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# Short-term responses to ocean acidification: effects on relative abundance of eukaryotic plankton from the tropical Timor Sea

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#### ABSTRACT

Anthropogenic carbon dioxide (CO<sub>2</sub>) emissions drive climate change and pose one of the major challenges of our century. The effects of increased CO<sub>2</sub> in the form of ocean acidification (OA) on the communities of marine planktonic eukaryotes in tropical regions such as the Timor Sea are barely understood. Here, we show the effects of high CO<sub>2</sub> (pCO<sub>2</sub>=1823±161 µatm,  $pH_T=7.46\pm0.05$ ) versus in situ CO<sub>2</sub> (pCO<sub>2</sub>=504±42 µatm, pH<sub>T</sub>=7.95±0.04) seawater on the community composition of marine planktonic eukaryotes immediately and after 48 hours of treatment exposure in a shipboard microcosm experiment. Illumina sequencing of the V9 hypervariable region of 18S rRNA (gene) was used to study the eukaryotic community composition. Down-regulation of extracellular carbonic anhydrase occurred faster in the high CO<sub>2</sub> treatment. Increased CO<sub>2</sub> significantly suppressed the relative abundances of different eukaryotic operational taxonomic units (OTUs), including important primary producers. These effects were consistent between abundant (DNA-based) and active (cDNA-based) taxa after 48 hours, e.g., for the diatoms Trieres chinensis and Stephanopyxis turris. Effects were also very species-specific among different diatoms. Planktonic eukaryotes showed adaptation to the CO<sub>2</sub> treatment over time, but many OTUs were adversely affected by decreasing pH. OA effects might fundamentally impact the base of marine biodiversity, suggesting profound outcomes for food web functioning in the future ocean.

**KEYWORDS:** high CO<sub>2</sub>, low pH, phytoplankton, protists, tropics, copepods, 18S rRNA sequencing

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#### **1. INTRODUCTION**

The world's oceans are a major sink for anthropogenic carbon dioxide (CO<sub>2</sub>) (Friedlingstein et al. 2019, Gruber et al. 2019). Absorption of atmospheric CO<sub>2</sub>, which is currently at 414 µatm (February 2020, https://www.co2.earth/) and driven by ongoing anthropogenic emissions, decreases pH while increasing the bicarbonate ion concentration and partial pressure of CO<sub>2</sub>  $(pCO_2)$  of seawater. Man-made CO<sub>2</sub> emissions are expected to lead to ocean pH changes greater than any experienced in the last 300 million years, with a maximum predicted pH reduction of 0.77 in surface water for the year 2300 (Caldeira & Wickett 2003). The decrease in seawater pH, known as ocean acidification (OA) or "the other CO<sub>2</sub> problem" (Doney et al. 2009), affects ocean biogeochemistry and the physiology of a plethora of marine organisms, especially the calcifying ones (Orr et al. 2005, Kaniewska et al. 2012, Schlüter et al. 2016). The fifth assessment report of the IPCC suggests that net primary productivity in the tropical ocean will most likely decline by 7-16 % by 2081-2100 assuming the worst case "business as usual" scenario (Representative Concentration Pathway=RCP8.5), which predicts an atmospheric CO<sub>2</sub> concentration of >1000 µatm by the end of this century (Pachauri et al. 2014). Moreover, profound implications of OA effects on community and ecosystem processes can be expected but are difficult to predict (Riebesell et al. 2007, Fabry et al. 2008).

Changes in seawater pH and  $pCO_2$  may have critical effects on the primary producers, i.e., phytoplankton that convert  $CO_2$  to organic carbon during photosynthesis, and thus affect the base of the marine food web. Increased  $CO_2$  is known to promote phytoplankton growth and photosynthetic rates (Wolf-Gladrow et al. 1999, Kroeker et al. 2013), although increases in the latter are assumed to be small and dependent on the phytoplankton species under investigation (reviewed by Raven et al. (2005)). Typical acclimation effects of phytoplankton to elevated  $CO_2$  include the down-regulation of carbon concentrating mechanisms, e.g., of the enzyme

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carbonic anhydrase (Mustaffa et al. 2017, Deppeler et al. 2018), demonstrating direct effects of OA on intracellular pH and enzyme activities. Furthermore, CO<sub>2</sub>-driven changes in element stoichiometry (King et al. 2015), community structure (Tortell et al. 2002, Feng et al. 2010), and size classes (Suffrian et al. 2008) of primary producers can alter food availability and quality for grazers and cascade up the whole food web (Rossoll et al. 2012, Cripps et al. 2016).

Copepods (phylum Arthropoda, class Copepoda (Pancrustacea)) are the most abundant metazoans on Earth (Humes 1994). They form the major dietary link between phytoplankton and ichthyoplankton as well as larger fish (Turner 2004). Copepods and other crustaceans were assumed to be more resilient towards OA effects compared to other marine organisms (Kurihara & Ishimatsu 2008, Whiteley 2011). However, this assumption mainly results from studies on adult copepod females and has been questioned (Cripps et al. 2014) supporting the lack of conclusive knowledge about OA effects on copepods that exist in the literature.

Insights on the effects of lowered pH on natural marine nano- to meso-sized planktonic organisms from the tropics are scarce. Hence, our objective was to study the impact of OA on a natural plankton community from the Timor Sea, tropical Indian Ocean in a microcosm experiment that was conducted onboard the R/V Falkor in October 2016. Probable OA effects on planktonic communities were investigated in one treatment with reduced pH  $(pCO_2=1823\pm161 \ \mu atm, \ pH_T=7.46\pm0.05)$  compared to an unaltered control  $(pCO_2=504\pm42 \ \mu atm, \ pH_T=7.95\pm0.04)$  at two consecutive time points. The effects of OA on abundant and active planktonic organisms were investigated as inferred from DNA and cDNA-based V9 hypervariable region of the 18S rRNA (gene) sequencing, respectively. Since literature on the topic reports inconclusive results of OA effects sometimes even for the same organism (Ridgwell et al. 2009, Meyer & Riebesell 2015, Thor & Oliva 2015), we expected to see

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significant short-term OA effects after different exposure times on the relative abundance of some taxa as well as missing responses in others.

#### 2. MATERIALS AND METHODS

#### 2.1 Sampling and microcosm setup

During a research cruise on R/V Falkor (FK161010), sample water containing planktonic organisms was collected from the chlorophyll maximum at 65 m depth using a Conductivity, Temperature, Depth (CTD) unit at 19:20, 20th October 2016 (UTC) from the Timor Sea (11°51.25'S, 127°15.23'E). The region is tropical with summer and winter sea surface temperature >26 and >22 °C, respectively, and experiences periodic cyclone-generated storm currents (James et al. 2004). Water temperature and salinity at the time of sampling were at 28 °C and 35, respectively. Sampled seawater was randomly filled from CTD bottles into 24 x 1-L Nalgene polycarbonate bottles (Thermo Fisher Scientific, Darmstadt, Germany), which were wrapped into a foil (No 298, Lee Filters, Andover, UK) filtering daylight to approach light conditions at the sampling depth. Half of the bottles were acidified using an equimolar addition of strong acid (1 M HCl) and HCO<sub>3</sub> (1 M NaHCO<sub>3</sub>) to simulate effects of low pH/high CO<sub>2</sub> (treatment HICO) according to conditions expected in a future ocean (Pachauri et al. 2014). The other half of samples was kept at *in situ* conditions (treatment ISCO). All bottles were placed into a baby pool (99 x 99 x 23 cm, Wehnke, Germany). During sample incubations, fresh oceanic water was constantly pumped into the pool to keep the temperature as similar as possible to *in situ* conditions. Pool temperature was monitored by a portable PCD650 (EuTech Instruments, Singapore, Supplementary Figure S1). Incubations were carried out from the 20<sup>th</sup>  $(t_0)$  to the 23<sup>th</sup> October, 00:40 UTC  $(t_2)$ , and for each day beginning at  $t_0$  four replicate bottles of each the HICO and the ISCO treatment were processed. The time period from acidification

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until water filtration for t<sub>0</sub> samples was three hours. Before water sample processing, the pH was measured in each bottle to monitor that the manipulation was effective.

### 2.2 Chlorophyll a extraction

Discrete chlorophyll *a* (chl *a*) analysis was performed on sampling water (initial samples after CTD, HICO, ISCO, n=4), using the fluorometer JENWAY 6285 (Bibby Scientific Ltd., Felsted, Essex, UK). Readings on standards were taken by using commercial pure chl *a* (Sigma Aldrich, Taufkirchen, Germany). Water samples (160–300 mL of the 1L) were filtered onboard the *R/V* immediately onto glass microfiber filters (GF/F, diameter: 25 mm, Whatman, UK), were stored at -20 °C, and shipped on dry ice for further analysis in the home laboratory. The filtered samples were extracted in 90% ethanol for 24 hours in the dark, before being measured fluorometrically in triplicates and according to the EPA Method 445.0 (Arar & Collins 1997).

#### 2.3 Analysis of carbonate chemistry

Dissolved inorganic carbon (DIC) and total alkalinity (TA) samples were analyzed at the University of Southampton using a VINDTA 3C (Marianda, Kiel, Germany). The DIC was measured by coulometric titration, and TA by potentiometric titration and calculated using a modified Gran plot approach as implemented by Calkulate (version 1.0.2) (Bradshaw et al. 1981, Humphreys 2015). Measurements were calibrated using certified reference material (batches 144, 151 and 160) obtained from A.G. Dickson (Scripps Institution of Oceanography, USA). The 1 $\sigma$  measurement precision was ±3 and ±2 µmol kg<sup>-1</sup> for DIC and TA, respectively. The remaining carbonate chemistry variables were calculated with PyCO2SYS (version 1.3.0) (Lewis & Wallace 1998, Van Heuven et al. 2011, Humphreys et al. 2020) using the carbonic acid dissociation constants of Lucker et al. (2000).

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The dissociation constant for bisulphate dissociation was chosen according to Dickson (1990), and the ratio of total borate to salinity was in accordance with Lee et al. (2010).

Discrete samples for inorganic nutrients (nitrate and nitrite, phosphate and silicate) were collected from the same intervals as the DIC and TA samples. Inorganic nutrients were measured using a continuous flow analyzer (Quattro, CFA), Reader, Easychem (Strickland 1968, Fanning & Pilson 1973). The detection limit was 0.4  $\mu$ M for nitrate and nitrite, 0.04  $\mu$ M for phosphate and 0.1  $\mu$ M for silicate, and the limit of determination was 1.0  $\mu$ M, 0.13  $\mu$ M, and 0.3  $\mu$ M respectively.

#### 2.4 DNA extraction and cDNA synthesis

For DNA and RNA extraction, a two-step filtration of sampling water (150-400 mL from each bottle) was conducted, namely water was filtered through a 3.0  $\mu$ m pore size membrane filter. The flow-through was further filtered onto a 0.2  $\mu$ m polycarbonate membrane filter (47 mm diameter, Whatman, Maidstone, UK). All filters were initially stored at -80°C prior analysis and shipped on dry ice to the home laboratory. Extraction of DNA and RNA from t<sub>0</sub> and t<sub>2</sub> filters for three biological replicates (half of a 0.2 and a 3  $\mu$ m filter were pooled in each case) was performed by using the DNA + RNA + Protein Extraction Kit (Roboklon, Berlin, Germany) and a modified protocol as described in Rahlff et al. (2017). Remaining DNA in RNA samples was digested on-column using 3 U of DNase, and all samples were subsequently checked for genomic DNA contaminations by polymerase chain reaction (PCR). A quantity of 10 ng RNA was reversely transcribed to cDNA using the NG dART Kit and therein included random hexamer primers (Roboklon, Berlin, Germany). Negative controls without reverse transcriptase and without RNA were included. The reaction was incubated for 60 minutes at 50°C followed by 5 minutes at 85°C.

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# 2.5 Library preparation from V9-18S rRNA (gene) amplicons, sequencing and bioinformatics

Two independent PCRs were carried out to attach the Nextera XT compatible Illumina adapter overhangs at the 5' ends of each amplicon following the Illumina 16S Metagenomic Sequencing Library Preparation guide (15044223Rev.B). The first PCR was conducted on a 3PrimeG thermocycler (Techne, Staffordshire, UK) using Phusion High Fidelity DNA Polymerase (Thermo Fisher Scientific, Darmstadt, Germany), 10 mM dNTPs, 3% DMSO and 0.5 µM of the Tara Oceans eukaryote primer set 1389F/1510R targeting the V9 hypervariable region of the 18S ribosomal RNA gene (Amaral-Zettler et al. 2009). The primers were tagged with part of the Nextera compatible Illumina adapter overhang at the 5' ends of each primer. The PCR-cycling program was modified from a previously described one for this primer set (Alberti et al. 2017), namely 27 and 30 cycles instead of 25 were chosen for DNA and cDNA template, respectively.

Five µL of the first PCR products have been purified using 2 µL of ExoSAP-IT PCR Product Cleanup Reagent (Thermo Fisher Scientific) following the manufacturer's protocol. A second PCR using Nextera XT V2 Indexed primers (dual indexing approach) has been performed to bind the Illumina overhang adapters to the product of the first PCR and was conducted at 98 °C for 2 min, 7 cycles of 15 s at 98 °C, 30 s at 62°C, 30 s at 72 °C and 72 °C for 2 min. The PCR products' cleanup was performed using 60% of the PCR products volume of Agencourt® Ampure® XP magnetic beads (Beckmann Coulter, Brea, CA, USA), and concentrations were measured using Qubit Fluorometric Quantitation (Thermo Fisher Scientific, Darmstadt, Germany). All amplicon concentrations were subsequently adjusted to 8 nM, pooled and purified from a 2% agarose gel stained with 2% Gel Red using Monarch® Nucleic Acid

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Purification kit (New England Biolabs, Ipswich, MA, USA). The library was sequenced in two independent runs on an Illumina MiSeq platform using the MiSeq Reagent Nano Kit v2 (150 paired-end cycles). The resulted reads have been demultiplexed by MiSeq considering Nextera index sequences for both forward and reverse strands.

BBMap tool (Bushnell 2018) has been used for trimming the primers. The Vsearch pipeline (Rognes et al. 2016) was used for making contigs from each sample using the following settings: expected mean size of 170 bp with +/- 30 bp intervals, minimum contig overlap at 25 bp, and maximum allowed differences in the contig at 15 bp. Contigs have been filtered with maximum expected errors of 0.5 and maximum of 0 ambiguities in the contig and de-replicated. All de-noised and de-replicated samples from libraries have been initially clustered at 98% similarity to produce the initial operational taxonomic units (OTUs). After excluding chimeras, the initial OTUs have been clustered at a similarity threshold of 97 % (species level) for each sample, which has been recently recommended for PCR-based high-throughput sequencing data to obtain more realistic richness and Shannon diversity data from 18S amplicon data (Wylezich et al. 2018). The generated OTUs have been blasted against GenBank to assign taxonomies. Sequence reads were deposited at NCBI's sequence read archive (SRA) under Bioproject ID PRJNA623264.

#### 2.6 Data analysis

A total number of 2435 OTUs resulted after implementing all filtrations, in which the number of unique reads per library ranged from 20616 to 41687. After de-noising, chimera detection and de-replicating this has been reduced to 3051 to 5673. Number of resulted OTUs were between 503 and 1136 for the different samples (Supplementary Table S1).

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Sequence data were analyzed using the "phyloseq" package (McMurdie & Holmes 2013) in R version 3.6.1. (Team 2017). Reads were normalized to minimum sequencing depth, which corresponded to 18056 reads. One replicate of the HICO  $t_0$  cDNA samples was removed from the dataset due to a too small sequencing depth of 334.

For all calculations and plots we focused on the following phylogenetic groups: apicomplexans, brown-algae, cercozoans, choanoflagellates, ciliates, crustaceans, cryptomonads, ctenophores, diatoms, dinoflagellates, euglenoids, eukaryotes (no further classification), forams, golden\_algae, green\_algae, haptophytes, kinetoplastids and pelagophytes. Relative abundances were calculated and used for the following analyses. Shannon-Wiener diversity indices ( $\alpha$ -diversity) were calculated using the estimate\_richness function of the "phyloseq" package. We visualized the differences ( $\beta$ -diversity) of the eukaryotic planktonic community composition of samples through a Nonmetric Multidimensional Scaling (NMDS) plot based on Bray–Curtis dissimilarity indices.

For finding significant differences for extracellular carbonic anhydrase (eCA) concentrations and Shannon-Wiener indices between treatments, a one-way analysis of variances (one-way ANOVA) with 95 % confidence level was performed using R. The prerequisites for parametric tests, i.e., normal distribution of data and variance homogeneity were previously confirmed using Shapiro and Bartlett tests in R, respectively. A Tukey HSD *post-hoc* test was used to make multiple comparisons of means to find significant differences.

A two-sided t-test assuming unequal variances (Welch t-test) was conducted to find significant differences ( $\alpha \le 0.05$ ) between relative abundances of individual OTUs in the ISCO and HICO treatments (n=3). Since the sample HICO t0 cDNA lacked one replicate, the comparison ISCO/HICO for t0 cDNA was omitted.

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To elucidate the abundance of significantly CO<sub>2</sub>-affected taxa in both treatments., the proportion of significantly CO<sub>2</sub>-affected OTUs was calculated and assigned to one of five abundance categories (Table 2).

#### **3. RESULTS**

#### 3.1 Carbonate chemistry and temperature

Temperature in the pool varied between 27.0 and 34.0 °C from  $t_0$  to  $t_2$  (Supplementary Figure S1). Seawater pH<sub>T</sub>, i.e., pH on the Total scale, was consistent at  $t_0$  and  $t_1$  under both *p*CO<sub>2</sub> conditions with average values (mean ± standard deviation) of 7.44±0.03 and 7.93±0.02 for the HICO and ISCO treatment, respectively (Figure 1A). We measured a small increase in pH<sub>T</sub> of circa 0.06 in both experiments at  $t_2$ , to 7.50±0.06 and 7.99±0.03. Dissolved inorganic carbon (DIC) was relatively stable throughout all experiments under both *p*CO<sub>2</sub> conditions (Figure 1B), with an average of 2018±14 µmol kg<sup>-1</sup> and 2213±14 µmol kg<sup>-1</sup> in the ISCO and HICO treatment, respectively.

The seawater  $pCO_2$  was calculated from measured pH<sub>T</sub> and DIC along with relevant metadata, using the mean DIC values for each CO<sub>2</sub> treatment where those measurements were missing (Figure 1C). Like pH<sub>T</sub>, seawater  $pCO_2$  was consistent at t<sub>0</sub> and t<sub>1</sub> under both  $pCO_2$  conditions with average values of 1910±73 µatm and 526±30 µatm for the HICO and ISCO treatments respectively, but these values decreased to 1648±144 µatm and 460±20 µatm at t<sub>2</sub>. The calculated decrease in  $pCO_2$  at t<sub>2</sub> was driven by the measured increase in pH<sub>T</sub>, which could be due to organic alkalinity degradation during the experiments (Martín Hernández-Ayon et al. 2007).

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#### 3.2 Chlorophyll *a*, nutrients, extracellular carbonic anhydrase

Discrete chl *a* analyses revealed concentrations <0.4 µg L<sup>-1</sup> (n=4) with no consistent difference between HICO and ISCO treatment (Figure 2A). Nutrient levels were overall low. PO<sub>4</sub> and NOX were below the detection limit (0.04 µM for PO<sub>4</sub> and 0.4 µM for NOX). Silicate levels dropped from t<sub>0</sub> to t<sub>2</sub> from maximum 7.20 µM to minimum 5.52 µM with no major differences between ISCO and HICO treatments (Figure 2B). Concentration of eCA in seawater was measured in quadruplicates from t<sub>0</sub> to t<sub>2</sub>. Methods and data for this experiment were reported in the paper and supplement material of Mustaffa et al. (2017). Indeed, the concentration of eCA decreased faster in the HICO treatment compared to the ISCO treatment (Supplementary Figure 2 of Mustaffa et al. (2017)). ANOVA demonstrated significant differences for the eCA concentration (*F*=9.51, *df*=4, *p*=0.00049). Means of eCA concentrations were not significantly different between ISCO and HICO treatment at t<sub>0</sub>, but weakly significant at t<sub>1</sub> (Tukey HSD, *p*=0.025). Missing data for the HICO treatment at t<sub>2</sub> were removed for the statistics (and not assumed to be equal to zero).

#### 3.3 Plankton alpha and beta diversity in response to acidification

The one-way ANOVA indicated weak significant differences between the mean species diversities of the different sample types (F=4.62, df=5, p=0.014). Species diversity analyses based on Shannon-Wiener index (Figure 3) showed that at t<sub>0</sub> DNA-based OTUs showed a higher diversity measure in the ISCO (mean: 3.6 ± standard deviation: 0.98, n=3) compared to the HICO treatment (2.3 ± 0.53, n=3), which was not significant (Tukey HSD, p=0.1). Such differences were less pronounced for the cDNA-based OTUs, where both measures were equally high (ISCO: 4.5 ± 0.16, n=3; HICO: 4.4, n=2). For t<sub>2</sub>, the Shannon-Wiener index

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differences between the acidification treatments based on DNA-based OTUs were less strong compared to  $t_0$  (ISCO:  $3.8 \pm 0.46$ , n=3; HICO:  $3.2 \pm 0.44$ , n=3) and even more similar for cDNA-based OTUs (ISCO:  $4.3 \pm 0.05$ , n=3; HICO:  $3.8 \pm 0.53$ , n=3). It follows that OTUs based on DNA sequencing (reflecting abundant taxa) seem slightly more prone to acidification effects because they have a lower Shannon-Wiener index compared to those based on cDNA sequencing (reflecting active taxa), though all differences between treatment pairings were not significant based on Tukey HSD test.

The community composition of each sample (Figure 4) revealed that the most striking differences in cDNA- and DNA-based OTUs between HICO and ISCO treatments at t<sub>0</sub> were based on the different relative abundance of crustaceans. Overall, diversity of phyla appeared comparable between replicates (Figure 4). However, the community composition plots of the individual groups (diatoms, dinoflagellates etc.) showed profound differences in relative abundances between t<sub>0</sub> and t<sub>2</sub> (Supplementary Figures S2 – S12). Many OTUs that were undetectable at t<sub>0</sub> reached considerable abundances at t<sub>2</sub>. For instance, after 48 hours, active and abundant OTUs assigned to uncultured kinetoplastids and especially the genus *Bodonidae* sp., the ciliate *Eutintinnus fraknoi*, the cercozoan *Massisteria marina* and the choanoflagellate *Calliacantha* sp., strongly increased in relative abundance from t<sub>0</sub> to t<sub>2</sub> (Supplementary Figures S4, S5, S6, S12). On the other hand, OTUs assigned to dinoflagellates and some haptophytes, e.g., *Phaeocystis antarctica* decreased in relative abundance, whereas other haptophytes, e.g., an OTU related to *Pavlova gyrans*, became more abundant during the incubation period (Supplementary Figures S3 and S7).

In agreement with these abundance changes of different taxa, the NMDS plot (Figure 5, stress= 0.070) indicated that four distinct community cluster were formed. The clusters could be mostly distinguished based on the nucleic acid type used as PCR template (DNA or cDNA) and the

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incubation period ( $t_0$  versus  $t_2$ ). Differences between HICO and ISCO treatment did not reveal strong effects at the phylum level, although at  $t_0$ , the two treatments showed some separation among the DNA-based OTUs. Communities based on cDNA and DNA resembled each other more at  $t_2$  than at  $t_0$ , and this observation was independent from the acidification treatment.

#### 3.4 OA effects on the OTU level

Despite the fact that the acidification treatment induced no major differences at the phylum level (Figure 4), significant differences ( $p \le 0.05$ ) were detectable at the OTU-level (Table 1). In most cases, the HICO treatment was associated with a lower relative abundance of significantly affected OTUs compared to the ISCO treatment.

For crustaceans, a higher relative abundance of the copepod species *Acartia longiremis* and *Calanus finmarchicus* was found among the DNA-based OTUs in the HICO treatment compared to the ISCO treatment (Supplementary Figure S8). For instance, the increase in relative abundance of an OTU assigned to *Calanus finmarchicus* was statistically significant in the HICO treatment at t<sub>0</sub> (ISCO: DNA: mean=0.002 %, STD=±0.004 %, HICO: DNA: mean=0.02 % ±0.0007 %, *p*=0.015).

At t<sub>2</sub>, we sometimes found the same OTUs to be significantly affected by the OA treatment among the DNA and cDNA-based OTUs, e.g. for the diatoms *Trieres chinensis, Stephanopyxis turris* (OTU\_54) or *Synedra* sp. (Table 1). All significantly affected taxa (based on Welch t-test) are shown in Table 1.

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#### 4. DISCUSSION

#### 4.1 OA effects are species or even ecotype-specific

Increased atmospheric CO<sub>2</sub> burden due to enhanced anthropogenic fossil fuel emissions and associated OA is one of the major global issues of our time. The effects of OA on natural eukaryotic plankton communities, especially from the tropical Timor Sea are poorly understood, and our data show that the HICO treatment influenced the relative abundance of certain abundant and active taxa from this oceanic region. Since the copy number of the small subunit rRNA genes is known to correlate with the biovolume of marine diatoms and can reach considerable numbers per cell in this group (Godhe et al. 2008) and also in ciliates (Gong et al. 2013), we are aware about the difficulties of using this gene for the interpretation of single-celled protist numbers. However, it is impossible to know the exact copy number of the 18S rRNA (gene) for each OTU belonging to different protists of a mixed community of which many species are uncultured or even unknown. We made comparisons on the relative abundance of an OTU by comparing two treatments, meaning that even if an OTU species-specifically contains multiple 18S rRNA gene copies, it will probably do so in both treatments and is thus comparable.

While significant CO<sub>2</sub> effects were not immediately apparent at higher taxonomic levels (Figure 4), they became visible at the OTU level. We expected that phytoplankton OTUs would be most sensitive to changes in CO<sub>2</sub> due to their inorganic carbon requirements for primary production and found responses to OA in diatoms to be highly species- or even ecotype specific. Many diatoms affected by the HICO treatment decreased in relative abundance as a response including *Trieres chinensis, Synedra* sp., *Fragilariopsis doliolus, Pseudonitzschia australis, Pseudo-nitzschia subcurvata* (OTU\_210), *Stephanopyxis turris* (OTU\_54), and *Pleurosigma* 

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sp. The diatom *Stephanopyxis turris* (OTU\_54) showed a decrease in relative abundance in response to the HICO manipulation both in the DNA-and the cDNA-based investigation after 48 hours, most likely indicating that abundance and activity were both reduced by the treatment. This further confirms that the significant effect was real and not a false-positive random finding. A few diatoms also increased in abundance in the HICO treatment such as *Cocconeis placentula, Chaetoceros* sp., *Corethron criophilum, Grammonema striatula,* another *Pseudo-nitzschia subcurvata* (OTU\_659) and another *Stephanopyxis turris* (OTU\_505), matching the generally constant chl *a* concentration. Our results are in agreement with Schulz et al. (2017) postulating that CO<sub>2</sub> effects on diatoms are more likely to become visible on the species rather than on the phylum- or other higher taxonomic level.

Many studies of OA effects on natural microbial communities focus on polar waters complicating direct comparison to our findings from a tropical region. By investigating phytoplankton populations from the Ross Sea, Southern Ocean, Tortell et al. (2008) reported that growth of larger chain-forming diatoms such as *Chaetoceros* spp. increased in abundance in response to high CO<sub>2</sub> (800 µatm), while the small pennate diatom *Pseudo-nitzschia subcurvata* decreased in this treatment. This finding matches our results apart from the fact that we had two different OTUs assigned to *Pseudo-nitzschia subcurvata*, and both had contrasting responses to the treatment at t<sub>0</sub> and that our HICO treatment was much higher ( $pCO_2=1823$  µatm). Also, we cannot conclude from our results that different morphologies of diatoms (centric vs. pennate) have different advantages in facing high and low CO<sub>2</sub> environments as proposed by Tortell et al. (2008), because in our study OTUs related to both morphology types decreased in relative abundance.

We also found contrasting responses to OA by two OTUs being both assigned to *Stephanopyxis turris*. This leads to the conclusion that very closely related diatom species (97% sequence

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identity) can have very different responses to OA. Different responses of the same model species *Emiliania huxleyi* (but from different ecotypes) to carbonate chemistry manipulations were previously described (Ridgwell et al. 2009). An ecotype refers to a genetically distinct geographic population within a species. Ecotype-specific OA responses might lead to conflicting results between studies and explain the contrasting responses we found for OTUs assigned to *Stephanopyxis turris* and *Pseudo-nitzschia subcurvata*.

Immediate responses to OA were also observed for the dinoflagellate *Vulcanodinium rugosum*, which decreased in relative abundance at t<sub>0</sub>. *V. rugosum* is known for being a harmful algae and producing pinnatoxins causing shellfish poisoning (Rhodes et al. 2011). While it is known that *V. rugosum* is thermophile and euryhaline (Abadie et al. 2016, Abadie et al. 2018), nothing is known about its response to OA. The immediate response and decrease in relative abundance among DNA-based OTUs from mean  $\pm$  STD=ISCO DNA mean: 1.9%  $\pm$  0.30 to HICO DNA mean: 0.76%  $\pm$  0.22 (n=3, Welch t-test, *p*=0.0083) suggests that this dinoflagellate was sensitive to the HICO treatment. Effects of CO<sub>2</sub>-induced OA on dinoflagellates that are able to form (harmful) algal blooms have been previously studied, again showing very species-specific responses. For instance, biomass production in calcareous *Scrippsiella trochoidea* decreased, while toxic *Alexandrium tamarense* did not respond to an OA treatment (Eberlein et al. 2014).

Two copepod OTUs were BLAST-assigned to *Acartia longiremis* and *Calanus finmarchicus*. Although both species have their respective main distribution in the North Atlantic, we decided to keep these names here because there are some records of these species in the southern hemisphere in OBIS (<u>https://obis.org</u>). Both copepods showed sudden increases in relative abundance in response to the HICO treatment at t<sub>0</sub>. The increases were striking but mostly not significant, most likely due to the variation between replicates. We can only speculate about the reason for the immediate response. It has been shown that lowered seawater pH can

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influence copepod reproduction, hatching and development in some species (Kurihara et al. 2004, Mayor et al. 2007) whereas it had no significant effect on physiological parameters or grazing of copepods in other studies (Kurihara & Ishimatsu 2008, Mayor et al. 2012, Isari et al. 2015, Hildebrandt et al. 2016). This implies that OA effects are very species and context-specific and also dependent on the geographic origin of the copepod population (Thor & Oliva 2015), the life stage (Cripps et al. 2014), gender (Cripps et al. 2016), and even food quality provided (McLaskey et al. 2019). Samples that were most affected by the HICO treatment at t<sub>0</sub> could have randomly contained a particular CO<sub>2</sub>-vulnerable life stage or gender of copepods. Or copepod carcasses and fecal pellets present in the samples were more slowly degraded in the HICO compared to the ISCO treatment.

By assigning affected taxa to different OTU abundance ranges (Table 2), we found that the proportion of HICO-treatment affected OTUs was lower compared to the ISCO treatment, especially in the high abundance ranges. This suggests that the HICO treatment overall had a suppressive effect on the relative abundance of taxa. For instance, OTUs that reached an abundance of 9.4% in the ISCO treatment in the 1-100% abundance range, were undetectable at such a high abundance in the HICO treatment. Not surprisingly, more OTUs were found in the lowest abundance range in the HICO treatment, which might be attributable to the pH manipulation.

#### 4.2 Adaptations to the ocean acidification treatment

The NMDS plot revealed that eukaryotic plankton community differences between DNA and cDNA-based OTUs became smaller at  $t_2$  compared to  $t_0$  (Figure 5). The incubation conditions (heat, low oxygen, and grazing pressure) likely selected for specific taxa that could resist those conditions and became abundant and active, while it led to a decreased abundance or even

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extinction of others. The "bottle enclosure effect" exerted by incubations in closed systems (Gieskes et al. 1979) includes decreasing biomass of picophytoplankton, increasing biomass of heterotrophic bacteria (Calvo-Diaz et al. 2011), and increases the likelihood for host cell-virus encounters with potential to alter the community composition (Haro-Moreno et al. 2019). However, containment of natural seawater for 48 hours led to only minor changes in species richness (Figure 3, Shannon-Wiener Index) as previously reported (Countway et al. 2005). We found an increase in the relative abundance of certain taxa from t<sub>0</sub> to t<sub>2</sub>, mainly of kinetoplastids, cercozoans, cryptomoands and choanoflagellates (Supplementary Figures S5, S6, S9, S12), while other taxa such as dinoflagellates might be due to increased grazing or to enhanced vulnerability towards incubation conditions and associated community changes.

The fact that abundant and active eukaryotic communities became more similar with increased incubation time emphasizes that several members of the planktonic community were strongly affected by the incubation conditions. Since the effect occurred after 48 hours ( $t_2$ ) and independent of the OA treatment, our data show that longer bottle incubations to test for OA effects have to be conducted with great caution. If a reliable presentation of the *in situ* community is needed, direct processing at  $t_0$  is required as has been recently suggested for prokaryotic communities (Haro-Moreno et al. 2019). On the other hand, the "bottle enclosure effect" can also be systematically used to allow rare and underrepresented populations to flourish. OA studies often apply acclimation periods with gradually increasing CO<sub>2</sub> to let organisms slowly adjust to new pH conditions (Hancock et al. 2018), which is probably a more realistic representation of OA than sudden pH manipulations. However, here we used acute elevations of seawater  $pCO_2$  to obtain a better understanding of the short-term sensitivity of planktonic organisms and because rapid (within a few days) shifts in picoeukaryote communities due to OA were previously detected (Meakin & Wyman 2011).

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Adaptation of the planktonic eukaryotes to the OA treatment also became visible at the enzymatic level. The down-regulation of carbon concentrating mechanisms processes in response to high CO<sub>2</sub> serves as an energy saving mechanism, which has been previously observed in model phytoplankton species (Wu et al. 2010, Moroney et al. 2011) and natural phytoplankton communities from Antarctica (Young et al. 2015, Deppeler et al. 2018). Carbonic anhydrases, zinc-containing metallo-enzymes that catalyze the hydration of CO<sub>2</sub> into bicarbonate, are important components for the functioning of carbon concentrating mechanisms under CO<sub>2</sub>-limiting conditions (Mondal et al. 2016). Likewise, our data show a rapid decline in the concentration of extracellular carbonic anhydrase (eCA) in response to the HICO treatment in the (autotrophic) plankton community indicating an adaptation to the manipulation. The eCA concentration dropped more slowly in the ISCO compared to the HICO treatment (Supplementary Fig.2 of Mustaffa et al. (2017)). Dropping of eCA concentration in the ISCO treatment probably happened because of a depletion in phytoplankton biomass due to grazing in closed bottles and associated accumulation of respired CO<sub>2</sub> suppressing carbon concentrating mechanisms.

Despite constant pumping of fresh ocean water into the sampling pool, heating of incubation bottles during the day because of the tropical climate (Supplementary Figure S1) was a clear limitation of our experiment. Previous studies did not detect a synergetic effect of OA and warming on plankton communities (Paul et al. 2015, Horn et al. 2016). But to exclude additional effects by warming, e.g., the decrease of heterotrophic flagellate biomass (Moustaka-Gouni et al. 2016), we decided to compare effects of OA treatment only within but not between t<sub>0</sub> and t<sub>2</sub>. However, we assume that planktonic organisms were probably not adversely affected, i.e. killed, by the temperature at least until t<sub>2</sub> because of the positive development of some groups

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and ongoing transcription of rRNA indicating active cell growth and activity (Schaechter et al. 1958, Poulsen et al. 1993, Lanzén et al. 2011).

OA effects on natural planktonic communities especially from tropical regions are poorly understood and require further investigation. Incubations over several days provide the advantage that they sometimes allow rare taxa to flourish (even if artificially), hence allowing effects of OA to be discovered that would have stayed obscure otherwise. OA might particularly affect the rare and some thermophilic taxa believed to cope well with climate change without considering their actual response to OA. Although being rare, some taxa could be of human interest, because they produce important metabolites or are somehow involved in the complex interactions of the oceanic food web. Diatoms and dinoflagellates as important primary producers and base of the food web seemed particularly vulnerable to OA and incubationinduced community changes. Overall, unravelling the interactive or opposed effects of a warming and simultaneously acidifying ocean on different eukaryotic species from natural planktonic communities is a difficult task due to differential intra- and interspecies responses. The consequences for whole trophic networks of a future ocean will consequently be even harder to predict.

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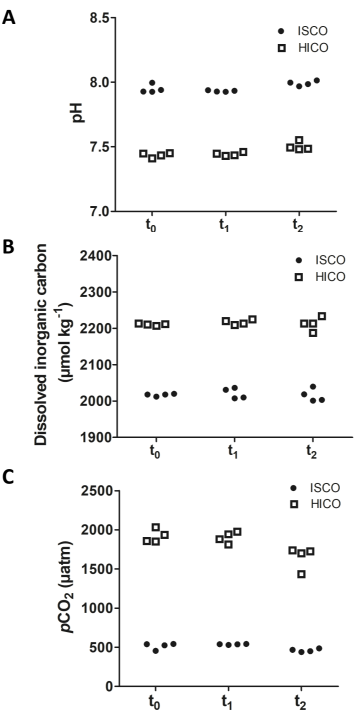
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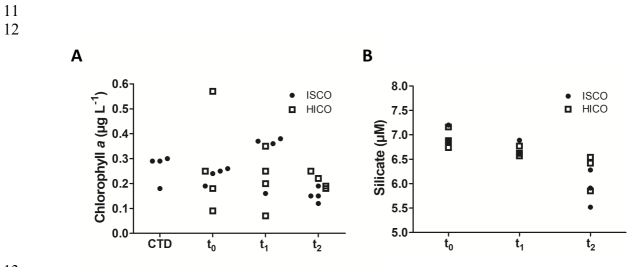
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- 2 Figures

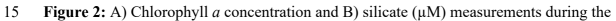


**Figure 1:** Carbonate chemistry measurements during the experiments. A) Seawater pH (Total scale) measured directly during the experiments at *in situ* temperature (c. 25 °C). B) Dissolved inorganic carbon (DIC) measured later from subsamples of the experiment media. C) Seawater  $pCO_2$  at *in situ* temperature calculated from the pH and DIC measurements shown in A) and B). For samples with no DIC measurement, the average DIC across all timepoints for all experiments at a similar CO<sub>2</sub> level was used instead (i.e. 2018 µmol kg<sup>-1</sup> for ISCO experiments, and 2213 µmol kg<sup>-1</sup> for HICO experiments).

Ocean acidification effects on planktonic eukaryotes

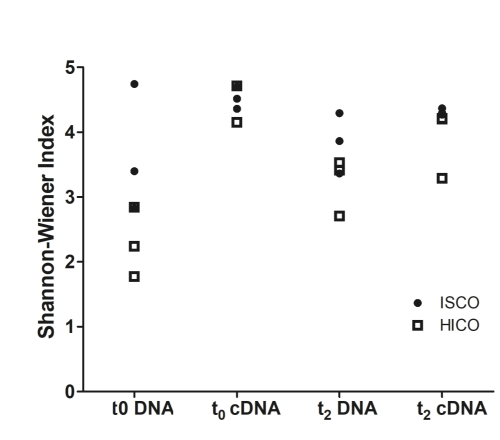


## 13 14



16 experiments (n=4).

#### Ocean acidification effects on planktonic eukaryotes



18 19

**Figure 3:** Calculated Shannon-Wiener indices (species richness) of ISCO and HICO-treated

20 samples. HICO= high  $pCO_2$ , ISCO= in situ  $pCO_2$ 

### Ocean acidification effects on planktonic eukaryotes

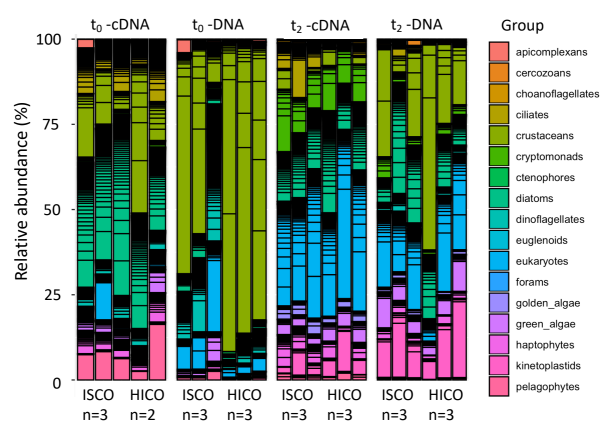
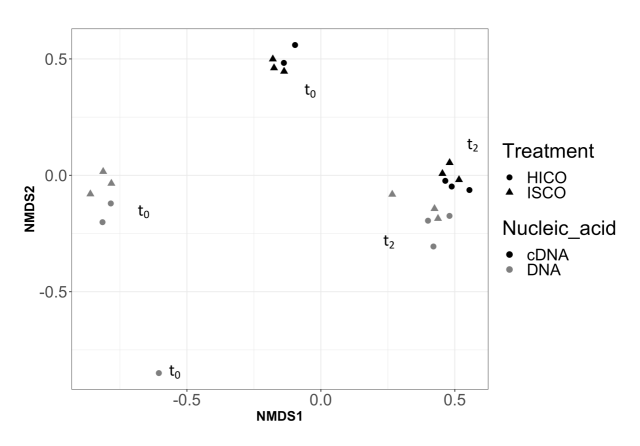
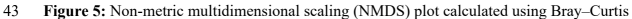


Figure 4: Community composition of taxa expressed as relative abundance (%) of operational
 taxonomic units (OTUs) for ISCO and HICO-treated samples of three replicates (with
 exception of one sample).

Ocean acidification effects on planktonic eukaryotes





- 44 dissimilarity. Circles indicate sample clusters. Stress=0.070, HICO= high  $pCO_2$ , ISCO= in
- 45 *situ p*CO<sub>2</sub>

Ocean acidification effects on planktonic eukaryotes

- 46 **Table 1:** Operational taxonomic units (OTUs) that are significantly affected ( $p \le 0.05$ ) by the
- 47 ocean acidification treatment (n=3 for each group). The value in the  $ISCO_m$  and  $HICO_m$
- 48 column represents the mean of relative abundance (%) for three biological replicates.
- 49 Asteriks indicate significance (Sig.) level (\*=  $p \le 0.05$ , \*\*=  $p \le 0.01$ , \*\*\*=  $p \le 0.001$ ).
- 50 CV=coefficient of variation, STD=standard deviation

#### DNA t<sub>0</sub>

DIVINU										
OTU	Group	Species	ISCO <sub>m</sub>	STD	CV (%)	HICO <sub>m</sub>	STD	CV (%)	p-value	Sig.
OTU_418	ciliates	Eutintinnus_cfapertus	0.015	0.002	16.6	0	0.000		0.009	**
OTU_490	ciliates	Rimostrombidium_lacustris	0.010	0.003	35.3	0	0.000		0.039	*
OTU_431	ciliates	Aristerostoma_marinum	0.022	0.008	34.5	0.004	0.007	173.2	0.045	*
OTU_888	crustaceans	Calanus_finmarchicus	0.002	0.004	173.2	0.019	0.001	4.0	0.015	*
OTU_967	cryptomonads	Geminigera_cryophila	0.007	0.001	16.6	0	0.000		0.009	**
OTU_793	cryptomonads	Goniomonas_spSH8	0.032	0.004	13.9	0.006	0.000	4.0	0.010	**
OTU_61	cryptomonads	uncultured_Proteomonas	0.012	0.003	22.2	0	0.000		0.016	*
OTU_25	diatoms	Fragilariopsis_doliolus	0.436	0.021	4.9	0.117	0.075	63.8	0.013	*
OTU_143	diatoms	Grammonema_striatula	0.076	0.019	25.7	0.008	0.004	47.3	0.023	*
OTU_34	diatoms	Bacillaria_paxillifer	0.254	0.058	23.0	0.064	0.012	18.9	0.026	*
OTU_659	diatoms	Pseudonitzschia_subcurvata	0.002	0.004	173.2	0.012	0.000	4.0	0.042	*
OTU_138	diatoms	Pseudonitzschia_australis	0.078	0.025	31.8	0.014	0.004	24.6	0.045	*
OTU_210	diatoms	Pseudonitzschia_subcurvata	0.052	0.016	30.5	0.011	0.019	173.2	0.046	*
OTU_174	diatoms	Pleurosigma_sp102	0.097	0.030	31.2	0.031	0.013	43.0	0.047	*
OTU_312	dinoflagellates	Herdmania_litoralis	0.327	0.036	11.0	0.056	0.006	10.0	0.005	**
OTU_84	dinoflagellates	Vulcanodinium_rugosum	1.880	0.302	16.1	0.757	0.221	29.2	0.008	**
OTU_365	dinoflagellates	Gyrodinium_spHJ2011	0.093	0.013	13.6	0.035	0.005	13.4	0.009	**
OTU_1708	dinoflagellates	uncultured_Syndiniales	0.012	0.003	22.2	0	0.000		0.016	*
OTU_801	dinoflagellates	Herdmania_litoralis	0.029	0.008	27.9	0.004	0.007	173.2	0.018	*
OTU_1195	dinoflagellates	uncultured_Amoebophrya	0.032	0.010	30.7	0.002	0.004	173.2	0.024	*
OTU_321	dinoflagellates	Amphidinium_belauense	0.146	0.040	27.6	0.036	0.021	58.8	0.024	*
OTU_527	dinoflagellates	Scrippsiella_cferinaceus	0.083	0.012	14.4	0.027	0.021	77.8	0.027	*
OTU_716	dinoflagellates	Peridinium_sociale	0.021	0.005	22.2	0.006	0.006	101.1	0.030	*
OTU_713	dinoflagellates	Scrippsiella_hangoei	0.021	0.005	21.9	0.006	0.006	98.5	0.031	*
OTU_1209	dinoflagellates	Karenia_mikimotoi	0.032	0.010	33.0	0.004	0.004	86.7	0.034	*
OTU_794	dinoflagellates	uncultured_Amoebophrya	0.065	0.024	36.9	0.004	0.007	173.2	0.038	*
OTU_988	dinoflagellates	uncultured_dinoflagellate	0.010	0.003	35.3	0	0.000		0.039	*
OTU_1764	dinoflagellates	uncultured_Amoebophrya	0.010	0.003	35.7	0	0.000		0.040	*
OTU_797	dinoflagellates	uncultured_Syndiniales	0.024	0.010	41.6	0.002	0.004	173.2	0.049	*
OTU_450	dinoflagellates	Islandinium_tricingulatum	0.044	0.015	34.5	0.008	0.003	39.6	0.049	*
OTU_356	eukaryotes	Vacuolaria_virescens	0.032	0.004	14.2	0.002	0.004	173.2	0.001	**
OTU_1217	eukaryotes	Larcopyle_butschlii	0.012	0.003	22.2	0	0.000		0.016	*
OTU_1725	eukaryotes	Picobiliphyte_spMS58411	0.010	0.003	35.3	0	0.000		0.039	*
OTU_373	eukaryotes	Artostrobus_sp2014	0.088	0.025	27.9	0.034	0.020	59.9	0.043	*
OTU_445	golden_algae	Spumellalike_flagellate_JBNZ43	0.007	0.001	16.6	0	0.000		0.009	**
OTU_792	green_algae	uncultured_Chlorophyta	0.021	0.005	22.2	0.002	0.004	173.2	0.006	**
OTU_1809	green_algae	uncultured_Chlorophyta	0.007	0.001	16.6	0	0.000		0.009	**
OTU_2192	green_algae	uncultured_Chlorophyta	0.007	0.001	16.6	0	0.000		0.009	**
OTU_795	green_algae	uncultured_Chlorophyta	0.027	0.008		0	0.000		0.025	*
OTU_112	green_algae	uncultured_Chlorophyta	0.067	0.021	30.6	0.015	0.016	110.8	0.028	*
OTU_524	kinetoplastids	Kentomonas_sorsogonicus	0.015	0.002	16.6	0.002	0.003	173.2	0.009	**
OTU_305	pelagophytes	Coccoid_pelagophyte_CCMP1395	0.070	0.025	35.9	0.011	0.014	126.5	0.036	*

OTU	Group	Species	ISCO <sub>m</sub>	STD	CV (%)	HICOm	STD	CV (%)	p-value	Sig.
OTU_251	ciliates	Eutintinnus_tenuis	0.034	0.009	27.1	0.000	0.000		0.024	*
OTU_222	ciliates	Laboea_strobila	0.064	0.016	25.5	0.023	0.008	36.3	0.031	*
OTU_1046	ciliates	uncultured_Tintinnida	0.013	0.005	37.9	0	0.000		0.045	*
OTU_61	cryptomonads	uncultured_Proteomonas	0.652	0.228	35.0	0.010	0.005	49.9	0.039	*
OTU_793	cryptomonads	Goniomonas_spSH8	0.027	0.010	37.9	0	0.000		0.045	*
OTU_145	diatoms	Synedra_spp517	0.543	0.125	23.0	0.163	0.111	67.7	0.017	*
OTU 54	diatoms	Stephanopyxis turris	1.153	0.413	35.8	0.209	0.272	130.0	0.037	*
OTU 209	diatoms	Trieres chinensis	0.099	0.033	32.9	0.024	0.025	102.4	0.037	*
OTU 52	diatoms	Trieres chinensis	1.674	0.136	8.1	0.385	0.493	128.2	0.038	*
OTU 505	diatoms	Stephanopyxis turris	0.005	0.004		0.017	0.005		0.042	*
OTU 713	dinoflagellates	Scrippsiella hangoei	0.013	0.005		0.003		173.2	0.050	*
OTU 2077	eukaryotes	Bicosoeca vacillans	0.008	0.000		0	0.000	17012	0.001	***
OTU_115	eukaryotes	Rhizochromulina cf. marina	0.299	0.094		0.078	0.034	44 3	0.042	*
DTU 62	eukaryotes	Pirsonia guinardiae	0.592	0.196		0.100	0.047		0.042	*
DTU_401	eukaryotes	Rhizidiomyces apophysatus	0.010	0.004		0	0.000	17.1	0.046	*
OTU 442	golden algae	Spumella_spGOT220	0.010	0.004		0.002		173.2	0.037	*
_										*
OTU_602	golden_algae	Hibberdia_magna	0.040	0.016		0.007		106.3	0.050	***
DTU_1946	green_algae	uncultured_Chlorophyta	0.008	0.000		0	0.000		0.001	***
DTU_2405	green_algae	Micromonas_commoda	0.008	0.000		0	0.000	100 6	0.001	
OTU_851	green_algae	uncultured_Chlorophyta	0.032	0.008		0.008		100.6	0.018	*
DTU_523	green_algae	Prasinoderma_coloniale	0.048	0.009		0.015	0.015		0.041	*
DTU_192	haptophytes	Haptophyta_environmental_samples	0.114	0.024		0.010	0.009		0.010	**
DTU_168	haptophytes	Haptophyta_environmental_samples	0.096	0.028	29.7	0.005	0.004	87.7	0.029	*
OTU_412	haptophytes	Haptophyta_environmental_samples	0.065	0.021	31.8	0.007	0.007	99.4	0.029	*
DTU_394	haptophytes	prymnesiophyte_symbiont_4	0.037	0.013	35.1	0.005	0.004	86.6	0.040	*
DTU_246	haptophytes	Haptophyta_environmental_samples	0.043	0.018	41.4	0.005	0.009	173.2	0.047	*
DNA t <sub>2</sub>										
DTU	Group	Species	ISCO <sub>m</sub>	STD	CV (%)	HICOm	STD	CV (%)	p-value	Sig
DTU 222	ciliates	Laboea strobila	0.315	0.044	14.0	0.093	0.065	69.7	0.011	*
DTU 349	ciliates	uncultured Tintinnida	0.038	0.011	28.0	0	0.000		0.025	*
DTU 351	ciliates	Strombidium biarmatum	0.087	0.028	32.2	0.013	0.013	100.7	0.029	*
DTU 1822	ciliates	Eutintinnus tenuis	0.011	0.004		0.000	0.000		0.036	*
DTU 331	ciliates	uncultured ciliate	0.080	0.023		0.025	0.007	27.1	0.046	*
DTU_54	diatoms	Stephanopyxis_turris	2.612	0.242		0.293		128.2	0.002	
DTU 52					2.0	0.275				**
_			1.031		46	0.240	0.151	62.9		**
TII 1492	diatoms	Trieres_chinensis Coccouries_placentula	1.031	0.047		0.240	0.151		0.007	**
_	diatoms	Cocconeis_placentula	0.002	0.047 0.004	173.2	0.017	0.004	21.7	0.007 0.008	** **
DTU_475	diatoms diatoms	Cocconeis_placentula Chaetoceros_sp	0.002 0.004	0.047 0.004 0.004	173.2 86.6	0.017 0.025	0.004 0.006	21.7 25.9	0.007 0.008 0.015	** ** *
DTU_475 DTU_169	diatoms diatoms diatoms	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum	0.002 0.004 0.034	0.047 0.004 0.004 0.003	173.2 86.6 10.3	0.017 0.025 0.141	0.004 0.006 0.028	21.7 25.9 19.6	0.007 0.008 0.015 0.020	** ** *
DTU_475 DTU_169 DTU_145	diatoms diatoms diatoms diatoms	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517	0.002 0.004 0.034 0.770	0.047 0.004 0.004 0.003 0.161	173.2 86.6 10.3	0.017 0.025 0.141 0.303	0.004 0.006 0.028 0.156	21.7 25.9 19.6 51.5	0.007 0.008 0.015 0.020 0.023	** ** * *
DTU_475 DTU_169 DTU_145 DTU_1078	diatoms diatoms diatoms diatoms dinoflagellates	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium	0.002 0.004 0.034 0.770 0	0.047 0.004 0.004 0.003 0.161 0.000	173.2 86.6 10.3 20.9	0.017 0.025 0.141 0.303 0.010	0.004 0.006 0.028 0.156 0.004	21.7 25.9 19.6 51.5 36.1	0.007 0.008 0.015 0.020 0.023 0.041	** ** * * *
DTU_475 DTU_169 DTU_145 DTU_1078 DTU_2303	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans	0.002 0.004 0.034 0.770 0 0.036	0.047 0.004 0.003 0.161 0.000 0.004	173.2 86.6 10.3 20.9 10.7	0.017 0.025 0.141 0.303 0.010 0.006	0.004 0.006 0.028 0.156 0.004 0.006	21.7 25.9 19.6 51.5 36.1 100.8	0.007 0.008 0.015 0.020 0.023 0.041 0.004	** ** * * * * * *
0TU_475 0TU_169 0TU_145 0TU_1078 0TU_2303 0TU_242	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes	0.002 0.004 0.034 0.770 0 0.036 0.032	0.047 0.004 0.003 0.161 0.000 0.004 0.006	173.2 86.6 10.3 20.9 10.7 19.1	0.017 0.025 0.141 0.303 0.010 0.006 0.006	0.004 0.006 0.028 0.156 0.004 0.006 0.006	21.7 25.9 19.6 51.5 36.1 100.8 100.7	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007	** ** * * * * * * * * * * * *
DTU_475 DTU_169 DTU_145 DTU_1078 DTU_2303 DTU_742 DTU_765	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004	173.2 86.6 10.3 20.9 10.7 19.1 173.2	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.006	0.004 0.006 0.028 0.156 0.004 0.006 0.006 0.004	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008	** ** ** ** ** **
DTU_475         DTU_169         DTU_145         DTU_1078         DTU_2303         DTU_742         DTU_765         DTU_113	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360	0.004 0.028 0.156 0.004 0.006 0.006 0.004 0.162	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017	***
DTU_475 DTU_169 DTU_145 DTU_1078 DTU_2303 DTU_742 DTU_765 DTU_113 DTU_62	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581	0.004 0.006 0.028 0.156 0.004 0.006 0.006 0.004	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008	* * * * * * * * * *
DTU_475 DTU_169 DTU_145 DTU_1078 DTU_2303 DTU_742 DTU_765 DTU_113 DTU_62	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360	0.004 0.028 0.156 0.004 0.006 0.006 0.004 0.162	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017	*** * * * * * * * * *
DTU_475 DTU_169 DTU_145 DTU_1078 DTU_2303 DTU_742 DTU_765 DTU_113 DTU_62 DTU_187	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581	0.004 0.006 0.028 0.156 0.004 0.006 0.006 0.004 0.162 0.249 0.066	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031	* * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_1078         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_62         DTU_187         DTU_2047	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.006 0.017 0.360 0.581 0.177	0.004 0.006 0.028 0.156 0.004 0.006 0.006 0.004 0.162 0.249 0.066	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040	* * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_1078         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_62         DTU_187         DTU_2047         DTU_1358	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 35.1	0.017 0.025 0.141 0.303 0.010 0.006 0.017 0.360 0.581 0.177 0.002	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048	* * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_1078         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_62         DTU_187         DTU_2047         DTU_1358         DTU_145	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 35.1 30.5	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581 0.177 0.002 0.006	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.004	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2 18.4	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.000	* * * * * * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_145         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_62         DTU_187         DTU_2047         DTU_1358         DTU_74         DTU_74	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0 0.089	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.0027	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 35.1 30.5 22.1	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581 0.177 0.002 0.006 0.213	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.000 0.000 0.039	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2 18.4 43.4	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.000 0.048	* * * * * * * * * * * * * * * * * * * *
DTU_475 DTU_169 DTU_169 DTU_145 DTU_2303 DTU_742 DTU_765 DTU_765 DTU_113 DTU_62 DTU_187 DTU_2047 DTU_1358 DTU_74 DTU_712 DTU_519	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0 0.089 0.057	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 35.1 30.5 22.1 10.9	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581 0.177 0.002 0.006 0.213 0.008	0.004 0.006 0.028 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.000 0.039 0.004	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2 18.4 43.4 89.6	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.040 0.048 0.000 0.014 0.016	* * * * * * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_145         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_2047         DTU_1358         DTU_742         DTU_745         DTU_765         DTU_113         DTU_2047         DTU_1358         DTU_74         DTU_74         DTU_519         DTU_264	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta uncultured_Chlorophyta Haptophyta_environmental_samples uncultured_marine_haptophyte	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0 0.089 0.057 0.095 0.131	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.024	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 35.1 30.5 22.1 10.9 18.0	0.017 0.025 0.141 0.303 0.010 0.006 0.010 0.360 0.360 0.581 0.077 0.002 0.006 0.213 0.008 0.015 0.027	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.000 0.039 0.004 0.013 0.014	$\begin{array}{c} 21.7\\ 25.9\\ 19.6\\ 51.5\\ 36.1\\ 100.8\\ 100.7\\ 21.7\\ 44.9\\ 42.8\\ 37.0\\ 173.2\\ 2.2\\ 18.4\\ 43.4\\ 89.6\\ 53.4 \end{array}$	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.007 0.007 0.017 0.031 0.040 0.048 0.000 0.048 0.000 0.014 0.001 0.001 0.001 0.001 0.000 0.023 0.041 0.004 0.004 0.007 0.007 0.007 0.004 0.007 0.004 0.007 0.001 0.004 0.007 0.001 0.004 0.007 0.001 0.004 0.007 0.001 0.004 0.007 0.001 0.004 0.004 0.004 0.007 0.001 0.004 0.007 0.004 0.004 0.004 0.007 0.007 0.004 0.004 0.007 0.007 0.004 0.000 0.004 0.004 0.000 0.004 0.000 0.004 0.000 0.001 0.005 0.001 0.005 0.	* * * * * * * * * * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_145         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_2047         DTU_2047         DTU_21358         DTU_742         DTU_2047         DTU_159         DTU_519         DTU_264         DTU_2047	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae haptophytes haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta uncultured_Chlorophyta Haptophyta_environmental_samples uncultured_marine_haptophyte Haptophyta_environmental_samples	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 0.324 0.011 0 0.089 0.057 0.095 0.131 0.338	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.024 0.065	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 35.1 30.5 22.1 10.9 18.0 19.2	0.017 0.025 0.141 0.303 0.010 0.006 0.010 0.360 0.360 0.581 0.177 0.002 0.006 0.213 0.008 0.015 0.027 0.080	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.000 0.039 0.004 0.013 0.014 0.064	$\begin{array}{c} 21.7\\ 25.9\\ 19.6\\ 51.5\\ 36.1\\ 100.8\\ 100.7\\ 21.7\\ 44.9\\ 42.8\\ 37.0\\ 173.2\\ 2.2\\ 18.4\\ 43.4\\ 89.6\\ 53.4\\ 80.2 \end{array}$	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.003 0.040 0.040 0.048 0.000 0.048 0.000 0.014 0.001 0.005 0.008	* * * * * * * * * * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_145         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_62         DTU_187         DTU_2047         DTU_1358         DTU_112         DTU_519         DTU_264         DTU_911	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae green_algae haptophytes haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta uncultured_Chlorophyta Haptophyta_environmental_samples uncultured_marine_haptophyte Haptophyta_environmental_samples Haptophyta_environmental_samples	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 0.324 0.011 0 0.089 0.057 0.095 0.131 0.338 0.099	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.024 0.024 0.065 0.019	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 35.1 30.5 22.1 10.9 18.0 19.2 19.2	0.017 0.025 0.141 0.303 0.010 0.006 0.017 0.360 0.581 0.177 0.002 0.006 0.213 0.008 0.015 0.027 0.080 0.006	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.000 0.039 0.004 0.013 0.014 0.064 0.006	$\begin{array}{c} 21.7\\ 25.9\\ 19.6\\ 51.5\\ 36.1\\ 100.8\\ 100.7\\ 21.7\\ 44.9\\ 42.8\\ 37.0\\ 173.2\\ 2.2\\ 18.4\\ 43.4\\ 89.6\\ 53.4\\ 80.2\\ 100.7\\ \end{array}$	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.000 0.014 0.016 0.001 0.005 0.008 0.008	* * * * * * * * * * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_145         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_62         DTU_187         DTU_1358         DTU_12047         DTU_158         DTU_519         DTU_264         DTU_910         DTU_397         DTU_192	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae green_algae haptophytes haptophytes haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta uncultured_Chlorophyta Haptophyta_environmental_samples uncultured_marine_haptophyte Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 0.324 0.011 0 0.089 0.057 0.095 0.131 0.338 0.099 0.626	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.024 0.013 0.019 0.137	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 30.5 22.1 10.9 18.0 19.2 19.2 21.8	0.017 0.025 0.141 0.303 0.010 0.006 0.017 0.360 0.581 0.177 0.002 0.006 0.213 0.008 0.213 0.008 0.213 0.008 0.215 0.027 0.080 0.006 0.007	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.249 0.066 0.004 0.009 0.004 0.039 0.004 0.013 0.014 0.064 0.006 0.071	$\begin{array}{c} 21.7\\ 25.9\\ 19.6\\ 51.5\\ 36.1\\ 100.8\\ 100.7\\ 21.7\\ 44.9\\ 42.8\\ 37.0\\ 173.2\\ 2.2\\ 18.4\\ 43.4\\ 89.6\\ 53.4\\ 80.2\\ 100.7\\ \end{array}$	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.000 0.014 0.001 0.005 0.008 0.008 0.008 0.008 0.008 0.009	* * * * * * * * * * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_1078         DTU_2303         DTU_742         DTU_745         DTU_113         DTU_62         DTU_1078         DTU_2047         DTU_1358         DTU_74         DTU_1519         DTU_264         DTU_192         DTU_2037	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae green_algae haptophytes haptophytes haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta uncultured_Chlorophyta Haptophyta_environmental_samples uncultured_marine_haptophyte Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0 0.089 0.057 0.095 0.131 0.338 0.099 0.626 0.015	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.024 0.013 0.010 0.024 0.0137 0.004	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 30.5 22.1 10.9 18.0 19.2 19.2 21.8 24.1	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.360 0.360 0.213 0.008 0.015 0.027 0.008 0.015 0.027 0.080 0.006 0.077 0	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.249 0.066 0.004 0.009 0.004 0.039 0.004 0.013 0.014 0.064 0.006 0.071 0.000	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2 18.4 43.4 89.6 53.4 80.2 100.7 91.9	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.000 0.014 0.001 0.005 0.008 0.008 0.009 0.019	* * * * * * * * * * * * * * * * * * * *
DTU_475         DTU_169         DTU_145         DTU_145         DTU_1078         DTU_2303         DTU_742         DTU_745         DTU_165         DTU_1765         DTU_187         DTU_2047         DTU_113         DTU_187         DTU_1145         DTU_187         DTU_1158         DTU_112         DTU_519         DTU_264         DTU_191         DTU_2037         DTU_246	diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae green_algae haptophytes haptophytes haptophytes haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta Uncultured_Chlorophyta Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0 0.089 0.057 0.095 0.131 0.338 0.099 0.626 0.015 0.237	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.024 0.065 0.019 0.137 0.004 0.061	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 30.5 22.1 10.9 18.0 19.2 19.2 21.8 24.1 25.9	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581 0.177 0.002 0.006 0.213 0.008 0.015 0.027 0.080 0.015 0.027 0.080 0.006	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.003 0.004 0.013 0.014 0.013 0.014 0.064 0.071 0.000 0.006	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2 18.4 43.4 89.6 53.4 80.2 100.7 91.9 99.3	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.048 0.000 0.014 0.016 0.001 0.005 0.008 0.008 0.009 0.019 0.022	*** ***********************************
DTU_475         DTU_169         DTU_145         DTU_145         DTU_1078         DTU_2303         DTU_742         DTU_765         DTU_113         DTU_62         DTU_187         DTU_1358         DTU_12047         DTU_1519         DTU_264         DTU_192         DTU_2037         DTU_246         DTU_246         DTU_1044	diatoms diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae haptophytes haptophytes haptophytes haptophytes haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta Haptophyta_environmental_samples uncultured_marine_haptophyte Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0 0.089 0.089 0.089 0.095 0.131 0.338 0.099 0.626 0.015 0.237 0.023	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.022 0.013 0.010 0.024 0.065 0.019 0.137 0.004 0.004 0.004	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 35.1 30.5 22.1 10.9 18.0 19.2 19.2 21.8 24.1 25.9 16.3	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581 0.177 0.002 0.006 0.213 0.002 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.009 0.004 0.003 0.004 0.013 0.014 0.014 0.064 0.006 0.071 0.000 0.006 0.006	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2 18.4 43.4 89.6 53.4 80.2 100.7 91.9 99.3 99.3	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.000 0.014 0.016 0.001 0.005 0.008 0.008 0.009 0.019 0.022 0.022	* * * * * * * * * * * * * * * * * * * *
DTU_1482 DTU_475 DTU_169 DTU_145 DTU_145 DTU_2303 DTU_742 DTU_742 DTU_765 DTU_113 DTU_62 DTU_162 DTU_1358 DTU_2047 DTU_1358 DTU_74 DTU_1358 DTU_74 DTU_112 DTU_519 DTU_264 DTU_191 DTU_297 DTU_2037 DTU_2037 DTU_246 DTU_1044 DTU_112	diatoms diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae haptophytes haptophytes haptophytes haptophytes haptophytes haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta Haptophyta_environmental_samples uncultured_marine_haptophyte Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0 0.089 0.057 0.095 0.131 0.338 0.099 0.626 0.015 0.237 0.023 0.085	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.024 0.065 0.019 0.137 0.004 0.061 0.004 0.004	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 30.5 22.1 10.9 18.0 19.2 19.2 21.8 24.1 25.9 16.3 20.0	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581 0.177 0.002 0.006 0.213 0.002 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.003 0.004 0.003 0.004 0.013 0.014 0.064 0.006 0.071 0.006 0.006 0.006 0.006	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2 18.4 43.4 89.6 53.4 80.2 100.7 91.9 99.3 99.3 87.6	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.000 0.014 0.016 0.001 0.005 0.008 0.008 0.009 0.019 0.022 0.022 0.022 0.032	***************************************
DTU_475 DTU_169 DTU_169 DTU_145 DTU_2303 DTU_742 DTU_765 DTU_113 DTU_62 DTU_187 DTU_2047 DTU_1358 DTU_74 DTU_1358 DTU_74 DTU_120 DTU_120 DTU_120 DTU_264 DTU_191 DTU_2037 DTU_2037 DTU_246 DTU_1044	diatoms diatoms diatoms diatoms diatoms dinoflagellates eukaryotes eukaryotes eukaryotes eukaryotes eukaryotes green_algae green_algae green_algae haptophytes haptophytes haptophytes haptophytes haptophytes	Cocconeis_placentula Chaetoceros_sp Corethron_criophilum Synedra_spp517 uncultured_Gymnodinium Bicosoeca_vacillans haptophytes Pseudobodo_tremulans golden_algae Pirsonia_guinardiae Telonema_subtile Ciliophrys_infusionum Tetraselmis_spGSL018 uncultured_Chlorophyta Haptophyta_environmental_samples uncultured_marine_haptophyte Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples Haptophyta_environmental_samples	0.002 0.004 0.034 0.770 0 0.036 0.032 0.002 0.843 2.133 0.324 0.011 0 0.089 0.089 0.089 0.095 0.131 0.338 0.099 0.626 0.015 0.237 0.023	0.047 0.004 0.003 0.161 0.000 0.004 0.006 0.004 0.107 0.595 0.036 0.004 0.000 0.027 0.013 0.010 0.022 0.013 0.010 0.024 0.065 0.019 0.137 0.004 0.004 0.004	173.2 86.6 10.3 20.9 10.7 19.1 173.2 12.7 27.9 11.1 30.5 22.1 10.9 18.0 19.2 21.8 24.1 25.9 16.3 20.0 33.5	0.017 0.025 0.141 0.303 0.010 0.006 0.006 0.017 0.360 0.581 0.177 0.002 0.006 0.213 0.002 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027 0.027	0.004 0.006 0.028 0.156 0.004 0.006 0.004 0.162 0.249 0.066 0.004 0.009 0.004 0.003 0.004 0.013 0.014 0.014 0.064 0.006 0.071 0.000 0.006 0.006	21.7 25.9 19.6 51.5 36.1 100.8 100.7 21.7 44.9 42.8 37.0 173.2 2.2 18.4 43.4 89.6 53.4 80.2 100.7 91.9 99.3 99.3 87.6 67.3	0.007 0.008 0.015 0.020 0.023 0.041 0.004 0.007 0.008 0.017 0.031 0.040 0.048 0.000 0.014 0.016 0.001 0.005 0.008 0.008 0.009 0.019 0.022 0.022	*** * * * * * * * * * * * * * * * * * *

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- **Table 2:** Proportion (%) of significantly CO<sub>2</sub>-affected OTUs within the given relative abundance range for each sample

Rel. Abundance range of OTUs	ISCO-DNA-t0	HICO-DNA-t0	ISCO-DNA-t2	HICO-DNA-t2	ISCO-cDNA-t2	HICO-cDNA-t2
1-100%	9.4	0.0	12.0	0.0	13.4	0.0
0.1-1%	6.1	8.3	7.6	8.1	9.5	9.6
0.01-0.1%	6.4	5.1	6.9	3.3	7.8	6.8
0.001-0.01%	3.2	6.3	2.3	6.4	2.5	7.9
0-0.001%	0.0	0.9	0.0	0.5	0.1	0.2

Ocean acidification effects on planktonic eukaryotes

#### **Supplementary Material**

# Short-term responses to ocean acidification: effects on relative abundance of eukaryotic plankton from the tropical Timor Sea

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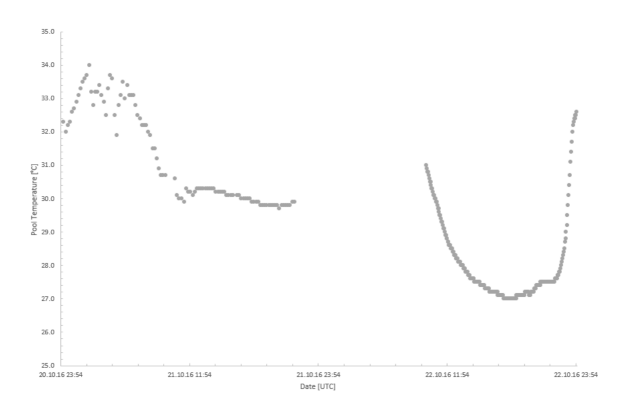
and Utrecht University, 1790 AB Den Burg, Texel, the Netherlands

<sup>5</sup>Leibniz Institute for Baltic Sea Research (IOW), 18119 Rostock, Germany

#Present address: Project management Jülich, 18069 Rostock, Germany

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#### Ocean acidification effects on planktonic eukaryotes



**Supplementary Figure S1:** Temperature of pool water (°C) surrounding incubated bottles. Dates refer to  $t_0$  to  $t_2$ , respectively. The data gap happened due to an unrecognized full memory of the measuring device.

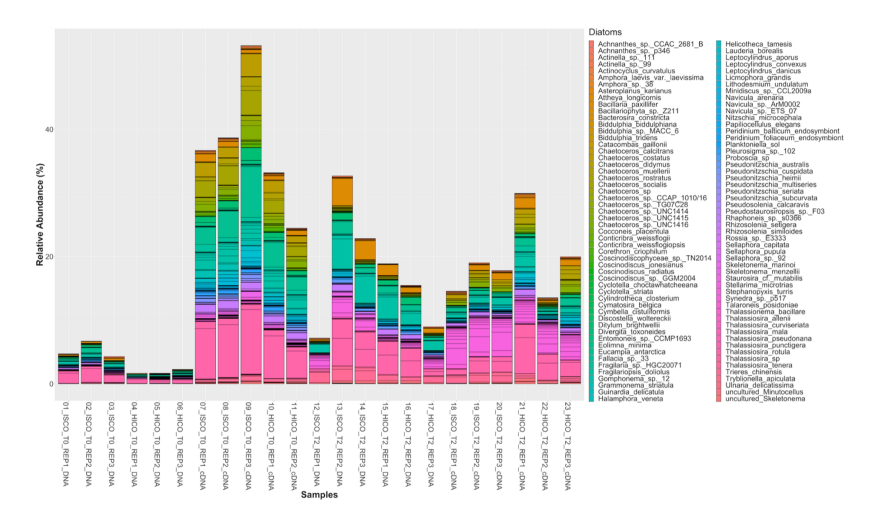
bioRxiv preprint doi: https://doi.org/10.1101/2020.04.30.068601; this version posted May 3, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

Ocean acidification effects on planktonic eukaryotes

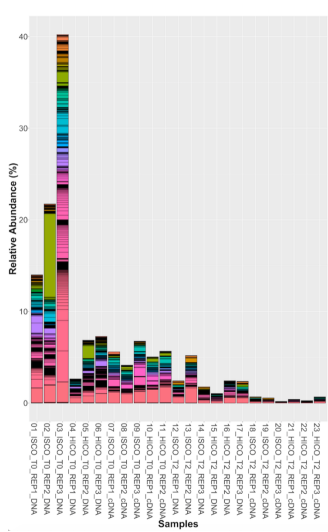
#### Sample OR NC NFC NDC NOTU ISCO T0 Rep1 DNA ISCO T0 Rep2 DNA ISCO T0 Rep3 DNA HICO T0 Rep1 DNA HICO T0 Rep2 DNA HICO T0 Rep3 DNA ISCO T0 Rep1 cDNA ISCO T0 Rep2 cDNA ISCO T0 Rep3 cDNA HICO T0 Rep1 cDNA HICO T0 Rep2 cDNA ISCO T2 Rep1 DNA ISCO T2 Rep2 DNA ISCO T2 Rep3 DNA HICO T2 Rep1 DNA HICO T2 Rep2 DNA HICO T2 Rep3 DNA ISCO T2 Rep1 cDNA ISCO T2 Rep2 cDNA ISCO T2 Rep3 cDNA HICO T2 Rep1 cDNA HICO T2 Rep2 cDNA HICO T2 Rep3 cDNA

#### Supplementary Table S1: Sequencing statistic

OR= original reads NC= number of contigs NFC= number of contigs after filtering NDC= number of dereplicated contigs NOTU= number of OTUs



**Supplementary Figure S2:** Relative abundance (%) of diatoms in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and 48 hours (t<sub>2</sub>) treatment exposure.



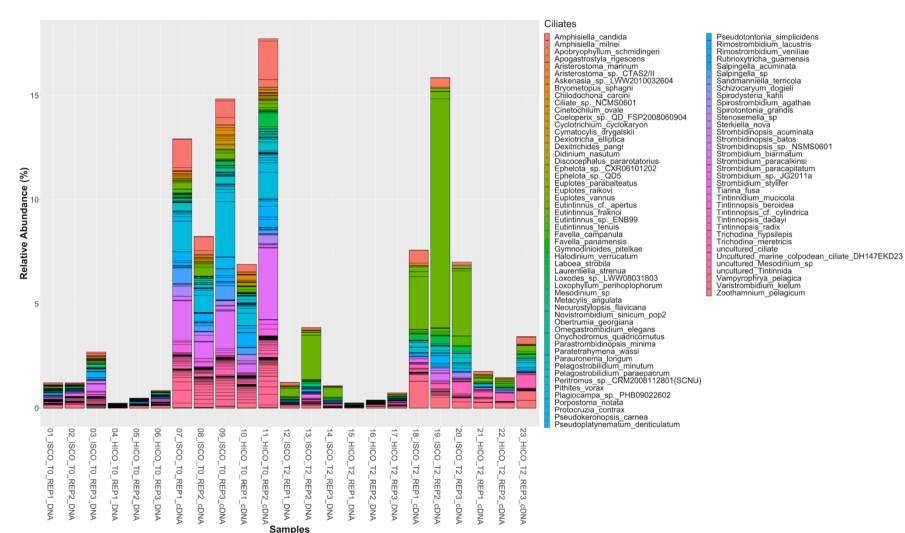
#### Dinoflagellates

Adenoides eludens Adunoodinium\_glandula Alexandrium\_beruvianum Alexandrium\_lamarense Arnoebophrya\_sp\_ex\_Akashiwo\_sanguineaum Arnoebophrya\_sp\_ex\_Caratium\_tripos Amoebophrya\_sp\_ex\_Conyaulax\_polygramma Amoebophrya\_sp\_ex\_Gonyaulax\_polygramma Amoebophrya\_sp\_ex\_Karlodinium\_instriatum Amoebophrya\_sp\_ex\_Prorocentrum\_minimum Amoebophrya\_sp\_ex\_Prorocentrum\_minimum Amoebophrya\_sp\_ex\_Prorocentrum\_minimum Amoebophrya\_sp\_ex\_Cropsiella\_sp Amoebophrya\_sp\_ex\_CC 1626 Amphidiniopsis\_uroënsis Amphidiniopsis\_uroënsis Amphidinium\_belauense Amphidinium\_carterae Aduncodinium glandula Amphidinium\_carterae Amphidinium\_sp.\_D1CMSTAC020 Archaeperidinium\_sp.provide Azadinium\_cf.\_poporum\_HJ2011 Azadinium\_dexteroporum Biecheleria brevisulcata Blastodinium\_contortum Blastodinium\_inornatum Blastodinium mangini Blastodinium navicula Blastodinium\_oviforme Blastodinium\_spinulosum Brandtodinium nutricula Bysmatrum caponii Calciodinellum\_operosum Chytriodinium\_sp.\_Atlantic Coolia\_malayensis Cryptoperidiniopsis brodyi cryptoperidiniopsid\_sp.\_Folly\_C5 cryptoperidiniopsoid\_sp.\_FL0Y\_C5 cryptoperidiniopsoid\_sp.\_FL0Z1 Dinophyceae\_sp.\_et.Thalassicolla\_nucleata Dinophyceae\_sp.\_UDMS0803 Dinophysis\_acuminata Dinophysis\_acuta Diplopsalis\_caspica Diplopsalopsis\_bomba Durinskia\_baltica Durinskia oculata Ensiculifera imariensis Erist-Quillera\_interensis Esoptrodinium\_sp.\_RP Euduboscquella\_sp.\_ex\_Favella\_markusovszkyi Euduboscquella\_sp.\_ex\_Favella\_panaemensis Euduboscquella\_sp.\_ex\_Tintjnnopsis\_cf.\_subacuta Fragilidium\_duplocampanaeforme Fragilidium\_sp. CCMP1920 Gloeodinium\_viscum

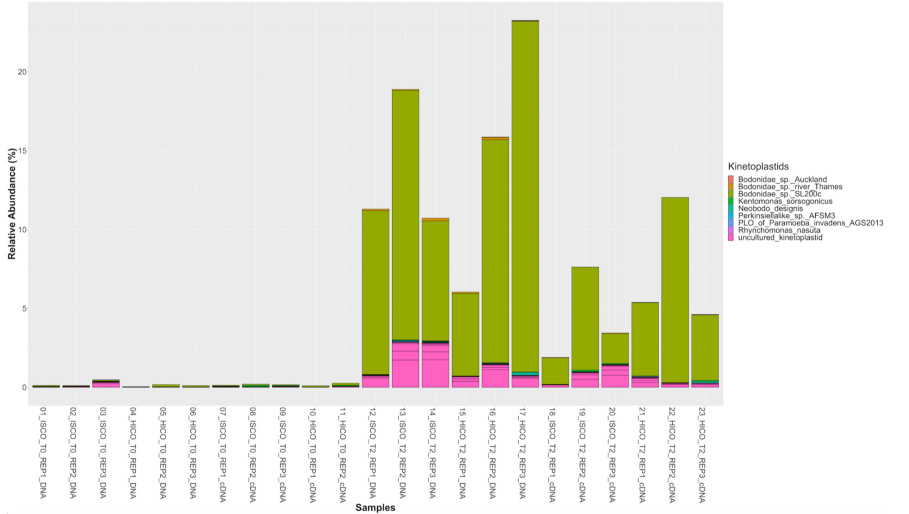
Goniodoma polvedricum Gonyaulax\_polygramma Gymnodinium\_cf.\_mikimotoi Gymnodinium\_impudicum Gymnodinium limneticum Gymnodinium litoralis Gymnodinium\_sp.\_ex\_Spongotrochus\_glacialis Gymodinium\_sp.\_ex\_Spongotrochus\_glacialis Gyrodinium\_sp.\_H2011 Hematodinium\_sp.\_ex\_2011 Hematodinium\_sp.\_ex\_Crangon\_crangon Herdmania\_litoralis Heterocapsa niei leterocapsa rotundata Heterocapsa\_triquetra Histioneis\_sp\_FTL62 Islandinium\_tricingulatum Karenia\_mikimotoi Karenia selliformis Karlodinium veneficum Kryptoperidinium foliaceum Lepidodinium viride evanderina fissa Lingulodinium\_polyedra Margalefidinium\_fulvescens Margalefidinium\_polykrikoides Nusuttodinium amphidinioides Oodinium\_pouchetii Ornithocercus\_magnificus Paragymnodinium\_shiwhaense Parvodinium\_inconspicuum Pelagodinium\_beii Pentapharsodinium tvrrhenicum Peridinium sociale Peridinium\_sociale Peridinium\_sociale Pfiesteria sp. B112456 Pfiesterialike\_dinoflagellate\_Lucy Pfiesterialike\_sp.\_CCMP1838 Pfiesterialike\_sp.\_NC\_LucyV27 Pheloregroup\_contended Phalacroma\_oxytoxoides Phalacroma rapa Pheopolykrikos hartmanni Piscinoodinium\_sp.\_RWL2007 Posoniella\_tricarinelloides Prorocentrum\_emarginatum Prorocentrum\_mexicanum Prorocentrum minimum Prorocentrum\_sipadanense Prorocentrum triestinum Proterythropsis\_sp.\_BSL2009a Protoceratium reticulatum Protodinium\_simplex Protoperidinium\_americanum Protoperidinium\_bipes

Protoperidinium conicum Protoperidinium depressum Protoperidinium\_divergens Protoperidinium\_excentricum Protoperidinium monovelum Protoperidinium monovelum Pyrocystis\_sp.\_CCAP\_1131/2 Pyrophacus\_steinii Scrippsiella\_cf.\_erinaceus Scrippsiella\_hangoei Scrippsiella trochoidea Sinophysis\_stenosoma Stoeckeria\_sp.\_SSSC09 Symbiodinium\_microadriaticum Syndinium\_sp.\_ex\_Corycaeus\_sp Syndinium\_turbo Takayama\_cf.\_pulchellum Theleodinium\_calcisporum Thoracosphaera\_heimii Torodinium teredo Tripos\_tenuis uncultured Amoebophrya uncultured dinoflagellate uncultured Duboscquella uncultured\_Gymnodinium uncultured\_Heterocapsa uncultured\_marine\_dinoflagellate uncultured marine Syndiniales uncultured\_Syndiniales unidentified\_syndiniales unidentified\_symbiotic\_dinoflagellate\_BBSR\_159 Vulcanodinium\_rugosum Warnowia\_sp.\_BSL2009a Yihiella\_yeosuensis

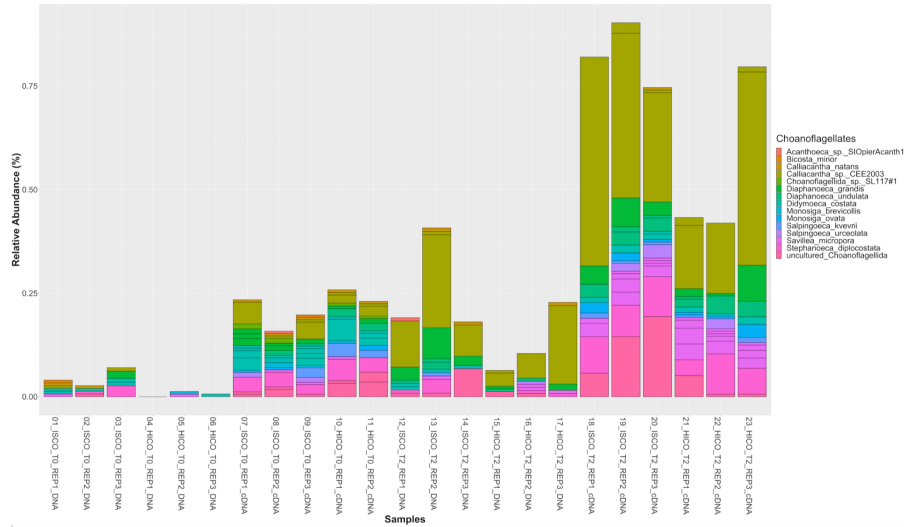
**Supplementary Figure S3:** Relative abundance (%) of dinoflagellates in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and 48 hours (t<sub>2</sub>) treatment exposure.



**Supplementary Figure S4:** Relative abundance (%) of ciliates in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and 48 hours (t<sub>2</sub>) treatment exposure.

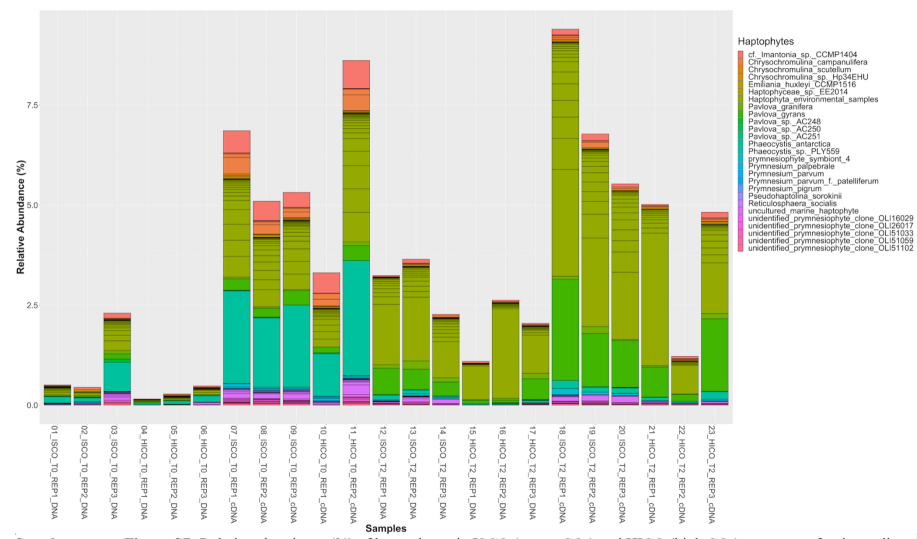


**Supplementary Figure S5:** Relative abundance (%) of kinetoplastids in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and 48 hours (t<sub>2</sub>) treatment exposure.



**Supplementary Figure S6:** Relative abundance (%) of choanoflagellates in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and 48 hours (t<sub>2</sub>) treatment exposure.

14



15 16

Supplementary Figure S7: Relative abundance (%) of haptophytes in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and

17 48 hours (t<sub>2</sub>) treatment exposure.

#### Crustaceans Acartia Longiremis Amphiascoides atopus Anthessius sp. New CaledoniaRJH2004 Calanus finmarchicus Centropäges furcatus Clausidium vancouverense Conchoecia sp. OC2001 Corycaeus speciosus Cyclopina gracilis Euchirella rostrata 75 Euchirella rostrata Eudiaptomus\_environmental\_sample Euderpina\_acutifrons Exumella\_mediterranea Macrocyclops\_albidus Macrocyclops\_albidus Meganycliphanes\_norvegica Oithona\_simplex\_ Oithona\_sp.\_2\_New\_CaledoniaRJH2004 Oithona\_sp.\_Roadstead\_of Brest Oncaea\_sp.\_Roadstead\_of\_Brest Pareucalanus\_attenuatus Subeucalanus\_pileatus Relative Abundance (%) Tessarabrachion\_oculatum Thysanopoda pectinata 25 0 01\_ISCO\_T0\_REP1\_DN/ 02\_ISCO\_T0\_REP2\_DNA 03\_ISCO\_T0\_REP3\_DNA 04\_HICO\_T0\_REP1\_DNA 05\_HICO\_T0\_REP2\_DNA 07\_ISCO\_T0\_REP1\_cDN/ 06\_HICO\_T0\_REP3\_DNA 08\_ISCO\_T0\_REP2 09\_ISCO\_T0\_REP3\_cDNA 10\_HICO\_T0\_REP1\_cDNA 11\_HICO\_T0\_REP2\_cDNA 12\_ISCO\_T2\_REP1\_DNA 13\_ISCO\_T2\_REP2\_DNA 14\_ISCO\_T2\_REP3\_DNA 15\_HICO\_T2\_REP1\_DNA 16\_HICO\_T2\_REP2\_DNA 17\_HICO\_T2\_REP3\_DNA 18\_ISCO\_T2\_REP1\_cDN/ 20\_ISCO\_T2\_REP3 21\_HICO\_T2\_REP1\_cDNA 22\_HICO\_T2\_REP2\_cDNA 23\_HICO\_T2\_REP3\_cDN# 19\_ISCO\_T2\_REP2\_cDN/ \_cDNA CDN/A

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20 21 22

Supplementary Figure S8: Relative abundance (%) of crustaceans in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and

Samples

48 hours (t<sub>2</sub>) treatment exposure.

Cryptomonads Cryptomonads Cryptophyta\_sp\_CRMAL01 Geminigera\_cryophila Goniomonas\_sp\_SH3 Goniomonas\_sp\_SH3 Goniomonas\_sp\_SH4 Hemiselmis\_andersenii Hemiselmis\_virescens Proteomonas\_sulcata Rhinomonas\_pauca Teleaulax\_amphioxeia uncultured\_Cryptophyta uncultured\_Geminigera uncultured\_Geniomonas uncultured\_Proteomonas 15 Relative Abundance (%) 5 01\_ISCO\_T0\_REP1\_DNA 02\_ISCO\_T0\_REP2\_DNA 03\_ISCO\_T0\_REP3\_DNA 04\_HICO\_T0\_REP1\_DNA 05\_HICO\_T0\_REP2\_DNA 07\_ISCO\_T0\_REP1\_cDNA 09\_ISCO\_T0\_REP3\_cDNA 21\_HICO\_T2\_REP1\_cDNA 22\_HICO\_T2\_REP2\_cDNA 06\_HICO\_T0\_REP3\_DNA 08\_ISCO\_T0\_REP2\_cDNA 10\_HICO\_T0\_REP1\_cDNA 11\_HICO\_T0\_REP2\_cDNA 17\_HICO\_T2\_REP3\_DNA 20\_ISCO\_T2\_REP3\_cDNA 23\_HICO\_T2\_REP3\_cDNA 12\_ISCO\_T2\_REP1\_DNA 13\_ISCO\_T2\_REP2\_DNA 14\_ISCO\_T2\_REP3\_DNA 15\_HICO\_T2\_REP1\_DNA 16\_HICO\_T2\_REP2\_DNA 18\_ISCO\_T2\_REP1\_cDNA 19\_ISCO\_T2\_REP2\_cDNA

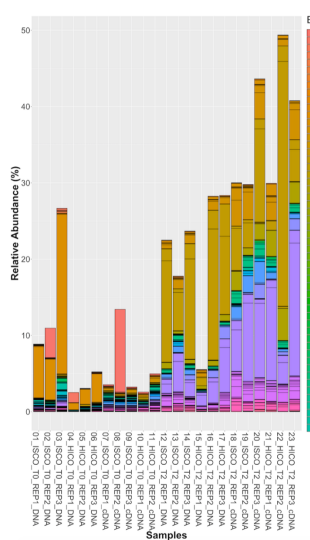
25 26

27

Samples Supplementary Figure S9: Relative abundance (%) of cryptomonads in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and 48 hours (t<sub>2</sub>) treatment exposure.

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#### Eukarvotes

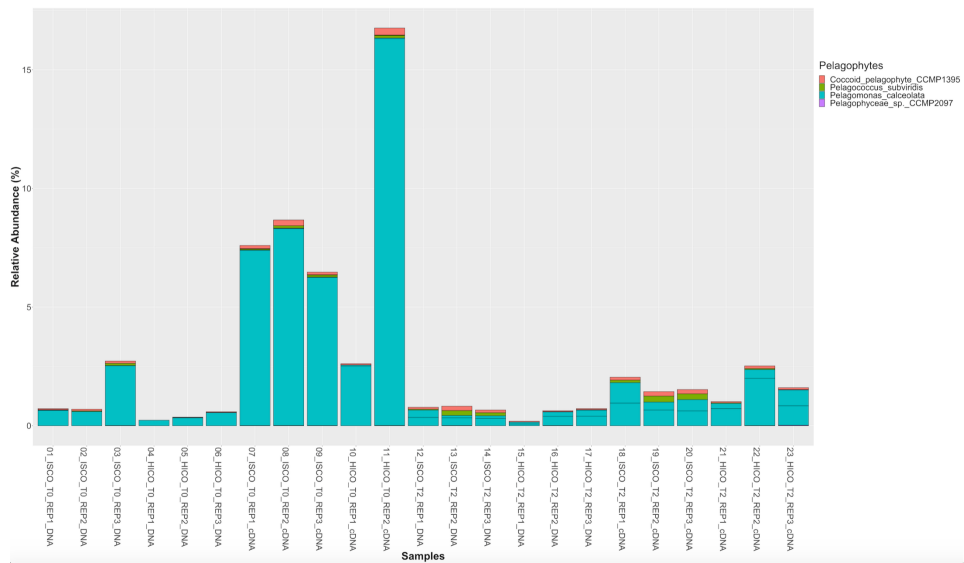
Acanthocolla cruciata Acanthometra\_sp Acanthometra sp. 2 7202 Acanthometra\_sp.\_3\_813 Acavomonas peruviana Acrispumella\_msimbaziensis Amastigomonas\_sp.\_2\_Bamfield Amphiacon\_denticulatus Amphibelone\_anomala Anurofeca sp. LAH2003 Artostrobus\_sp.\_\_2014 Astrosphaera\_hexagonalis Bicosoeca\_petiolata Bicosoeca\_vacillans Blastocystis\_sp Blastocystis\_sp. subtype\_3 Blastocystis\_sp.\_subtype\_4 Caecitellus parvulus Caecitellus pseudoparvulus Cafeteria minima Chattonella\_marina Chattonella\_subsalsa Chaunacanthid 217 Chaunacanthid sp. 6200 Chlamydaster sterni Chrysowaernella\_hieroglyphica Ciliophrys\_infusionum Colponema\_edaphicum Copromyxa\_protea Cyanophora paradoxa Dictyocha\_speculum Dictyocha\_speculum Diplonema\_ambulator Diplonemida\_sp\_\_13\_RG2016 Diplonemida\_sp\_\_13\_RG2016 Diplonemida\_sp\_\_15b\_RG2016 Diplonemida\_sp\_\_215b\_RG2016 Diplonemida\_sp\_\_37\_RG2016 Diplonemida\_sp\_\_37\_RG2016 Diplonemida\_sp.\_47\_RG2016 Diplonemida\_sp.\_9sb\_RG2016 chinamoeba\_exundans Ellobiopsis\_chattonii Eucyrtidium hexastichum eukaryotic\_picoplankton\_environmental\_sample Fabornonas\_tropica Giraudyopsis\_stellifera golden\_algae Gromia melinus Halocafeteria\_sp.\_WVII\_10/2\_320 haptophytes Hartmannella abertawensis Heliodiscus\_asteriscus lemistasia phaeocysticola

Heterophrys\_marina Heterosigma akashiwo Hexacontium gigantheum Hexaconus\_serratus Incisomonas marina Larcopyle\_butschlii Leucocryptos\_marina Lithomelissa\_setosa Lithomelissa sp. 8012 Mastigamoeba guttula Multimonas\_koreensis Ochromonadaceae\_sp. CCMP2298 Paramoeba aestuarina Paramoeba branchiphila Paramoeba\_eilhardi Percolomonas cosmopolitus Percioiomonas\_cosmopolitus Perkinsus qugwadi phytoplankton\_environmental\_sample Picobiliphyte\_sp.\_MS58411 Picobiliphyte\_sp.\_MS58422 Picomonas\_judraškeda Pirsonia\_guinardiae Pirsonia\_guinardiae Pirsonia\_punctigera Pirsonia verrucosa Placidia\_cafeteriopsis Planomonas\_brevis Planomonas\_elongata Poterioochromonas\_malhamensis Pseudobodo tremulans Pseudochattonella\_verruculosa Pseudocubus obelīscus Pterocystis sp Pygsuia biforma Rhizcitomulina cf. marina Rhynchopus sp. ATCC\_50229 Snyderella kirbyi Solenicola\_setigera Squamamoeba japonica Sticholonche\_sp. JJP2003 Stramenopile sp. RCC822 Stramenopile sp. BCC822 Stramenopile sp. ZC822 Stramenopile sp. Z211 Synura\_curtispina Telonema\_subtile Tetrapyle octacantha Pseudocubus\_obeliscus Tetrapyle\_octacantha Thalassicolla\_pellucida Triastrum\_aurivillii Triparma pacifica Trizona brandti Uncultured marine\_alveolate\_Group\_I\_DH148EKD18 Uncultured marine\_alveolate\_Group\_II\_DH145EKD20 Uncultured\_marine\_alveolate\_Group\_II\_DH147EKD20 Vacuolaria\_viresceñs Vannella\_spipetala Vannella\_sp.\_ELH5 Vannella\_sp.\_ELH5 Vannella\_sp.\_KONT2Pe Vannella\_sp.\_S3M13

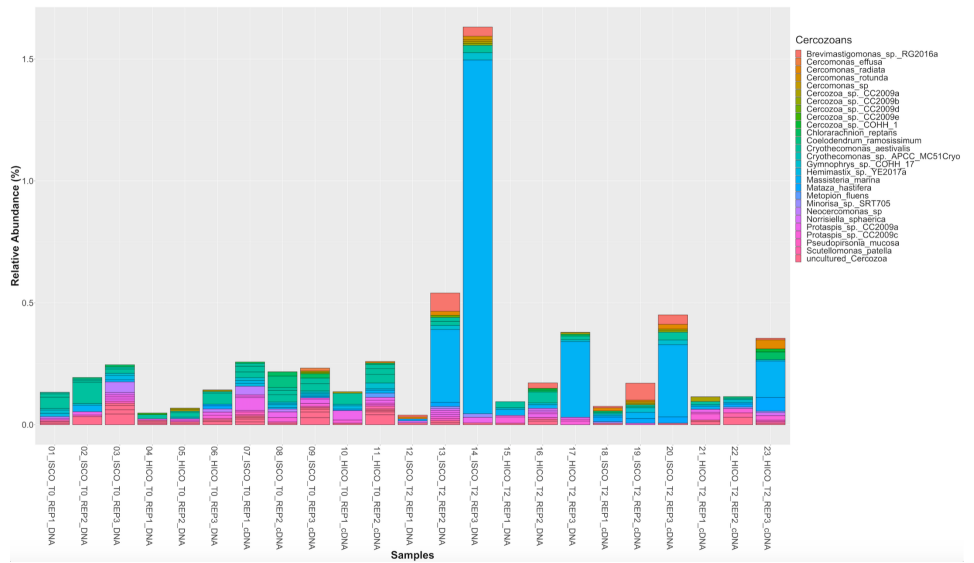
30 31

Supplementary Figure S10: Relative abundance (%) of eukaryotes in ISCO (*in situ* CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and

32 48 hours  $(t_2)$  treatment exposure.



Supplementary Figure S11: Relative abundance (%) of pelagophytes in ISCO (in situ CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) 35 and 48 hours  $(t_2)$  treatment exposure.



36 37

Supplementary Figure S12: Relative abundance (%) of cercozoans in ISCO (in situ CO<sub>2</sub>) and HICO (high CO<sub>2</sub>) treatment after immediate (t<sub>0</sub>) and

38 48 hours (t<sub>2</sub>) treatment exposure.