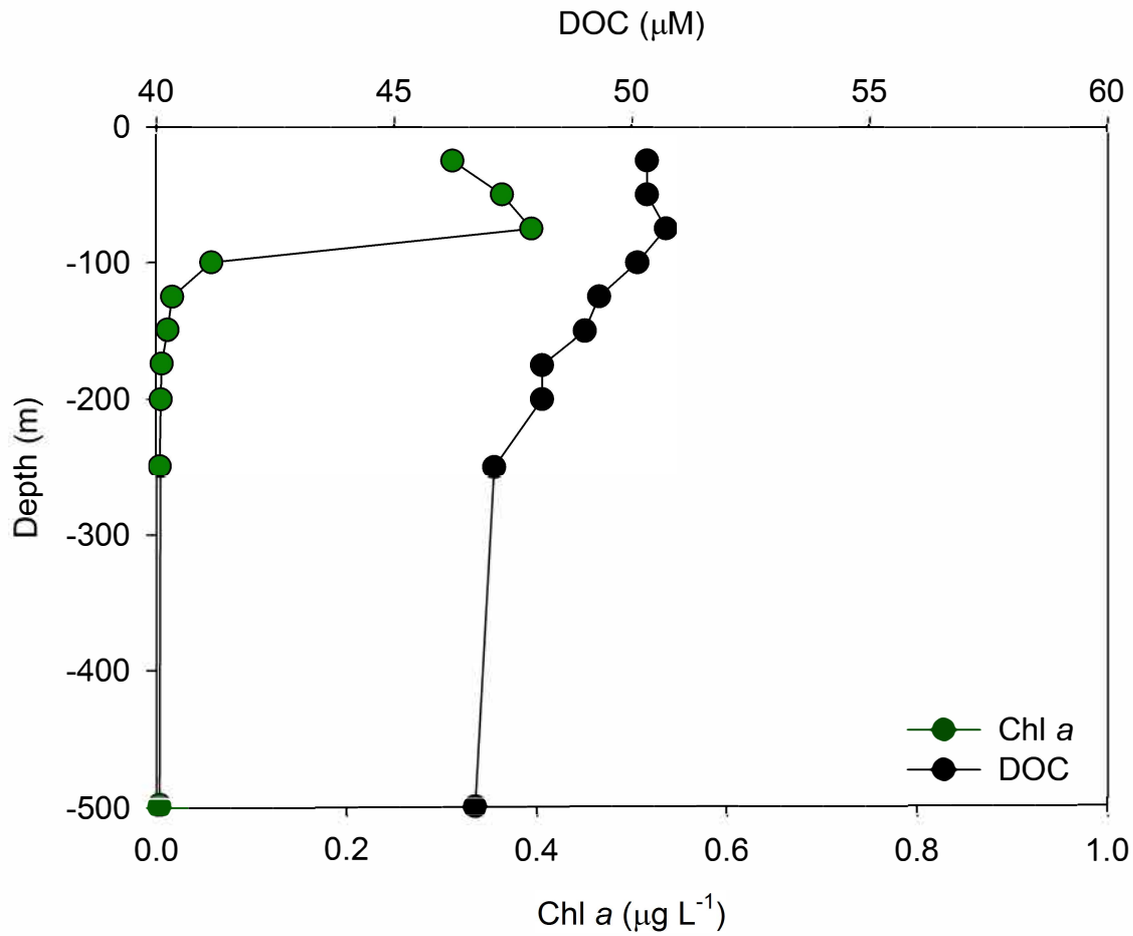


Fig. S3

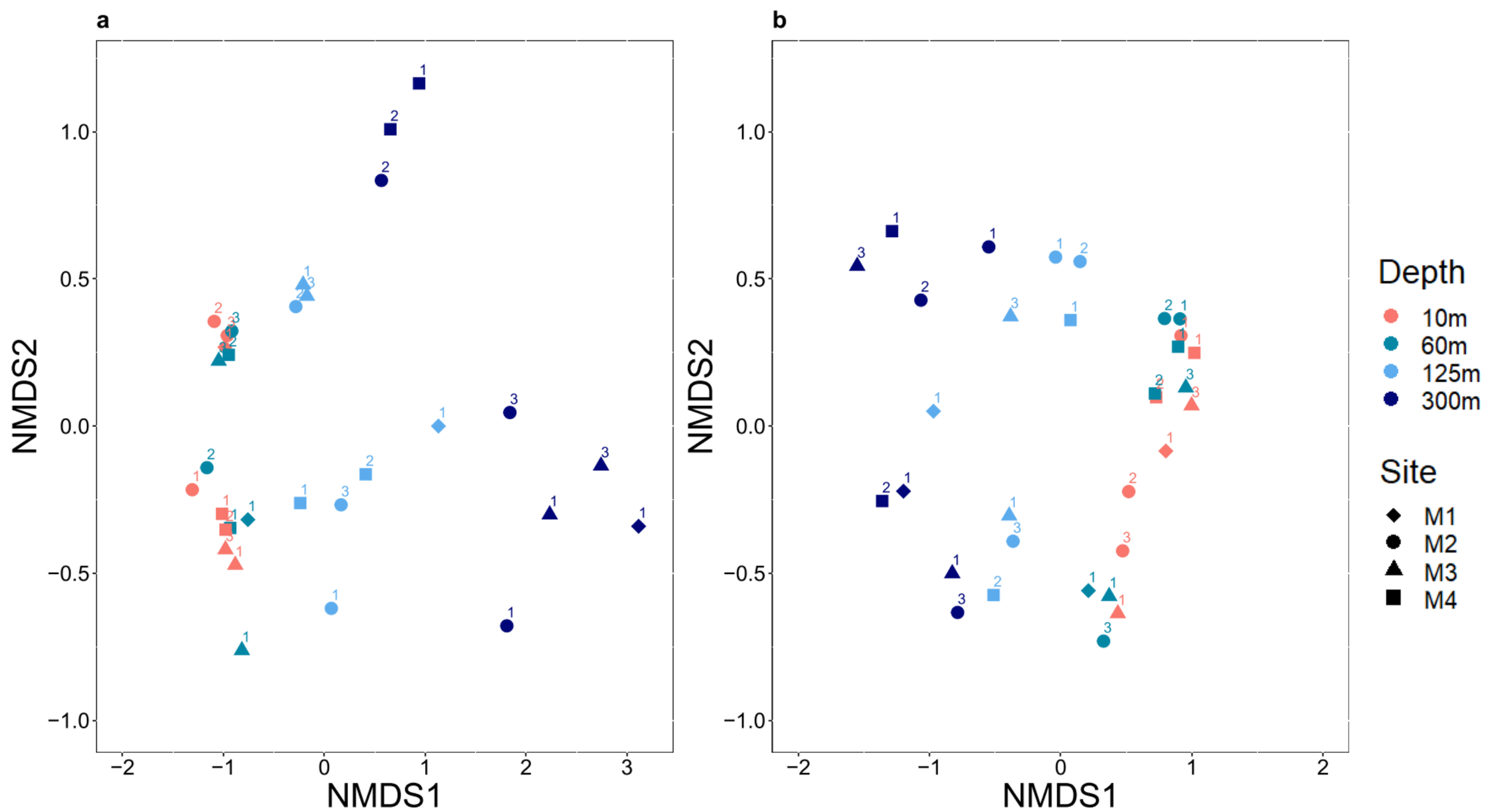


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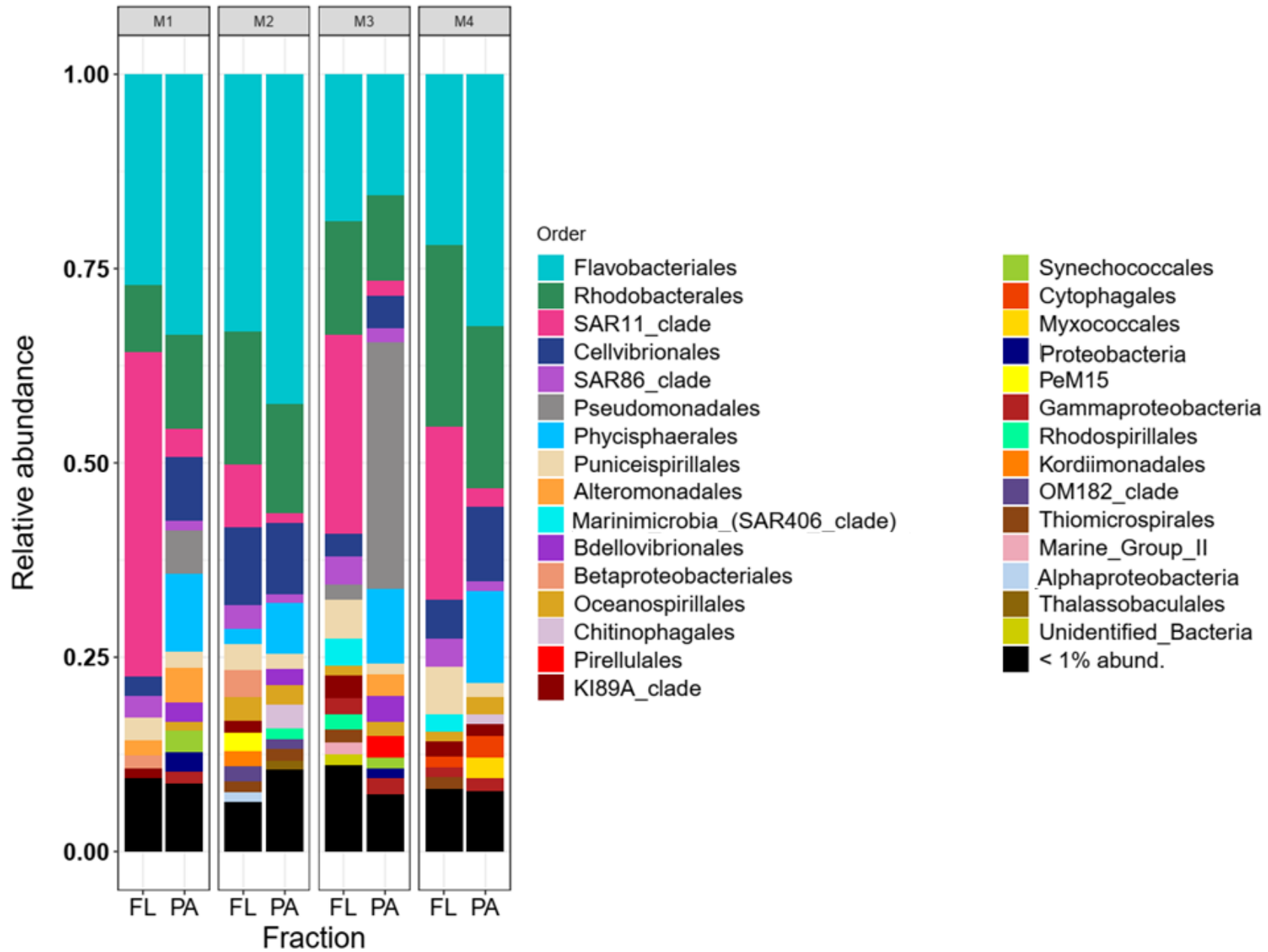


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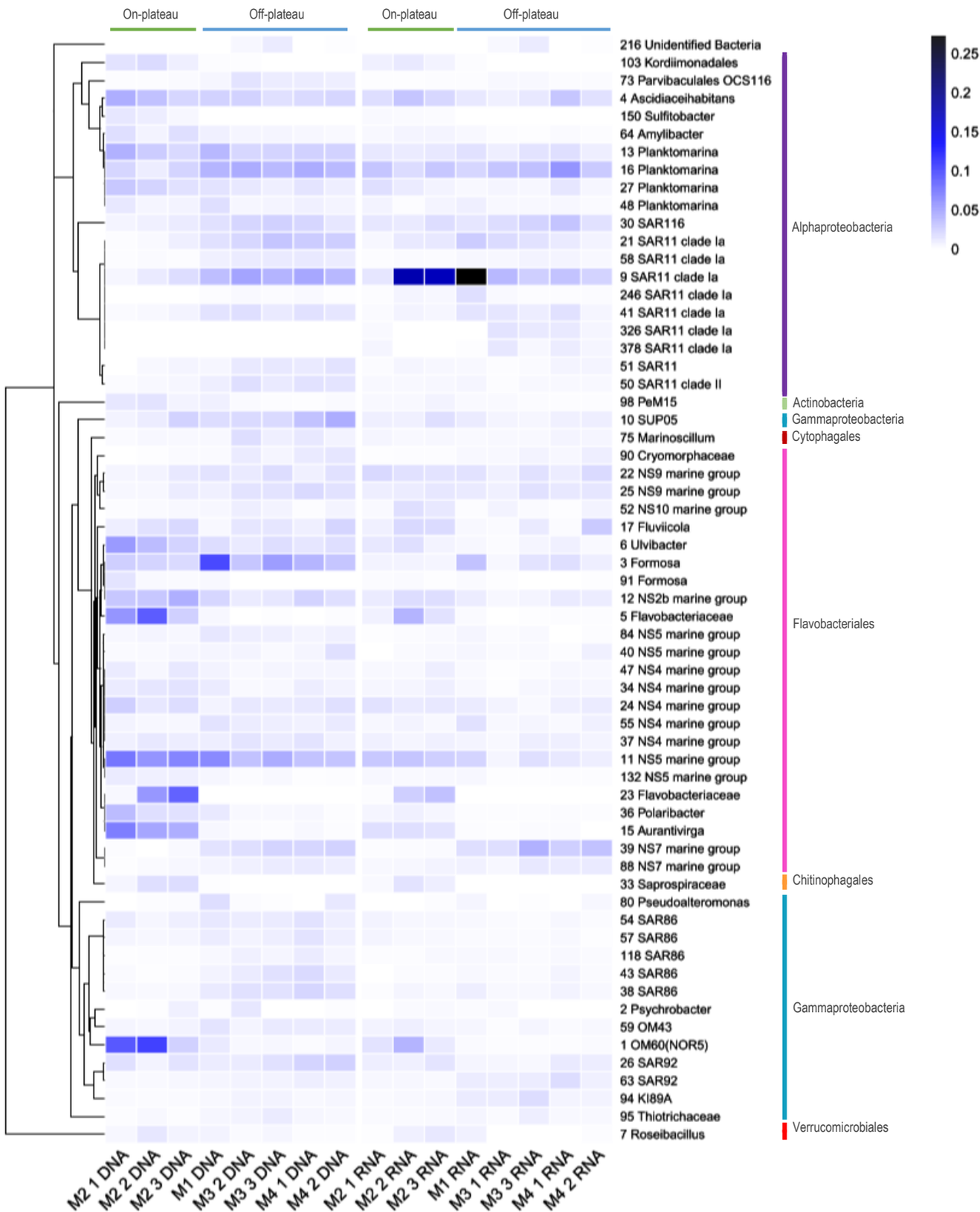


Fig. S6





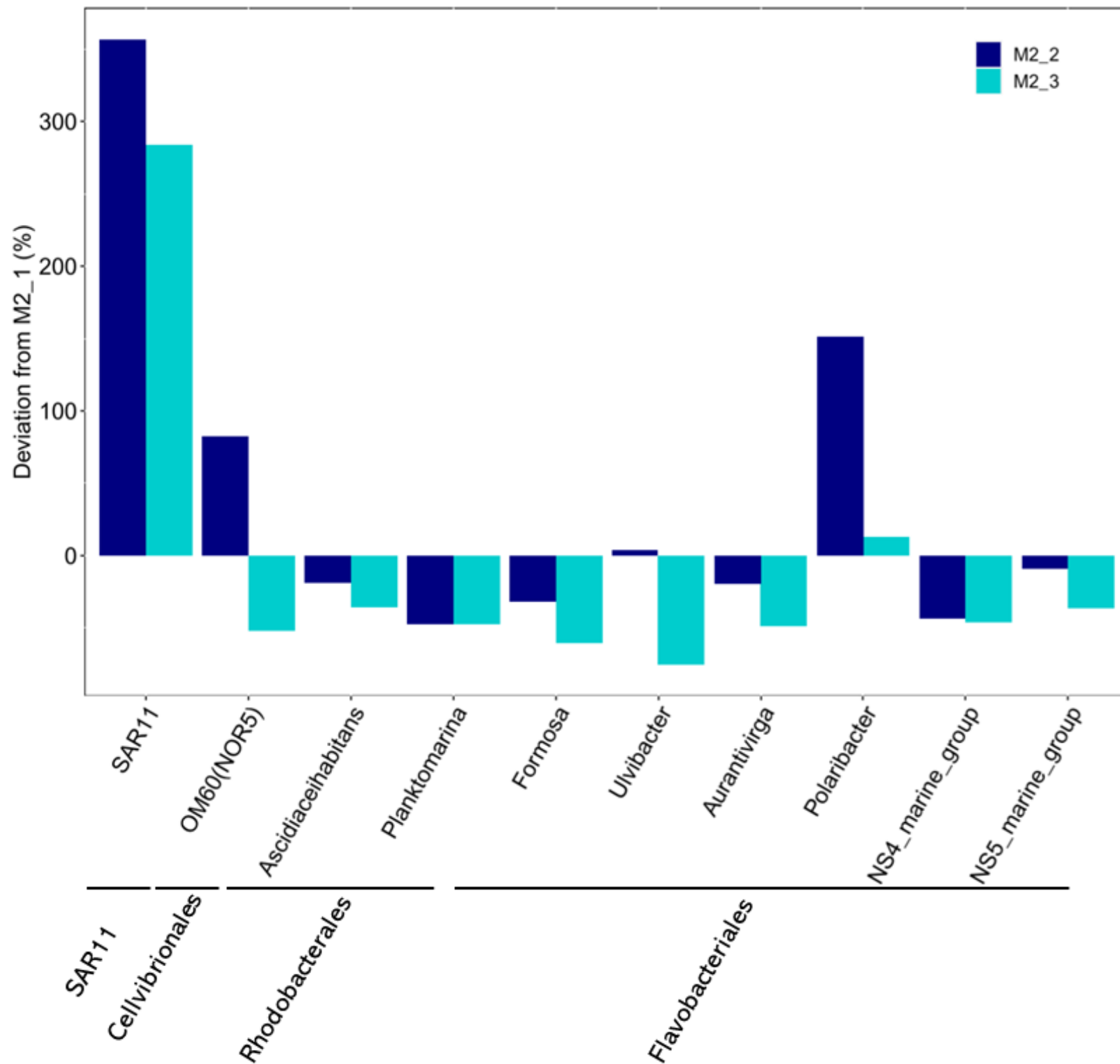


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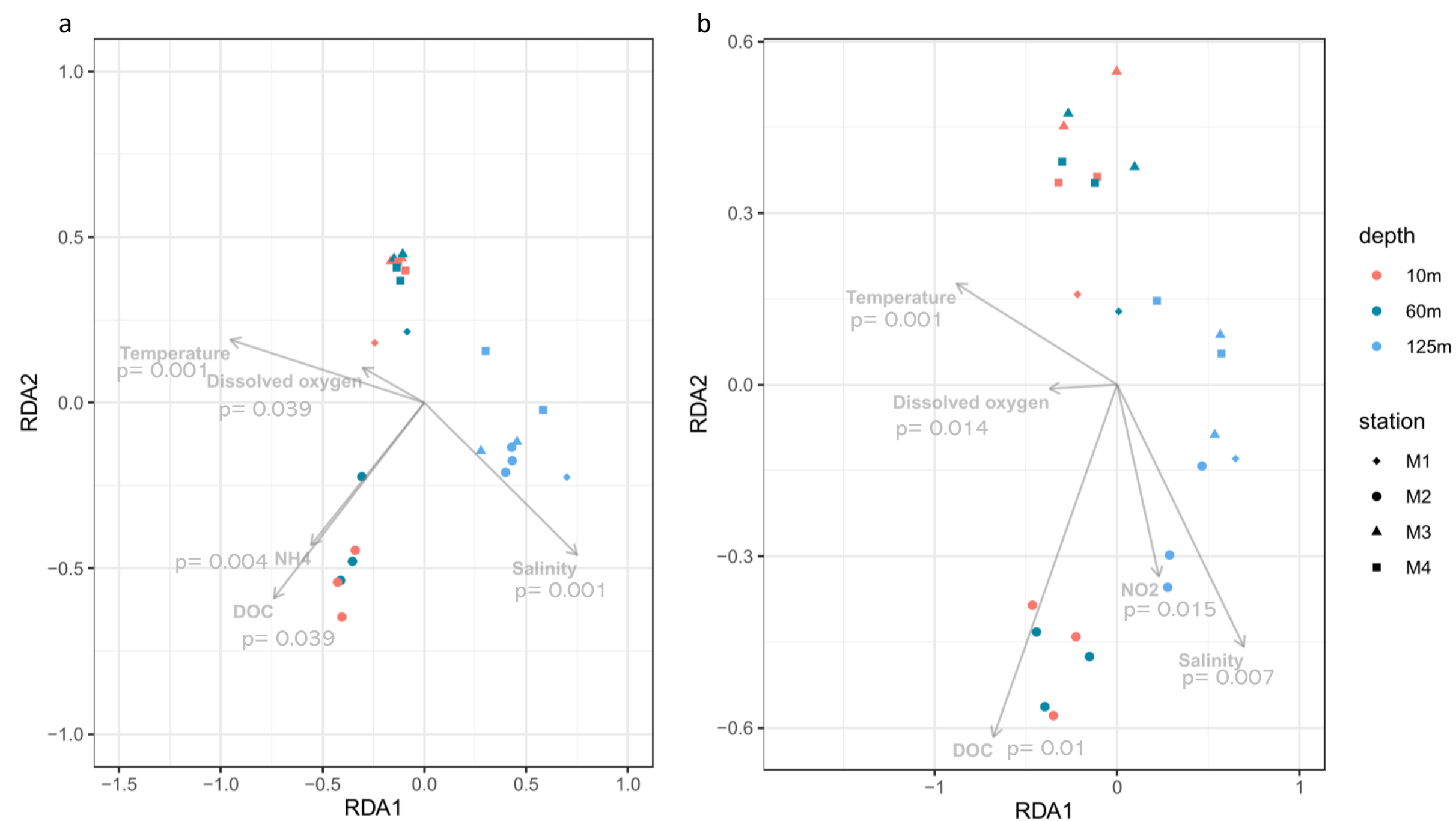


Fig. S10

## Supplementary Figures

**Figure S1.** Seasonal bloom dynamics for on-plateau station M2 and off-plateau stations M3 and M4. The green line indicates the 10-year climatological mean (2007-2017) of Chl *a* concentrations and its SD illustrated as the green shaded area. The blue line denotes Chl *a* concentrations for the year 2018 (MOBYDICK cruise). Chl *a* data are from Global Ocean Satellite Observations (Copernicus-GlobColour). Reprocessed data are provided by Copernicus Marine Service. The time period of the MOBYDICK cruise is marked with a rectangle. Note different scales of y-axis.

**Figure S2.** Depth profiles of temperature and density anomaly for the repeated visits at the 4 stations.

**Figure S3.** Depth profiles of DOC, prokaryotic abundances and Chl *a* at station M1.

**Figure S4.** Non-Metric Multidimensional Scaling (NMDS) of potentially active (RNA) prokaryotic communities in the (a) free-living and (b) particle-attached fraction from all depth layers based on Bray-Curtis Dissimilarity.

**Figure S5.** Relative abundance of potentially active (RNA) free-living (FL) and particle-attached (PA) taxa grouped at order level in surface waters (10 m). For stations M2, M3 and M4 the mean relative abundances of the repeated visits are shown.

**Fig. S6.** Relative abundance of ASVs with corresponding phylum (right colored sidebar) in the free-living fraction in surface waters (10 m). Only ASVs with >1% relative abundance in at least one sample are shown. On-plateau in green; off-plateau in blue. DNA: total community, RNA: potentially active community.

**Fig. S7.** Relative abundance of ASVs with corresponding phylum (right colored sidebar) in the particle-attached fraction in surface waters (10 m). Only ASVs with >1% relative abundance in at least one sample are shown. On-plateau in green; off-plateau in blue. DNA: total community, RNA: potentially active community.

**Fig. S8.** Relative abundance of ASVs that contribute significantly ( $p < 0.05$ ) to the dissimilarity between stations M2 and M3 (SIMPER analysis) at 60m. Asterisk highlight significant differences for a given ASV between sites, in either the free-living or particle-attached fractions, for the total (DNA) or potentially active (RNA) communities. Only the ASVs with relative abundance >5% in at least one of the samples are shown. Note that the ASVs that contribute to the difference among sites are not always the same for the total and the active communities

**Fig. S9.** Changes in the absolute abundance of dominant taxa (RNA, free-living fraction) between the first and the second (M2\_3) and the first and the third (M2\_3) visit at station M2 at 10 m. Deviations are given as percent; positive and negative values indicate an increase and a decrease, respectively, in abundance over the repeated visits at station M2. The deviation was calculated based on a formula published in Agogué et al (2011).

**Fig. S10.** Redundancy analysis of the total community (DNA-based dataset) in the free-living fraction (a) and particle attached (b). The significant environmental response-vectors are shown. Colors indicate water depths and symbol shapes correspond to sites. The variables were selected prior the RDA analysis (with variance inflation <10) excluding the redundant constraints.

Table S1. Percent prokaryotic cells in the 0.8  $\mu\text{m}$  filtered seawater of cells in unfiltered seawater.

Station	Depth (m)	Fraction of cells in < 0.8 $\mu\text{m}$ size fraction (%)
M1	15	115
M2-1	10	84
M2-2	10	93
M2-2	60	82
M2-3	60	79
M3-1	10	91
M3-1	60	83
M3-3	10	86
M3-3	60	83
M4-2	15	96

Table S2. Prokaryotic diversity as illustrated by the Observed, Shannon and Inverse Simpson Index for all depths and repeated visits at the 4 sites. NA – Not Available

Station	M1				M2_1				M2_2				M2_3				
Fraction	free-living DNA																
Depth (m)	10	60	125	300	10	60	125	300	15	60	125	300	10	60	125	300	15
Observed	357	448	588	604	256	329	480	590	293	310	469	675	347	297	515	637	439
Shannon	4,630417587	4,911170603	5,016702863	5,1742284	4,050761314	4,519927552	4,973573443	5,324898909	4,199674721	4,321745361	4,923799894	5,39013288	4,557528066	4,376234884	5,036416351	5,372785124	5,005918554
InvSimpson	42,82129835	58,02194061	31,7209722	52,58309965	28,9092411	46,7586488	46,12942033	72,23921259	29,21192078	36,99498058	41,97304834	64,65604812	43,44739127	35,8288635	45,03269349	72,06032229	74,84599849
Station	M1				M2_1				M2_2				M2_3				
Fraction	attached DNA																
Depth (m)	10	60	125	300	10	60	125	300	15	60	125	300	10	60	125	300	15
Observed	282	283	545	866	259	231	290	439	273	231	334	704	347	297	515	637	439
Shannon	4,013059399	3,913617347	5,071673974	6,012926553	3,671495301	3,588997236	3,963424462	4,611143781	3,685029331	3,40815961	4,123871042	5,691295054	3,6529	3,538	3,796	3,9944	3,518
InvSimpson	18,41429539	19,21990533	47,68899127	178,9420594	14,55652021	13,95914847	20,17997005	32,94964122	12,69473407	10,4241389	20,2013473	118,7923767	14,21149175	10,4241389	20,2013473	118,7923767	8,177114762
Station	M1				M2_1				M2_2				M2_3				
Fraction	free-living RNA																
Depth (m)	10	60	125	300	10	60	125	300	15	60	125	300	10	60	125	300	15
Observed	510	1107	1279	857	930	NA	1066	1031	467	953	680	948	513	569	1274	1346	1015
Shannon	4,391351947	6,26176654	6,553091444	5,643766982	6,043665186	NA	6,373004977	5,967207732	4,587999483	6,038872206	5,208954219	5,991221674	4,794227976	4,936724393	6,56084046	6,661230197	6,147218663
InvSimpson	12,47474901	199,7510742	236,2879011	31,81886256	170,37426	NA	284,3064592	40,76355835	23,92997487	161,2016865	41,25067073	111,66426	27,0774122	28,72407112	256,0138486	202,2292605	171,5273223
Station	M1				M2_1				M2_2				M2_3				
Fraction	attached RNA																
Depth (m)	10	60	125	300	10	60	125	300	15	60	125	300	10	60	125	300	15
Observed	896	990	961	1152	896	990	961	1152	866	363	443	1188	1023	860	1053	795	995
Shannon	5,79465954	5,95422116	6,189837213	6,454896518	5,79465954	5,95422116	6,189837213	6,454896518	5,886758741	4,507552806	5,030836167	6,340782925	6,373348086	5,849730962	6,215210067	4,897418311	6,044759366
InvSimpson	85,19544355	93,97085242	203,6656298	281,132465	85,19544355	93,97085242	203,6656298	281,132465	131,207044	39,15112692	69,59025164	86,48967977	298,7720653	82,54598082	157,5615963	11,48080568	105,1908396

<b>M3_1</b>				<b>M3_2</b>				<b>M4_1</b>				<b>M4_2</b>			
free-living DNA				free-living DNA				free-living DNA				free-living DNA			
60	125	300	15	60	125	300	10	60	125	300	15	60	125	300	
457	438	774	416	396	519	659	382	388	495	684	412	419	618	694	
5,119911706	5,003944767	5,569760803	4,886634643	4,766888034	5,033568653	5,378535363	4,769761755	4,761656847	5,104457433	5,448336438	4,902835399	4,892809069	4,943056375	5,444155036	
86,28261896	72,41899328	79,1641964	65,26185117	59,56053611	47,57175899	66,88524437	61,40002813	61,24781921	69,85134114	63,62046237	69,68530427	70,01692317	27,79842907	65,40092231	
<b>M3_1</b>				<b>M3_2</b>				<b>M4_1</b>				<b>M4_2</b>			
attached DNA				attached DNA				attached DNA				attached DNA			
60	125	300	15	60	125	300	10	60	125	300	15	60	125	300	
457	438	774	288	298	315	771	243	267	366	669	275	306	267	707	
3,286	3,6634	4,77	3,997718952	4,08066732	3,954872015	5,750910961	3,717988494	3,884385475	4,404890291	5,647105594	3,969754376	4,070741559	2,928439517	5,557859019	
7,240784755	13,19206106	16,93033334	20,40416923	22,46182369	18,90960618	129,6652914	16,20826173	19,36629488	29,77360972	125,7471711	19,14907738	22,8616917	4,539264699	73,51675384	
<b>M3_1</b>				<b>M3_2</b>				<b>M4_1</b>				<b>M4_2</b>			
free-living RNA				free-living RNA				free-living RNA				free-living RNA			
60	125	300	15	60	125	300	10	60	125	300	15	60	125	300	
1103	680	1211	897	523	703	1033	816	945	1213	946	1027	572	1351	937	
6,443330058	4,863603132	6,369838161	5,872947044	4,541682703	5,04721143	6,157978297	5,645590785	5,936458808	6,5245001	6,0079269	6,114155036	4,721951749	6,647294232	6,070729353	
280,3100004	18,29747807	82,2504423	127,1842817	15,83392416	26,54139064	78,5280407	94,82600929	136,7852678	300,0499905	102,6859357	161,4269072	18,65007181	273,5372002	151,9662869	
<b>M3_1</b>				<b>M3_2</b>				<b>M4_1</b>				<b>M4_2</b>			
attached RNA				attached RNA				attached RNA				attached RNA			
60	125	300	15	60	125	300	10	60	125	300	15	60	125	300	
947	996	878	1088	1089	1116	1165	943	1038	1270	1098	1093	383	929	1243	
5,914238798	6,20376749	5,761394155	6,281651595	6,284950066	6,390307622	6,365012264	6,038599251	6,21247134	6,563289489	6,293072004	6,295741049	4,636097191	5,793395001	6,469338878	
81,74213719	141,1394363	75,05190811	191,8025779	173,4126956	181,1869336	103,1794949	130,9402697	177,3185809	247,1549593	95,5618546	196,6217623	40,27047017	67,49777923	159,5258776	

Table S3. Partial Mantel tests for total (DNA) and potentially active (RNA) prokaryotic community composition, diatom community composition and environmental parameters. Rho values are shown. \*\*  $p < 0.01$ , \*  $p < 0.05$ . Diatom community composition was determined by microscopic observations and the results are presented in Lafond et al. (2020)

	Prokaryotic Community Composition			
	Total		Active	
	Free-living	Particle-attached	Free-living	Particle-attached
Diatom community composition	0.6006 **	0.4498**	0.5257**	0.3682**
Combined environmental parameters	0.3525**	0.3003*	0.3969**	0.3126**
Temperature	0.2559**	0.3627**	0.5493**	0.3558**
Salinity	0.2553*	0.315**	0.4528**	0.2677**
Dissolved oxygen	0.2783*	0.257	0.3782*	0.2457*
NH <sub>4</sub> <sup>+</sup>	0.3668**	0.1354	0.1262	0.1222
PO <sub>4</sub> <sup>3-</sup>	0.08138	0.0796	0.114	0.1021
Si(OH) <sub>4</sub>	0.4228**	0.4413**	0.6532**	0.4747**
NO <sub>2</sub> <sup>-</sup>	0.1993	0.2615*	0.3636*	0.2568*
NO <sub>3</sub> <sup>-</sup>	0.05504	0.05192	0.07594	0.07463
DOC	0.3153**	0.07243	-0.06333	0.1312

Hernandez et al. (JMarSyst)																						
Table S4. Environmental parameters used for the RDA analyses.																						
Station	Depth (m)	Salinity	Temp (°C)	Oxygen (µM)	NH4+ (nM)	PO4 (µM)	Si(OH)4 (µM)	NO2 (µM)	NO3 (µM)	Chlorophyll c3 (µg/L)	Peridinin (µg/L)	Fucoaxanthin (µg/L)	Prasincoxanthin (µg/L)	19'-Hexanoyloxyfucoxanthin (µg/L)	Alloxanthin (µg/L)	Zeaxanthin (µg/L)	Lutein(µg/L)	Chlb (µg/L)	Chlorophyll a (µg/L)	Phaeophytin a (µg/L)	POC (µM)	PON (µM)
M1	25	33.89	5.08	322.04	421	1.63	6.49	0.28	24.76	0.0378	0.0045	0.0738	0.0021	0.1285	0.0023	0.0018	0.0008	0.0096	0.3111	0.0019	4.838	0.863
M1	50	33.89	4.90	320.48	712	1.67	6.9	0.29	24.55	0.0514	0.005	0.0939	0.0035	0.1428	0.0037	0.0016	0.0013	0.0178	0.3633	0.0025	4.964	0.855
M1	125	34.01	1.97	296.44	0	2.15	30.02	0.03	31.47	0.0015	0.0023	0.0116	0	0.0013	0.0004	0	0	0	0.0173	0.0036	1.511	0.234
M1	250	34.22	2.21	231.95	0	2.3	47.63	0.04	34.45	0.0004	0.0012	0.0025	0	0	0	0	0	0	0.0035	0.0022	0.907	0.174
M2_1	9	33.86	5.17	322.09	704	1.45	1.17	0.27	21.62	0.0346	0.0103	0.077	0.0027	0.0921	0.0031	0.0048	0.0015	0.0109	0.2795	0.003	5.531	0.88
M2_1	48	33.87	5.08	320.83	729	1.45	1.14	0.27	21.68	0.0416	0.0124	0.0854	0.0022	0.1013	0.0032	0.0046	0.002	0.0105	0.2886	0.0031	5.481	0.837
M2_1	124	33.93	1.88	325.59	642	2.07	22.85	0.37	28.96	0.0038	0.0057	0.0674	0	0	0	0.0006	0.0004	0	0.128	0.0149	3.447	0.382
M2_1	349	34.25	2.08	231.70	0	2.33	51.44	0.03	36.89	0.0003	0	0.0046	0	0	0	0	0	0	0.006	0.0048	3.662	0.288
M2_2	15	33.87	5.24	316.00	1090	1.47	1.29	0.3	21.25	0.0431	0.008	0.1124	0.0059	0.0743	0.0037	0.004	0.001	0.0223	0.3246	0.0025	4.437	0.691
M2_2	50	33.86	5.24	316.15	1106	1.48	1.35	0.27	21.38	0.0429	0.0083	0.114	0.0059	0.0751	0.0038	0.004	0.0011	0.0227	0.3336	0.0024	3.541	0.544
M2_2	125	33.93	2.23	319.25	297	2.05	21.69	0.32	28.95	0.0044	0	0.034	0	0.0037	0	0.0004	0.0007	0	0.0561	0.0118	2.178	0.315
M2_2	350	34.25	2.06	230.07	0	2.47	51.65	0.03	34.57	0.0003	0	0.0041	0	0	0	0.0002	0	0	0.005	0.0054	1.161	0.153
M2_3	15	33.86	5.18	318.82	899	1.51	2.6	0.3	21.75	0.0761	0.0125	0.216	0.0184	0.1033	0.0049	0.0047	0.0013	0.0519	0.575	0.0045	5.915	0.933
M2_3	50	33.86	5.07	317.59	957	1.49	2.67	0.3	21.61	0.0914	0.0173	0.2356	0.0167	0.1239	0.0063	0.0043	0.0015	0.0534	0.6218	0.0061	5.984	0.966
M2_3	127	33.95	2.11	315.71	263	2.06	23.5	0.32	28.91	0.0026	0.002	0.0212	0	0.0025	0	0.0003	0.0005	0	0.0347	0.0079	1.728	0.26
M2_3	351	34.31	2.15	214.34	0	2.34	56.46	0.04	34.56	0.0002	0	0.0033	0	0	0	0	0	0	0.0041	0.0045	1.034	0.14
M3_1	25	33.82	5.60	314.77	501	1.62	2.31	0.24	23.39	0.0256	0.0049	0.034	0	0.1041	0.0013	0.0014	0	0.0073	0.191	0	4.262	0.719
M3_1	50	33.82	5.60	314.67	508	1.62	2.31	0.24	23.27	0.0256	0.0051	0.0343	0	0.1013	0.0013	0.0014	0	0.0072	0.1892	0	4.05	0.704
M3_1	125	33.87	2.72	316.75	503	1.99	18.01	0.4	28.04	0.0281	0.0056	0.0372	0	0.0502	0	0.0007	0	0.0019	0.1152	0.0062	2.288	0.347
M3_1	250	34.06	2.23	276.04	0	2.15	34.58	0.02	32.05	0.0006	0	0.0027	0	0	0	0	0	0	0.0045	0.0018	0.998	0.187
M3_3	24	33.81	5.32	312.90	722	0.01	2.42	0.01	0.02	0.0189	0.0045	0.029	0	0.065	0.0012	0.0006	0.0003	0.0054	0.1402	0	3.297	0.547
M3_3	49	33.81	5.32	313.27	733	1.61	2.45	0.23	23.02	0.0193	0.0043	0.0292	0	0.0657	0.0012	0.0006	0.0003	0.0056	0.1413	0	3.117	0.525
M3_3	126	33.90	2.71	313.79	246	2.01	20.67	0.38	28.19	0.0119	0.0054	0.0222	0	0.0177	0	0.0004	0	0.001	0.0625	0.0052	1.644	0.254
M3_3	249	34.03	2.21	285.48	0	2.1	32.31	0.02	30.97	0.0009	0.0004	0.0046	0	0	0	0.0002	0	0	0.0067	0.0024	0.903	0.127
M4_1	23	33.85	4.50	324.05	354	1.72	4.13	0.25	25.5	0.0231	0.0039	0.0483	0	0.0832	0.0011	0.0014	0.0004	0.003	0.1801	0	3.853	0.646
M4_1	48	33.85	4.48	323.24	349	1.68	4.19	0.25	25.5	0.0233	0.0037	0.0547	0	0.079	0.0011	0.0016	0.0005	0.0029	0.1811	0	4.414	0.691
M4_1	124	33.91	2.83	324.06	812	1.94	18.85	0.28	27.57	0.0221	0.0024	0.0739	0	0.0154	0	0.0009	0.0007	0.001	0.1158	0.0064	2.932	0.43
M4_1	251	34.07	1.73	287.82	0	2.13	36.83	0.01	32.17	0.001	0	0.0043	0	0.0005	0	0.0003	0	0	0.006	0.0026	1.194	0.17
M4_2	24	33.85	4.46	320.13	481	1.71	4.8	0.27	24.79	0.0278	0.003	0.0658	0	0.0857	0.0012	0.0009	0.0006	0.0038	0.2151	0.0013	3.745	0.564
M4_2	52	33.85	4.46	320.23	495	1.71	4.8	0.27	24.27	0.0275	0.0033	0.0641	0	0.0847	0.0012	0.0009	0.0007	0.0037	0.2078	0	3.536	0.58
M4_2	125	33.91	2.56	320.07	930	1.99	20.54	0.35	26.91	0.0141	0.0015	0.0408	0	0.0186	0	0.0005	0.0003	0.0047	0.0785	0.0035	2.074	0.329
M4_2	246	34.07	1.96	282.39	0	2.13	34.66	0.04	30.94	0.0006	0.0012	0.0033	0	0	0	0	0	0	0.0056	0.0021	1.136	0.153

DOC (µM)	Prokaryotic Abundance (cells /mL)	Photosynthetic Pico-Nano-eukaryotes (cells/mL)*	Heterotrophic Nanoflagellates (cells /mL)*	Viral like particles (VLPs/mL)*	Bacterial Production (µg C/L/h)*
50,3	606000	8040	943	3,10E+06	21,54
50,3	619000	8010	964	3,69E+06	18,04
48,1	253000	38,5	366	2,05E+06	1,97
46,2	222000	25,2	168	1,28E+06	1
52,5	1180000	3190	1340	4810000	22,16
53,4	1150000	2770	1700	5850000	30,99
49,8	371000	41,5	466	2110000	5,01
46,8	269000	16,3	257	1170000	1,37
54,6	837000	3770	2840	8,02E+06	46,23
56,7	794000	3490	2910	7,11E+06	35,81
49	367000	80	726	2,94E+06	4,59
46,8	220000	13,3	278	1,31E+06	1,3
54,6	665000	8130	1730	7,40E+06	56,91
55	708000	8410	1890	7,27E+06	45,09
49	301000	56,1	551	2,43E+06	3,58
45,9	214000	14,8	300	1,09E+06	1,27
50,4	696000	3720	625	3,64E+06	15,59
50,2	665000	3590	763	3,65E+06	14,39
47	358000	42,8	655	2,40E+06	4,5
44,2	191000	7,38	222	1,52E+06	0,41
49,1	446000	1490	847	4,10E+06	13,18
52	445000	1520	798	3,71E+06	13,76
49,4	325000	33,9	931	2,83E+06	3,62
44,2	205000	10,3	362	2,25E+06	1,69
49,3	552000	3180	851	3,45E+06	13,98
50	550000	3160	965	3,80E+06	13,1
49,4	440000	220	655	3,00E+06	6,82
44,9	193000	8,4	178	1,43E+06	0,3
49,5	501000	3700	884	3320000	14,39
51,2	506000	3760	889	3060000	14,81
50	388000	104	658	2740000	5,56
44,9	199000	4,2	301	1390000	0,51