Sea-ice retreat may decrease carbon export and vertical microbial connectivity in the Eurasian Arctic basins

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Abstract

Arctic Ocean sea-ice cover is shrinking due to warming. Long-term sediment trap data show higher export efficiency of particulate organic carbon in regions with seasonal sea-ice compared to regions without sea-ice. To investigate this sea-ice enhanced export, we compared how different phytoplankton communities in seasonally ice-free and ice-covered regions of the Fram Strait affect carbon export and vertical dispersal of microbes. In situ collected aggregates, combined with microbial source tracking revealed that larger aggregates from sea-ice and under-ice diatom blooms were responsible for higher export efficiency and vertical microbial connectivity. During early summer, Phaeocystis aggregates dominated the ice-free regions and exported two-fold less carbon than diatom aggregates in ice-covered regions, and also less surface-born microbial clades to the deep-sea. This suggests that continuous ice-loss will further decrease pelagic-benthic coupling, impacting the quantity and quality of food input due to formation of slow-settling aggregates, with potential repercussions for Arctic deep-sea ecosystems.
Introduction

The Arctic Ocean is currently undergoing unprecedented changes due to ongoing climate warming. Ice coverage is declining by \(\sim 13\%\) per decade compared to the mean September extent for 1981-2010\(^1\), and climate models project that business-as-usual scenarios will result in seasonally ice-free conditions by 2050\(^2\). Increasing temperature, combined with declining sea-ice extent, ice thickness and multiyear ice\(^3\) are impacting the composition of primary producers in the Arctic Ocean\(^4\). For example, Atlantic phytoplankton species, such as *Phaeocystis* spp., are already seasonally reoccurring in the Fram Strait\(^5\), and were also recently observed during a phytoplankton bloom in the central Arctic Ocean\(^6\). It has been suggested that temperate phytoplankton species will become resident in the Eurasian basins of the Arctic Ocean if the intrusion of warming Atlantic waters continues\(^7\). As primary producers form the base of the food web, such shifts are likely to have drastic consequences, not just in the pelagic realm, but also for pelagic-benthic coupling and biogeochemical cycling in the Arctic Ocean\(^8\). However, the complexity of factors driving Arctic productivity regionally makes it difficult to generalize to the future carbon flux in the entire Arctic Ocean\(^9\).

Remote sensing of ocean color in the Arctic shelf seas suggests an increase in net primary production by \(57\%\) since 1998\(^10\), which likely enhances vertical carbon and nutrient fluxes\(^11\). Furthermore, the transformation from thick multiyear to thin first year ice is increasing light transmission through the ice\(^12\). Accordingly, field observations show an increased spatial and temporal extent of sea-ice algae and under-ice pelagic phytoplankton in the Arctic basins, for which ocean color based PP assessments are not available\(^6,13–15\). When the ice melts, such ice-algae and under-ice phytoplankton blooms that are mostly composed of diatoms can also deliver substantial pulses of carbon and nutrients to benthic ecosystems\(^8,16,17\). For example in 2012, the release of the fast-sinking ice-algae, *Melosira* spp., from melting sea-ice delivered up to \(9\) g of carbon per square meter of seafloor, which was more than \(85\%\) of the total carbon export that year\(^16\). Primary production models suggest that with the northward propagation of ice-edge blooms along leads and polynyas, the impact of ice-algae and under-ice phytoplankton blooms on productivity is likely to increase\(^18\), with ecological consequences for pelagic and benthic ecosystems\(^16,19\). However, it remains unclear what ecological effects an earlier and spatially substantial retreat of the ice-edge will have for carbon export on the Arctic shelf seas and margins. Key factors could be timing of stratification, e.g. by meltwater, the phytoplankton composition during the bloom, mismatches to grazers, and other effects on carbon export efficiency, including the microbial loop in the euphotic zone and remineralization of aggregates during sinking to deep-waters.

It is well established that sinking particles are an essential conduit of carbon and nutrients to heterotrophic organisms in the deep ocean\(^20\). The microbial loop retains most carbon and nutrients in the surface ocean, and can influence carbon export efficiency substantially, next to other factors such as particle sizes and the activities of grazers\(^21\). Recent studies have also revealed that particles are vectors that disperse microbes from the surface to the deep ocean some of which may carry specific heterotrophic functions in the remineralization of sinking matter\(^22–24\). Hence, sinking particles can play a key role in determining the structure and functioning of deep-sea microbial communities beyond supply of energy and nutrients\(^22,23\). To date, such so-called vertical microbial connectivity has mainly been demonstrated in temperate and tropical oceanic settings\(^22–26\). However, Rapp et al. (2018) found that sedimentation of sea-ice algae influences microbial community composition at the seafloor in the central Arctic. Considering the changes that the Arctic Ocean is currently undergoing, it is critical to understand how microbial connectivity and carbon export will be impacted by changing ice regimes and associated alterations in the composition and rates of primary producers.
Here, we assess the efficiency of particle export and vertical connectivity in the Fram Strait under seasonally ice-covered and ice-free conditions, which were defined based on temporal duration of sea-ice presence during the productive season. The Fram Strait represents the major deep-water gateway to the Arctic Ocean basins. Warm Atlantic water flows northwards via the West Spitzbergen Current (WSC) through the eastern part of Fram Strait, whereas cold Arctic water and sea-ice flows southwards into the Atlantic via the East Greenland Current (EGC) in its western part (Fig. 1a). The annual ice volume export through the western Fram Strait is currently increasing by 11% per decade during spring and summer due to ice thinning and increasing drift speed\(^{27}\). If Atlantic warming of the Arctic Ocean continues, it is projected that stronger ice-melt will occur in the western Eurasian Basin, eventually reducing ice export through Fram Strait\(^{28}\). Here we studied the effect of sea-ice distribution on phytoplankton community composition, carbon export efficiency and vertical microbial connectivity in the Fram Strait. We test the hypotheses 1) that settling aggregates formed in ice-covered regions sink faster than those formed in ice-free regions, which may result in lower export flux for ice-free regions. With regard to the effect on microbial community structure we postulated that 2) particle-associated microbes originate from surface waters and have little exchange with free-living microbes at depth, 3) that there is stronger vertical microbial connectivity in ice-covered compared to ice-free regions, and 4) that the stronger microbial connectivity leads to higher representation of surface-born microbes in the deep-sea including sediments. We found that large and fast-settling diatom aggregates in ice-covered regions resulted in higher carbon export efficiency near the ice and thus stronger vertical microbial connectivity compared to the seasonally ice-free region of the Fram Strait, which was dominated by smaller, slow-sinking *Phaeocystis* spp. aggregates. Our results suggest that with the ice-edge seasonally retreating from the Eurasian basins, carbon export efficiency and vertical connectivity may decline in large regions of the Arctic Ocean.

### Results

**Settling aggregates in ice-free and ice-covered regions**

We studied microbial communities associated with settling aggregates in contrasting sea-ice conditions between June 24\(^{th}\) and July 16\(^{th}\) 2016 at the Long-Term Ecological Research (LTER) observatory HAUSGARTEN in the Fram Strait (Expedition PS99.2 with RV POLARSTERN). We classified ice concentration >15% as ice-covered conditions and ice concentration <15% as ice-free conditions. According to this classification, the stations in the West Spitsbergen Current (‘HG’; seafloor depths ~2500 m) were seasonally ice-free stations, as the majority of the productive season in 2016 (March - July) they had no sea-ice (Fig. 1a; Table 1). On the other hand, in the East Greenland Current (‘EG’; seafloor depth: ~1000-2700 m) and the northern (‘N’; ~2500-2800 m) stations, sea-ice was present during most of the productive season, and these sites were thus defined as ice-covered stations (Fig. 1a; Table 1). Long-term sediment trap collections of particulate organic carbon (POC) fluxes at 200 m depth in the seasonally ice-free HG4 station have shown that the POC flux peaks early in the season due to ice-associated carbon export and again later in the season during the ice-free period due to POC export of pelagic production\(^{29}\). The POC export during the ice-associated flux peak (February to April) between 2001 and 2013 showed an inverse relationship between POC flux and distance to the ice-edge within 0-80 km, i.e. the zone most influenced by melt water (\(R^2=0.39, p<0.01\); Fig. 1b). This suggests that sea ice proximity can enhance POC fluxes. When the ice edge was beyond 80 km from the sediment traps, we observed no spatial effects between sea ice and POC export. To further test the hypotheses of sea-ice effects on carbon export efficiency, we assessed the exported organic matter and the vertical microbial connectivity at stations with contrasting ice-conditions in the Fram Strait during summer 2016.
First we used a Lagrangian Particle Tracking algorithm based on the observed aggregate sinking velocities (Table 1), to test whether our spatial classification scheme permitted the differentiation of particle origins. It showed less horizontal transport in the ice-covered regions compared to the ice-free regions (Supplementary Fig. 1), and differences in the origin of the particles. Particles in the ice-free region (‘HG’ stations) were primarily from the Atlantic waters south of the investigated region, and the majority of aggregates (82%) reaching the deep ocean (>1000 m) and seafloor originated from ice-free surface waters (Supplementary Fig. 1). In the ice-covered region (‘EG’ and ‘N’ stations) below 1000 m, a 60% of the aggregates originated from the ice-covered surface waters (Table 1).

All sampled stations were at the later stage of the phytoplankton bloom, based on the rate of consumed nitrate, silica and phosphate above the seasonal pycnocline (50 m depth, Supplementary Table 1). Microscopic analyses of water samples revealed that phyto- and protozooplankton communities in the chlorophyll \(a\) maximum (10 – 28 m depth) in the ice-free regions were dominated by \textit{Phaeocystis} spp., heterotrophic dinoflagellates and ciliates, while the ice-covered regions were dominated by planktonic diatoms and \textit{Phaeocystis} spp. (Supplementary Table 1). This was reflected in the composition of \textit{in situ} formed aggregates collected using a marine snow catcher (MSC) directly below the chlorophyll \(a\) maximum (60 m depth), where \textit{Phaeocystis} spp. dominated aggregates of the ice-free region and planktonic diatoms those of the ice-covered regions (Fig. 2). The aggregates from the ice-covered regions were two-fold larger (Wilcoxon Signed-Ranks Test; \(p<0.01\); Table 2) and sank two-fold faster than the aggregates collected in the ice-free regions (Wilcoxon Signed-Rank Test; \(p<0.01\); Table 2). Half of the aggregates collected in the ice-free region (13 out of 24) were smaller than 512 µm in diameter, while almost all (33 out of 36) collected aggregates in the ice-covered regions were larger than 512 µm (Fig. 2).

In addition, drifting sediment traps were equipped with a viscous gel to capture and preserve the size and structure of intact settling aggregates. The gel traps confirmed the MSC observations of \textit{Phaeocystis} spp. dominated aggregates in the ice-free regions and planktonic diatom-dominated aggregates in the ice-covered regions. The gel traps showed similar numbers of particles exported in the ice-free and ice-covered regions, but confirmed that the aggregates in the ice-covered regions had on average two-fold larger diameters than those collected in the ice-free regions (Wilcoxon Signed-Ranks Test; \(p<0.01\); Table 2). The larger diameters in the ice-covered regions translated into an order of magnitude larger average volume compared to aggregates of the ice-free regions (Table 2). Hence, in the period assessed in early summer, larger and faster-settling aggregates in the ice-covered regions caused a two-fold higher carbon export compared to the ice-free regions (Table 2). The carbon to nitrogen ratios (C:N, mol:mol) were 11 in the ice-free regions and 8 in the ice-covered regions, indicating export of fresher material from under the ice (Table 2). Furthermore, at the ice-stations macroscopic strands of the sea-ice diatom \textit{Melosira arctica} were observed by sea ice sampling (Fig. 2e), as well as during high-resolution imaging of the seafloor in the ice-covered stations (doi:10.1594/PANGAEA.873926).

**Free-living and particle-associated microbial communities**

For both regions, free-living (FL) and particle-associated (PA) microbial communities of four distinct water layers: surface (10-30 m), epipelagic (100 m), mesopelagic (1000 m) and bathypelagic (~50 m above the seafloor), were characterized using Illumina 16S rRNA gene sequencing. For the analyses of the microbial community composition we chose amplicon sequence variants (ASVs) as the highest possible taxonomic resolution the method provides\(^\text{30}\). The final dataset consisted of 3,709,676 sequences from 66 samples that were assigned to 6,253 ASVs associated with bacterial and archaeal
lineages (Supplementary Table 2). Rarefaction curves did not reach a plateau in any of the sampled communities, however, estimated asymptotic extrapolation to double amount of sequences showed only few additional ASVs (Supplementary Figure 2). Thus, our sequencing depth was satisfactory to represent most of the bacterial and archaeal diversity in all sampled microbial communities. The classes Alphaproteobacteria, Bacteroidia, and Gammaproteobacteria dominated the microbial communities in both FL and PA fractions, with no differences between ice-free and ice-covered regions (Supplementary Fig. 3). Each of these classes comprised more than 15% of the sequences and more than 10% of the ASVs in the entire dataset (Supplementary Fig. 3). In the deep ocean communities (>1000 m) there was an increasing abundance of the clades SAR202 (class Dehalococcoidia), SAR324 (Marine group B), SAR406 (Marinimicrobia), and the archaeal class Nitrososphaeria, each comprising 1-6% of the sequences and 3-6% of the ASVs in the entire dataset (Supplementary Fig. 3).

With increasing depth in the water column, microbial communities showed an increase in both richness (based on Chao1 richness estimator; Kruskal-Wallis test; Chi square=37.24, df=3, p<0.01), and beta-diversity (based on Shannon’s diversity index; Kruskal-Wallis test; Chi square=39.89, df=3, p<0.01; Supplementary Fig. 4). In the FL communities, this trend was mostly caused by significant differences between the communities of surface and epipelagic, and epipelagic to mesopelagic layers (post-hoc Wilcoxon Signed-Ranks Test; p-adjusted<0.01). In contrast, in the PA communities the richness did not show significant changes from surface to epipelagic, and from epipelagic to mesopelagic layers, but significantly increased between meso- and bathypelagic layers (post-hoc Wilcoxon Signed-Ranks Test; p-adjusted<0.01). The beta-diversity of the PA communities did not change in the upper 100 m of the water column (i.e., from surface to epipelagic waters), but significantly increased with depth below the epipelagic layer (post-hoc Wilcoxon Signed-Ranks Test; p-adjusted<0.01; Supplementary Fig. 4).

The composition of microbial communities showed clear separation between the FL and PA communities (PERMANOVA test; F1,64=10.14, R2=0.09, p<0.01). In both fractions the communities showed a specific clustering with the four distinct water layers (Fig. 3; PERMANOVA test; F3,64=10.11, R2=0.30, p<0.01). Compared to the surface water-originating FL and PA communities, the dissimilarity between deeper FL and PA communities increased with depth (Kruskal-Wallis test; Chi square=54.94, df=7, p<0.01). There was no difference in depth-related dissimilarity of FL communities between ice-free and ice-covered regions (Fig. 3; Wilcoxon Signed-Ranks Test; p>0.01). However, in the PA fraction the communities of the ice-free region had significantly higher dissimilarity along the water column, compared to the PA communities of the ice-covered region (Fig. 3; Wilcoxon Signed-Ranks Test; p<0.01).

**Vertical connectivity and shifts in particle-associated communities**

Many free-living microbes are adapted to colonize particles in the water column. Thus, the observed vertical dissimilarity pattern of the PA communities could be associated with the changing diversity of the FL communities. In order to test this hypothesis and to estimate the extent of colonization, we applied a microbial source tracking (MST) Bayesian algorithm ‘SourceTracker’. This MST approach assumes that ASVs diversity in various ‘source’ (i.e. FL) and corresponding ‘sink’ (i.e. PA) communities allows identification of statistically probable links between them (for detailed explanation see Methods section). The MST analysis showed a strong effect of the surface and epipelagic FL microbes on the composition of PA communities along the entire water column (Fig. 4). Within the surface and epipelagic layers, a particularly high proportion (84±5%) of the PA communities was associated with surface and epipelagic FL communities. In contrast, at meso-
bathypelagic depths the PA communities showed only a weak link to meso- and bathypelagic FL communities (ca. 2 and 8% of the communities, respectively), and a large fraction (72±5%) was not linked to any FL community. However, at meso- and bathypelagic depths, 27±6% of the PA communities in ice-covered and 11±2% of PA communities in ice-free regions were linked to surface and epipelagic FL communities (Fig. 4; Supplementary Table 3).

By statistical tests of comparative sequence enrichment, we identified the microbial taxonomic groups that became significantly more abundant on sinking particles as a function of depth. The ASVs within the PA communities were defined as enriched when they had a log$_2$ fold change of absolute value higher than 1 (i.e., double the amount of sequences) and an adjusted $p$ value lower than 0.1 (Fig. 5). This test looked at consecutive pelagic layers: surface-epipelagic, epipelagic-mesopelagic and mesopelagic-bathypelagic. In both ice-free and ice-covered regions PA communities became enriched with increasing depth in the classes Gammaproteobacteria (with 40 and 18 enriched ASVs, respectively), Planctomycetes (with 37 and 27 enriched ASVs, respectively), Bacteroidia (43 and 8 enriched ASVs, respectively), and the poorly characterized class OM190 (with 34 and 19 enriched ASVs, respectively). The enriched ASVs of these classes reached up to 5% of the sequences in the PA communities of the ice-covered regions and up to 10% of the sequences in the ice-free region. However, while the enriched ASVs of the classes Gammaproteobacteria and Bacteroidia were present also in the FL communities, the enriched ASVs of the classes Planctomycetes and OM190 were absent from the FL fraction (<0.5% of the sequences). Overall, we observed larger changes with depth in the PA communities of the ice-free region (where sinking speed was lower), resulting in more than double the amount of PA-enriched ASVs, in comparison to the ice-covered regions (348 and 158 ASVs, respectively; Supplementary Table 4).

**Transport of surface water-originating microbes to the bathypelagic: water column vs. seafloor**

Some of the vertically PA-enriched ASVs were also present in the FL communities along the water column (Fig. 6). In the bathypelagic, the PA-enriched ASVs comprised 17±2% of the sequences in the FL communities of the ice-covered region, and 47±4% of the sequences in the FL communities of the ice-free region. The most abundant family that consisted of such ASVs was the archaeal family Nitrosopumilacea, which comprised 3-4% and 6-19% of sequences in FL communities of the ice-covered and ice-free regions, respectively.

The seeding of the deep-sea sediment by microbes on sinking particles was tested using 7 deep-sea sediment samples (uppermost centimeter) collected at the same stations as the water column communities across the Fram Strait. This dataset consisted of 1,209,785 sequences that were assigned to 11,145 ASVs associated with bacterial and archaeal lineages (Supplementary Table S2; Supplementary Fig. S2). The sediment microbial communities were mainly affiliated to the classes Alphaproteobacteria, Gammaproteobacteria, and Nitrososphaeria (Supplementary Fig. S3).

The vertically PA-enriched ASVs were also identified in the deep-sea sediment communities of both ice-covered and ice-free regions. These shared ASVs between the PA and the sediment communities were associated mainly with the archaeal family Nitrosopumilaceae (17 ASVs) and the bacterial family Woeseiaceae (8 ASVs; class Gammaproteobacteria), each comprised ca. 2-4% of the sequences in the sediment communities (Fig. 6). Interestingly, in contrast to the PA-enriched ASVs of the family Nitrosopumilaceae that were also abundant in the FL communities of the bathypelagic, the shared ASVs of the family Woeseiaceae were absent from the FL communities (<0.3% of sequences in all FL communities). Overall, in the ice-free region, 31% of the PA-enriched ASVs were present in the sediment and comprised ca. 17% of the sequences in the sediment communities.
In contrast, in the ice-covered regions 39% of the PA-enriched ASVs were present in the sediment and comprised ca. 11% of the sequences in the sediment communities (Fig. 6).

Discussion

Throughout the world’s oceans, settling particles export organic matter and nutrients, as well as microbes and their enzymes, to the deep ocean\textsuperscript{20,22,23,26}. However, little is known about the influence of sea-ice on the dynamics on the composition and sinking velocities of settling particles and hence on export flux and efficiency in the Arctic. This is due to difficulties with the exchange of sediment traps and appropriate year-round upper ocean observations in ice-covered regions, limiting microbiological and biogeochemical deep-sea studies in the Arctic Ocean\textsuperscript{32,33}. From what is known, snow and ice cover affect productivity by light limitation, and thereby carbon export is relatively low under the ice\textsuperscript{29,34}. However, recently, the ice cover thinned substantially, so that it has lesser effect on light limitation and rafting of particles in the Central Arctic basins and in the Fram Strait\textsuperscript{27,28}. Furthermore, it was found that the ice-margin can have stimulating effects on primary production early in the season, by meltwater-induced stratification and through seeding with ice-associated primary producers\textsuperscript{18,35}. Spatially and temporarily this can lead to higher export efficiencies and stronger pelagobenthic coupling in regions with seasonal presence of the ice-margin, or covered partially by thinning sea-ice, including in the Fram Strait\textsuperscript{36}, in the regions north of Svalbard\textsuperscript{37}, as well as in the Central Arctic\textsuperscript{16}.

Here, we studied the role of sea ice on settling particle characteristics and vertical microbial connectivity, and postulate links to carbon export efficiency in the Fram Strait. Our long-term assessment of the role of ice-coverage on particle export during periods with sea ice near the HG4 station suggested an important function of sea ice-distance on export fluxes early in spring during the ice-influenced phytoplankton bloom period. This encouraged us to assess the underlying principles of this connection between ice-associated export and the fate of microbial communities attached to particles close to the ice margin. At the time of sampling in June-July 2016, the late stage of the ongoing phytoplankton bloom was dominated by diatoms in the ice-covered region while in the adjacent ice-free region it was dominated by \textit{Phaeocystis} spp. To test how ice-coverage impacts vertical connectivity and export of organic matter, we compared characteristics of sinking marine aggregates from ice-covered and ice-free regions of the Fram Strait during the productive period. In the ice-covered region we found larger diatom aggregates, with two-fold higher size-specific sinking velocities compared to the smaller \textit{Phaeocystis} spp. aggregates that dominated ice-free regions. This caused almost two-fold higher carbon export rates under the ice, compared to adjacent ice-free waters during the same period. The long-term record in the Fram Strait also shows that annual particle flux is lower during warm water phases with less ice\textsuperscript{38}, and characterized by a shift from diatom to coccolithophorid and \textit{Phaeocystis} spp. dominated phytoplankton during summertime at HG4 station\textsuperscript{39}. This is similar to observations north of Svalbard where ice-associated diatom production resulted in higher export than that observed for ice-free regions dominated by \textit{Phaeocystis} spp.\textsuperscript{37}. Taken together, this suggests that in the early Arctic summer, fast settling diatom aggregates drive export in ice-covered regions, whereas in warming, Atlantic-water influenced regions, the slower settling \textit{Phaeocystis} spp. aggregates dominate and will lead to more pelagic recycling. In this study, this also affected carbon to nitrogen ratios of the sinking matter, which were lower for the settling particles collected by the drifting traps in the ice-covered regions compared to the ice-free regions of the Fram Strait (Table 1). Hence, a potential future shift to Atlantification of the Eurasian Arctic basins\textsuperscript{40}, with larger areas of thermally stratified open waters, flagellate-dominated phytoplankton blooms, slower settling aggregates and stronger grazing pressure may lead to higher degradation and
transformation of organic matter during its journey through the water column, thus, resulting in lower amounts and less labile organic matter reaching the seafloor.

In this study we tested, for the first time in Arctic deep waters, the previously established hypothesis that vertical microbial connectivity is stronger in ecosystems dominated by fast-settling aggregates\textsuperscript{22,23}, due to the shorter transit time through the water column. In both ice-covered and ice-free regions of the Fram Strait, free-living (FL) pelagic microbial communities from different depths had greater dissimilarities to each other than the particle-associated (PA) communities from the same depths. This suggests a stratified water column with distinct microbial communities in the different water layers, as well as a vertical dispersal of microbial communities between surface ocean and deep-sea via sinking particles. In this context, settling particles are not only important for the export of organic matter to the deep ocean, but they also promote microbial heterotrophic activity and seeding\textsuperscript{22,23,33}, and thereby shape microbial biogeography and biogeochemical functioning in meso- and bathypelagic realms.

The surface water-originating microbial families that were significantly enriched on particles collected at depth, such as various members of the class \textit{Bacteroidia}, are associated with phytoplankton blooms in the region\textsuperscript{32,41}, and are known to be highly active organic matter degraders\textsuperscript{42}. Furthermore, it has recently been shown that there is a dominance of enzymatic activity phylogenetically linked to these taxonomic groups in the bathypelagic\textsuperscript{43} and that this enzymatic activity is predominantly linked to a particle-associated lifestyles\textsuperscript{44}. This indicates that active microbes originating from surface waters and associated with sinking particles continue to process organic matter while they sink to the deep ocean, and thus may remain key players in the biogeochemical cycling in the deep ocean. The cold water-adapted (i.e., psychrophilic) microbes of polar waters may potentially thrive in the deep Arctic ocean characterized by a relatively uniform temperature. Our results show that almost half of the bathypelagic FL communities consisted of vertically PA-enriched ASVs. This suggests that surface water-originating microbes may realize an ecological niche in bathypelagic waters. Evidence for this is provided by the archaeal family \textit{Nitrosopumilaceae}, which was the most abundant among taxonomic groups with PA-enriched ASVs in bathypelagic FL communities. Previous analyses showed that epipelagic and bathypelagic members of this family are phylogenetically closer to each other than those found in the intermediate mesopelagic waters\textsuperscript{45}. Based on our results, this pattern may be explained by a niche realization in the bathypelagic of \textit{Nitrosopumilaceae} family members exported from the surface ocean on sinking particles.

In both ice-free and adjacent ice-covered regions, the high similarity between surface PA and FL communities suggested that particles were colonized in surface waters, similar to other oceanic regions\textsuperscript{22,23,25,26}. Even in the bathypelagic, a substantial proportion of the PA community was still comprised of microbes recruited in the surface ocean. Notably, particles in the meso- and bathypelagic contained a high proportion of sequences that were not linked to the FL community at any depth, indicating a shift in population densities in the particle, e.g. by growth of otherwise rare types in the particles. Alternatively, the relatively long sinking time of days to weeks could have led to PA microbial communities at depth containing an imprint of surface water’s microbial communities that were no longer present or horizontally offset during our sampling. Overall, we conclude from our data that the observed vertical changes in the PA communities are substantially affected by sinking speed, causing different encounter rates and colonization in the surface ocean\textsuperscript{46}, and differences in time for ecological succession within the particles\textsuperscript{47}. The succession can result from transformations in the aggregate composition during aging and turnover\textsuperscript{48}, grazing by protozoans\textsuperscript{49}, viral infection\textsuperscript{50}, or changing environmental conditions, such as increasing hydrostatic
Common to all these processes is that slower sinking will enhance selection of some taxonomic groups and lead to the demise of others, potentially allowing rare taxa to become abundant at depth while those that were abundant at shallower depths become rarer. Furthermore, a large fraction of sinking particles remains suspended in the bathypelagic. In this way, settling aggregates should be viewed as constantly changing microcosms that have some exchange with their surroundings in the deep ocean, but where particle sinking speed is an important driver of succession. In Arctic waters, it seems that fast aggregate sinking speed is strongly related to ecological impact from sea-ice cover.

Since the seafloor is the final destination for those particles that make the journey through the water column, we tested whether the vertical microbial connectivity extends to deep-sea sediment. We found that ca. 10-20% of the sequences in the deep-sea sediment were related to PA-enriched microbes originating from the surface waters and deposited via sinking particles, in both ice-covered and ice-free regions. These results are comparable to observations in the Central Arctic Ocean after deposition of ice algae on the deep seafloor, and are higher than the global average of <10%. Interestingly, the family Woeseiaceae (class Gammaproteobacteria) showed the strongest benthic-pelagic connectivity via sinking particles, indicating its export from surface to bathypelagic waters via sinking particles. Recent genomic characterization of this largely unknown taxonomic group (which was conducted using sediment samples collected in the Fram Strait) suggests their involvement in the cycling of detrital proteins in marine benthic environments. Using targeted fluorescence microscopy of the total pelagic microbial communities (based on samples collected in parallel to this study), Hoffmann et al. (2020) also showed, that cells of this taxonomic group were present throughout the entire water column, comprising <1% of the community. We found that pelagic members of the Woeseiaceae were associated with sinking particles, but not free-living, suggesting that this important benthic heterotroph is one of the few types of bacteria that cover all water depths by a particle-associated life style.

In conclusion, our study supports the notion that sea-ice retreat can have an important ecological impact on carbon flux characteristics, and on long term potentially affect the deep-ocean microbial diversity. Fast settling ice-associated diatom aggregates drive higher export efficiency and cause stronger pelagic-benthic coupling including the transport of functionally important microbial groups, whereas slow settling Phaeocystis spp. aggregates associated with seasonally ice-free regions may lead to more pelagic recycling and less connectivity. These changes may substantially alter deep water and seafloor communities in the Arctic.

**Methods**

**Water sampling and metadata collection**

The sampling was performed during RV Polarstern expedition PS99.2 to the LTER observatory HAUSGARTEN in the Fram Strait (June 24th – July 16th 2016). Water samples were collected using 12 L Niskin bottles mounted on a CTD rosette (Sea-Bird Electronics Inc. SBE 911 plus probe) equipped with double temperature and conductivity sensors, a pressure sensor, chlorophyll a fluorometer, and transmissometer. At all stations water samples were collected from surface at 10-30 m, 100 m, 1000 m and ~50 m above the seafloor (Supplementary Table S2). For assessing archaeal and bacterial community composition, 4 L in epipelagic (<100 m) and 8-12 L in meso- and bathypelagic waters were filtered with a peristaltic pump (Masterflex; Cole Parmer) through successive membrane filters of 5 μm (Whatman Nucleopore, 47 mm polycarbonate), and 0.22 μm (Millipore Sterivex™ filters). In addition, deep-sea sediment cores were collected with a TV-guided
multicorer, and subsamples of the uppermost centimeter of the cores were collected with syringes (Supplementary Table S2). All samples were stored at -20 °C until DNA isolation.

Hydrographic data of the seawater including temperature and salinity (doi:10.1594/PANGAEA.871952), as well as the inorganic nutrient concentrations (doi:10.1594/PANGAEA.906132) were retrieved from PANGAEA. The map in Fig. 1a was generated using ArcMap (v10.5) with Esri world countries dataset (www.esri.com) in a WGS 1984 Arctic Polar Stereographic map projection. The mean monthly sea-ice concentrations for Fig. 1a were retrieved from http://data.seaiceportal.de 56, and sea surface temperature was obtained from NOAA NCEP real-time analysis (http://polar.ncep.noaa.gov/sst/rtg_high_res/).

Long-term sediment trap deployment and sea-ice distance estimation

The long-term moored KIEL sediment trap (sampling area 0.5 m² and 20 collection cups) was deployed and recovered yearly from 2001 to 2013 at the central LTER observatory HAUSGARTEN station (HG4 - 79.01 °N, 4.20 °E; Fig. 1). The deployment depths of the sediment trap was ~ 200 m. Sampling cups were filled with filtered seawater adjusted to a salinity of 40 and fixed with 0.14% final solution of HgCl₂. The opening time of the sampling cups varied between 7 and 59 days, depending on the season with short opening time during the polar day and long opening time during the polar night. Swimmers were removed after recovery and triplicate subsamples were measured for particulate organic carbon (POC) by filtering the material onto pre-combusted Whatman GF/F filters, soaking them in 0.1N HCl, and drying at 60°C before analyzing with a CHN elemental analyzer14,36. The data was retrieved from PANGAEA (doi: 10.1594/PANGAEA.855473).

To evaluate the impact of the sea ice concentration on the POC flux during periods when peak POC flux was associated with sea ice (February to April), we used POC flux collected by the long-term sediment trap of the central station HG4. The daily distance between the sea-ice edge and HG4 was estimated using daily sea-ice concentration satellite images from NSIDC/NOAA (http://nsidc.org/data/5051). The images were generated using the NASA Team algorithm 57 and mapped to a 25 x 25 km grid. This satellite data set was derived from brightness and temperature data generated from Scanning Multichannel Microwave Radiometer and Sensor Microwave Imager and Sounder equipped on the Nimbus-7 satellite and the Defence Meteorological Satellite Program, respectively. The distance to the ice-edge was defined at the position with 15% sea-ice concentration. The ice-edge nearest the HG4 position was used to calculate the daily ice-distance and averaged for each opening time of the collection cups (~14 days) on the long-term moored sediment traps.

Microscopic analysis of phyto- and protozooplankton

The plankton community composition at the chlorophyll a maximum was identified and the phytoplankton abundance was counted using light microscopy. Seawater samples were preserved in hexamethylenetetramine-buffered formalin (final concentration 0.5-1%) and stored in brown glass bottles. For microscopic analyses an aliquot of 50 mL was transferred to Utermöhl settling chambers where the cells were allowed to settle for 48 hours. At least 500 cells of the dominant phytoplankton species or groups were counted with an inverted microscope at three different magnifications using phase contrast according to Utermöhl (1958) and Edler (1979).
On-board characterization of marine aggregates and sinking velocity measurements

Using a marine snow catcher (MSC, OSIL, United Kingdom) we sampled intact aggregates from 60 m at ice-free and ice-covered regions, and measured their size, composition, and sinking velocities. The aggregates were individually transferred to a vertical flow chamber filled with Whatman GF/F filtered seawater collected from the same MSC and kept at in situ temperature. The x-, y-, and z-axes of each aggregate were measured in the vertical flow system using a horizontal dissection microscope and an ocular with a scale. The aggregate volume was thereafter calculated assuming an ellipsoidal shape and the equivalent spherical diameter (ESD) was calculated from the aggregate’s volume. The sinking velocity was measured by increasing the upward flow in the flow-chamber until the aggregate was floating one diameter above the net. The sinking velocity was thereafter calculated by determining the volumetric flow rate three times, and dividing the average of these measurements by the area of the flow chamber. The composition of the aggregates was determined with an inverted light microscope using Utermöhl chambers (Fig. 2).

Aggregate and carbon export to 100 m

Aggregate and carbon export to 100 m depth was measured using the free-drifting surface-tethered sediment traps in the ice-free and ice-covered regions. The drifting traps consisted of a drifting array attached to a surface buoy equipped with a GPS satellite transmitter, two surface floats and 12 small buoyancy balls that served as wave breakers to reduce hydrodynamic mixing effects on the sediment traps. The 100 m collection depth was equipped with four gimbal-mounted cylinders, each 1 m tall and 10.4 cm in inner diameter. Three of the cylinders collected samples for biogeochemical measurements and the last collection cylinder contained 200 ml of a viscous gel, which intercepted and preserved settling particles without destroying their original sizes and structures. Upon recovery, the material collected for biogeochemical fluxes was fixed with HgCl₂ and stored at 4°C until further analyses in the home laboratory. The particles collected in the gels were photographed using a stereo microscope equipped with a 3.1 megapixel digital camera and a 105 mm macro lens, resulting in a pixel size of 12 μm. The image analyses were performed with a routine written in MATLAB (The MathWorks) using the image analysis toolbox. Each image was converted into grey scale and the background was removed by applying a threshold value. The calibrated pixel area (mm²) in each projected particle was converted into the equivalent spherical diameter (ESD).

DNA isolation and 16S rRNA amplicon sequencing

Genomic bacterial and archaeal DNA was isolated from size-fractionated filtration through 5 µm and 0.22 µm filters membranes to analyze the particle-associated (PA, >5 µm) and the free-living (FL, >0.22 µm and <5 µm) community. The isolations were conducted by a combined chemical and mechanical procedure using the PowerWater DNA Isolation Kit and PowerSoil DNA Isolation Kit for the sediment samples (MO BIO Laboratories, Inc., Carlsbad, CA, USA). Prior to DNA isolation the Sterivex™ cartridges of the 0.22 µm membranes were opened in order to place the filters in the kit-supplied bead beating tubes. The isolation was continued according to the manufacturer’s instructions, and DNA was stored at -20 °C. Library preparation was performed according to the standard instructions of the 16S Metagenomic Sequencing Library Preparation protocol (Illumina, Inc., San Diego, CA, USA). The hyper variable V4–V5 region of the 16S rRNA gene was amplified using bacterial primers 515F-Y (5’-GTGYCAGCMGCCGCGGTAA-3’) and 926R (5’-CCGYCAATTYMTTTRAGTTT-3’). Sequences were obtained on the Illumina MiSeq platform in a 2 × 300 bp paired-end run (CeBiTec Bielefeld, Germany), following the standard instructions of the 16S Metagenomic Sequencing Library Preparation protocol (Illumina, Inc., San Diego, CA, USA). Raw paired-end, primer-trimmed reads were deposited in the European Nucleotide Archive (ENA).
under accession number PRJEB30254. The data were archived using the brokerage service of the German Federation for Biological Data (GFBio).  

**Bioinformatics and statistical analyses**

The raw paired-end reads were primer-trimmed using cutadapt. Further analyses were conducted using R (v3.6.3; [http://www.Rproject.org/](http://www.Rproject.org/)) in RStudio (v1.2.5033; [http://www.rstudio.com/](http://www.rstudio.com/)). The trimmed libraries were processed using DADA2 (v1.14.1) following the suggested tutorial ([https://benjjneb.github.io/dada2/tutorial.html](https://benjjneb.github.io/dada2/tutorial.html)). Briefly, chimeras and singletons were filtered out. The produced amplicon sequence variants (ASVs) were taxonomically classified against the Silva reference database (release 138). The ASVs that were taxonomically unclassified on domain level, or not assigned to bacterial or archaeal lineages, were excluded from further analysis. Furthermore, all ASVs which were taxonomically assigned to mitochondria and chloroplast were removed from the dataset.

Sample data matrices were managed using the R package ‘phyloseq’ (v1.28.0) and plots were generated using R package ‘ggplot2’ (v3.3.0). The sample rarefaction analyses were conducted using R package ‘iNEXT’ (v2.0.20). Prior to beta-diversity analyses, a prevalence threshold (i.e., in how many samples did an ASV appear at least once) of 4% was applied on the ASV abundance table. Principal component analysis (PCA) and dissimilarity comparisons between FL and PA communities were conducted on a stabilized ASV abundance table based on the geometric mean. The fold-change in abundance of each ASV between the water layers was calculated using the R package ‘DEseq2’ (v1.24.0). The method applies a generalized exact binomial test on variance stabilized ASV abundance.

Based on the assumption that the particle-associated microbial communities (i.e., ‘sink’ communities) are the result of various events of colonization of marine aggregates by free-living microbes (i.e., ‘source’ communities); a Bayesian microbial source tracking algorithm ‘SourceTracker’ (v1.0) was applied on the ASV abundance table. The algorithm performance was validated using a ‘leave-one-out’ approach, in which each ‘source’ (i.e., FL) community was hidden, in turn, from the training dataset, and its origin was predicted based on the rest of the source samples in the dataset. The entire analysis was conducted under default conditions: burn-in period - 100, restarts - 10, dirichlet hyperparameters (α, β) - 0.001. All samples were randomly sub-sampled to 5,000 sequences. Scripts for the molecular data processing and statistical analyses can be accessed at [https://github.com/edfadeev/Vertical_connectivity_Arctic_Ocean](https://github.com/edfadeev/Vertical_connectivity_Arctic_Ocean).

**Modeled aggregates sinking trajectories**

A Lagrangian particle tracking algorithm was used to back-track particles from the sampling depth to the surface. A detailed description of the model can be found [Wekerle et al, 2018](https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7725388/). Briefly, the backward particle computation is done by reversing the flow field, i.e. particles are treated as if they were rising from the sampling depth to the surface with a negative sinking speed, being horizontally displaced with the reversed horizontal velocity. Particles were advected with daily averaged 3D model velocities from the ocean general circulation model FESOM (an ocean-sea ice model based on unstructured meshes). The particle sinking speed was computed by adding a constant sinking speed to the modelled vertical velocity. In this study, we used a FESOM configuration optimized for the Fram Strait, applying a mesh resolution of 1 km. The performance of the model was validated for the sampling time period by oceanographic observations (Supplementary Fig. 5).
The backward trajectory calculation was performed for all three sampled regions (ice-free ‘HG’ stations, and ice-covered ‘EG’ and ‘N’ stations), using on-board measurements of aggregate sinking velocities (Table 1). Trajectories were released around 300 m above the seafloor once per day during the year March - July 2016, however we restricted the analysis to particles that reached the ocean surface between March and July 2016. A time step of 30 min was used for the trajectory calculation, and bi-hourly positions and corresponding temperature and salinity values were stored. To quantify the vertical distribution of particles, their positions were binned into a grid with bin sizes of 25 m depth x 0.05° Longitude/Latitude and then divided by the total number of particles to determine the fraction of particles originating from each grid box (Table 2). The daily concentrations of sea-ice were retrieved from Centre d’Exploitation et de Recherche SATellitaire (CERSAT; http://cersat.ifremer.fr/).

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Authors contribution

EF analyzed the data and wrote the manuscript with guidance from MHI, AB, CB and IS. MHI conducted the aggregate characterization on board. SR conducted the long-term sediment traps analysis. AR and AMW contributed to sampling, and data analysis of an early version of the manuscript. EMN conducted the microscopic analysis of phyto- and protozooplankton and was responsible for POC flux analysis of long-term moorings. CW modeled the aggregates sinking trajectories. All authors contributed to the final version of the manuscript.

Competing interests

The authors declare no competing interests.

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Figure 1. Overview of sampling area in the Fram Strait. (a) Monthly average of sea-ice cover and sea surface temperature during July 2016. The arrows represent general directions of the WSC (in red) and the EGC (in blue). The yellow edge of the HG4 station indicates the position of a long-term sediment trap deployment. (b) Particulate Organic Carbon (POC) flux collected at 200 m by a long-term moored sediment trap between 2001 and 2013 during the spring period between February and April, plotted as a function of the distance to the ice-edge. The regression shows that there was a significantly negative relationship between the distance to the ice-edge and the magnitude of POC flux ($R^2=0.39$, $p <0.01$).
Figure 2. Exemplary light microscopy images of marine aggregates from MSC deployments in Fram Strait. (a,b) Aggregates dominated by diatoms from ice-covered region - ‘EG’, where (a) is mainly diatoms and (b) is diatoms together with Phaeocystis spp. (c,d) Aggregates dominated by flagellates in the ice-free region - ‘HG’, where (c) is a copepod fecal pellet composed of flagellates and (d) is a marine snow aggregate formed from Phaeocystis spp. colonies. (e) Chains of Melosira spp., diatoms growing under the sea-ice. (f) Calanus spp. (copepod) fecal pellets collected at the ice-covered region - ‘N’ and formed from mainly Phaeocystis spp. colonies with a few diatoms. All scale bars indicate 200 µm long. White arrows point towards diatom chains and black arrows point towards colonies of flagellates.
Figure 3. Free-living and particle-associated community patterns throughout the water column of the Fram Strait. (a) Principal component analysis (PCA) of microbial communities. Ellipses encompass clustering of each of the fractions by water layer (S-surface, E-epipelagic, M-mesopelagic, B-bathypelagic), with normal confidence of 0.95. The percentages on both axes represent the explained variance of the axis. (b) Euclidean distances between microbial communities in each fraction along the entire water column. The colors represent different geographic origins: ice-covered (blue) and ice-free (red) regions. (NS) – not significant (Wilcoxon signed-rank test; \(p\)-adjusted>0.01). (*** – significant (Wilcoxon signed-rank test; \(p\)-adjusted<0.001).
Figure 4. Proportion estimates of the source water masses for microbial communities in different water layers, using 'SourceTracker'. The source estimates for each free-living (FL) community was conducted estimated using leave-one-out approach (i.e., based on all other FL communities; see methods), and the sources of the particle-associated (PA) communities were estimated based on the FL communities. The ice-covered stations are marked with an asterisk.
Figure 5. Differences in particle-associated (PA) community composition between the distinct water layers, in ice-covered and ice-free regions. Enriched taxonomic families between each two consecutive depths (surface-epipelagic, epipelagic-mesopelagic and mesopelagic-bathypelagic), ordered according to labels between the panels. The y-axis represents the mean log$_2$ fold change for microbial families with more than 3 ASVs with log$_2$ fold change absolute value higher than 1 (standard error is smaller than the point). Positive value represent enrichment in deeper water layers and negative value represents enrichment in shallower water layer. The numbers near the symbols represent the number of ASVs enriched in the depth. The x-axis is ordered according to the different taxonomic classes, represented by the color code.
Figure 6. Overview of the sequence proportion of enriched ASVs in the free-living, particle-associated, and uppermost centimeter of deep-sea sediment microbial communities. The classes represented by colors according to the legend, all classes with sequence proportion below 2% were classified as “Other classes”. The ice-covered stations are marked with an asterisk.
Table 1: Sinking aggregate trajectories characteristics of particles reaching the surface ocean between March 1\textsuperscript{st} 2016 and July 31\textsuperscript{st} 2016 in different regions of Fram Strait. The sinking trajectories were modeled using the measured in situ aggregates sinking velocities in each region, as well as using hypothetical low (20 m d\textsuperscript{-1}) and high (60 m d\textsuperscript{-1}) velocities. The values after ± represent standard deviation.

|                                      | EG       | N       | HG       |
|--------------------------------------|----------|---------|----------|
| Station coordinates                  | 78.81° N / 2.729° W | 79.74° N / 4.185° E | 79.06° N / 4.51° E |
| Starting depth (m) of trajectory calculation | 2350 | 2350 | 1950 |
| Number of days during 2016 with ice-coverage >15% | 197 | 107 | 78 |
| Number of days during March - July 2016 with ice-coverage >15% | 92 | 38 | 29 |
| Measured sinking velocity (m d\textsuperscript{-1}) | 52 | 52 | 29 |
| Median catchment radius (km)          | 78 ± 45 | 74 ± 53 | 118 ± 97 |
| Aggregates originated from ice-covered waters (% of total)* | 72 | 44 | 16 |
| Median sinking trajectory length (km)  | 194 ± 62 | 181 ± 64 | 392 ± 111 |
| Low sinking velocity (m d\textsuperscript{-1}) | 20 | 20 | 20 |
| Median catchment radius (km)          | 142 ± 79 | 132 ± 115 | 124 ± 124 |
| Aggregates originated from ice-covered waters (% of total)* | 50 | 36 | 15 |
| Median sinking trajectory length (km)  | 536 ± 124 | 527 ± 159 | 572 ± 175 |
| High sinking velocity (m d\textsuperscript{-1}) | 60 | 60 | 60 |
| Median catchment radius (km)          | 70 ± 46 | 70 ± 53 | 94 ± 76 |
| Aggregates originated in ice-covered waters (% of total)* | 74 | 41 | 6 |
| Median sinking trajectory length (km)  | 179 ± 55 | 161 ± 63 | 233 ± 71 |
Table 2: Vertical fluxes and marine aggregates characteristics in the epipelagic waters (0-100 m) of ice-covered and ice-free regions. ESD: equivalent spherical diameter. The values after ± represent standard errors.

|                                | ice-covered region | ice-free region |
|--------------------------------|--------------------|-----------------|
| **Drifiting traps fluxes**     |                    |                 |
| Particulate organic carbon (POC; mg m⁻² d⁻¹) | 60                 | 32              |
| Particulate organic nitrogen (PON; mg m⁻² d⁻¹) | 9                  | 3               |
| POC to PON ratio (mol:mol)     | 8                  | 12              |
| **Gel trap fluxes**            |                    |                 |
| Number of particles            | 2003               | 1399            |
| Total particle number flux (# m⁻² d⁻¹) | 36 x 10⁴           | 37 x 10⁴        |
| Total particle volume flux (mm³ m⁻² d⁻¹) | 11 x 10³           | 1.5 x 10³       |
| Average ESD of particles (mm)  | 0.2 ± 0.2 (range: 0.1 – 2) | 0.1 ± 0.1 (range: 0.1 – 1) |
| Average volume of particles (mm³) | 0.03 ± 0.2         | 0.003 ± 0.02    |
| **Vertical flow chamber**      |                    |                 |
| Number of collected particles  | 36                 | 24              |
| Average ESD of particles (mm)  | 0.9 ± 0.1 (range: 0.3 – 2.4) | 0.6 ± 0.1 (range: 0.3 – 1.3) |
| Average sinking velocity of particles (m d⁻¹) | 52.8 ± 0.9         | 29.5 ± 0.7      |
| Dominant phytoplankton in particles | Diatoms            | Flagellates (Phaeocystis spp.) |