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Earlier sea-ice melt extends the oligotrophic summer period in the Barents Sea with low algal biomass and associated low vertical flux



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ABSTRACT

The decrease in Arctic sea-ice extent and thickness as a result of global warming will impact the timing, duration, magnitude and composition of phytoplankton production with cascading effects on Arctic marine food-webs and biogeochemical cycles. Here, we elucidate the environmental drivers shaping the composition, abundance, biomass, trophic state and vertical flux of protists (unicellular eukaryotes), including phytoplankton, in the Barents Sea in late August 2018 and 2019. The two years were characterized by contrasting sea-ice conditions. In August 2018, the sea-ice edge had retreated well beyond the shelf break into the Nansen Basin (>82°N), while in 2019, extensive areas of the northwestern Barents Sea shelf (>79°N) were still ice-covered. These contrasting sea-ice conditions resulted in marked interannual differences in the pelagic protist community structure in this area. In August 2018, the protist community was in a post-bloom stage of seasonal succession characterized by oligotrophic surface waters and dominance of small-sized phytoplankton and heterotrophic protists (predominantly flagellates and ciliates) at most stations. In 2019, a higher contribution of autotrophs and large-celled phytoplankton, particularly diatoms, to total protist biomass compared to 2018 was reflected in higher chlorophyll a concentrations and suggested that the protist community was still in a late bloom stage at some stations. It is noteworthy that particularly diatoms contributed a considerably higher proportion to the protist biomass at the ice-covered stations in both years compared to the open-water stations. This pattern was also evident in the higher vertical protist biomass flux in 2019, dominated by dinoflagellates and diatoms, compared to 2018. Our results suggest that the predicted transition toward an ice-free Barents Sea will lengthen the oligotrophic summer period with low algal biomass and associated low vertical flux.

1. Introduction

Ecosystem structures, ecological interactions and dynamics in the

Barents Sea are strongly driven by water-mass characteristics and seasonality in this high-latitude environment. Separated by the Polar Front (Parsons et al., 1996), the northern (Arctic) part of the Barents Sea with

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Fig. 1. Sampling station locations and sea-ice conditions in the northwestern Barents Sea during the Nansen Legacy cruises JC1_2 in August 2018 and Q3 in August 2019. The blue line indicates the position of the Polar Front (as indicated by the 0 °C surface isotherm) and was provided by the Institute of Marine Research. Ice data are from 15 August 2018 and 15 August 2019 and were provided by the Norwegian Ice Service - MET Norway.

cold and low-salinity Arctic water entering from the northeast is typically ice-covered in winter (Smedsrud et al., 2013), while the southern (subarctic) part remains ice-free for the majority of the year due to the inflow of warm, saline and nutrient-rich Atlantic water masses (Loeng, 1991; Loeng et al., 1997; Wassmann et al., 2006; Loeng and Drinkwater, 2007; Hunt et al., 2013). The Barents Sea is a highly productive shelf sea, particularly the subarctic part, supporting large zooplankton and commercial fish stocks, which in turn represent important prey for seals, whales, and seabirds (Wassmann et al., 2006; Loeng and Drinkwater, 2007; Sakshaug et al., 2009).

Apparent from the large-scale environmental changes observed over the past decades in the study region, the Barents Sea is particularly sensitive to climate warming (Skagseth et al., 2020; Isaksen et al., 2022; Smedsrud et al., 2022). Increasing volumes of inflowing warm Atlantic water masses (Oziel et al., 2016; Lind et al., 2018) lead to the continuous decrease in sea-ice extent and cause thinning of the ice cover during all seasons (Årthun et al., 2012, 2019, 2021; Onarheim and Årthun, 2017). While gross pelagic primary production is predicted to increase with declining sea-ice cover and consequently longer open-water seasons (Ellingsen et al., 2008; Drinkwater, 2011; Arrigo and van Dijken, 2015), increasing stratification of the surface water column in summer can amplify nutrient limitation reducing phytoplankton growth (Tremblay et al., 2009; Slagstad et al., 2015) and favour regenerated production and communities (Tremblay et al., 2015). Additionally, rising atmospheric CO₂ levels are resulting in acidification of the ocean, which is expected to affect calcifying organisms negatively (Kroeker et al., 2013; Skogen et al., 2014).

Superimposed on climate-change trends, multidecadal and interannual variations in ocean heat transport caused by large-scale climatic forcing (Vinje, 2001; Smedsrud et al., 2013, 2022; Oziel et al., 2016) have a marked impact on primary and secondary production, and subsequently ecosystem dynamics in the Barents Sea (Reigstad et al., 2002; Dalpadado et al., 2014; Ingvaldsen et al., 2021). Extended periods of open water, as the direct result of sea-ice loss caused by warming, but also interannual variability in sea-ice extent will impact the timing and magnitude of the primary production as well as the composition of phytoplankton communities in the Barents Sea (Neeley et al., 2018; Dalpadado et al., 2020; Dong et al., 2020). Earlier ice break-up and melt lead to earlier phytoplankton blooms in the northern Barents Sea and a longer open-water season to an increase in pelagic annual net primary productivity (Kahru et al., 2016; Dalpadado et al., 2020). However, the length of the highly productive period has not increased, which is likely the result of nutrient depletion as the season progresses (Kahru et al., 2016; Lewis et al., 2019). This will have consequences for vertical flux and the energy transfer to the seafloor during the extended lowproductive summer season.

Atlantic and Arctic water masses in the Barents Sea, characterised by different properties and varying sea-ice conditions, host distinctly different phytoplankton communities (Owrid et al., 2000; Metfies et al., 2016; Downes et al., 2021). The composition of marine protist plankton communities in the Barents Sea changes seasonally. In spring, diatoms often dominate algal biomass (von Quillfeldt, 2000; Degerlund and Eilertsen, 2010; Vodopyanova et al., 2020), while after the spring diatom bloom, motile taxa, such as flagellates belonging to the prymnesiophytes, are a prominent component of the phytoplankton community in the subarctic part in this period (Giraudeau et al., 2016; Silkin et al., 2020). Also, other flagellates and heterotrophic protists reach their peak (Ratkova and Wassmann, 2002). Physical factors, such as the presence of sea ice, temperature and salinity of seawater, light conditions and mixed layer depth, chemical factors, including nutrient availability as well as biological factors, such as top-down control by pathogens, parasites and grazers have decisive impacts on the composition of phytoplankton and protist communities (Neeley et al., 2018; Sugie et al., 2020). In addition, the fact that the spring bloom is usually dominated by a few diatom species belonging to the genera Thalassiosira and Chaetoceros and the haptophyte Phaeocystis pouchetii points to lifecycle strategies, including defence against grazers, that have evolved to optimize resource uptake after the end of winter (Hegseth et al., 2019; Assmy et al., 2023).

Climate-driven changes in algal communities at the base of the

Table 1

Stations in the northwestern Barents Sea sampled for environmental variables (sections 2.3, 2.4, 3.2 and 3.3), protist community composition (microscopy; sections 2.5.1 and 3.4), pico- and nanoplankton (flow cytometry; sections 2.5.2 and 3.5) and protist vertical flux (sections 2.5.3 and 3.6) in August 2018 and 2019.

2018	18					2019					
Station #	Date of sampling	Latitude °N	Longitude °E	Bottom depth (m)	Standard sampling depths (m)*	Station #	Date of sampling	Latitude (°N)	Longitude (°E)	Bottom depth (m)	Standard sampling depths (m)*
P1 ⁺	08/08	75.997	31.229	325	5, 10, 20, 30, 40, 60, 90	P1 ⁺	08/08	76.000	31.219	325	5, 10, 30, 45, 60, 90
P2 ⁺	10/08	77.502	33.996	192	5, 10, 20, 30, 40, 60, 90	P2	12/08	77.501	33.987	186	10, 30, 50, 60
Р3	12/08	78.751	33.998	305	5, 10, 20, 30, 40, 60, 90	Р3	13/08	78.749	34.001	307	10, 30, 60, 90
P4 ⁺	14/08	79.749	34.001	335	5, 10, 20, 30, 40, 60, 90	P4 ⁺	14/08	79.693	34.230	353	10, 30
Р5	15/08	80.501	34.006	163	5, 20, 30, 40, 60, 90	P5 ⁺	15/08	80.497	33.989	163	20, 30, 90
PICE1 ⁺	17/08 & 18/08	83.332/ 83.349	31.540/ 31.577	3930	5, 10, 30, 40, 60, 90	P6 ⁺	18/08	81.585	31.519	1100	10, 30, 90
SICE3	20/08	83.226	26.125	3911	5, 10, 20, 30	P7 ⁺	21/08	81.926	29.139	3300	15, 30, 60, 90
						SICE4	24/08	81.978	24.473	3600	10, 20, 30, 60, 90

*Deviations are noted in respective sections.

⁺ Sampling stations for vertical particle flux.

pelagic food web will have cascading effects on the entire Arctic marine ecosystem (Hays et al., 2005; Post et al., 2013; Thingstad, 2020). Assessing qualitative and quantitative changes in protist plankton (especially primary producers) under contrasting environmental scenarios is therefore essential for developing a better understanding of ecosystem characteristics in the decades to come (Acevedo-Trejos et al., 2014; Lyon and Mock, 2014; Barton et al., 2016). While some microalgae seem to be able to acclimate to their changing habitat (Pančić et al., 2015; Rivero-Calle et al., 2015; Hoppe et al., 2018), others are likely to be impacted negatively (Hare et al., 2007; Coello-Camba et al., 2014; Kvernvik et al., 2018). As a result of a prolonged openwater season, warming waters, increasing CO₂ concentrations and reduced nutrient availability, the trend toward small-sized algae (picoplankton) has been reported for different parts of the Arctic Ocean (Li et al., 2009; Morán et al., 2010; Coello-Camba et al., 2014; Zhang et al., 2015; Sugie et al., 2020), where they succeed over large-sized diatoms (Marinov et al., 2010).

In this study, we assessed and compared the composition of planktonic protist communities and their vertical flux along a south-north transect in the northwestern Barents Sea during two years with contrasting sea-ice conditions. We related their taxonomic compositions and size structures, from pico- to microplankton, to environmental conditions (sea-ice cover, temperature, salinity, nutrients) in order to reveal the drivers of interannual variability in biomass, abundance and diversity of the Barents Sea microbial food web. The contrasting two years with different ice conditions provided a unique opportunity to test the hypothesis that earlier ice melt would lead to an earlier spring bloom and an extended oligotrophic summer season dominated by flagellates.

2. Material and methods

2.1. Sampling campaigns

Seawater samples were collected in August of two consecutive years (2018, 2019) within the framework of the Norwegian Nansen Legacy project (arvenetternansen.com). Both sampling campaigns were conducted onboard RV *Kronprins Haakon* during a similar summer period (2018: 6 to 23 August, seven stations, joint cruise JC1_2; 2019: 5 to 27 August, eight stations, seasonal cruise Q3). The northward transects covered areas from 76°N to 83°N in the northwestern Barents Sea (Fig. 1, Table 1). Details on individual sampling procedures can be found in the project sampling protocols (https://doi.org/10.7557/nlrs.5719).

2.2. Sea-ice conditions

Sea-ice conditions along the entire cruise track were assessed using the standardized IceWatch ASSIST protocol (https://icewatch.met.no/d ata_guide). Briefly, visual observations conducted regularly provided information on ice concentration, primary ice type, snow depth as well as melt-pond coverage. The entire data for both expeditions is available online (https://icewatch.met.no/) and only ice conditions within the vicinity of our biological stations are reported in this study (Table A.1).

2.3. Temperature, salinity and inorganic nutrients

Water-column temperature and salinity measurements were obtained using a conductivity-temperature-depth (CTD) sensor (Sea-Bird SBE 911plus) mounted onto a rosette equipped with 24 10L-Niskin bottles. All CTD data can be found in Ingvaldsen (2022) and Reigstad (2022). Seawater samples for the determination of nitrate [NO₃], phosphate [PO₄³⁻] and silicic acid [Si(OH)₄] were drawn from the Niskin bottles into 20 mL pre-rinsed plastic HDPE vials and were preserved with 250 µL chloroform and stored in the dark at 4 °C. Post-cruise analysis was performed using a Flow Solution IV analyzer (O.I. Analytical, USA) following standard procedures (Grasshoff et al., 2009) at the Institute of Marine Research, Bergen, Norway. The analyser was calibrated by routine measurements of reference seawater obtained from Ocean Scientific International Ltd., UK. Detection limits were 0.5 $\mu mol \ L^{-1}$ for [NO_3], 0.06 $\mu mol \ L^{-1}$ for [PO_4^3-] and 0.7 $\mu mol \ L^{-1}$ [Si (OH)₄], respectively. All nutrient data can be found in Chierici et al. (2021a,b).

Photosynthetically active radiation intensity (μ mol m⁻² s⁻¹) was retrieved from a Biospherical Licor Chelsea sensor mounted onto the rosette. The direct output of the sensor was used at all stations in 2018, and the P1 station in 2019. At stations P2-SICE4 in 2019, the rosette was deployed through the vessel's moonpool and the shadow of the vessel impacted the light measurements down to 25–30 m. For those stations, a Beer-Lambert curve was fitted on the data beyond this depth in order to model what the surface irradiance would have been. For the sea icecovered stations, a light transmittance of 4 % under summer first-year ice with <10 cm of snow covered was assumed (Nicolaus et al., 2012) in order to estimate the irradiance incident on the sea-ice surface. An average between the above numbers, using the ice-cover proportion as weighing factor, was calculated to obtain a representative surface irradiance at each station.



Fig. 2. A) Temperature (°C), B) salinity, concentrations of C) nitrate (μ mol L⁻¹), D) silicate (μ mol L⁻¹), E) phosphate (μ mol L⁻¹) and F) Chlorophyll (chl) *a* concentration (μ g L⁻¹) including the euphotic zone depth (white line) represented by the depth where photosynthetic available radiation (PAR) is 1% of PAR incident at the ocean surface, along the sampling transects in the northwestern Barents Sea in August 2018 (left panels) and August 2019 (right panels). The sampled depths (black dots) are plotted along the station (black line).



Fig. 3. Satellite-derived monthly composites of surface chl a (μ g L⁻¹) in the Barents Sea for 2018 (top row), 2019 (middle row) and differences between 2019 and 2018 (bottom row). Sampling stations during both years are represented by the red dots.

Table 2

Loss rate (% d⁻¹) and flux (mg C m⁻² d⁻¹) at 90 m for the different protist groups along the sampling transects in the northwestern Barents Sea in August 2018 and August 2019 (note: the loss rate calculation at P4 in 2019 is based on integration from 0 to 30 m). 0 = below detection limit.

Parameter	Group	2018		2019	2019					
		P1	P2	P4	PICE1	P1	P4	Р5	P6	P7
Loss rate (% d ⁻¹)	Diatoms	0.2	0	0	2.2	0.6	3.2	65.3	1.2	0.5
	Dinoflagellates	1.2	0.3	4.7	0.8	2.8	2.7	9.5	2.2	0.3
	Other flagellates	4.8	2.2	29.8	1.8	0.9	5.9	10.5	2.0	3.4
	Ciliates	0.2	2.2	0	0	0.3	30.6	0.6	0.1	4.3
Flux (mg C $m^{-2} d^{-1}$)	Diatoms	0.8	0	0	10.1	1.7	3.3	186.5	16.4	3.2
	Dinoflagellates	37.1	6.2	9.2	6.2	20	20.6	68.4	31.1	1.9
	Other flagellates	20.4	7.4	4.1	12.6	10.2	21	44.9	22.1	20.6
	Ciliates	4.3	18.5	0	0	5.0	104.7	1.0	0.2	20.3

2.4. Chlorophyll (chl) a concentrations

Water subsamples for chl *a* concentration measurements were collected at discrete depths (Table 1) from casts taken by a CTD rosette. Water was collected into plastic bottles and stored in a dark and cold location until further processing (within 1 h). Between 0.15 and 1 L of water was filtered through 25 mm Whatman GF/F filters under low vacuum pressure (~30 kPa). Filters were stored in polypropylene tubes with 5 mL of methanol added for chl *a* extraction (overnight at 0–4 °C). Chlorophyll *a* concentrations were measured in the dark according to Holm-Hansen and Riemann (1978) procedure with a Turner Design AU10 fluorometer. The entire chl *a* datasets can be found in Vader (2022).

In order to extend the spatial and temporal coverage of our study, chl *a* surface concentrations (as a proxy for algal biomass) were also estimated by using satellite imagery provided by MODIS-Aqua (Moderate-resolution Imaging Spectroradiometer; 4.6 km resolution data). We used

L3 monthly averaged data for the area of interest $(71^{\circ}-80^{\circ} \text{ N}, 16^{\circ}-52^{\circ} \text{ E})$ for the period of April to September in 2018 and 2019. Near-surface chl *a* concentrations were estimated by using bio-optical models for certain chl *a* ranges. For very clear oligotrophic waters, when chl *a* concentration was below 0.15 mg m⁻³, the CI (Colour Index) algorithm was applied (Hu et al., 2012), for concentrations above 0.2 mg m⁻³, the OC3 band ratio model was used (O'Reilly et al., 2000) and between 0.15 and 0.2 mg m⁻³, a combination of the aforementioned methods was applied. The data can be assessed and downloaded from the NASA (National Aeronautics and Space Administration) OBPG (Ocean Biology Processing Group) archive data website (<u>https://oceancolor.gsfc.nasa.gov/13/order/</u>). It should be noted that the satellite-derived chl *a* estimates are only valid for the upper surface (~10 m) of the ocean, and are unable to capture the presence of subsurface/deep chl *a* maxima.



Fig. 4. A) Protist abundances (10^3 cells L⁻¹) and B) biomass (μ g C L⁻¹) along the sampling transects in the northwestern Barents Sea in August 2018 (top panels) and August 2019 (bottom panels). Note the difference on the x-axis for the stations PICE1 and SICE3 in 2018.

2.5. Microbial community composition

2.5.1. Light microscopy: Nano- and microplankton communities

Seawater was collected at discrete depths (Table 1) with Niskin bottles attached to a CTD rosette. For microscopic analysis, 190 mL of seawater from each depth was filled into 200 mL brown glass bottles directly from the Niskin bottles. Samples were fixed with 0.8 mL of 25 % glutaraldehyde and 10 mL of 20 % hexamethylenetetramine-buffered formalin solutions to yield final concentrations of 0.1 and 1 %, respectively. Samples were stored cool (ca. 15 °C) and dark until further processing.

Identification and quantification (cells L^{-1}) of protists were carried out with a Nikon inverted light microscope equipped with phase and differential interference contrasts and objectives 10–60× (resulting in 100–600 × magnification) following the Utermöhl method (Utermöhl, 1958; Edler et al., 2010). Different volumes of sedimentation columns were used (50 mL or 55 mL in 2018; 10 mL, 50 mL or 55 mL in 2019) according to the algal abundance in the samples. Abundances were converted to carbon biomass based on published geometric relationships for biovolume conversion (Hillebrand et al., 1999) and biovolume to carbon conversion factors (Menden-Deuer and Lessard, 2000). Protist community compositions are reported as abundances (cells L⁻¹) and biomass (μ g C L⁻¹), and both numerically integrated with a trapezoidal formula over the uppermost 90 m of the water column (standing stocks as cells m⁻² and g C m⁻²; note that station SICE3 [0–30 m] in 2018 and stations P2 [0–60 m] and P4 [0–30 m] in 2019 were integrated over shallower depths due to the absence of samples from greater depths). Organisms were identified to the lowest possible taxonomic level and



Fig. 5. Depth-integrated stocks of A) protist abundances $(10^9 \text{ cells m}^{-2})$, B) biomass (g C m⁻²), C) trophic mode (g C m⁻²) and D) chl *a* (mg m⁻²) for the upper 90 m surface layer along the sampling transects in the northwestern Barents Sea in August 2018 (left panels) and August 2019 (right panels). *For stations SICE3 in 2018, and P2 and P4 in 2019, protist stocks were integrated over shallower depth. Blue lines indicate stations that were sea ice-covered during the time of sampling.

were named according to the World Register of Marine Species (WoRMS). Protist taxonomy data can be found in Assmy et al. (2022a) for JC1_2 and Assmy et al. (2022b) for Q3.

2.5.2. Flow cytometry: Pico- and nanophytoplankton communities

Seawater was collected at discrete depths (Table 1) with Niskin bottles attached to a CTD rosette. Samples for flow cytometric abundance analysis were prepared in triplicates by fixing 1.8 mL of seawater with 36 μL 25 % glutaraldehyde (0.5 % final concentration) at 4 °C in the dark for a minimum of 2 h, then flash-frozen in liquid nitrogen and stored at -80 °C until further processing.

The abundances of pico- and nano-sized phytoplankton and heterotrophic nanoflagellates (HNF) were determined using an Attune® NxT, Acoustic Focusing Cytometer (Invitrogen by Thermo Fisher Scientific) with a syringe-based fluidic system and a 20 mW 488 nm (blue) laser. Pico- and nanophytoplankton were counted after thawing the sample and the various groups of protists were discriminated based on their red fluorescence (BL3) vs. orange fluorescence (BL2), red fluorescence (BL3) vs. side scatter (SSC) and orange fluorescence (BL3) vs. side scatter (SSC) (Paulsen et al., 2016). For HNF analysis, samples were stained with SYBR Green I for 2 h in the dark and subsequently 1-2 mL was measured at a flow rate of 500 μ L min⁻¹ following the protocol of Zubkov et al. (2007). Pico- and nanophytoplankton community compositions are reported as cells mL^{-1} and abundance was converted into biomass (µg C L^{-1}). HNF biomass was estimated using 0.220 pg C μm^{-3} as conversion factor (Børsheim and Bratbak, 1987). Biomass of pico- and nanophytoplankton was calculated using equivalent spherical diameter (ESD) to estimate cell volume (CV) and the formula Carbon per cell [pg C $cell^{-1}$] = 0.216 × CV ^{0.939} to calculate carbon content (Mullin et al.,

1966; HELMCOM-Guidelines for monitoring of phytoplankton species composition, abundance, and biomass 2021). ESDs for the three phytoplankton size groups were estimated using a bead mix (1, 2, 4, 6 and 15 μ m) during flow cytometry and are given as average based on the estimated upper and lower limits of the gates of the different phytoplankton groups (picophytoplankton [size range 0.2–2 μ m]: r = 0.55 μ m; small nanophytoplankton [2–5 μ m]: r = 1.75 μ m; larger nanophytoplankton [5–10 μ m]: r = 3.75 μ m).

2.5.3. Vertical flux

Short-term sediment traps (KC-Denmark, 2-4 cylinders, aspect ratio > 6) were deployed at six different depths at stations P1, P2, P4 and PICE1 in August 2018 and at stations P1, P4, P5, P6 and P7 in August 2019 (Table 1). Before the deployment, the cylinders were filled with high-salinity deep water that was collected at the station and filtered through GF/F filters in order to minimize flushing during deployment and recovery. The sediment traps were freely drifting in open water or anchored to an ice floe in ice-covered waters to facilitate a semi-Lagrangian drift. Deployment times varied from 19 to 41 h, with shorter deployment periods in order to minimize particle degradation when a high amount of sinking material was expected, and longer periods when little material was expected to sink out. No preservatives were used during deployment. After retrieval, contents of cylinders from the same depth were pooled in one carboy. The samples were stored at 4 °C until further processing within 20 h after the retrieval. After homogenizing, a 100 mL water sample was fixed with a glutaraldehydelugol solution (Rousseau et al., 1990) and stored at 4 °C in the dark. Protist communities were analysed microscopically as described in section 2.5.1 and daily flux rates are reported as cells $m^2 d^{-1}$ and





C - Other flagellates

D - Ciliates



Fig. 6. Depth-integrated stocks of protist biomass (g C m^{-2}) for the upper 90 m surface layer representing A) diatoms, B) dinoflagellates, C) other flagellates, and D) ciliates along the sampling transects in the northwestern Barents Sea in 2018 (left) and 2019 (right). * For stations SICE3 in 2018, and P2, P4 in 2019, protist stocks were integrated over shallower depth. Blue lines indicate stations that were sea ice-covered during the time of sampling.

biomass as mg C m⁻² d⁻¹.

The relative daily loss rates at 90 m depth were calculated as:

% Loss rate
$$d^{-1} = \left(\frac{\text{flux at 90m } [\text{mg } C \text{ } m^{-2} \text{ } d^{-1}]}{\text{integrated standing stock } (0 - 90m) [\text{mg } C \text{ } m^{-2}]}\right) * 100$$

3. Results

3.1. Sea-ice conditions

The ice conditions in August 2018 and August 2019 differed substantially (Fig. 1, Table A.1). During the sampling in 2018, sea ice occurred mainly in the Arctic Basin, north of the shelf break, while in 2019, also large parts of the northwestern Barents Sea shelf were icecovered.

According to sea ice observations from the bridge, areas represented by sampling stations P1 to P5 were ice-free in 2018. The ice edge (10 % ice cover) was encountered at ca. 82.9° N and 31.4° E. All the remaining stations were covered by ice from 70 to 100 %. First-year ice (FYI) with a thickness of around 1.0 m dominated at station PICE1, while secondyear ice with a thickness of 1.7–1.8 m dominated at station SICE3. Sea ice was covered by a thin, around 2 cm thick snow cover, and melt ponds covered 10–30 % of the surface of the ice floes. At the ice-covered stations, open water areas were limited to leads of 50 to 200 m width.

In 2019, ice extended much further south and only stations P1 to P3 were ice free. The ice edge was encountered between two ice

observations: 79.1°N and 34.0°E (0 % ice cover) and 79.8°N and 34°E (60 % ice cover). Ice concentration, dominated by FYI < 0.7 m thickness and narrow leads of open water <200 m wide, increased from 60 % to 90 % between P4 and P6. While areas at P7 had similar conditions to those represented by station P6, less ice (70 % cover) occurred further west, where sampling station SICE4 was located, with open water in narrow leads of 50–200 m width. FYI < 0.7 m dominated in the entire area (for more details see Van Engeland et al., submitted for publication, same issue).

3.2. Temperature, salinity, and inorganic nutrients

In 2018 and 2019, Atlantic water dominated the water column south of the Polar Front (station P1) while surface waters further north were fresher polar water masses (Fig. 2A, B). Temperature-salinity distribution at stations P2-P5 indicated that the relatively warm upper ocean north of the Polar Front was a result of surface warming rather than intrusion of Atlantic Water (Fig. A.1). Coldest and least saline surface water was observed at the northernmost stations, most evident in 2019. Shoaling of warm Atlantic water was observed at the shelf break (station P6) and in the southern Nansen Basin (station P7) indicative of the Svalbard branch of Atlantic water entering the Arctic Ocean along the shelf break north of Svalbard. The nutricline of nitrate, silicate and phosphate was generally deeper in 2018 (50 m) compared to 2019 (20–30 m), except at the permanently open water station P1 and station P2 just north of the polar front (Fig. 2C-E). In 2018, surface nutrient concentrations were still elevated (nitrate > 4 μ M) at the two



Fig. 7. Cell biomass of small phytoplankton and heterotrophic nanoflagellates $(0.2 - 10 \ \mu\text{m})$ along the sampling transects in the northwestern Barents Sea in 2018 (left panels) and 2019 (right panels) divided according to trophy and size into A) picophytoplankton $(0.2-2 \ \mu\text{m})$, B) nanophytoplankton $(2-10 \ \mu\text{m})$ and C) heterotrophic nanoflagellates.

northernmost ice stations. However, it needs to be noted that those stations were about 1° latitude further north than the sea ice stations P6 and P7 in 2019 due to the northward retreat of the ice edge in August 2018 and hence not directly comparable.

3.3. Chlorophyll a concentrations

Chlorophyll *a* concentrations and standing stocks were overall lower in 2018 compared to 2019, reaching maximum values in the upper 10 m at station PICE1 in 2018 (0.87 to $1.09 \ \mu g \ L^{-1}$) and peaked at station P5 with 2.57 $\ \mu g \ L^{-1}$ at 20 m in 2019 (Fig. 2F). In both years, the values were the highest in the top 60 m of the water column. In 2019, chl *a* concentrations at the ice-covered stations reached their maximum in surface waters (Fig. 2F).

Monthly chl *a* climatologies were derived from remote sensing data for 2018 (top row), 2019 (middle row) as well as differences between 2019 and 2018 (bottom row; Fig. 3). The grey and white colours correspond to the land and cloud masks, respectively. An extensive phytoplankton bloom was observed in May 2018. Aggregation and decaying phytoplankton in large bloom events will result in increased backscatter, which impacts the chl *a* retrieval models used here. Hence, the actual chl *a* values might have been lower, but our data suggest a significant bloom in May. The difference map (bottom row) for May shows markedly lower chl *a* concentrations in 2019 compared to 2018 (dark blue colour). In the area of interest ($71^{\circ} - 80^{\circ}$ N, $16^{\circ} - 52^{\circ}$ E), in May 2018, the average chl *a* value was 1.7 ± 2.1 mg m⁻³. This shows that on average the difference in chl *a* content corresponds to 3.5 mg m⁻³

(Table A.2). Besides this event in May 2018, a general increase in certain areas can be observed (red colour in Fig. 3, bottom row) or no difference (green colour in Fig. 3, bottom row) in the chl *a* content over the entire seasonal production cycle (Table 2).

3.4. Protist community composition

All identified protist taxa from Niskin bottle samples are presented in Table A.3.

Protist abundances were overall higher in August 2019 compared to August 2018, except at the ice-covered stations PICE1 and SICE3, where abundances in 2018 exceeded those in 2019 and were about one order of magnitude higher compared to values observed at open-water areas in 2018 (Fig. 4A). In contrast, the overall biomass was higher in 2018 compared to 2019 (Fig. 4B). In both years, (depth-integrated) algal abundances and biomass were the lowest at stations P3 and P4 (note: in 2019, data only available from 0 to 30 m; Figs. 4B, 5A, B). In 2018, heterotrophic taxa clearly dominated the protist community, except for the two ice-covered stations where autotrophic taxa prevailed (Fig. A.2), while in 2019, the majority of the community was auto- or mixotrophic (Fig. 5C, A.2).

In 2018, diatoms numerically predominated the protist community at the southernmost station (P1), dinoflagellates dominated further north (stations P2 to P4), whereas other flagellates (primarily prymnesiophytes) dominated at the northernmost stations (P5 to SICE3). In turn, in 2019, flagellates dominated numerically the entire study area (Fig. 4A, 5A). Chrysophyte flagellates were highly abundant at station P5 in 2018 and P1 in 2019 but contributed less to the biomass at these





Fig. 8. Vertical flux of protist A) abundances (10^6 cells m⁻² d⁻¹) and B) biomass (mg C m⁻² d⁻¹) at selected depths for the upper 200 m of the water column along the sampling transects in the northwestern Barents Sea in August 2018 (top panels) and August 2019 (bottom panels). Note: data not available for all stations.

stations (Fig. A.3). In terms of biomass, dinoflagellates and ciliates dominated in the ice-free areas (stations P1 to P5) and other flagellates dominated at the two ice-covered stations in 2018 (PICE1 and SICE3; Fig. 4B, 5B). In 2019, dinoflagellates and oligotrich ciliates dominated protist biomass in open-waters at stations P1 to P3 and in the proximity

of the ice edge (station P4), while diatoms were a prominent component of the protistan community in the ice-covered areas (stations P5 to P7; Fig. 4B, 5B). Diatom biomass was the highest in the uppermost 40 m at the ice-covered stations during the two years of observations.



Fig. 9. Biomass distribution in the microbial food web (Conceptual model after Thingstad et al., 2020) and vertical particle flux in the northwestern Barents Sea in **A**) August 2019 considering stations P2 to P5 representative of the interior shelf (north of the polar front and south of the continental slope). Box sizes indicate differences in biomass (for bacteria, protists, copepods) and nutrient concentrations (small = low, large = high). Mixotrophic nanoflagellates are included under autotrophic flagellates. Blue arrows indicate flow between the different components of the microbial food web where the thickness of the arrows indicates the strength of the flow from food to consumer. Vertical flux of protists is represented by red arrows (thicker arrow indicates higher protist flux). Future longer open water oligotrophic seasons (as seen in 2018) are likely to strengthen the left-hand side of the food web model comprising smaller forms relative to the more energy-efficient pathway of larger diatoms to the *Calanus* food web (as seen in 2019). Lower biomass and dominance of small-celled protists result in lower vertical flux. Our data indicate a strong trophic relationship between mixotrophic ciliates (*Strombidium, Mesodinium*) and autotrophic flagellates in both years. HNF were main grazers of bacteria.



Fig. A1. Distribution of Conservative Temperature and Absolute Salinity below 10 m depth from CTD profiles collected at the station locations in the northwestern Barents Sea (Fig. 1) during the two cruises. Water mass categories from Sundfjord et al. (2020) are shown as black boxes. Blue circles highlight measurements from < 50 m depth and between 77°N and 81°N (stations P2-P5). Due to intermittent sensor issues, the profile from station P7 in 2019 was replaced by a profile taken 7.5 km to the northeast.

3.4.1. Diatoms

Diatoms contributed to overall biomass to a larger extent in 2019 than in 2018 (Fig. 4B). In both years, the highest diatom standing stocks were observed at the ice-covered stations (Fig. 5B). It is noteworthy that diatom standing stocks were similar in both years at the ice-free station P1, dominated by Atlantic water masses, and at the same time considerably higher than at the open-water stations (P2 and P3) north of the Polar Front (Fig. 5B).

In 2018, the diatom community was primarily represented by Leptocylindrus minimus in Atlantic waters (station P1), followed by Pseudonitzschia spp., which dominated also in ice-covered waters (station SICE3). Interestingly, both taxa did not contribute largely to the diatom community in 2019 (Fig. 6A, Fig. A.4A). In comparison to 2018, species of the genus Fragilariopsis were abundant in 2019, particularly at the icecovered stations P4 and P5. Important contributors to phytoplankton biomass in 2019 were species of Thalassiosira, especially in ice-covered waters represented by stations P4 to SICE4 (Fig. 6A). A species commonly found in both pelagic and sea ice-associated communities, Shionodiscus biocolatus (formerly Thalassiosira bioculata), contributed largely to the diatom biomass at the northernmost ice-covered stations in both years (SICE3 in 2018 as well as P7 and SICE4 in 2019). The oceanic diatom Rhizosolenia hebetata f. semispina was a dominant component of the diatom community at the open-water station P1 in 2018 and at the ice-covered stations P5 to P7 in 2019, while Rhizosolenia hebetata f. hebetata dominated at P1 in 2019 (Fig. 6A). Chaetoceros species contributed moderately to the protistan biomass at the ice station PICE1 in 2018 and P6 and SICE4 in 2019.

3.4.2. Dinoflagellates

Highest dinoflagellate standing stocks were found at the southernmost stations on either side of the Polar Front (P1 and P2) in 2018 (Fig. 5B). In other areas, interannual differences were not very pronounced and dinoflagellate standing stocks varied between 0.19 and 3.21 g C m⁻². Athecate dinoflagellates (naked or unarmoured) dominated dinoflagellate biomass in both years, except for station P1 in 2019. Thecates were also important at station P5 in 2019 (Fig. A.5). Among the athecate dinoflagellates, gymnodinoid species were high in numbers and biomass in both years (Fig. 6B, Fig. A.4B). Another numerically important component were thecate dinoflagellates of the genus *Heterocapsa*, which were more abundant in 2019 than in 2018, particularly at station P2 (Fig. A.4B). Peridiniales had an overall higher biomass in 2019 vs. 2018.

3.4.3. Other flagellates

Dominance in the phylogenetically diverse flagellate community was quite variable across the transect in both years (Fig. 6C, Fig. A.4C). In 2018, the flagellate community at the ice stations PICE1 and SICE3 was dominated by the prymnesiophyte *Phaeocystis pouchetii*, both in numbers and biomass, which was distinctly less abundant in 2019 (Fig. 6C, Fig. A.4C). The raphidophyte *Heterosigma* sp. dominated the flagellate biomass at the ice-covered stations P6 to SICE4 in 2019 and at station P2 in 2018. At station P5, the prymnesiophyte *Chrysochromulina* sp. dominated in 2018. At station P1, the prasinophyte *Pterosperma* sp. dominated in 2018 while the chrysophyte *Dinobryon* sp. was a dominant component in 2019 (Fig. 6C).



Trophic mode Autotroph Mixotroph Heterotroph Undefined

Fig. A2. Protist trophic mode (μ g C L⁻¹) per depth (m) along the sampling transects in the northwestern Barents Sea in August 2018 (top panels) and August 2019 (bottom panels).

3.4.4. Ciliates

Ciliate biomass was generally higher in 2018 than in 2019, except at the southernmost station P1 and the northernmost ice-covered areas, where ciliate standing stocks were similar in both years (Fig. 5B, 6D). The genus *Strombidium* dominated the ciliate biomass at most stations in both years (Fig. 6D), and also in numbers at stations P2, P4 and P5 in 2018, as well as SICE4 in 2019 (Fig. A.4D). *Parafavella* sp. contributed largely to the ciliate biomass at station P1, particularly in 2019. Mixotrophic *Mesodinium rubrum* dominated ciliate standing stocks at stations P5 and P6 in 2019.

3.5. Pico- and nanophytoplankton community composition

Abundances of autotrophic organisms in the size range of $0.2 - 10 \,\mu m$ were determined using flow cytometry, with highest abundances measured at depths corresponding to the chl a maximum and above. Data were grouped into pico- and small nano-sized phytoplankton $(0.2-2 \,\mu\text{m} \text{ and } 2-10 \,\mu\text{m}, \text{ respectively})$ and visualized as cell biomass (μg $C L^{-1}$) for the upper 100 m of the transects in 2018 and 2019 in Fig. 7. Dominating size groups were different during the contrasting years and picophytoplankton biomass was the highest in 2018, with up to 2.5 µg C L^{-1} at the ice-covered station SICE3 (Fig. 7A). Picophytoplankton abundances in 2018 increased gradually from south to north, with 2000 cells mL⁻¹ at station P1, 4000 cells mL⁻¹ at station P2, 7000 cells mL⁻¹ at station P3 until 12,000 cells mL^{-1} at station P4 and decreasing thereafter before highest abundances were detected at station SICE3 (19,000 cells mL⁻¹; Fig. A.6). Picophytoplankton abundances measured in 2019 were comparable in terms of spatial pattern, but overall values were lower than in 2018.

Nanophytoplankton was generally more abundant in 2019 than 2018, with biomass concentrations of up to 80 μ g C L⁻¹ at station P7 in 2019 (Fig. 7B). Highest biomass in 2018 was measured furthest south (station P1) with around 30 μ g C L⁻¹. In 2019, the nanoplankton

biomass pattern corresponded to the chl a maximum.

Biomass of heterotrophic nanoflagellates (HNF) was comparable in 2018 and 2019 with peaks at around 4–5 μ g C L⁻¹ (Fig. 7C). In 2018, highest concentrations were at stations P3 and P4, while in 2019, highest biomass coincided with the chl *a* maximum, peaking at stations P2, P4, P5 and P6.

3.6. Vertical flux of protist communities

All identified protist taxa from sediment trap samples are presented in Table A.4.

Both protist cell and biomass vertical flux were higher in 2019 than in 2018 (Fig. 8). The vertical flux of cells ranged from 195 to 2184 10^6 cells m⁻² d⁻¹ in 2019 and from 215.7 to 1306 10^6 cells m⁻² d⁻¹ in 2018 across all depths. The only exception was P1 in 2019, where cell fluxes were lower than in 2018, and a strong flux attenuation from 932 10^6 cells m⁻² d⁻¹ at 30 m to 54.1 10^6 cells m⁻² d⁻¹ at 200 m depth. The vertical flux of protist biomass ranged from 4.6 to 198 mg C m⁻² d⁻¹, except for one station in 2019 with high flux dominated by diatoms (station P5 at 90 m with 296 mg C m⁻² d⁻¹; Fig. 8B).

In general, flagellates dominated cell fluxes at all stations in both years (Fig. 8A, Fig. A.7A). This reflected mostly the pattern in the suspended communities; except for stations P1 to P4 in August 2018, where diatoms and/or dinoflagellates dominated the suspended communities, whereas the sinking cells were dominated by flagellates. Despite the dominance of flagellates in terms of cell flux, dinoflagellates contributed more to the carbon flux. Diatom carbon flux was higher in 2019 than in 2018, particularly at stations P5 and P6, both in numbers and biomass (Fig. 8A, B).

Interestingly, the stations with the highest biomass in the suspended communities were not always dominated by diatoms but rather by dinoflagellates and flagellates, although this was not reflected in the protist carbon flux. The highest protist carbon flux was measured in



Fig. A3. Protist A) abundances (10^9 cells m⁻²) and B) biomass μ g (C L⁻¹) per depth (m) of all identified taxa along the sampling transects in the northwestern Barents Sea in August 2018 (top panels) and August 2019 (bottom panels). Note the difference on the x-axis for the stations PICE1 and SICE3 in 2018.

August 2019 at station P5 at 90 m and was dominated by diatoms of the genus *Thalassiosira* (Fig. 8B, Fig. A.7B). In 2018, cell flux at the southernmost station P1 and the northernmost ice-covered station PICE1 was dominated by prymnesiophytes (particularly *Phaeocystis pouchetii*; Fig. A.7A), however, in terms of carbon flux, dinoflagellates dominated at P1 and PICE1, represented by a mixed community (Fig. 8B). *Fragilariopsis* spp. and *Chaetoceros* spp. dominated diatom cell flux at stations P1 and PICE1, respectively, but due their relatively small size, diatoms contributed little to carbon flux at those stations.

Loss rates at 90 m were the highest in 2019 at station P4, dominated by ciliates (>30 % relative to suspended communities) and station P5, dominated by diatoms (>60 %), and in 2018 at P4, dominated by

flagellates (>30 %, Table 2).

Although average total particulate organic carbon (POC) flux was comparable in both years (Amargant-Arumí et al. in prep., same issue), protist carbon flux in 2018 contributed much less to total POC flux compared to 2019, with a mean of 25.8 % \pm 18.9 in 2018 vs. a mean of 54.6 % \pm 25.1 in 2019 considering all stations and depths.

4. Discussion

4.1. Contrasting late summer environmental conditions

Large interannual variability in sea-ice cover and oceanographic



Fig. A4. Depth-integrated stocks of protist abundances $(10^9 \text{ cells m}^{-2})$ for the upper 90 m surface layer representing A) diatom species, B) dinoflagellate species, C) other flagellates, and D) ciliate species along the sampling transects in the northwestern Barents Sea in 2018 (left) and 2019 (right) *For stations SICE3 in 2018, and P2 and P4 in 2019 integrated stocks were integrated over shallower depth. Note the difference on the y-axes. Blue lines indicate stations that were sea ice-covered during the time of sampling.



Fig. A5. Depth-integrated stocks of dinoflagellate A) abundances (10^9 cells m⁻²) and B) biomass μ g (C L⁻¹) per depth classified in athecates and thecates along the sampling transects in the northwestern Barents Sea in August 2018 and August 2019. *For stations SICE3 in 2018, and P2 and P4 in 2019 integrated stocks were integrated over shallower depth. Blue lines indicate stations that were sea ice-covered during the time of sampling.

conditions is typical for the Barents Sea (Falk-Petersen et al., 2000; Furevik, 2001; Serykh and Kostianoy, 2019). The two sampling years 2018 and 2019 offer a prime example for these annual fluctuations, characterized by contrasting environmental conditions, including seaice cover, sea-water temperature, and salinity as well as nutrient availability.

Despite sampling during the same time of the year, the earlier sea-ice retreat and longer open water summer season in 2018 caused differences in the timing of seasonal succession and thus in the composition of the pelagic protist communities in late summer (Figs. 4-6). This is corroborated by previous studies from other seasonally ice-covered regions of the Arctic Ocean showing a link between seawater temperature, sea-ice cover and protist phenology (Nöthig et al., 2015; Dabrowska et al., 2020). In the Barents Sea, the spring bloom typically occurs in May, starting in the permanently ice-free areas with elevated concentrations along the ice-edge rather than in ice-free areas (Engelsen et al., 2002), indicating that meltwater-induced stabilization of the water column has a positive effect on primary production (Qu et al., 2006). The earlier onset and more intense spring bloom in 2018 as evidenced from satellite-derived chl *a* (Fig. 3) can be attributed to the earlier ice retreat compared with 2019. Hence, the protist community in August 2018 was in a seasonally later stage of its succession than in August 2019. The interannual differences in chl *a* revealed by our *in situ* observations are not reflected in the satellite data which show similar chl *a* values in August for both years. However, it needs to be considered that satellites do not capture the subsurface chl *a* maximum that largely accounted for



Fig. A6. Abundances (10^3 cells m⁻²) of small phytoplankton (0.2–10 μ m) along the sampling transects in the northwestern Barents Sea in 2018 (left panels) and 2019 (right panels) divided according to size into A) picophytoplankton, B) nanophytoplankton and C) heterotrophic nanoflagellates.

the higher chl *a* standing stocks in 2019. Indeed, satellites can underestimate Arctic Ocean primary production (based on satellite-derived chl *a*) by up to 40 % during the post-bloom period with a pronounced subsurface chl *a* maximum (Ardyna et al. 2017). The suggested higher chl *a* values in the coastal areas recorded from satellites in both years were likely an artefact caused by sediment-rich waters that interfere with the chl *a* retrieval algorithm (Blondeau-Patissier et al., 2014; Blix et al., 2018). However, this did not compromise the comparison of satellite-derived and *in situ* chl *a* observations as we were focusing on the more off-shore parts of the northwestern Barents Sea in this study.

In 2018, the protist community was dominated by small-sized autotrophic (Fig. 7) and heterotrophic protists, mainly flagellates and ciliates (Fig. 4), indicative of a late-summer oligotrophic state (see 4.2 and 4.3). This is in line with other studies investigating interannual variability in taxonomic compositions of planktonic communities (Wiktor and Wojciechowska, 2005; Kubiszyn et al., 2014; Hegseth et al., 2019; Dabrowska et al., 2020, 2021; Dybwad et al., 2021; Assmy et al., 2023). In 2018, the relatively warm polar surface layer (>2.5 °C) extended further north (to at least station P5) and had a larger vertical extent (ca. 50 m) than in 2019, indicative of longer exposure to surface insolation. This was reflected in the deeper nutricline of nitrate, silicate and phosphate and much less pronounced subsurface chl a maxima in 2018 compared to 2019 (Fig. 2C--F-). In 2019, the subsurface chl a maximum was clearly following the shoaling of the nutricline from south to north along the transect and was well within the euphotic zone (Fig. 2F). Although the euphotic zone extended to greater depths in 2018 due to lower particle concentrations and no sea-ice cover, except at the two northernmost ice stations (Fig. 2F), the deeper nutricline in 2018 likely limited phytoplankton growth. The water column at stations P1

and P2 south and north of the Polar Front showed similar nutrient concentrations in both years, likely due to the absence of sea ice over a longer period at these southernmost locations of the study area. There was no distinct surface low-salinity layer in 2018 compared to 2019 suggesting that wind events had likely mixed the surface layer although not sufficiently deep to entrain nutrients from sub-surface waters. The permanently open-water station P1 sustained higher late-summer protist standing stocks than meltwater-stratified previously ice-covered stations (P2 to P5 in 2018 and P2 to P4 in 2019) and likely showed less pronounced seasonality.

The more extensive ice cover in 2019 during the time of sampling (southern limit at ca. 79.8°N) resulted in a colder and fresher surface layer than in 2018. Autotrophic protists, including ice-associated taxa, had a higher contribution to protist standing stocks in August 2019, as suggested by the higher chl *a* concentrations compared to August 2018. In August 2019, chl *a* concentrations peaked at station P5 with 2.6 μ g L⁻¹, indicating that the pelagic ecosystem was in a late-bloom situation. During the spring bloom chl *a* concentrations can reach up to 20 μ g L⁻¹ (Engelsen et al., 2002), while in summer, concentrations typically do not exceed 2 μ g L⁻¹ in Atlantic waters (Piwosz et al., 2009). In August 2018, chl *a* concentrations were very low (<0.5 μ g L⁻¹), typical for summer conditions in the study area (Dalpadado et al., 2014). This was also reflected in 2–22-fold higher chl *a* standing stocks in 2019 (range from 34 – 67 mg chl *a* m⁻²) compared to 2018 (range from 3 to 18 mg chl *a* m⁻²).



Fig. A7. Vertical flux of protist A) abundances $(10^6 \text{ cells m}^{-2} \text{ d}^{-1})$ and B) biomass (mg C m⁻² d⁻¹) per depth (m) of all identified taxa along the sampling transects in the northwestern Barents Sea in August 2018 (top panels) and August 2019 (bottom panels). Note: data not available for all stations.

4.2. Differences in protist composition and consequences for vertical carbon flux

which was in line with the shallower silicline as diatoms require a minimum of 2 μ M to dominate phytoplankton communities (Egge and Aksnes, 1992). This can be largely attributed to the above-mentioned differences in seasonal succession stage linked to the onset of ice melt

Diatoms had a higher contribution to protist standing stocks in 2019,

Table A1

Sea-ice concentration and properties along the sampling transects in the northwestern Barents Sea in the two sampling years 2018 and 2019. Nd: data not available.

Station	Lat (°N)	Lon (°E)	Sea-ice concentration	Primary ice type	Ice thickness (cm)	Snow depth (cm)	Melt pond (tenths)
2018							
P1	75.9966	31.2299	0				
P2	77.5015	33.9962	0				
P3	78.7508	33.9978	0				
P4	79.6932	33.9961	0				
P5	80.5006	34.0064	0				
PICE1	83.3321	31.5402	8	FYI	100	1	3
SICE3	83.2321	25.6739	9	SYI	180	2	1
2019							
P1	76.0000	31.2194	0				
P2	77.5006	33.9865	0				
P3	78.7498	34.0008	0				
P4	79.6932	34.2300	6	FYI < 70 cm	nd	nd	nd
P5	80.5289	33.9602	8	FYI < 70 cm	nd	nd	nd
P6	81.585	31.5195	9	FYI < 70 cm	nd	nd	nd
P7	81.9262	29.1396	9	FYI < 70 cm	nd	nd	nd
SICE4	81.9957	24.9952	7	$FYI < 70 \ cm$	nd	nd	nd

Table A2

Descriptive statistics for estimated monthly chlorophyll (chl) *a* climatologies considering the area of interest ($71^{\circ} - 80^{\circ}$ N latitude and $16^{\circ} - 52^{\circ}$ E longitude).

Month	Mean chl a		Standard	d deviation	Variance		
	2018	2019	2018	2019	2018	2019	
April	0.77	0.97	1.37	2.07	1.87	4.27	
May	5.16	1.72	6.48	2.11	41.93	4.47	
June	1.23	0.91	1.50	0.82	2.25	0.67	
July	0.53	0.66	0.41	0.86	0.17	0.74	
August	0.62	0.60	0.71	0.83	0.50	0.70	
September	0.91	1.14	0.73	1.32	0.53	1.74	

as diatom standing stocks were generally elevated at the ice-covered stations. It is interesting to note that in both years, diatom standing stocks were extremely low at open-water stations that were previously ice-covered, while at the permanently open-water station P1, diatom standing stocks were comparable to those in ice-covered waters. Diatoms showed the strongest relationship to sea ice at all locations, except for the permanently open-water station P1, suggesting strong ice-ocean feedbacks on nutrient and light availability *via* sea-ice meltwater stratification as well as snow and sea-ice light attenuation, respectively.

Species of the ice-associated diatom genus Fragilariopsis (Ratkova and Wassmann, 2002; Lundholm and Hasle, 2010; Szymanski and Gradinger, 2016), were found at high abundance near the ice edge (stations P4 and P5) in 2019 (Fig. A.4), while the centric diatom Shionodiscus bioculatus contributed a large share of diatom standing stocks at the northernmost ice stations in 2018 and 2019 (Fig. 6A). This species has been shown to accumulate in sea-ice ridges (Syvertsen 1991; Fernández-Méndez et al., 2018), hence its dominance in the water column likely reflects cryo-pelagic coupling caused by ice melt. The elevated vertical carbon flux of 296 mg C m⁻² d⁻¹ (station P5) observed in 2019 (Fig. 8B) was dominated by pelagic diatoms of the genus Thalassiosira and roughly 2-fold higher than the flux dominated by flagellates and dinoflagellates ($<160 \text{ mg C m}^{-2} \text{ d}^{-1}$). Thalassiosira species are known to dominate Arctic spring blooms (von Quillfeldt, 2000; Degerlund and Eilertsen, 2010; Hegseth et al., 2019; Assmy et al., 2023) and significantly contribute to vertical carbon flux (Rynearson et al., 2013; Dybwad et al., 2021) and were likely late remains of the preceding bloom. It needs to be noted that many of the dominant diatoms in both years were species with a relatively wide geographic distribution and are known to often occur later in the season, such as Thalassiosira nordenskioeldii (von Quillfeldt, 2000), Chaetoceros debilis/curvisetus (Tomas, 1997) and the Rhizosolenia species. The genus Rhizosolenia hosts large cylindrical species that have been associated with a shade flora growing at low light levels, likely facilitated by buoyancy regulation to allow them to migrate

between the euphotic zone and the deep nutricline (Kemp et al., 2000).

Athecate dinoflagellates of the genus *Gymnodinium* made a large contribution to dinoflagellate standing stocks (Fig. 6B, Fig. A.5). This diverse but understudied genus (Kubiszyn and Wiktor, 2016) is known to include mixotrophic and heterotrophic species (Tomas, 1997; Stoecker and Lavrentyev, 2018). The obligatory heterotrophic dinoflagellate genus *Protoperidinium* was found predominantly at station P1 in 2018 and is known to feed on large diatoms *via* their pallium feeding mode (Jacobson and Anderson, 1986). The prominent role of dinoflagellates is also reflected in their high contribution to biomass flux (Fig. 8B).

Within the flagellate community, the prymnesiophyte Phaeocystis pouchetii, an important bloom former in the Atlantic sector of the Arctic (Degerlund and Eilertsen, 2010; Ardyna et al., 2020; Dabrowska et al., 2020), sticks out with the highest abundances at the two northernmost ice-covered stations in 2018 (Fig. A.4), likely fuelled by the still elevated surface nutrient concentrations. This species often occurs at a later successional stage after the diatom spring bloom (von Quillfeldt, 2000; Hegseth et al., 2019), which is consistent with the generally low diatom biomass in 2018 (Fig. 4B). Deep mixing at station P1, indicated by elevated temperatures down to > 100 m, might have facilitated downward transport of Phaeocystis single cells, but overall, the carbon flux at station P1 and PICE1 was relatively low ($<150 \text{ mg C m}^{-2} \text{ d}^{-1}$). This is in agreement with previous vertical flux studies from the Barents Sea and north of Svalbard that have shown that a shift from a diatom to a Phaeocystis dominance could weaken the biological carbon pump (Wiedmann et al., 2020; Dybwad et al., 2021). The observed increase of Phaeocystis pouchetii in the Barents Sea (Orkney et al., 2020) could hence have a negative impact on strength of the biological carbon pump.

Altered temperature and seawater properties have further been suggested to favour the coccolithophore Emiliania huxleyi (Smyth et al., 2004; Giraudeau et al., 2016), which has shown a poleward expansion (Hegseth and Sundfjord, 2008; Winter et al., 2014; Oziel et al., 2017; Neukermans et al., 2018). Somewhat surprisingly, E. huxleyi contributed very little to protist biomass in 2018 and was virtually absent in 2019. The bloom of this species had likely been terminated by the time of sampling. Silkin et al. (2020) reported E. huxleyi blooms in the Barents Sea every summer between 2014 and 2018, typically being most prevalent in August (Signorini and McClain, 2009; Hovland et al., 2014). Our sampling stations, except P1, were likely located above its northernmost distribution limit, as this species is associated with Atlantic water masses (Silkin et al., 2020). This is supported by metabarcoding data which detected this species predominantly at station P1 in both years (B. Edvardsen, pers. comm.). Furthermore, it cannot be excluded that coccolithophores lost their scales, to some extent, during sample fixation and were consequently categorised as unidentified flagellates. Both P. pouchetii and E. huxleyi are predicted to increase in abundance and

Table A3

Overview of station presence of all identified protists from Niskin bottle samples collected during August 2018 and August 2019 in the northwestern Barents Sea.

Таха	Class	Species	August 2018	August 2019	Carbon conversion factor (pg C cell $^{-1}$)
Diatoms	Bacillariophyceae	Attheya longicornis	SICE3	P6, P7, SICE4	12
Diatoms	Bacillariophyceae	Bacterosira bathyomphala	P1, SICE3	P6	383
Diatoms	Bacillariophyceae	Chaetoceros borealis	P1, PICE1, SICE3	P4, P5, P6, P7, SICE4	403
Diatoms	Bacillariophyceae	Chaetoceros cf. atlanticus	SICE3		276
Diatoms	Bacillariophyceae	Chaetoceros cf. borealis/convolutus/	P1, P3		270
D	D 111 1 1	concavicornis	25		000
Diatoms	Bacillariophyceae	Chaetoceros cf. borealis/eibenii	P5 DICE1 CICE2	DF D7	339
Diatoms	Bacillariophyceae	Chaetoceros cf. debilis/curvisetus	PICE1, SICES	P5, P7 P6	204
Diatoms	Bacillariophyceae	Chaetoceros concavicornis f trisetosa	TIGET	P1 P7 SICE4	177
Diatoms	Bacillariophyceae	Chaetoceros debilis	PICE1, SICE3	SICE4	208
Diatoms	Bacillariophyceae	Chaetoceros decipiens	P3, PICE1, SICE3	P4, P6, P7, SICE4	563
Diatoms	Bacillariophyceae	Chaetoceros furcellatus	P2, PICE1, SICE3	P3, P5, P6, SICE4	24
Diatoms	Bacillariophyceae	Chaetoceros gelidus (formerly C. socialis)		P6, SICE4	21
Diatoms	Bacillariophyceae	Chaetoceros laciniosus		P3, P7, SICE4	144
Diatoms	Bacillariophyceae	Chaetoceros similis		P1, SICE4	102
Diatoms	Bacillariophyceae	Chaetoceros sp.	P1, PICE1, SICE3	P2	25
Diatoms	Bacillariophyceae	Chaetoceros subtilis	01070	P1 Pí	27
Diatoms	Bacillariophyceae	Chaetoceros tenuissimus	SICE3	P6	6.0
Diatoms	Bacillariophyceae	Chaetoceros teres	D4 DICE1	SICE4	4//
Diatoms	Bacillariophyceae	Chaeloceros wighama	P4, PICEI	P7, SIGE4	91
Diatoms	Bacillariophyceae	Coscinouiscus sp. 60–70 µm	D1 D2 D5 DICE1	PO, P7 D1 D3 D6 D7 SICE4	23
Diatoniis	Dacmanophyceae	Symmotica closter ann	SICE3	11,10,10,17,31024	20
Diatoms	Bacillarionhyceae	Diploneis littoralis	PICE1		413
Diatoms	Bacillariophyceae	Entomoneis kiellmanii	SICE3		951
Diatoms	Bacillariophyceae	Eucampia groenlandica	PICE1, SICE3	P4, P6, P7, SICE4	966
Diatoms	Bacillariophyceae	Fragilariopsis cf. nana		P1, P5	3.5
Diatoms	Bacillariophyceae	Fragilariopsis cylindrus	PICE1	P4, P5, P6, P7, SICE4	12
Diatoms	Bacillariophyceae	Fragilariopsis nana		P4, P5, P7	3.5
Diatoms	Bacillariophyceae	Fragilariopsis oceanica	PICE1	P7, SICE4	17
Diatoms	Bacillariophyceae	Gyrosigma fasciola	PICE1		801
Diatoms	Bacillariophyceae	Lennoxia faveolata		P3, P4, P7	3.0
Diatoms	Bacillariophyceae	Leptocylindrus danicus	P1, SICE3		95
Diatoms	Bacillariophyceae	Leptocylindrus minimus	P1	P1, SICE4	22
Diatoms	Bacillariophyceae	Licmophora gracilis	P1, P2, SICE3	P5	683
Diatoms	Bacillariophyceae	Melosira arctica	PICE1, SICE3		199
Diatoms	Bacillariophyceae	Navicula sp. 50–60 µm	PICEI	DA CICEA	891 540
Diatoms	Bacillariophyceae	Navicula directa	DE	P4, SICE4	100
Diatoms	Bacillariophyceae	Navicula transitans	PJ	D4 D6 D7 SICE4	109
Diatoms	Bacillariophyceae	Nitzschia 150–160 um	SICE3	r4, r0, r7, 516E4	411
Diatoms	Bacillariophyceae	Nitzschia 50–60 µm	51020	P6, SICE4	152
Diatoms	Bacillariophyceae	Nitzschia 70–80 µm	PICE1	,	207
Diatoms	Bacillariophyceae	Nitzschia 90–150 µm	P2		321
Diatoms	Bacillariophyceae	Nitzschia frigida	PICE1	P4, P6, P7, SICE4	29
Diatoms	Bacillariophyceae	Nitzschia promare	PICE1	SICE4	89
Diatoms	Bacillariophyceae	<i>Opephora</i> 60–70 μm		P6	279
Diatoms	Bacillariophyceae	Pennales 20–30 µm	P4, P5		13
Diatoms	Bacillariophyceae	Pennales 40–50 µm	P5	P7	65
Diatoms	Bacillariophyceae	Pennales 50–60 µm	PICE1		279
Diatoms	Bacillariophyceae	Pleurosigma stuxbergii	P1		539
Diatoms	Bacillariophyceae	Porosira glacialis		SICE4	3388
Diatoms	Bacillariophyceae	Prodoscia alata	DI DO DICEI CICEO	P1, P7, SICE4	1876
Diatoms	васшатюрпусеае	Pseudo-nitzscnia ci. delicatissima/	P1, P2, PICE1, SICE3	P1, P4, P5, P6, P7,	18
Diatoms	Bacillarionbyceae	Pseudo-nitzschia cf. granii		D1 SICE4	38
Diatoms	Bacillariophyceae	Pseudo-nitzschia cf. pungens	SICES	P1, 51014	147
Diatoms	Bacillariophyceae	Pseudo-nitzschia pungens	51615	SICE4	147
Diatoms	Bacillariophyceae	Rhizosolenia hebetata f. hebetata	P1, PICE1. SICE3	P1	1679
Diatoms	Bacillariophyceae	Rhizosolenia hebetata f. semispina	P1, PICE1, SICE3	P1, P6, P7, SICE4	1260
Diatoms	Bacillariophyceae	Shionodiscus bioculatus (formerly	PICE1, SICE3	P5, P6, P7, SICE4	2279
		Thalassiosira bioculata)		·	
Diatoms	Bacillariophyceae	Skeletonema cf. costatum		P5, SICE4	
					33
Diatoms	Bacillariophyceae	Skeletonema marinoi		P3, P5	33
Diatoms	Bacillariophyceae	Synedropsis hyperborea	PICE1	P6, P7, SICE4	18
Diatoms	Bacillariophyceae	Thalassiosira 10–20 µm		P1, P2, P5, P7, SICE4	121
Diatoms	Bacillariophyceae	Thalassiosira 20–30 µm	P3, P5, PICE1	P1, P2, P4, P5, P6, P7,	340
Distores	Pagillarianharasa	The lassing 20, 40	DI DO DE DICEI	SICE4	770
Diatoms	васшатюрпусеае	1 natassiosira 30–40 µm	гі, гэ, гэ, PIGEI	r'/	//0

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Таха	Class	Species	August 2018	August 2019	Carbon conversion factor (pg $C \text{ cell}^{-1}$)
Diatoms	Bacillariophyceae	Shionodiscus bioculatus var. exigua (formerly Thalassiosira bioculata var.	P1, SICE3	P7	2279
	N 111 1 1	exigua)		DI DC DE CICEA	
Diatoms	Bacillariophyceae	Thalassiosira cf. gravida/antarctica	P1, PICE1, SICE3	P1, P6, P7, SICE4	762
Diatoms	Bacillariophyceae	Thalassiosira cf. pacifica/hyalina		P4, P5	746
Diatoms	Bacillariophyceae	Thalassiosira gravida		P6	762
Diatoms	Bacillariophyceae	Thalassiosira nordenskioeldii	PICE1, SICE3	P4, P6, P7, SICE4	427
Dinoflagellates	Dinophyceae	Alexandrium 20–30 µm	SICE3	P1	1860
Dinoflagellates	Dinophyceae	Alexandrium 30–40 µm		P3, P7	2563
Dinoflagellates	Dinophyceae	Amphidinium longum	P1, P2, P3	P1, P2, P3, P4, P6, SICE4	189
Dinoflagellates	Dinophyceae	Amphidinium sphenoides	P2, P3, P4, P5, PICE1, SICE3	P1, P2, P5, P6, P7, SICE4	99
Dinoflagellates	Dinophyceae	Amphidoma acuminata	SICE3		52
Dinoflagellates	Dinophyceae	Amylax triacantha	P1, P3		815
Dinoflagellates	Dinophyceae	Cochlodinium 70–80 um		SICE4	2925
Dipoflagellates	Dinophyceae	Dicrogrisma peilopargialla	SICES	DIGET	126
Dinoflagellates	Dinophyceae	Dirophysege 20, 40 um	DICE1		2022
Dinonagenates	Dinophyceae	Dinophycede 30–40 µm	PICEI		3832
Dinoflagellates	Dinophyceae	Dinophysis acuta	P1		0408
Jinoflagellates	Dinophyceae	Gonyaulax spinifera	P1, P5		2123
Dinoflagellates	Dinophyceae	Gymnodiniales indet.	P2		1849
Dinoflagellates	Dinophyceae	Gymnodiniales indet. 10–20 µm	P5		130
Dinoflagellates	Dinophyceae	Gymnodiniales indet. 20-30 µm	P1, P2, P3, P5	P2, P4, P6, SICE4	484
Dinoflagellates	Dinophyceae	Gymnodiniales indet. 30–40 um	P3	P1, P3, SICE4	1223
Dinoflagellates	Dinophyceae	Gymnodiniales indet, 40–50 um	P3. PICE1	P2	2243
Dinoflagellates	Dinophyceae	Gymnodiniales indet. 50, 60 um	10,11011	D6	2210
Dinoflagellates	Dinophyceae	Gymnodinium 5–10 µm	P1, P2, P3, P4, P5,	P6 P4, P5, P6, P7	49
Dinoflagellates	Dinophyceae	Gymnodinium 10–20 µm	P1, P2, P3, P4, P5, PICE1_SICE3	P1, P2, P3, P4, P5, P6, P7, SICE4	130
Dinoflagellates	Dinophyceae	Gymnodinium 20–30 µm	P1, P2, P3, P5, PICE1, SICE3	P1, P2, P5, P7, SICE4	484
Dinoflagellates	Dinophyceae	Gymnodinium arcticum	P5 PICE1	P6 SICF4	121
Dipoflagellates	Dinophyceae	Cymnodinium of filum	SICE2	10,01011	1056
	Dinophyceae	Gynnodinium Ci. Juum	DI DO DO DA DE	D1 D0 D0 D4 D5 D6	1950
	Dinophyceae		P1, P2, P3, P4, P5, PICE1, SICE3	P1, P2, P3, P4, P5, P6, P7, SICE4	200
Dinoflagellates	Dinophyceae	Gymnodinium gracilentium	PI, P2, P3, P4, P5, PICE1	P1, P2, P3, P4, P7, SICE4	266
Dinoflagellates	Dinophyceae	Gymnodinium ostenfeldii 10–20 µm	PICEI		57
Dinoflagellates	Dinophyceae	Gymnodinium	P1, P2, P5	P5, P6, SICE4	1849
Dinoflagellates	Dinophyceae	Gymnodinium wulffii	P1, P2, P3, P4, P5, PICE1, SICE3	P1, P3, P5, P6, P7, SICE4	182
Dinoflagellates	Dinophyceae	Gyrodinium	P3		2567
Dinoflagellates	Dinophyceae	Gyrodinium 10–20 µm	P2	P1, SICE4	91
Dinoflagellates	Dinophyceae	Gyrodinium 20–30 µm	P1, P2, P3, P4, P5, PICE1	P3, P6, P7	432
Dinoflagellates	Dinophyceae	Gyrodinium cf. fusiforme 20–30 um	P2		2487
Dinoflagellates	Dinophyceae	Gyrodinium 30–40 µm	P1, P2, P3, P4, P5, PICE1	P2, P4, P5, P6	1164
Dinoflagellates	Dinophyceae	Gyrodinium 40–50 um	P3	P5. SICE4	2243
Dinoflagellator	Dinophyceae	Gyrodinium 50, 60 um	D4 DICE1	D3	2802
vinoflagellates	Dinophyceae	Curodinium 60, 70 ·····	F7, FIGE1	1.5	2002
	Dinophyceae	Gyrounium $60-70 \ \mu m$	SIGES DE		5/51 5/51
Jinoflagellates	Dinophyceae	Gyrodinium ct. gravida/antarctica	P5		826
Dinoflagellates	Dinophyceae	Gyrodinium cf. pepo	P1		2567
Dinoflagellates	Dinophyceae	Gyrodinium flagellare	Р5	P2, P3, P4, P5, P6, P7, SICE4	8
Dinoflagellates	Dinophyceae	Gyrodinium fusiforme	P1, P2, P3, P4, P5, PICE1, SICE3	P2, P3, P4, P5, P6, P7, SICE4	2429
Dinoflagellates	Dinophyceae	Gyrodinium grave	P1, P2, P3, SICE3	P2, P3, P6, P7, SICE4	2567
Dinoflagellates	Dinophyceae	Gyrodinium spirale	P1, P3, P5, PICE1	P1, P2, P3, P4, P6, P7, SICE4	4392
Dinoflagellates	Dinophyceae	Heterocapsa 30–40 µm		SICE4	856
Dinoflagellates	Dinophyceae	Heterocapsa rotundata	P2, P3, SICE3	P1, P2, P3, P4, P6, P7	36
Dinoflagellates	Dinophyceae	Heterocapsa sp.		P6	178
Dinoflagellates	Dinophyceae	Heterocapsa triauetra		P2, P5	225
Dinoflagellates	Dinophyceae	Lebouridinium glaucum	P1, P2, P3, P4, P5, PICE1, SICE3	P1, P2, P3, P5, P6, P7, SICE4	439
Dinoflagellates	Dinonhycese	Lessardia elopaata	SICE3	D1 D4 D6 D7 CICEA	166
Dinoflagellates	Dinophyceae	Micracanthodinium claytonii	P2, P5, PICE1, SICE3	P1, P2, P3, P4, P6, P7,	546
				SICE4	
Dinoflagellates	Dinophyceae	Nematopsides vigilans		P4, P6, SICE4	499
Dinoflagellates	Dinophyceae	Peridiniales	P1, P2		1900
Dinoflagellates	Dinophyceae	Peridiniales 10–20 µm	P1, P2, P3, P4, P5, PICE1, SICE3	P2, P3, P4, P5, P6, P7, SICE4	128

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Table A3 (continued)

Table AS (continu	leu)				
Таха	Class	Species	August 2018	August 2019	Carbon conversion factor (pg C cell $^{-1}$)
Dinoflagellates	Dinophyceae	Peridiniales 20–30 µm	P1, P2, P3, P4, P5, PICE1, SICE3	P1, P2, P3, P4, P5, P6, P7, SICE4	506
Dinoflagellates	Dinophyceae	Peridiniales 30–40 µm	P1, P2, P4	P1, P3, P6	950
Dinoflagellates	Dinophyceae	Phalacroma rotundatum (formerly Dinophysis rotundata)	P1	P1, P4, P5, SICE4	2479
Dinoflagellates	Dinophyceae	Polarella glacialis	P1	P3, P4, P7, SICE4	51
Dinoflagellates	Dinophyceae	Pronoctiluca pelagica	P1, P3, P4, PICE1,	P3, P5, P7, SICE4	297
Dinoflagellates	Dinophyceae	Prorocentrum cordatum	P1, P2, P3, P5, PICE1	P1, P3, P4, P5, P6, SICE4	173
Dinoflagellates	Dinophyceae	Protoperidinium	P1, P3		6473
Dinoflagellates	Dinophyceae	Protoperidinium 20–30 um	P1	P7	517
Dinoflagellates	Dinophyceae	Protoperidinium 30–40 um	P1, P3		1383
Dinoflagellates	Dinophyceae	Protoneridinium 40–50 um	P1		2867
Dinoflagellates	Dinophyceae	Protoperidinium 50–60 um	P1		5116
Dipoflagellates	Dinophyceae	Protoperidinium bines	D1 D2 SICE3	D1 D4 D5 D7 SICE4	172
Dinoflagenates	Dinophyceae	Protopertaintain opes	P1, P3, SICES	P1, P4, P3, P7, SIGE4	1/2
Dinonagenates	Dinophyceae	Protopertaintain brevipes	PI, SICES	P1, P4, P0	1280
Dinoflagellates	Dinopnyceae	Protoperialnium cerasus	P3, SICE3	PI	3380
Dinoflagellates	Dinophyceae	Protoperidinium cf. depressum/oceanica	PI		12,388
Dinoflagellates	Dinophyceae	Protoperidinium depressum	P1		12,388
Dinoflagellates	Dinophyceae	Protoperidinium islandicum		P1	3116
Dinoflagellates	Dinophyceae	Protoperidinium oceanicum		P1	12,388
Dinoflagellates	Dinophyceae	Protoperidinium pallidum	P1	P1, P6	9764
Dinoflagellates	Dinophyceae	Protoperidinium pellucidum	P1, P2, P5, SICE3	P1, P6, P7, SICE4	3273
Dinoflagellates	Dinophyceae	Protoperidinium pyriforme	P1	P6	3380
Dinoflagellates	Dinophyceae	Scrippsiella trochoidea	P1, P4	P1, P6	695
Dinoflagellates	Dinophyceae	Torodinium robustum	P1, P2, P3, PICE1, SICE3	P1, P2, P4, P5, SICE4	569
Dinoflagellates	Dinophyceae	Tripos arcticus (formerly Ceratium arcticum)	P1	P1 P2	8128
Dinoflagellates	Dinophyceae	Tripos fusus	P1	,	2058
Dinonagenates	Chlorodendronbycese	Pachysphaera pelagica	D1 SICE2	D2 D4 D5 D7	63
Other	Chlorophygoag	Chlamudomonae on	FI, SIGES	r2, r4, r3, r7	71
Oulei As sallatas	Chlorophyceae	Chianyaomonas sp.		P1, P3, P7	/1
Other	Choanoflagellatea	Bicosta spinifera		P3, P4, P6, SICE4	5.1
flagellates Other	Choanoflagellatea	Calliacantha natans		P1 P2 P3 P4 P5 P7	39
flagellates	Choanoflagellatea	Monosiaa marina	D5 DICE1 SICE3	SICE4 P2 P3 P4 P5 P6 P7	9
flagellates	Chosen flagellates		13, 11021, 51025	SICE4	10
flagellates	Choanoriageilatea	Saipingoeca inquillata		P2, P4, P7	10
Other flagellates	Chrysophyceae	Chrysophyceae sp.	P3	P2, P3, P4, P5, P6, P7, SICE4	77
Other flagellates	Chrysophyceae	Dinobryon balticum	P1, P2, P3, P4, P5, PICE1, SICE3	P1, P2, P3, P4, P6, P7, SICE4	18
Other flagellates	Chrysophyceae	Dinobryon faculiferum		P1, P3, P5, P6, P7, SICE4	11
Other flagellates	Chrysophyceae	Dinobryon sp.		P1, P5, P6, P7, SICE4	20
Other	Chrysophyceae	Ochromonas sp.	PICE1		7.0
Other	Cryptophyceae	Cryptophyceae indet.	P1, P2, P3, P5, SICE3	P1, P2, P3, P4, P5, P6,	34
Other	Cryptophyceae	Leucocryptos marina	P1, P2, P3, P4, P5,	P1, P2, P3, P4, P5, P6,	24
Other	Cryptophyceae	Plagioselmis prolonga	P5	P2, P4, P7	10
Other	Cryptophyceae	Rhodomonas sp.		P5, SICE4	16
Other	Cryptophyceae	Teleaulax amphioxeia		P2	21
Other	Cryptophyceae	Teleaulax sp.	P2, SICE3	P2, P3, P4, P5, P6, P7,	27
Other	Dictyochophyceae	Apedinella radians	P1, P3	SICE4 P1, P3, P6, SICE4	50
tiagellates Other	Dictyochophyceae	Octactis speculum (formerly Dictyocha	P1, PICE1, SICE3	P3, P4, P5, P6, P7,	461
Other	Dictyochophyceae	speculum) Pseudopedinella pyriformis		SICE4 P4	30
tiageilates Other	Euglenoidea	Euglenoidea indet.	P4	P4, P7	233
flagellates Other	Euglenoidea	Eutreptiella sp.		P7	489
tlagellates Other flagellates	Eukaryota incertae sedis	Flagellates indet.	PICE1, SICE3	P2	67

Table A3 (continued)

Таха	Class	Species	August 2018	August 2019	Carbon conversion factor (pg C cell $^{-1}$)
Other	Eukaryota incertae sedis	Flagellates indet. 0–3 µm	PICE1		1.2
Other	Eukaryota incertae sedis	Flagellates indet. 3–7 µm	P1, P2, P3, P4, P5,	P1, P2, P3, P4, P5, P6,	10
flagellates Other	Eukaryota incertae sedis	Flagellates indet. 7–10 µm	PICE1, SICE3 P1, P2, P5, PICE1,	P7, SICE4 P1, P2, P3, P5, P6, P7,	39
flagellates Other	Eukaryota incertae sedis	Flagellates indet. 10–20 µm	SICE3 P5, PICE1	SICE4 P2, P3, P5, P6, SICE4	205
flagellates Other	Pyramimonadophyceae	Halosphaera sp.		P5, SICE4	40
flagellates Other	Prasinophyceae	Pterosperma	P1		6005
flagellates Other	Prasinophyceae	Pterosperma parallelum		P1	6005
flagellates Other	Pyramimonadophyceae	Pyramimonas nansenii	SICE3	P7	4.2
flagellates Other	Pyramimonadophyceae	Pyramimonas virginica		P2, P4, P6	4.6
flagellates Other	Prymnesiophyceae	Chrysochromulina spp.	P2, P5, PICE1	P2, P3, P6, P7	38
flagellates Other	Prymnesiophyceae	Phaeocystis pouchetii	P1, P2, P4, PICE1,	P4, P5, P6, P7, SICE4	9.4
flagellates	Prymnesionhyceae	Prympesionhyceae 10–20 um	SICE3	P3	62
flagellates	Prympesiophyceae	Dynamimonas ef virainica		P6	4.6
flagellates	Drymposionhyceae	Pyramimonas en	D1 CICES		90 90
flagellates	Prynnesiopnyceae	Pyruminonus sp.	FI, SICES	P2, P4, P3, P7, SICE4	02
flagellates	Raphidophyceae			SICE4	8.0
Other flagellates	Raphidophyceae	Heterosigma sp.	P2, P3	P1, P2, P6, P7, SICE4	242
Other flagellates	Telonemea	Telonema subtile	PICE1	P2, P6, P7	7.9
Other flagellates	Xanthophyceae	Meringosphaera mediterranea		P2, P3	89
Ciliates	Ciliophora	Ciliophora		P3	245
Ciliates	Ciliophora	Ciliophora 30–40 µm	P1, P2, P4	P5, P7	1072
Ciliates	Ciliophora	Ciliophora 40–50 µm	P1	P5	1135
Ciliates	Ciliophora	Ciliophora 50–60 µm	P1		1298
Ciliates	Ciliophora	Ciliophora 60–70 um	P1 P3		1661
Ciliatea	Cilianhara	Cilianhara 70, 90 um	D1		1001
Cillates	Chiophora		PI		1924
Ciliates	Ciliophora	Ciliophora 80–90 µm	P1, P5		2910
Ciliates	Gymnostomatea	Didinium gargantua		P2, P7, SICE4	4048
Ciliates	Gymnostomatea	Didinium sp.	P5, PICE1, SICE3	P2	4171
Ciliates	Gymnostomatea	Mesodinium rubrum	P1, P2, P3, P4, P5, PICE1, SICE3	P3, P4, P5, P6, P7, SICE4	3248
Ciliates	Gymnostomatea	Mesodinium ruhrum 40-50 um	P1	broet	3248
Ciliates	Limetrichee	Finlates on	11	D6	066
Cillates	Hypotricilea	Euplotes sp.		PO	966
Ciliates	Oligohymenophorea	Scuticociliatia indet. 10–20 µm	P3		532
Ciliates	Oligohymenophorea	Scuticociliatia indet.	P1, P2, P3, P5, PICE1	P2, P3, P6, P7, SICE4	532
Ciliates	Oligotrichea	Acanthostomella norvegica	P2, P3	P2, P3, P4, P5, P6, P7, SICE4	312
Ciliates	Oligotrichea	Laboea strobila	P1, P2, P3, P4, P5, SICE3	P1, P2, P3, P4, P6	2711
Ciliates	Oligotrichea	Leegaardiella ovalis	P1, P3, P5, PICE1, SICE3	P2, P5, SICE4	672
Ciliates	Oligotrichea	Leevaardiella sol	P1	SICE4	456
Ciliates	Oligotrichea	Laprotintinnus en	D1 D2	DIGET	1660
Ciliates	Oligotrichea	Leprotitututus sp.	P1, P2, D2, D4, D5	D2 D2 D4 D5 D6 D7	204
Cillates	Oligotricliea	Lonmanniella ovijormis	P1, P2, P3, P4, P5, PICE1	F2, F3, F4, F5, F6, F7, SICE4	204
Ciliates	Oligotrichea	Lohmanniella oviformis 20–30 µm	P2		204
Ciliates	Oligotrichea	Parafavella gigantea	P5, SICE3		28,883
Ciliates	Oligotrichea	Parafavella obtusangula	P1	P1, P3	26,503
Ciliates	Oligotrichea	Paratontonia gracillima		P4, P7	1660
Ciliates	Oligotrichea	Ptychocylis obtusa	P1, P2, P3, P5	P2, P3, P4, P6, P7,	1660
Ciliates	Oligotriches	Salningella secata	P1	JIGLT	1660
Cilictor	Oligotrishas	Supilizena secun	r 1	D7	1072
Cillates	oligouricnea	Saroomanni striatum		P/	10/2
Ciliates	Oligotrichea	Strombidium	P2, PICE1	P2, P6, P7	1010
Ciliates	Oligotrichea	Strombidium 10-20 µm	P2, P4, PICE1, SICE3	P2, P6, P7	185
Ciliates	Oligotrichea	Strombidium 20-30 µm	P1, P2, P3, P4, P5, PICE1, SICE3	P2, P3, P4, P5, P6, P7, SICE4	779
Ciliates	Oligotrichea	Strombidium 30-40 µm	P1, P2, P3, P4, P5, PICE1, SICE3	P2, P3, P6, P7, SICE4	2009

Table A3 (continued)

Таха	Class	Species	August 2018	August 2019	Carbon conversion factor (pg C cell $^{-1}$)
Ciliates	Oligotrichea	Strombidium cf. acutum 30–40 µm	P1		2009
Ciliates	Oligotrichea	Strombidium 40–50 µm	P1, P2, P3, P5		4077
Ciliates	Oligotrichea	Strombidium 50–60 µm	P3		7176
Ciliates	Oligotrichea	Strombidium 60–70 μm	P1, P2	P7	11,488
Ciliates	Oligotrichea	Strombidium cf. acutum	P1, P2, P3		2009
Ciliates	Oligotrichea	Strombidium cf. scutellum	P3		2009
Ciliates	Oligotrichea	Strombidium cf. sulcatum	P5, SICE3	P4, P5, P6	4077
Ciliates	Oligotrichea	Strombidium cf. sulcatum/vestitum		P6	2762
Ciliates	Oligotrichea	Strombidium cf. vestitum	P4, P5	P2	1403
Ciliates	Oligotrichea	Strombidium conicum	P1, P2, P3, P4, P5,	P1, P2, P3, P4, P5, P6,	11,170
			PICE1, SICE3	P7, SICE4	
Ciliates	Oligotrichea	Strombidium constrictum	P1, P3, P4, P5, SICE3	P1, P2, P3, P4, P6, P7,	6377
				SICE4	
Ciliates	Oligotrichea	Tintinnus inquilinus	P1, P2, P5	P1	1660
Ciliates	Prostomatea	Balanion comatum	P1		245
Ciliates	Prostomatea	Prorodon ovum	P1		2910

extend poleward as a result of Atlantification (Neukermans et al., 2018; Orkney et al., 2020).

The dominance of Dinobryon, Heterosigma and Chrysochromulina at the other stations both in 2018 and 2019 (Fig. 6C) can likely be explained by their mixotrophic feeding mode, which will give them a competitive advantage in the oligotrophic surface layer since they can supplement their diet by feeding on bacteria, which has not been shown for Phaeocystis. Indeed, under-ice blooms of Chrysochromulina under low nutrient conditions have been attributed to mixotrophy (Søgaard et al., 2021) and Dinobryon is known to thrive under low nutrient and light conditions late in the season (McKenrie et al., 1995; Hegseth et al., 2019). The kleptoplastidic ciliates Strombidium and Mesodinium rubrum, which keep the plastids of their prey for a while to perform photosynthesis, dominated ciliate standing stocks (Fig. 6D), supporting the general finding that mixotrophy is a prominent trophic mode in the Arctic Ocean, especially during summer (Stoecker and Lavrentyev, 2018). Interestingly, at open water station P1, in both 2018 and 2019, protist standing stocks were among the highest, particularly if flagellates are excluded, but abundances of ciliates and dinoflagellates were rather low, illustrating that particularly large taxa, such as thecate dinoflagellates of the genus Protoperidinium and tintinnid ciliates of the genus Parafavella, were largely restricted to this open water station. Overall, it can be concluded that the relative contribution of heterotrophic and mixotrophic protists to total protist standing stocks was higher in 2018 than in 2019, reflecting the more oligotrophic status and more advanced successional stage of the protist community in August 2018. This is also confirmed by the higher contribution of protist carbon flux to total POC fluxes in 2019 compared to 2018, suggesting that vertical flux in 2018 had a higher share of regenerated material and detritus.

4.3. Relationship between protist size structure and nutrient concentrations

The most pronounced differences in the protist communities between the late summers were observed for nanophytoplankton. Its biomass was considerably higher in August 2019 than in August 2018 (Fig. 7B), reflected in the higher chl *a* standing stocks in 2019. Indeed, much of the difference in chl *a* standing stocks can likely be explained by this group as the interannual differences in larger protists identified under light microscopy were less pronounced, also indicating that nanoplanktonic protists were likely not fully captured by the latter method. This is further supported by the close match of nanophytoplankton biomass, as measured by flow cytometry, and the subsurface chl *a* maximum in both years. This match with the subsurface chl *a* maximum was particularly evident for the larger of the two nanophytoplankton size fractions (5–10 µm; data not shown), indicating a close association with the nutricline. Biomass of the smaller nanophytoplankton size fraction (2–5 μ m) and picophytoplankton (Fig. 7A) on the other hand showed an inverse pattern with chl *a* concentrations and were generally most abundant where nutrient concentrations were the lowest. The biomass of HNF largely matched the subsurface chl *a* maximum (Fig. 7C), particularly in 2019, suggesting active feeding on smaller phytoplankton. However, they also matched the abundance of large bacteria (Amargant-Arumí et al. in prep., same issue), which was likely their main food source since HNF are major consumers of bacteria in the ocean (Sherr et al., 1997; Kopylov et al., 2016). Thus, the most likely scenario is that elevated levels of labile organic carbon in the subsurface chl *a* maximum sustained bacterial growth, which again supported their HNF grazers.

In the context of environmental change, the sea-ice conditions and consequently protist community structure in the northwestern Barents Sea during 2018 can give us a glimpse into the future when the Barents Sea is predicted to be sea-ice free during the summer (Onarheim and Årthun, 2017; Serreze and Meier, 2019), and longer open-water periods are likely to ultimately favour mixo- and heterotrophic flagellates and their protozoan grazers due to higher surface temperatures and decreased nutrient availability as outlined in Fig. 9. It is suggested that small-celled species, covering pico- and smaller nanophytoplankton fractions, will benefit from oligotrophic environments with longer openwater seasons (Nöthig et al., 2015; Zhang et al., 2015; Wang et al., 2019) as they can adapt to low nutrient conditions (Zhang et al., 2016) due to their larger surface-to-volume-ratio and thus superior nutrient uptake capability (Litchman et al., 2007). In comparison to e.g. (large-celled) diatoms, which are in the preferred prey-size class of Calanus copepods (Mullin, 1963), an essential link between primary production and higher trophic levels, smaller cells are suboptimal for efficient grazing by mesozooplankton (Levinsen et al., 2000; Kiørboe, 2011). While picophytoplankton is significant for the ocean's carbon cycle (Buitenhuis et al., 2012; Richardson, 2019), changes in protist community composition with the warming of the ocean are expected to have direct consequences for copepod grazers and the associated food web (Li et al., 2009; Finkel et al., 2010; Dąbrowska et al., 2020; Thingstad, 2020).

5. Conclusions

Our study shows interannual variability in summer protist communities during two years with contrasting sea-ice conditions in the northwestern Barents Sea. The environmental conditions encountered in August 2018 are likely representative of the future Barents Sea, with earlier sea-ice melt and a longer open water season (Onarheim and Årthun, 2017). Annual primary production in the Arctic Ocean has been shown to increase as a result of a longer open water season (Arrigo and van Dijken, 2015), and these changes in the physical

Table A4

Overview of all identified species of sinking protists from sediment trap deployments collected during August 2018 and August 2019 in the Barents Sea. *species not observed in sediment trap samples.

Johane Datame	Таха	Class	Species	August 2018	August 2019	Carbon conversion factor (pg C $cell^{-1}$)
DistanceBeditatiophysicsAddres squares interainsPIC PIPICPICPICPICDistanceBeditatiophysicsAddres squares interainsPIC PIPIC PIPICPICDistanceBeditatiophysicsConserver convolutePIC PIPIC PIPICPICDistanceBeditatiophysicsConserver convolutePIC PIPICPICPICDistanceBeditatiophysicsConserver convolutePIC PIPICPICPICDistanceBeditatiophysicsConserver on forelinePICPICPICPICDistanceBeditatiophysicsConserver on forelinePICPICPICPICDistanceBeditatiophysicsConserver on forelinePICPICPICPICDistanceBeditatiophysicsConserver on forelinePICPICPICPICDistanceBeditatiophysicsConserver on forelinePICPICPICPICDistanceBeditatiophysicsConserver on forelinePICPICPICPICPICDistanceBeditatiophysicsConserver on forelinePICPICPICPICPICPICDistanceBeditatiophysicsConserver on forelinePIC<	Diatoms	Bacillariophyceae	Actinocyclus sp.*		P4	2460
Distance Distance Distance Distance BeciliariophysicsAstronuc subsymptials Cancourse break Cancourse break Cancou	Diatoms	Bacillariophyceae	Actinocyclus tenuissimus*		P4	1794
Diskow Diskow	Diatoms	Bacillariophyceae	Attheva septentrionalis*	P1. PICE1	P5. P6	82
DistantBetHardphysicsChemicronic barrellsPAL <td>Diatoms</td> <td>Bacillariophyceae</td> <td>Bacterosira bathyomphala</td> <td>PICE1</td> <td>P6, P7</td> <td>383</td>	Diatoms	Bacillariophyceae	Bacterosira bathyomphala	PICE1	P6, P7	383
Distant Distant Distant Bellinicity Distant Bellin	Diatoms	Bacillariophyceae	Chaetoceros borealis		P4, P5, P6, P7	403
Distance Di	Diatoms	Bacillariophyceae	Chaetoceros concavicornis	PICE1	P5. P6	177
Dialon: Data Data BacillarisphyseCharacorus despina (ancorus despina)PICHN. N. P.S.3Data Data Data BacillarisphyseCharacorus spika (ancorus despina)PICHN. N. P.21Data Data Data BacillarisphyseCharacorus spika (ancorus)PICHN. N. P.21Data Data Data BacillarisphyseCharacorus spika (ancorus)PICHN. P. S. N.30Data Data Data Data BacillarisphyseCharacorus calcians (ancorus)PICHN. P. S. N.30Data Data Data Data Data BacillarisphyseCostonolics spi (ancorus)PI CHN. N. N.30.323Data Data Data Data BacillarisphyseCostonolics spi (ancorus)PI CHN. N. N.30.323Data Data Data BacillarisphysePropiorispi (anan (ancorus)PI CH <td>Diatoms</td> <td>Bacillariophyceae</td> <td>Chaetoceros convolutus</td> <td>11021</td> <td>P1 P7</td> <td>231</td>	Diatoms	Bacillariophyceae	Chaetoceros convolutus	11021	P1 P7	231
Discours Discours Bacillarisphyrese Bacillarisphyrese Chanceres sp.Chanceres sp. PCL3PCL3 PCL4 PCL3P, F7 PCL3 PCL324Discours Bacillarisphyrese Chanceres sp. Bacillarisphyrese Chanceres sp. Chanceres sp. Cha	Diatoms	Bacillariophyceae	Chaetoceros deciniens	PICF1	P6 P7	563
DiscoseBacillarispicesConserver splats (formatly Coscials)PIC1 <td>Diatoms</td> <td>Bacillariophyceae</td> <td>Chaetoceros furcellatus</td> <td>D2 D4 DICE1</td> <td>D4 D5 D6 D7</td> <td>24</td>	Diatoms	Bacillariophyceae	Chaetoceros furcellatus	D2 D4 DICE1	D4 D5 D6 D7	24
Distance Distance Distance Distance Bardiariophyce Bardiariophyce Chancers sp.PICE1 PICE1 PICE1 PI, P4, P5, P014 PA PICE1 PI, P4, P5, P0144 PICE1 PI, P1, P1, P1, P1, P1, P1, P1, P1, P1, P1	Diatoms	Bacillariophyceae	Chaetoceros galidus (formerly C socialis)	DICE1	D6 D7	21
Dialons Dialons Bendlariophysics Dialons Bendlariophysics Dialons Bendlariophysics Concords/ansignmuPICEL P1, P2, PICEL P1, P2, PICEL P1, P4, P5, P5P1 P2 P2 P2 P1, P4, P5, P5P1 P2 P2 P2, PICEL P1, P4, P5, P5P1 P2 P2, PICEL P1, P4, P5, P5P1 P2 P2, PICEL P1, P4, P5, P5P1 P2 P2, PICEL P1, P4, P5, P5P1 P2 P2 P2, PICEL P1, P2, PICEL P1, P2, PICEL P1, P2, PICEL P1, P4, P5, P5P1 P2 P	Diatoms	Pacillariophyceae	Chaetoceros genuus (tormerry C. sociaits)	DICE1	r0, r7	21
Diacons Diacons Bacillariophyces Bacillariophyces Chastores mainstrans Chastores mainstrans Chastores mainstrans Philes Philes	Diatoms	Bacillariophyceae	Chaetoceros sp	DICE1	D1	25
Jackman (P)Jackman (P) <th< td=""><td>Diatoms</td><td>Pacillariophyceae</td><td>Chaetoceros sp.</td><td>D1 D2 DICE1</td><td></td><td>25</td></th<>	Diatoms	Pacillariophyceae	Chaetoceros sp.	D1 D2 DICE1		25
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Distors Distors Distors Balancian physics Distors Balancian physics Balancian physics Distors Balancian physics Balancian physics Distors Balancian physics Balancian physics Distors Balancian physics Balancian physics Distors Balancian physics Balancian Balancian physics Balancian	Diatome	Bacillariophyceae	Coscinodiscus controlis*		D5	36 320
Diacons BacillariophycesSolidariophyces ProglarepatricPL2 IPCLI PL <b< td=""><td>Diatoms</td><td>Pacillariophyceae</td><td>Coscinodiscus centralis</td><td></td><td>FJ DE D6 D7</td><td>7067</td></b<>	Diatoms	Pacillariophyceae	Coscinodiscus centralis		FJ DE D6 D7	7067
Datama Datama BacillariophycesDistanta Equipation 	Diatoms	Bacillarianhusses	Cosciliouiscus sp.	D1 D2 DICE1	P3, P0, P7	22
DatasetDatasetDatasetDatasetDatasetDatasetPressPressDistanceBacillanciphyceePressPressPressPressPressDistanceBacillanciphyceePressPressPressPressPressDistanceBacillanciphyceeNoricel optagiaPressPressPressPressDistanceBacillanciphyceeNoricel optagiaPressPressPressPressPressDistanceBacillanciphyceeNoricel optagiaPressPressPressPressPressDistanceBacillanciphyceeNoricel optagiaPress<	Diatoms	Bacillarianhusses	Cyunaromeca closterium	PI, PZ, PICEI	P1, P4, P3, P7	23
JackanowsBacklinkolaytyckeeProglampies LymnusPL PLSPL PLSPL PLSPL PLSPL PLSPL PLSPL PLSPL PLSPL	Diatoms	Bacillariophyceae	Eucampia groenianaica Enceileriopeis culindrus	PICEI DI DICEI	P4, P5, P0, P7	900
Bacharson DistoresReallistrophyceseProglaringsis accandingProglaringsis Proglaringsis accandingProglaring Proglaringsis Proglaringsis accandingProglaring Proglaringsis Proglaringsis accandingProglaring 	Diatonis	васшанорнусеае	Fraguariopsis Cyunarias	PI, PIGEI	P1, P4, P5, P0,	12
Data Data Data Data Data DataData Data DataProperty as a Data Data DataProperty as Data Data Data DataDeterm Data <br< td=""><td>Distants</td><td>Desillarianhusses</td><td>Tanailanian sia mana</td><td></td><td>P7</td><td>2 5</td></br<>	Distants	Desillarianhusses	Tanailanian sia mana		P7	2 5
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Data Data Data Data 	Diatoms	Bacillariophyceae	Fraguariopsis oceanica	PICEI	P4	1/
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Diatoms Diatoms BacillariophyceeAdvicul splantans	Diatoms	Bacillariophyceae	Navicula pelagica	PICEI	P4, P5, P6	109
Dutons Dutons BacillanophyceaeNakila sys.P5240Dutons BacillanophyceaeNakila transitorsPICEP4P7Dutons BacillanophyceaeNakila transitorsPICEP4, P5, P629DutonsBacillanophyceaeNitzschia frigdaPICEP428DutonsBacillanophyceaeNitzschia frigdaPICEP4539DutonsBacillanophyceaePeterosigna sy.P4539DutonsBacillanophyceaePeterosigna sy.P4539DutonsBacillanophyceaePeterosigna sy.P1P1, P1P1, P5, P6, P718DutonsBacillanophyceaePeterosinschia synauP1, P1P1, P1, P5, P6, P718DutonsBacillanophyceaePeterosinschia synauP1, P1P1, P1, P5, P6, P718DutonsBacillanophyceaePeterosinschia synauP1P1P2P3DutonsBacillanophyceaeNakoselna hobetan L hobetanP1P1P2P3DutonsBacillanophyceaeShorodocus hobetan CP1P2P3P3DutonsBacillanophyceaeShorodocus hobetan CP1P2P3P3DutonsBacillanophyceaeThalassiofra QangP1P2P3P3DutonsBacillanophyceaeThalassiofra QangP1P3P3P4DutonsBacillanophyceaeThalassiofra QangP1P2P3P4P3DutonsBacillanophyceaeTh	Diatoms	Bacillariophyceae	Navicula septentrionalis*		P5	87
Datoms Datoms BacillariophyceeNorkula vankedfanit*Pro12/Datoms BacillariophyceeNitzskia frigidaPICE1P4, P5, P629Datoms BacillariophyceeNitzskia frigidaPICE1P4, P5, P628Datoms BacillariophyceePauloila tominat*PICE1P1, P5, P6, P718Datoms BacillariophyceeProvinsi gloidailsP6338Datoms BacillariophyceeProvinsi gloidailsP1, PICE1P1, P5, P6, P718Datoms BacillariophyceeProvinsi gloidailsP1, PICE1P1, P5, P6, P718Datoms BacillariophyceePseudo-nitzskin graniP1, PICE1P1, P5, P6, P718Datoms BacillariophyceeRhizosoleni hobetara 1. sensipinaP1, PICE1P1, P5, P6, P718Datoms BacillariophyceeRhizosoleni hobetara 1. sensipinaP1, PICE1P1, P5, P6, P718Datoms BacillariophyceeRhizosoleni hobetara 1. sensipinaP1, PICE1P1, P5, P6, P718Diatoms DatomsBacillariophyceeThalassioaira 20-30 µnP1P1220Diatoms DatomsBacillariophyceeThalassioaira 20-30 µnP1P274Diatoms DatallariophyceeThalassioaira 4, revisflogis*P1P1P3260Diatoms DatallariophyceeThalassioaira 4, revisflogis*P1P2P1P1Diatoms DatallariophyceeThalassioaira 4, revisflogis*P1P2P4260Diatoms DiatomsBacillariophycee<	Diatoms	Bacillariophyceae	Navicula sp.		P6	260
Datoms Datoms Bacillariophyce DiatomsNurclai vandergima"PICE1PI77Diatoms Bacillariophyce DiatomsBacillariophyce Bacillariophyce Plansing globalPICE1PI28Diatoms Bacillariophyce DiatomsBacillariophyce Plansing globalPICE1PI59Diatoms Bacillariophyce DiatomsBacillariophyce Planda-mitsching grani princePIPICE1PI, PCE1PI, PF, P6, P718Diatoms Bacillariophyce DiatomsBacillariophyce Planda-mitsching grani princePIPICE1PI, PF, P6, P718Diatoms Bacillariophyce DiatomsBacillariophyce Bacillariophyce Plansichin stratating grani Bacillariophyce PlantomsPIPICE1PI, PF, P6, P718Diatoms DiatomsBacillariophyce BacillariophyceNincoolenia hebetari I. hebrain BacillariophycePIPICE1PI, P5, P6, P718Diatoms DiatomsBacillariophyce BacillariophyceShinondscub hebrain I. hebrains Bacillariophyce Diatosistic 2-30 µmPICE1P1, P5, P6, P718Diatoms DiatomsBacillariophyce Diatassistic 2-30 µmPICE1P1, P2, P5, P6, P7150Diatoms DiatomsBacillariophyce Diatassistic a cl, weisfigit*PICE1P1, P2, P5, P6, P7150Diatoms DiatomsBacillariophyce Diatassistic a rankelodidiPICE1P1, P2, P5, P6, P7150Diatoms DiatomsBacillariophyce Diatassistic a rankelodidiPICE1P1, P2, P5, P6, P7150Diatoms <b< td=""><td>Diatoms</td><td>Bacillariophyceae</td><td>Navicula transitans</td><td></td><td>P6</td><td>127</td></b<>	Diatoms	Bacillariophyceae	Navicula transitans		P6	127
Datoms Datoms BacillariophyceeNitzsche Ingission* Nitzsche Ingission*PICE1P4, P5, P6, 29Diatoms DatomsBacillariophyceePaulella tartistatia P1FA539Diatoms DiatomsBacillariophyceePorosina glocialisP63388Diatoms DiatomsBacillariophyceePorosina glocialisP1, PICE1P1, P5, P6, P718Diatoms DiatomsBacillariophyceePaudo-nitzsche grani P1, PICE1P1, P5, P6, P718Diatoms DiatomsBacillariophyceePaudo-nitzsche grani Statosoleria hebetata 1. hebetataP1, PICE1P1, P5, P6, P718Diatoms DiatomsBacillariophyceeRhizosoleria hebetata 1. sensipinaP1P11260Diatoms DiatomsBacillariophyceeSkeletonema costatum bioculariaP1P11260Diatoms DiatomsBacillariophyceeThalassioira 20-30 µmP1P11260Diatoms DiatomsBacillariophyceeThalassioira 20-30 µmP1P492Diatoms DiatomsBacillariophyceeThalassioira C, gravida/antarcicaP1P19292Diatoms DiatomsBacillariophyceeThalassioira C, weisflögi*P6P774Diatoms DiatomsBacillariophyceeThalassioira C, weisflögi*P1P2P4260Diatoms DiatomsBacillariophyceeThalassioira C, weisflögi*P1P2P4260Diatoms DiatomsBacillariophyceeThalassioira C, weisflögi*P1<	Diatoms	Bacillariophyceae	Navicula vanhoeffenii*	B10004	P4	77
Diatoms Bacillariophyceae Nitschial ongissima* PI CEI 28 Diatoms Bacillariophyceae Plearosigma sp. P4 539 Diatoms Bacillariophyceae Perado mizschia granii P1, PCEI P1, PS, P6, P7 18 Diatoms Bacillariophyceae Peado mizschia granii P1, PCEI P1, PA, P5, P5, P7 38 Diatoms Bacillariophyceae Peado mizschia granii P1, PCEI P1, P5, P6, P7 18 Diatoms Bacillariophyceae Rheado-mizschia granii P1 268 Diatoms Bacillariophyceae Rheado-mizschia granii P1 1260 Diatoms Bacillariophyceae Sheadomizschia granii P1 276 33 Diatoms Bacillariophyceae Sheadomizschia granii P1 P1, P5, P6, P7 18 Diatoms Bacillariophyceae Sheadomizschia granii dramizschia P1CEI P1, P4, P5, P6, P7 2279 Diatoms Bacillariophyceae Thalassisniar 1, Palassianiar P1CEI P1, P4, P5, P6, P7 2279 Diatoms	Diatoms	Bacillariophyceae	Nitzschia frigida	PICE1	P4, P5, P6	29
Diatoms DiatomsBacillariophyceae Berussian glaciallisPrice Persian glaciallisPrice Persian glaciallisPrice Persian glaciallisPersian Persian Persian PersianPersian Persian Persian PersianPersian Persian Persian PersianPersian Persian Persian PersianPersian Persian Persian PersianPersian Persian Persian Persian PersianPersian Persian PersianPersian Persian PersianPersian Persian PersianPersian Persian Persian PersianPersian Persian Persian PersianPersian Persian Persian PersianPersian Persian Persian PersianPersian Persian Persian PersianPersian Persian Persian Persian Persian PersianPersian Persian Persian Persian Persian Persian Persian Persian PersianPersian Persian Persian Persian Persian Persian Persian PersianPersian Persian<	Diatoms	Bacillariophyceae	Nitzschia longissima*	P1		28
Diatoms DiatomsBacillariophyceae BacillariophyceaePetrologing ap.P4S39DiatomsBacillariophyceaePetrologing aprilP1, PICEIP1, P5, P6, P718DiatomsBacillariophyceaePetrologing aprilP1, PICEIP1, P4, P5, P738DiatomsBacillariophyceaePetrologing aprilP1, PICEIP1, P5, P6, P718DiatomsBacillariophyceaeRhizoschnia hebetata 1. thebatanaP1P1, P5, P6, P71670DiatomsBacillariophyceaeShitomschnia hebetata 1. thebatanaP1P1, P5, P6, P718DiatomsBacillariophyceaeShitomschnia hebetata 1. fubratanaP1P1, P5, P6, P718DiatomsBacillariophyceaeShitomschnia hebetata 1. fubratanaP1P1, P5, P6, P72279DiatomsBacillariophyceaeShitomschnia hebetata 1. fubratanaP1CEIP1, P4, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira of gravida/antarcticaP1CEIP1, P4, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira of gravida/antarcticaP1CEIP1, P4, P5, P6, P7159DiatomsBacillariophyceaeThalassiosira of gravida/antarcticaP1CEIP1, P4, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira of gravida/antarcticaP1CEIP1, P4, P5, P6, P7231DiatomsBacillariophyceaeThalassiosira of gravida/antarcticaP1CEIP5, P6, P7230DiatomsBacillariophyceaeThalassiosira on diatarctic	Diatoms	Bacillariophyceae	Pauliella taeniata*	PICE1		69
Diatoms DiatomsBacillariophyceae Peudo-nizschia cf. delicatissima/pseudodelicatissima PI, PICE1PI, P5, P7, P738Diatoms DiatomsBacillariophyceaePeudo-nizschia cf. delicatissima/pseudodelicatissima PI, PICE1P1, P5, P7, P738Diatoms DiatomsBacillariophyceaePeudo-nizschia erita/*P1268Diatoms DiatomsBacillariophyceaeRhizosolenia hebetara f. hebetaraP1, PICE1P1, P5, P6, P718Diatoms DiatomsBacillariophyceaeSkeletomena costatumP1126033DiatomsBacillariophyceaeSkeletomena costatumP11260DiatomsBacillariophyceaeThalassiosira 20-30 µnP191, P5, P6, P7279DiatomsBacillariophyceaeThalassiosira 20-30 µnP1, P5, P6, P7279DiatomsBacillariophyceaeThalassiosira (Grmerly ThalassiosiraP1CE1P1, P5, P6, P7279DiatomsBacillariophyceaeThalassiosira ordenskiedaliP1P1, P5, P6, P7762DiatomsBacillariophyceaeThalassiosira nordenskiedaliP1P5, P6, P7762DiatomsBacillariophyceaeThalassiosira nordenskiedaliP1P5, P6, P7762DiatomsBacillariophyceaeThalassiosira nordenskiedaliP1P5, P6, P7764DiatomsBacillariophyceaeThalassiosira nordenskiedaliP1P6, P7764DiatomsBacillariophyceaeThalassiosira nordenskiedaliP1P2P6P7Diatoms <t< td=""><td>Diatoms</td><td>Bacillariophyceae</td><td>Pleurosigma sp.</td><td></td><td>P4</td><td>539</td></t<>	Diatoms	Bacillariophyceae	Pleurosigma sp.		P4	539
DiatomsBacillariophyceaePeaudo-nitschia cf. delicatistma/pseudodelicatistmaP1, PCR1P1, P4, P5, P6, P718DiatomsBacillariophyceaePicho-nitschia seriata*P1, PCR1P1, P4, P5, P6, P738DiatomsBacillariophyceaeRhizosolenia hebetata f. hebetataP1, PCR1P1, P5, P6, P7167DiatomsBacillariophyceaeRhizosolenia hebetata f. hebetataP1, PCR1P1, P5, P6, P718DiatomsBacillariophyceaeSkelcanema costamaP1P1, P5, P6, P718DiatomsBacillariophyceaeShinodicus biosolatusP1P1, P5, P6, P718DiatomsBacillariophyceaeShinodicus biosolatusP1P1, P5, P6, P718DiatomsBacillariophyceaeShinodicus biosolatus (Grmentery ThalassiosiraP1CE1P1, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira cl. gravida/antarcticaP1CE1P1, P4, P5, P6, 7627279DiatomsBacillariophyceaeThalassiosira cl. gravida/antarcticaP1P5, P6, P7159DiatomsBacillariophyceaeThalassiosira rolemskiedetiP1P1, P4, P5, P6, 77746DiatomsBacillariophyceaeAlexandrium sp.P1, P2, P4, P1, P5, P6, P71301DiatomsBacillariophyceaeAlexandrium sp.P1, P2, P4, P1, P4, P51301DinofagellatesDinophyceaeAlexandrium sp.P1, P2, P4, P1, P4, P5126DinofagellatesDinophyceaeDinophyceaeAnghidiniam indensidesP1, P2, P4, P728	Diatoms	Bacillariophyceae	Porosira glacialis		P6	3388
DiatomsBacillariophyceaePseudo-nitschia graniPi (Pi ChPi (Pi Pi, Pi, Pi	Diatoms	Bacillariophyceae	Pseudo-nitzschia cf. delicatissima/pseudodelicatissima	P1, PICE1	P1, P5, P6, P7	18
DiatomsBacillariophyceaePasudo-mizschia seriara*P1268DiatomsBacillariophyceaeRhizosolani hebetara f. semispinaP1, PIC.1P1, PS, P6, P71679DiatomsBacillariophyceaeSkeleronera costatumP1P1C.1P1, PS, P6, P718DiatomsBacillariophyceaeSyndropis hyperboreaPICE1P1, PS, P6, P718DiatomsBacillariophyceaeShinondiscus bioculatus (formerly ThalassiosiraPICE1P1, PS, P6, P72279DiatomsBacillariophyceaeShinondiscus bioculatus (formerly ThalassiosiraPICE1P1, P4, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira cf. gravida/antarcticaPICE1P1, P4, P5, P6, P7267DiatomsBacillariophyceaeThalassiosira f. gravida/antarcticaPICE1P1, P4, P5, P6, P7159DiatomsBacillariophyceaeThalassiosira f. gravida/antarcticaPICE1P5, P6, P7159DiatomsBacillariophyceaeThalassiosira hydina*P1, P2, P4, P1, P1, P4, P51301DiatomsBacillariophyceaeThalassiosira pnetholesP1, P2, P4, P1, P4, P51301DinoflagellatesDinophyceaeAmphidmini ngenoP1P2P699DinoflagellatesDinophyceaeCochodmium sp.P1P2P1, P2, P12380DinoflagellatesDinophyceaeDinophyceaeDinophyceae99373DinoflagellatesDinophyceaeDinophyceaeP1, P2P1P14693Dinofla	Diatoms	Bacillariophyceae	Pseudo-nitzschia granii	P1, PICE1	P1, P4, P5, P7	38
DiatomsBacillariophyceaeRhizosolenia hebetata i. hebetataPI, PICEIPI, PICEIPI, PICEIPI, PICEIPI, PICEI </td <td>Diatoms</td> <td>Bacillariophyceae</td> <td>Pseudo-nitzschia seriata*</td> <td></td> <td>P1</td> <td>268</td>	Diatoms	Bacillariophyceae	Pseudo-nitzschia seriata*		P1	268
Diatoms DiatomsBacillariophyceaeRhiesolenia hebetari f. semispinaPI1260Diatoms DiatomsBacillariophyceaeSyledcomac costatumPICE1PI, PS, P6, P718Diatoms DiatomsBacillariophyceaeSyledcomac costaturPICE1PI, PS, P6, P72279DiatomsBacillariophyceaeShionofistus bioclants (formerly Thalassiosira costaturPICE1PI, P4, PS, P6, P72279DiatomsBacillariophyceaeThalassiosira c1: gravida/antarcticaPICE1PI, P4, PS, P6, P7746DiatomsBacillariophyceaeThalassiosira tordinskieddiiPICE1P5, P6, P7746DiatomsBacillariophyceaeThalassiosira nordenskieddiiPICE1P5, P6, P7427DiatomsBacillariophyceaeThalassiosira nordenskieddiiPICE1P5, P6, P7427DiatomsBacillariophyceaeAlexandrium sp.PICE1P1, P2, P4, P1, P2, P4, P1, P2, P4,P1, P4, P51301DinoflagellatesDinophyceaeAmphidinium sp.P1P1, P2P699DinoflagellatesDinophyceaeAmphidinium sp.P2P4, P5, P6, P72814DinoflagellatesDinophyceaeDinophyceaeAmphidinium sp.P1P14633DinoflagellatesDinophyceaeDinophyceaeP1P14633DinoflagellatesDinophyceaeDinophyceaeP1P14633DinoflagellatesDinophyceaeDinophyceaeP1P14633DinoflagellatesDi	Diatoms	Bacillariophyceae	Rhizosolenia hebetata f. hebetata	P1, PICE1	P1, P5, P6, P7	1679
DiatomsBacillariophyceaeSkeltonema costaumP533DiatomsBacillariophyceaeSynedrojski hyperboreaPICE1P1, P5, P6, P718DiatomsBacillariophyceaeThalassiosira 0.30 µmPICE1P1, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira cf. gravida/antarcticaPICE1P1, P4, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira cf. gravida/antarcticaPICE1P1, P4, P5, P6, P7762DiatomsBacillariophyceaeThalassiosira valuesPICE1P5, P6, P7159DiatomsBacillariophyceaeThalassiosira valuesPICE1P5, P6, P7427DiatomsBacillariophyceaeThalassiosira valuesPICE1P4260DinofingellatesDinophyceaeAlexandrium sp.P1, P2, P4,P1, P4, P51301DinofingellatesDinophyceaeAlexandrium sp.P1, P2, P4,P1, P4, P52814DinofingellatesDinophyceaeOcorlodinium sp.P2P4189DinofingellatesDinophyceaeOcorlodinium sp.P2733DinofingellatesDinophyceaeDinophyceae inder.P1P1, P4, P52814DinofingellatesDinophyceaeDinophyceae inder.P1P14693DinofingellatesDinophyceaeDinophyceae inder.P1P14693DinofingellatesDinophyceaeDinophyceaeOrganula vasP1P14693DinofingellatesDinophyceaeOrganu	Diatoms	Bacillariophyceae	Rhizosolenia hebetata f. semispina		P1	1260
DiatomsBacillariophyceaeSynchroyis hyperboreaPICE1PI, PS, P6, P718DiatomsBacillariophyceaeShionodiscus bioculatus (formerly Thalassiosira 20-30 µmPICE1P1, PS, P6, P72279DiatomsBacillariophyceaeThalassiosira 1: gravida/antarcticaPICE1P1, PS, P6, P7762DiatomsBacillariophyceaeThalassiosira 1: gravida/antarcticaPICE1P1, P4, P5, P6, P7762DiatomsBacillariophyceaeThalassiosira nordenskioeldiiPICE1P5, P6, P7746DiatomsBacillariophyceaeThalassiosira nordenskioeldiiPICE1P5, P6, P7427DiatomsBacillariophyceaeThalassiosira nordenskioeldiiPICE1P5, P6, P7427DiatomsBacillariophyceaeThalassiosira nordenskioeldiiP1, P2, P4,P1, P4, P51301DinoflagellatesDinophyceaeArghidrinium longumP4189DinoflagellatesDinophyceaeCocholnium sp.P1P2P699DinoflagellatesDinophyceaeDinophyceae 10-20 µmP2373373DinoflagellatesDinophyceaeDinophyceae 10-20 µmP1P1P14693DinoflagellatesDinophyceaeGornaulax graciits*P1P193363DinoflagellatesDinophyceaeGornaulax graciits*P1P1P14693DinoflagellatesDinophyceaeGornaulax graciits*P1P1P14693DinoflagellatesDinophyceaeGornaul	Diatoms	Bacillariophyceae	Skeletonema costatum		P5	33
DiatomsBacillariophyceaeThalassisira 2.0-30 µmPICE1P4340DiatomsBacillariophyceaeShionodicus bioculatus (formerly ThalassiosiraPICE1P1, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira cf. gravida/antarcticaPICE1P1, P4, P5, P6, P7762DiatomsBacillariophyceaeThalassiosira fuxilina*PICE1P5, P6, P7159DiatomsBacillariophyceaeThalassiosira nordenskioeldiiPICE1P5, P6, P7427DiatomsBacillariophyceaeTrachyneis sp.*P4260DinoflagellatesDinophyceaeAlexandrium sp.P1, P2, P4, P1, P4, P51301DinoflagellatesDinophyceaeAmphidinium longumPICE1P1146DinoflagellatesDinophyceaeCochlodrium sp.P2P499DinoflagellatesDinophyceaeDinophyceae indet.P1, P2, P4, P1, P4, P72380DinoflagellatesDinophyceaeDinophyceae indet.P1P14693DinoflagellatesDinophyceaeDinophyceae indet.P1, P4, P1, P4, P5, P6, P7130DinoflagellatesDinophyceaeDinophycia argunita*P1, P4, P5, P6, P72380DinoflagellatesDinophyceaeDinophycia argunita*P1P1P1DinoflagellatesDinophyceaeDinophyceaeP1, P4, P72380DinoflagellatesDinophyceaeDinophyceaeP1P1, P4, P5, P6, P7130DinoflagellatesDinophyceaeGoryandax sp.P	Diatoms	Bacillariophyceae	Synedropsis hyperborea	PICE1	P1, P5, P6, P7	18
DiatomsBacillariophyceaeShionodiscub hoculatus (formerly Thalassiosira bioculatus)PICE1P1, P4, P5, P6, P72279DiatomsBacillariophyceaeThalassiosira cf. gravida/antarcticaPICE1P1, P4, P5, P6, P7762DiatomsBacillariophyceaeThalassiosira cf. weissflögit*PICE1P5, P6, P7762DiatomsBacillariophyceaeThalassiosira nordenskioeldiiPICE1P5, P6, P7427DiatomsBacillariophyceaeThalassiosira nordenskioeldiiPICE1P4260DinoflagellatesDinophyceaeAlexandrium sp.P1, P2, P4, PICE1P1, P4, P51301DinoflagellatesDinophyceaeAmphidinium sp. pentoidesP1P2P699DinoflagellatesDinophyceaeAmphidinium sp. pentoidesP1P2126373DinoflagellatesDinophyceaeCochodinium sp. pentoidesP1P2124933DinoflagellatesDinophyceaeDinophyceae inder.P12380373DinoflagellatesDinophyceaeDinophyceae inder.P1P14693DinoflagellatesDinophyceaeOmphysis acuminata*P1P14693DinoflagellatesDinophyceaeGoryaulax gracifis*P1P14693DinoflagellatesDinophyceaeGoryaulax gracifis*P1P1P44880DinoflagellatesDinophyceaeGoryaulax gracifis*P1P1P44893DinoflagellatesDinophyceaeG	Diatoms	Bacillariophyceae	Thalassiosira 20–30 µm		P4	340
Diatoms DiatomsBacillariophyceae Fulassioira cf. weissflogii*PICE1P1, P4, P5, P6, P7762 P7DiatomsBacillariophyceae BacillariophyceaeThalassioira ch. weissflogii*PG, P7746DiatomsBacillariophyceae BacillariophyceaeThalassioira ch. weissflogiiPICE1P5, P6, P7427DiatomsBacillariophyceaeThalassioira nordenskioeldiiPICE1P5, P6, P7427DiatomsBacillariophyceaeTrachyneis sp.*P4260DinoflagellatesDinophyceaeAtexandrium sp.P1, P2, P4, P1, P2, P4,P1, P4, P51301DinoflagellatesDinophyceaeAmphidinium sphenoidesP1, P2P4189DinoflagellatesDinophyceaeCocholdnium sp.P2P4260DinoflagellatesDinophyceaeOcholdnium sp.P2P1, P2, P4, P1, P2, P4,P1, P2, P4,DinoflagellatesDinophyceaeOcholnium sp.P2P4260DinoflagellatesDinophyceaeDinophyceaeP1, P2P493DinoflagellatesDinophyceaeDinophyceaeP1, P2P4P1DinoflagellatesDinophyceaeDinophyceaeP1, P2P4P3DinoflagellatesDinophyceaeDinophyceaeP1, P4P3P3DinoflagellatesDinophyceaeDinophyceaeP1, P4, P5130DinoflagellatesDinophyceaeOmophysis norregica*P1P14693DinoflagellatesDinophyceaeGoryaulax sp. </td <td>Diatoms</td> <td>Bacillariophyceae</td> <td>Shionodiscus bioculatus (formerly Thalassiosira</td> <td>PICE1</td> <td>P1, P5, P6, P7</td> <td>2279</td>	Diatoms	Bacillariophyceae	Shionodiscus bioculatus (formerly Thalassiosira	PICE1	P1, P5, P6, P7	2279
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DinoflagellatesDinophyceaeDinophyceae 10-20 μmP2373DinoflagellatesDinophyceaeDinophyceae indet.P1, P2P1, P4, P72380DinoflagellatesDinophyceaeDinophycia acuminata*P41921DinoflagellatesDinophyceaeDinophysis norvegica*P1P14693DinoflagellatesDinophyceaeGonyaulax gracilis*P1, P4, PICE1P54880DinoflagellatesDinophyceaeGonyaulax gracilis*P1, P4, PICE1P7130DinoflagellatesDinophyceaeGymnodiniales 10-20 μmP1P1, P4, P5, P6, PICE1130DinoflagellatesDinophyceaeGymnodiniales 20-30 μmP1P1, P4, P5, P6, PICE14880DinoflagellatesDinophyceaeGymnodinium 10-20 μmP1P1, P4, P5, P6, PI130DinoflagellatesDinophyceaeGymnodinium 10-20 μmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium 10-20 μmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium 10-20 μmP2, P4, PICE1P1, P4, P5, P6, P7121DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1, P2, P4, P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182	Dinoflagellates	Dinophyceae	Dicroerisma psilonereiella		P5	126
DinoflagellatesDinophyceaeDinophyceae indet.P1, P2P1, P4, P72380DinoflagellatesDinophyceaeDinophysis acuminata*P41921DinoflagellatesDinophyceaeDinophysis norvegica*P1P14693DinoflagellatesDinophyceaeGonyaulax gracilis*P1, P4, PICE1P54880DinoflagellatesDinophyceaeGonyaulax sp.P1, P2, P4,P1, P4, P5, P6,130DinoflagellatesDinophyceaeGymnodiniales 10–20 μmP1P1P14693DinoflagellatesDinophyceaeGymnodiniales 20–30 μmP1, P2, P4,P1, P4, P5, P6,130DinoflagellatesDinophyceaeGymnodinium 10–20 μmP1P1, P4, P5, P6,484DinoflagellatesDinophyceaeGymnodinium 10–20 μmP1P2P4, P5130DinoflagellatesDinophyceaeGymnodinium 10–20 μmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium 10–20 μmP2, P4, PICE1P1, P4, P5121DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, PICE1P1, P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182	Dinoflagellates	Dinophyceae	Dinophyceae 10–20 µm	P2		373
DinoflagellatesDinophyceaeDinophysis acuminata*P41921DinoflagellatesDinophyceaeDinophysis norvegica*P1P14693DinoflagellatesDinophyceaeGonyaulax gracilis*P1, P4, PICE1P54880DinoflagellatesDinophyceaeGonyaulax gracilis*P1, P2, P4,P1, P4P8DinoflagellatesDinophyceaeGymnodiniales 10–20 µmP1, P2, P4,P1, P4, P5, P6,130DinoflagellatesDinophyceaeGymnodiniales 20–30 µmP1P1, P4, P5, P6,484DinoflagellatesDinophyceaeGymnodiniumP1P1, P4, P5, P6,484DinoflagellatesDinophyceaeGymnodinium 10–20 µmP1P1, P4, P5, P6,484DinoflagellatesDinophyceaeGymnodinium 10–20 µmP1P1, P4, P5130DinoflagellatesDinophyceaeGymnodinium 10–20 µmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium 10–20 µmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium 10–20 µmP2P4, P5, P6, P7121DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, P1CE1P1, P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffitP1P4, P5182	Dinoflagellates	Dinophyceae	Dinophyceae indet.	P1, P2	P1, P4, P7	2380
DinoflagellatesDinophyceaeDinophysis norvegica*P1P14693DinoflagellatesDinophyceaeGonyaulax gracilis*P1, P4, PICE1P54880DinoflagellatesDinophyceaeGonyaulax sp.P1, P4, PICE1P4, P5, P6,130DinoflagellatesDinophyceaeGymnodiniales 10–20 µmP1, P2, P4,P1, P4, P5, P6,484DinoflagellatesDinophyceaeGymnodiniales 20–30 µmP1P1, P4, P5, P6,484DinoflagellatesDinophyceaeGymnodinium 10–20 µmP1P1, P4, P5, P6,484DinoflagellatesDinophyceaeGymnodinium 10–20 µmP1P1P4, P5130DinoflagellatesDinophyceaeGymnodinium 10–20 µmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, P1CE1P1, P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1, P2, P4, P5182DinoflagellatesDinophyceaeGymnodinium cf. wulffitP1P4, P5182	Dinoflagellates	Dinophyceae	Dinophysis acuminata*		P4	1921
DinoflagellatesDinophyceaeGonyaulax gracilis*P1, P4, PICE1P54880DinoflagellatesDinophyceaeGonyaulax sp.P1, P4, P5, P6, P1, P4, P5, P6, P14880DinoflagellatesDinophyceaeGymnodiniales 10–20 μmP1, P2, P4, P1, P4, P5, P6, P7130DinoflagellatesDinophyceaeGymnodiniales 20–30 μmP1P1P7DinoflagellatesDinophyceaeGymnodinium 10–20 μmP1CE1P7DinoflagellatesDinophyceaeGymnodinium 10–20 μmP1CE11849DinoflagellatesDinophyceaeGymnodinium 10–20 μmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, PICE1P1, P4, P5121DinoflagellatesDinophyceaeGymnodinium cf. arcticumP1, P2, P4, P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffitP1P4, P5182	Dinoflagellates	Dinophyceae	Dinophysis norvegica*	P1	P1	4693
DinoflagellatesDinophyceaeGonyaulax sp.P1, P2, P4, P1, P2, P4, P1, P2, P4, P1, P4, P5, P6, P7130DinoflagellatesDinophyceaeGymnodiniales 10–20 μmP1 P1CE1P7DinoflagellatesDinophyceaeGymnodiniales 20–30 μmP1 P1 P1 P1P1, P4, P5, P6, P1 P1849DinoflagellatesDinophyceaeGymnodinium 10–20 μmP1CE11849DinoflagellatesDinophyceaeGymnodinium 10–20 μmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium 10–20 μmP2P4, P5121DinoflagellatesDinophyceaeGymnodinium cf. arcticumP1, P2, P4, P1, P2, P4,P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1P4, P5182DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182	Dinoflagellates	Dinophyceae	Gonyaulax gracilis*	P1, P4, PICE1	P5	4880
DinoflagellatesDinophyceaeGymnodiniales 10–20 μmP1, P2, P4, P1, P2, P4, P1, P4, P5, P6, P7130DinoflagellatesDinophyceaeGymnodiniales 20–30 μmP1P1, P4, P5, P6, P7484DinoflagellatesDinophyceaeGymnodinium 10–20 μmPICE11849DinoflagellatesDinophyceaeGymnodinium 10–20 μmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, PICE111, P4, P5121DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1, P2, P4, P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182	Dinoflagellates	Dinophyceae	Gonyaulax sp.		P1, P4	4880
 Dinoflagellates Dinophyceae Gymnodiniales 20–30 µm PICE1 P1, P4, P5, P6, P1 P2 P4 P5 P5 P6 P7 P6 P6<!--</td--><td>Dinoflagellates</td><td>Dinophyceae</td><td>Gymnodiniales 10–20 μm</td><td>P1, P2, P4,</td><td>P1, P4, P5, P6,</td><td>130</td>	Dinoflagellates	Dinophyceae	Gymnodiniales 10–20 μm	P1, P2, P4,	P1, P4, P5, P6,	130
DinoflagellatesDinophyceaeGymnodiniales 20–30 μmP1P1, P4, P5, P6, P7484 P7DinoflagellatesDinophyceaeGymnodinium 10–20 μmPICE11849DinoflagellatesDinophyceaeGymnodinium 10–20 μmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, PICE1P1, P4, P5121DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1, P2, P4, P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffitP1P4, P5182				PICE1	P7	
Pionflagellates Dinophyceae Gymnodinium PICE1 1849 Dinoflagellates Dinophyceae Gymnodinium 10–20 µm P2 P4, P5 130 Dinoflagellates Dinophyceae Gymnodinium cf. arcticum P2, P4, PICE1 P1, P4, P5 121 Dinoflagellates Dinophyceae Gymnodinium cf. gracilentum P1, P2, P4, P4, P5, P6, P7 266 Dinoflagellates Dinophyceae Gymnodinium cf. wulffit P1 P4, P5 182	Dinoflagellates	Dinophyceae	Gymnodiniales 20–30 µm	P1	P1, P4, P5, P6,	484
DinoflagellatesDinophyceaeGymnodiniumPICE11849DinoflagellatesDinophyceaeGymnodinium 10–20 μmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, PICE1P1, P4, P5121DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1, P2, P4,P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182					P7	
DinoflagellatesDinophyceaeGymnodinium 10–20 µmP2P4, P5130DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, PICE1P1, P4, P5121DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1, P2, P4,P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182	Dinoflagellates	Dinophyceae	Gymnodinium	PICE1		1849
DinoflagellatesDinophyceaeGymnodinium cf. arcticumP2, P4, PICE1P1, P4, P5121DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1, P2, P4,P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182	Dinoflagellates	Dinophyceae	Gymnodinium 10–20 µm	P2	P4, P5	130
DinoflagellatesDinophyceaeGymnodinium cf. gracilentumP1, P2, P4, PICE1P4, P5, P6, P7266DinoflagellatesDinophyceaeGymnodinium cf. wulffiiP1P4, P5182	Dinoflagellates	Dinophyceae	Gymnodinium cf. arcticum	P2, P4, PICE1	P1, P4, P5	121
PICE1 Dinoflagellates Dinophyceae <i>Gymnodinium</i> cf. <i>wulffii</i> P1 P4, P5 182	Dinoflagellates	Dinophyceae	Gymnodinium cf. gracilentum	P1, P2, P4,	P4, P5, P6, P7	266
Dinoflagellates Dinophyceae Gymnodinium cf. wulffii P1 P4, P5 182				PICE1		
	Dinoflagellates	Dinophyceae	Gymnodinium cf. wulffii	P1	P4, P5	182

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Table A4 (continued)

Таха	Class	Species	August 2018	August 2019	Carbon conversion factor (pg C $cell^{-1}$)
Dinoflagellates	Dinophyceae	Gymnodinium galeatum	P1, P2, P4, PICE1	P1, P4, P5, P6, P7	266
Dinoflagellates	Dinophyceae	Gymnodinium gracilentum		P1	266
Dinoflagellates	Dinophyceae	Gymnodinium ostenfeldii		P5	57
Dinoflagellates	Dinophyceae	Gymnodinium simplex*	P1		182
Dinoflagellates	Dipophyceae	Gymnodinium wulffii	P2	P4	182
Dinoflagellates	Dinophyceae	Gyrodinium 30_40 um		P1	1164
Dinoflagellates	Dinophyceae	Gyrodinium cf. gracilantum		D4	266
Dinoflagellates	Dinophyceae	Cyrodinium of wulffii	D1	14	4201
Dinonagenates	Dinophyceae	Gyrodinium Ci. widjju	PI DI DICEI	D1 D4 D5 D(4391
Dinofiagellates	Dinopnyceae	Gyroainium Jiageilare	PI, PICEI	P1, P4, P5, P6, P7	8.0
Dinoflagellates	Dinophyceae	Gyrodinium fusiforme	P1, P2, PICE1	P1, P4, P5, P6, P7	2429
Dinoflagellates	Dinophyceae	Gyrodinium grave		P4	2567
Dinoflagellates	Dinophyceae	Heterocapsa arctica*	P2		77
Dinoflagellates	Dinophyceae	Heterocapsa rotundata	P4, PICE1	P1, P4, P5, P6	36
Dinoflagellates	Dinophyceae	Karlodinium sp.*		P4	99
Dinoflagellates	Dinophyceae	Katodinium glaucum	P1, P2	P1, P4, P5, P6,	439
Dipoflogollator	Dipophysooo	Lossandia alangata	D1 DICE1	P7	166
Dinonagenates	Dinophyceae		PI, PICEI	P1, P4, P5, P7	100
Dinofiageilates	Dinophyceae	Micracaninoanium claytonii	D1 D4 D1000	P4, P5, P6, P7	540
Dinoflagellates	Dinopnyceae	Oxyrrnis sp.*	P1, P4, PICE1	D1 54	45
Dinoflagellates	Dinophyceae	Peridiniales 20–30 µm		P1, P4	506
Dinoflagellates	Dinophyceae	Phalacroma rotundatum (formerly Dinophysis rotundata)		P1	2479
Dinoflagellates	Dinophyceae	Polarella glacialis	P1	P1, P4, P5	51
Dinoflagellates	Dinophyceae	Pronoctiluca pelagica		P1, P4, P6	297
Dinoflagellates	Dinophyceae	Pronoctiluca sp	PICE1	P1 P5 P6 P7	297
Dinoflagellates	Dipophyceae	Prorocentrum cordatum	D1	11,10,10,11	173
Dinoflagellates	Dinophyceae	Prorocentrum sp	11	D1 D5	173
Dinoflagellates	Dinophyceae	Protocenti un sp.	D1	F1, FJ	172
Dinonagenates	Dinophyceae	Protopertaintum otpes	P1 D1 D2	P4	1/2
Dinonagenates	Dinophyceae	Protopertaintum brevipes	P1, P2	P1, P0, P7	1280
Dinoflagellates	Dinophyceae	Protoperialnium cerasus		PI	3380
Dinoflagellates	Dinophyceae	Protoperidinium cf. ovatum*	P1		5116
Dinoflagellates	Dinophyceae	Protoperidinium depressum	P1	P1	12,388
Dinoflagellates	Dinophyceae	Protoperidinium pellucidum	P1	P1, P4, P5, P6	3273
Dinoflagellates	Dinophyceae	Protoperidinium sp.		P5, P7	6473
Dinoflagellates	Dinophyceae	Scrippsiella sp.	P1	P5, P7	816
Dinoflagellates	Dinophyceae	Tripos arcticus (formerly Ceratium arcticum)	P1	P1, P4	8128
Other flagellates	Chlorodendrophyceae	Pachysphaera pelagica		P4	63
Other flagellates	Chlorodendrophyceae	Pachysphaera sp.	P1, P2	P1, P5, P6, P7	63
Other flagellates	Chlorophyceae	Chlamydomonadaceae indet.	· · · · · · · · · · · · · · · · · · ·	P4	71
Other flagellates	Chlorophyceae	Chlamydomonas sp		P4	71
Other flagellates	Chlorophyceae	Dunaliella sp.*	P2	P7	36
Other flagellates	Choapoflagellatea	Calliacantha natans	D1 D2	D1 D4 D5 D6	30
Other hagenates	Ciloanonagenatea		F1, F2	P7	55
Other flagellates	Choanoflagellatea	Choanoflagellate indet.	P4		5.0
Other flagellates	Choanoflagellatea	Diaphanoeca pedicellata*		P1	21
Other flagellates	Choanoflagellatea	Monosiga marina	P1, P2, PICE1	P1, P4, P5, P6,	9
				P7	
Other flagellates	Choanoflagellatea	Pleurasiga echinocostata*		P1	5.0
Other flagellates	Choanoflagellatea	Salpingoeca sp.		P6, P7	10
Other flagellates	Chrysophyceae	Chrysophyceae sp.	P1, P2, PICE1	P1, P4, P5, P6,	77
				P7	
Other flagellates	Chrysophyceae	Dinobryon balticum	P1, P2, P4,	P1, P4, P5, P6,	18
0.1 7 7			PICE1	P7	22
Other flagellates	Chrysophyceae	Dinobryon divergens*	P2		32
Other flagellates	Chrysophyceae	Dinobryon faculiferum		P1, P4, P7	11
Other flagellates	Cryptophyceae	Cryptomonas sp.	P1, P2, P4, PICE1	P1, P4, P5, P6, P7	220
Other flagellates	Cryptophyceae	Cryptophyceae indet.	P2	P1, P4	34
Other flagellates	Cryptophyceae	Hemiselmis sp.*		P5	5
Other flagellates	Cryptophyceae	Leucocryptos marina	P1 P2	P1 P4 P5 P6	24
out a "			· -, - -	P7	
Other flagellates	Cryptophyceae	Rhodomonas sp.		P5, P6, P7	16
Other flagellates	Cryptophyceae	Teleaulax sp.	P1, P2	P1, P4, P5, P6, P7	27
Other flagellates	Dictyochophyceae	Octonaria speculum (formerly Dictyocha speculum)		P1, P4, P5, P6,	461
Other flagellates	Eukaryota incertae sedis	Flagellates indet.	P2	Р7 Р1, Р4, Р5. Р6.	67
				P7	
Other flagellates	Eukaryota incertae sedis	Flagellates indet. 0–3 µm	P2	P4, P6, P7	1.2
Other flagellates	Eukaryota incertae sedis	Flagellates indet. 20–30 µm		P4	205
Other flagellates	Eukaryota incertae sedis	Flagellates indet. 3–7 µm	P1, P2, P4, PICE1	P1, P4, P5, P6, P7	10

Table A4 (continued)

Таха	Class	Species	August 2018	August 2019	Carbon conversion factor (pg C $cell^{-1}$)
Other flagellates	Eukaryota incertae sedis	Flagellates indet. 7–10 µm	P1, P2, P4, PICE1	P1, P4, P5, P6, P7	39
Other flagellates	Prasinophyceae	Pterosperma sp.		P1, P4	6005
Other flagellates	Prymnesiophyceae	Chrysochromulina sp.	P1, P2, P4, PICE1	P1, P4, P7	38
Other flagellates	Prymnesiophyceae	Coccolithales 7–10 µm	P2		111
Other flagellates	Prymnesiophyceae	Coccolithales indet.		P1, P5	111
Other flagellates	Pyramimonadophyceae	Phaeocystis pouchetii	P1, P2, P4, PICE1	P1, P4, P5, P6, P7	9.4
Other flagellates	Prymnesiophyceae	Pyramimonas sp.	P1, P2, PICE1	P1, P4, P5, P6, P7	82
Other flagellates	Raphidophyceae	Commation sp.*	P2, P4	P4, P6, P7	8.0
Other flagellates	Raphidophyceae	Heterosigma sp.	P2	P1, P4, P5, P6, P7	242
Other flagellates	Raphidophyceae	Olisthodiscus sp.*	P2, P4, PICE1	P1	192
Other flagellates	Telonemea	Telonema sp.	P1, P4	P1, P4, P5	26
Other flagellates	Telonemea	Telonema subtile		P1	7.9
Ciliates	Ciliophora	Ciliophora	P2, P4, PICE1		245
Ciliates	Ciliophora	Ciliophora 10–20 µm	P1, P2, P4, PICE1		245
Ciliates	Ciliophora	Ciliophora 20–30 µm	P1, P2	P1, P5	608
Ciliates	Ciliophora	Ciliophora 30–40 um	P1	P6	1072
Ciliates	Ciliophora	Ciliophora 50–60 um	11	P1 P4	1298
Ciliates	Gymnostomatea	Cyclotrichium sp *		P4	244
Ciliates	Gymnostomatea	Didinium sp	P2	P4	4171
Ciliates	Gymnostomatea	Mesodinium ruhrum	D2 D4 DICE1	D1 D4 D5 D7	3248
Ciliates	Oligohymenophorea	Scuticociliatia indet	12,14,11011	D5 D6	532
Ciliates	Oligohymenophorea	Uronema marinum*	D1 D4	10,10	2163
Ciliates	Oligotrichea	Acanthostomella norvegica	P1 P2	P4 P5 P6 P7	312
Ciliates	Oligotrichea	Labora strohila	11,12	P1	2711
Ciliates	Oligotrichea	Leegaardiella ovalis		P1	672
Ciliates	Oligotrichea	Leegaardiella sol	P1, P2, PICE1	P1. P4. P6. P7	456
Ciliates	Oligotrichea	Leprotintinnus pellucidus*	,,	P1	1660
Ciliates	Oligotrichea	Leprotintinnus sp	P1	••	1660
Ciliates	Oligotrichea	Lohmanniella oviformis	P1, P2, P4,	P1, P4, P5, P6,	204
Ciliataa	Olizatriahaa	Demofectually abtuarded	PICEI	P7	26 502
Ciliates	Oligotrichea	Purujuvella oblusaligula	D1 D2	PI	1660
Ciliates	Oligotrichea	Strembidium	P1, P2 D1 D2 D4	D1 D4 D5 D7	1010
Gillates	Oligotricliea	Si onbiatant	PI, P2, P4, PICE1	F1, F4, F3, F7	1010
Ciliates	Oligotrichea	Strombidium 20–30 µm		P1, P4	779
Ciliates	Oligotrichea	Strombidium conicum		P1	11,170
Ciliates	Oligotrichea	Strombidium constrictum		P4	6377
Ciliates	Oligotrichea	Tintinnidae <i>indet</i> .	P1		1660
Ciliates	Oligotrichea	Tintinnopsis sp.		P5, P6, P7	1660

environment will likely change the timing of blooms, successional patterns and protist community structure (Li et al., 2009; Nöthig et al., 2015; Flores et al., 2019). In a low-ice scenario, the oligotrophic, low biomass summer season dominated by small phytoplankton as well as heterotrophic and mixotrophic protists will be prolonged and commences earlier, following an advanced spring bloom governed by an earlier sea-ice retreat. Considering the lower accessibility of small-sized protists for Calanus copepods, the question is how a shift in phytoplankton phenology as well as a longer oligotrophic open water season dominated by small-sized protists will impact the lipid storage capacity of the dominant copepod grazers, especially because metabolic demands are highest during the warmer summer season (Morata and Søreide, 2015). A trend towards smaller-sized algae is predicted to lengthen the food web (Vernet et al., 2019) and negatively impact the energy transfer through the classic food web leading from larger diatom-eating Calanus copepods (Fig. 9; Søreide et al., 2008; Campbell et al., 2009; Cleary et al., 2017) to e.g., pelagic fish. Long-term food web changes have been shown for the Northeast Atlantic as a result of an increased success of the picocyanobacteria Synechococcus (Schmidt et al., 2020). Besides the potential food-web impacts, a longer oligotrophic open water season dominated by small phytoplankton and hetero- and mixotrophic protists will likely also extend the period of low vertical carbon flux (Fig. 9; Wassmann and Reigstad, 2011) with unknown consequences for benthic

food webs, including the northern prawn (*Pandalus borealis*) (Carroll and Carroll, 2003; Dabrowska et al., 2020). Interestingly, the deeper mixed open-water station P1 sustained higher protist standing stocks, particularly diatoms, in late summer compared with the meltwater-stratified, previously ice-covered stations and likely shows less pronounced seasonality. This is consistent with findings from the Fram Strait that showed a reduction in the strength of the biological carbon pump as a result of sea ice-derived meltwater stratification (von Appen et al., 2021).

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Author Contributions

DK and PA contributed equally to the writing of the paper. Study design involved PA, RG, DK, YB, OM, MAA, GB, MR, and BE. Nutrient data were analysed by MC and EJ. KB and LMG analysed chlorophyll *a* satellite imagery. Microscopic analysis of protist community samples was carried out by AMD, AT and JMW. OM, GB and LMO contributed flow cytometry data on pico- and nanoplankton and heterotrophic

nanoflagellates. Vertical flux data was contributed by YB and MR. Data analyses and figures were done by LG with the help of the other authors. ØL provided the temperature-salinity plot (Fig. A.1). All authors contributed to data interpretation and to the writing of the manuscript.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Nutrient (Chierici et al., 2021a,b) and chlorophyll *a* (Vader, 2022) data have been published in the Norwegian Marine Data Centre. Phytoplankton biodiversity data (Assmy et al., 2022a,b) have been published in the Norwegian Polar Data Centre and GBIF.

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Appendix A

Figs. A1-A7 and Tables A1-A4.

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