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Does Arctic warming reduce preservation of organic matter in Barents Sea sediments?

Johan.C. Faust^{1*}, Mark A. Stevenson², Geoffrey D. Abbott², Jochen Knies^{3,4}, Allyson Tessin⁵, Isobel Mannion¹, Ailbe Ford¹, Robert Hilton⁶, Jeffrey Peakall¹ and Christian März¹

¹*School of Earth and Environment, University of Leeds, UK*

²*School of Natural and Environmental Sciences, Newcastle University, Newcastle upon Tyne, UK*

³*Geological Survey of Norway, Trondheim, Norway*

⁴*CAGE – Centre for Arctic Gas Hydrate, Environment and Climate, Department of Geology, UiT the Arctic University of Norway, Tromsø, Norway*

⁵*Department of Geology, Kent State University, Kent, OH, USA*

⁶*Department of Geography, Durham University, UK*

Keywords: Barents Sea, Geochemical sediment composition, Organic carbon bound to reactive iron, Carbon cycle, Arctic Ocean, Marine Surface Sediments

Summary

Over the last few decades, the Barents Sea experienced substantial warming, an expansion of relatively warm Atlantic water and a reduction in sea ice cover. This environmental change forces the entire Barents Sea ecosystem to adapt and restructure and therefore changes in pelagic-benthic coupling, organic matter sedimentation and long-term carbon sequestration are expected. Here we combine new and existing organic and inorganic geochemical surface sediment data from the western Barents Sea and show a clear link between the modern ecosystem structure, sea ice cover and the organic carbon and CaCO₃ contents in Barents Sea surface sediments. Furthermore, we discuss the sources of total and reactive iron phases and evaluate the spatial distribution of organic carbon bound to reactive iron. Consistent with a recent global estimate we find that on average 21.0±8.3 per cent of the total organic carbon is associated to reactive iron (fOC-Fe_R) in Barents Sea surface sediments. The spatial distribution of fOC-Fe_R, however, seems to be unrelated to sea ice cover, Atlantic water inflow or proximity to land. Future Arctic warming might, therefore, neither increase nor decrease the burial rates of iron-associated organic carbon. However, our results also imply that ongoing sea ice reduction and the

*Author for correspondence (J.Faust@Leeds.ac.uk).

†Present address: School of Earth and Environment (SEE), The University of Leeds, LS2 9JT Leeds, United Kingdom

19 associated alteration of vertical carbon fluxes might cause accompanied shifts in the Barents Sea surface
20 sedimentary organic carbon content, which might result in overall reduced carbon sequestration in the future.

21 Introduction

22 One of the most apparent signs of current global climate change is Arctic sea ice loss. Over the past four
23 decades, summer sea ice extent has drastically decreased by over 30% [1, 2] and the ongoing transformation of
24 the Arctic Ocean from an “icy land” into an open ocean forces the entire Arctic ecosystem to adapt and
25 restructure [3]. As the Arctic Barents Sea shelf area (Fig. 1) is a transition zone between the temperate North
26 Atlantic and the cold Arctic Ocean, it is climatically divided into two distinct regions. The northern area
27 experiences a cold and harsh Arctic climate and sustains an ice-associated ecosystem, while the southern part
28 has an Atlantic climate with a rich open water ecosystem and lucrative fisheries [4, 5]. During recent decades,
29 enhanced inflow of Atlantic water and atmospheric heat transport have dramatically warmed the Arctic, and in
30 particular the Barents Sea [6]. Sea ice loss and “Atlantification” of the northern Barents Sea are the consequences
31 [6-8]. Higher water temperatures and sea ice reduction modifies the Arctic marine ecosystem structure and,
32 therefore, changes the Arctic carbon cycle, i.e., atmospheric CO₂ uptake, pelagic-benthic coupling, organic
33 matter sedimentation and long term sequestration [3, 9-13]. An increase in the annual net primary production
34 in the Arctic and the Barents Sea has already been observed since the late 1990s and might rise in the future, due
35 to further summer sea ice reduction and longer phytoplankton growing seasons [10, 14]. However, these
36 environmental changes are complex and so far only a few studies link ongoing changes in the Arctic Ocean to
37 organic carbon burial, sedimentary biogeochemical cycles and the marine ecosystems [11, 15, 16]. Thus, there is
38 substantial uncertainty regarding current and future productivity and carbon burial estimates in the Arctic and
39 the Barents Sea.

40 The sequestration of organic matter in marine sediments is a fundamental mechanism for the removal of
41 carbon from the atmosphere and its storage over geological time periods [17]. Examining climatically induced
42 biogeochemical changes in Arctic marine sediments, is therefore, important for a better understanding of the
43 global carbon cycle. However, the processes that control organic carbon preservation in marine sediments,
44 including sedimentation rate [18, 19], presence and absence of oxygen [20-22], selective preservation of
45 biochemically unreactive compounds [23, 24], and protection of organic matter through interactions with a
46 mineral matrix [25-27] are complex and still not fully understood. A possible connection between iron and
47 organic carbon in marine sediments was already identified in 1970 [28], but only recently has the importance of
48 this relationship for organic matter preservation in marine sediments been recognised [29]. Due to their high
49 sorption capacity, iron oxides, in particular freshly precipitated and poorly crystalline iron (oxyhydr)oxides,

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4 50 like ferrihydrite, have a strong influence on organic carbon stabilization. During burial at the seafloor, organic
5 51 carbon adsorbed to these oxides is preserved against microbial degradation and can therefore bypass the
6 shallower oxic degradation regimes into, and possibly beyond, the zone of dissimilatory metal oxide reduction
7 52 [29]. Therefore, reactive iron phases may serve as an efficient shuttle to enhance organic carbon burial and
8 53 preservation in marine sediments. Lalonde et al. [29] investigated surface sediment samples from several marine
9 54 environments including the Southern Ocean, Mexican and Indian Margins, St. Lawrence estuary and gulf, and
10 55 the Black Sea. They proposed that on average 21.5% of the total organic carbon in marine surface sediments is
11 56 associated with reactive iron globally. Hence, Lalonde et al. [29] stated that “reactive iron phases serve as an
12 57 extremely efficient ‘rusty sink’ for organic carbon and are a key factor in the long-term storage of organic carbon
13 58 and the global cycles of carbon, oxygen and sulphur”. However, since this pioneering publication only a few
14 59 studies investigated the role of reactive iron on the preservation of organic carbon in natural marine sediments
15 60 [30-37]. And except for one study from the East Siberian Arctic Shelf [30], the type and amount of organic carbon
16 61 bound to iron oxides has not been examined in Arctic marine sediments. Moreover, there is still a general lack
17 62 of knowledge about reactive iron sources in relation to total iron content, the general sediment composition,
18 63 and the environmental setting. Making these mechanistic links is, however, necessary to evaluate the role of
19 64 organic carbon bound to iron phases and its role in the global carbon cycle, especially in a fast-changing
20 65 environment such as the Arctic Ocean.
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34 67 To better understand how ongoing “Atlantification” of the Barents Sea will change the organic and
35 68 inorganic sediment composition in the future, we combined new and existing surface sediment (0-1 cm) data of
36 69 organic carbon, total iron, calcium carbonate and grain size distribution of the seasonally ice-covered north and
37 70 permanently ice-free south western Barents Sea. Furthermore, to better constrain the controls on, and efficiency
38 71 of, carbon burial in the Arctic shelf seas we analysed the fraction of organic carbon bound to dithionite-
39 72 extractable iron phases (fOC-Fe_R).
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45 73 *Study Area*

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47 74 The Barents Sea is located between 70-81°N off the northern Norwegian coast and is bordered by the shelf
48 75 edge towards the Norwegian Sea in the west, the Norwegian archipelago Svalbard in the northwest and the
49 76 islands of Franz Josef Land and Novaya Zemlya (Russia) in the northeast and east. It is the largest pan-Arctic
50 77 shelf sea covering an area of 1.6 million square km with an average water depth of 230 m [38]. There are several
51 78 extensive overviews and reviews about the modern climate setting and ecosystem of the Barents Sea and we
52 79 refer to these references for a detailed description of the physical and ecological conditions [4, 10, 39-42]. In brief,
53 80 the present ecological setting as in all Arctic seas is characterized by very pronounced seasonal fluctuations in
54 81 insolation and, hence, primary production. However, despite the relatively short duration of the growing season
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4 82 in the Arctic, the Barents Sea is a high productivity shelf area where 40% of the total primary production of the
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6 83 Arctic Ocean takes place [43]. Water column primary productivity is generally inversely related to sea ice cover,
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8 84 i.e., lower rates occur in the north-east (30-70 g C m⁻²y⁻¹) and higher and less variable rates in the Atlantic water-
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10 85 influenced south-west (100-150 g C m⁻²y⁻¹) [39, 44]. The general oceanic circulation pattern of the western
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12 86 Barents Sea is dominated by the relatively warm northward flowing North Atlantic Current (temperature 2-
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14 87 8°C, salinity >35‰) which enters the Barents Sea from the southwest and the southward flowing cold Arctic
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16 88 currents (Spitsbergen and Persey; temperature <0°C, salinity <35‰) entering the Barents Sea from the northeast.
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18 89 The relatively sharp boundary between these water masses forms the oceanographic Polar Front (Fig. 1) [45]
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20 90 which is mainly determined by the bathymetry and is, therefore, relatively stable from year to year [46]. The
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22 91 northern Barents Sea is seasonally ice covered with maximum and minimum ice coverage in March-April and
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24 92 August-September, respectively. The heat content of the Atlantic water keeps the southern Barents Sea
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26 93 permanently ice-free. River runoff into the Barents Sea is very limited. Only one larger river, the Petchora River,
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28 94 enters directly into the south-eastern Barents Sea in Russia. Rivers on the Kola Peninsula, on Svalbard and in
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30 95 Norway are small and often drain into fjords. Thus, sediment discharge through river inflow is low and the
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32 96 main processes responsible for Barents Sea surface sediment distribution are re-deposition by winnowing from
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34 97 shallow banks into troughs and depressions, and deposition from sea ice. Hence, sedimentation rates are
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36 98 generally low, 0.04-2.1 mm/y since the last glacial period, but can be much higher proximal to glacier outlets
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38 99 e.g. close to Svalbard (Fig. 2; supplementary Tab. S1).

100 Material and Methods

101 *Surface sediments: sampling and preparation*

102 In July 2017, surface sediment samples were collected by using a multi-corer at 15 stations (supplementary
103 Tab. S2) along a general south-north gradient in the western Barents Sea (Fig. 1). The first 1 cm of an undisturbed
104 short sediment core at each station was sampled on-board the Royal Research Vessel James Clark Ross
105 immediately after core recovery. At seven stations (B3, B13-B18) samples were taken in 0.5 cm intervals and all
106 samples were stored in plastic bags at -20°C. Prior to any sediment analysis, except for grain size measurements,
107 all samples were freeze-dried and homogenized by gentle grinding using an agate mortar and pestle.

108 *Bulk elemental composition and grain size analysis*

109 Element composition of Barents Sea surface sediments was determined by wavelength dispersive X-ray
110 fluorescence (XRF). A sample split of 700 mg was mixed with 4200 mg di-lithiumtetraborate (Li₂B₄O₇,
111 Spectromelt A10), preoxidized at 500°C with 1.0 g NH₄NO₃ (p.a.) and fused to homogenous glass beads. The

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4 112 glass beads were analysed for 31 elements (Si, Ti, Al, Fe, Mn, Mg, Ca, Na, K, P, As, Ba, Co, Cr, Cu, Ni, Pb, Rb,
5 113 Sr, V, Y, Zn, Zr) using a Philips PW-2400 WD-XRF spectrometer calibrated with 53 geostandards at the
6 114 University of Oldenburg. Analytical precision and accuracy were better than 5% as checked by in-house and
7 115 international reference materials. Results are provided in the supplementary table S3.

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11 116 Grain size distribution was determined using a Mastersizer 2000E laser diffractometer at Leeds University,
12 117 UK. Samples taken in 0.5 cm intervals (stations B3, B13-B18) were mixed prior to grain size analysis. Sediment
13 118 samples were disaggregated in an ultrasonic bath for at least 15 min and grain size distribution of all samples
14 119 were analysed on bulk and on decarbonated samples, which were treated with 10% (vol.) HCl before analysis.
15 120 Grain size analysis was carried out on material within a particle diameter range of 0.1 to 1000 μm and results
16 121 are presented as cumulative volume percentages (supplementary Tab. S4 and S5).

22 23 122 *Organic carbon and reactive iron extraction and analysis (OC-Fe)*

24
25 123 To quantify the amount of organic carbon bound to iron oxides in Barents Sea surface sediment samples we
26 124 applied a citrate–dithionite iron reduction method which simultaneously dissolves all reactive iron
27 125 (oxyhydr)oxides and the organic carbon associated with these phases (OC-Fe). A detailed description of the
28 126 method can be found in Salvadó et al. [30]. Briefly, 0.25 g of each sample was transferred into 30 ml centrifuge
29 127 tubes. 15 ml of a solution containing 0.27M trisodium citrate ($\text{Na}_3\text{C}_6\text{H}_5\text{O}_7\cdot\text{H}_2\text{O}$) and 0.11M sodium bicarbonate
30 128 (NaHCO_3) was added, well mixed and heated up to 80°C in water bath. 0.1M sodium dithionite ($\text{Na}_2\text{S}_2\text{O}_4$) was
31 129 added to the mixture, maintained at 80°C and shaken every five minutes. After 15 min, the mixture was
32 130 centrifuged for 10 min at 4000 rpm and the supernatant was decanted and 200 μl of HCl were added to prevent
33 131 Fe precipitation. The sediment samples were rinsed three times with artificial seawater and then freeze-dried.
34 132 To quantify the organic carbon loss during the experiment, which was unrelated to iron oxides dissolution, a
35 133 control experiment was conducted. For the control experiment, a 0.25 g aliquot of each sample was treated the
36 134 same way as for the reduction experiment but the complexing and reducing agents (sodium citrate and sodium
37 135 dithionate) were replaced with sodium chloride to reach a solution of the same ionic strength. All samples were
38 136 weighed after the experiment to account for mass loss during the experiment. Dissolved iron in the supernatant
39 137 and rinse water of the control and reduction experiment was analysed using a Thermo Scientific iCE3000 Atomic
40 138 Absorption Spectrometer (AAS) at Leeds University, UK. Results are shown in the supplementary table S6 and
41 139 the relative error of the Fe analysis was $\pm 2.6\%$.

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44 140 Organic carbon (OC) content of the bulk sediment before and after the reduction and control experiments
45 141 was analysed on decarbonated samples using 10% (vol.) HCl, rinsed three times and dried overnight at 50°C.
46 142 OC content was determined with a LECO SC-144DR combustion analyser at Leeds University, UK

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4 143 (supplementary Tab. S6). The certified reference material LECO 502-062 and blanks were included in every
5 144 batch, and results are given in weight percentage. The relative error of the OC analysis was $\pm 1.7\%$.

145 *Sedimentary nitrogen and carbon isotope analysis*

10 146 Freeze dried sediments (~0.1 g) were acidified using 4 mol HCl (hydrochloric acid) to remove carbonates for 4
11 147 h, dried overnight at 60°C and analysed on a CS230 Carbon/Sulfur Determinator (Leco Corporation, Michigan,
12 148 USA) using porous crucibles to derive total organic carbon content (TOC). Precision/reproducibility was
13 149 $\pm < 0.1\%$. Total carbon (TC) and nitrogen were determined on a VarioMAX CNS Analyser (Elementar,
14 150 Langenselbold, Hesse, Germany) in at least duplicate (precision/reproducibility $\pm < 0.1\%$). Total inorganic
15 151 carbon (TIC) was calculated as the difference between the TC and TOC (TC-TOC). The calcium carbonate
16 152 (CaCO_3) content was estimated by multiplying TIC by 8.333. Bulk $\delta^{13}\text{C}_{\text{org}}$ was analysed at Elementex
17 153 Laboratories (Cornwall, UK) using IRMS on samples acidified three times using 4 mol HCL with drying at 60
18 154 °C between each acidification (precision/reproducibility to $\pm 0.2\%$).

155 Results and discussion

156 *Sources, spatial distribution and burial of organic carbon*

157 Compared to organic carbon cycling processes in the water column, there is generally a lack of knowledge
158 about the fate of sedimentary organic matter at and in the Arctic Barents Sea seafloor [47-50]. The link between
159 vertical carbon export and accumulation to primary productivity patterns and terrestrial sources is still not well
160 understood. Therefore, uncertainty remains about the origin of the sedimentary organic carbon, especially in
161 the northern Barents Sea. Based on $\text{C}_{\text{org}}/\text{N}_{\text{tot}}$ ratios, $\delta^{13}\text{C}_{\text{org}}$ signatures and pigment analysis, several studies argue
162 that the main source of sedimentary organic matter (OM) in Barents sea surface sediments is marine and derives
163 from productivity in the water column and ice-associated algae production [16, 47, 51-54]. However, by
164 accounting for the sedimentary inorganic nitrogen content, Knies et al. [55] showed that high amounts of
165 terrigenous OM (≥ 50 rel. %) can be present in the seasonally sea ice covered and coastal regions of the northern
166 Barents Sea, while high contributions of marine OM (> 60 rel. %) occur in the ice-free southwestern Barents Sea.
167 Our $\delta^{13}\text{C}_{\text{org}}$ values from the northern station B13-B17 vary between -21.35% to -23.08% and $\text{C}_{\text{org}}/\text{N}_{\text{tot}}$ values
168 range in all stations between 6 and 8.5 (supplementary Tab. S3), which indicates that these locations are strongly
169 influenced by marine OM.

170 The total organic carbon (OC) content of the Barents Sea surface sediments from this study, as well as
171 available OC data from the literature (Fig. 3) [47, 50, 56] show very similar trends. The OC content is higher in

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4 172 northern Barents Sea surface sediments and in coastal areas, whereas the ice-free southern areas show much
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6 173 lower OC contents (Fig. 3). Previous investigation of carbon burial rates in the northern Barents Sea show that
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8 174 carbon preservation in these sediments is considerably higher compared to other Arctic shelf areas [47]. A
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10 175 compilation of published linear sedimentation rates (Fig. 2; supplementary Tab. S1; adapted and extended from
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12 176 Pathirana et al. [50]) shows that sedimentation rates vary between 4 and 210 cm/kyr⁻¹ (average 64 cm/kyr⁻¹) for
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14 177 the entire western Barents Sea. They are lowest close to the western continental shelf edge, probably due to
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16 178 higher current velocities, and sedimentation rates in the seasonally ice covered northern Barents Sea (north of
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18 179 the median winter sea ice extent) are on average slightly higher (78.9 cm/kyr) than in the permanently ice-free
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20 180 southern regions (53.8 cm/kyr, south of the median winter sea ice extent, Fig. 2). This might be related to lower
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22 181 bottom current speed and higher sediment input from Svalbard and sea ice. The OC spatial distribution pattern
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24 182 could be related to different sedimentation rates and thus different oxygen exposure times as OC
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26 183 remineralization via oxygen reduction in marine sediments is the most effective process for OM degradation.
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28 184 However, investigations of sediment mixing and oxygen penetration depth of Barents Sea surface sediment
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30 185 show that at least the first centimetre is homogenised through physical and/or biological mixing [52, 57] and
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32 186 that the oxygen penetration depth in most locations of the Barents Sea is >1 cm [49, 58]. Hence, we assume that
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34 187 the overall OC decomposition is comparable between the northern and southern Barents Sea and that the spatial
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36 188 distribution of OC between the northern and southern Barents Sea is related to other controlling factors. Hence,
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38 189 we used the average sedimentation rates to estimate the average carbon burial rates north and south of the
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40 190 median winter sea ice extent (supplementary Tab. S7). In the seasonally sea ice covered northern area organic
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42 191 carbon burial rates are (6.3 gC/m²yr⁻¹) more than twice as high as in the ice-free southern region (2.4 gC/m²yr⁻¹).
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44 192 Even though these numbers present only an approximation derived from surface sedimentary OC, they are in
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46 193 relatively good agreement with carbon accumulation rate of 5.5 gC/m²yr⁻¹ published previously for the northern
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48 194 Barents Sea area [47]. Based on these findings, we suggest that carbon sequestration in the ice-free southern
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50 195 Barents Sea sediments is lower compared to the ice-covered northern region.

196 *Inverse relationship between total organic carbon content and calcium carbonate*

197 In pelagic sediments, variations in biogenic carbonate content are mainly controlled by dissolution, dilution,
198 and/or productivity changes. Hence, due to the strong relationship of CaCO₃ to marine productivity and, thus,
199 water temperature, salinity, nutrient supply and degree of ice coverage, CaCO₃ is often applied as a proxy to
200 reconstruct climate and environmental changes. Carbonate content in surface sediments from the eastern central
201 Arctic Ocean, north of the Barents Sea, were found to be mainly of biogenic origin [59] and CaCO₃ contents in
202 southern Barents Sea surface sediments show a good correspondence with planktonic foraminifera abundances
203 [60]. In agreement with these findings, our results show a strong relationship between CaCO₃ and Ca ($r = 0.99$)

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4 204 and both parameters are anti-correlated to terrigenous elements like Si, Fe, K, Ti and Al ($r \leq -0.49$; supplementary
5 205 Fig S1). This suggests that the carbonate content in Barents Sea sediments largely reflects the calcareous shell
6 206 fragments from either planktonic or benthic organisms and, that terrigenous CaCO_3 sources have only a very
7 207 minor effect on the composition of Barents Sea surface sediments.

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12 208 The variable carbonate content is also reflected in the grain size distribution in Barents Sea surface sediments
13 209 (Fig. 4). In the southern Barents Sea, bulk grain size distribution at stations B1 to B11 is much more
14 210 heterogeneous with higher contributions of coarse-grained material (35% $>63 \mu\text{m}$) compared to the clay and silt
15 211 fraction dominated northern stations B13 to B18 (87% $<63 \mu\text{m}$). The decarbonated grain size analyses, however,
16 212 show that the siliciclastic fraction is dominated by the silt fraction (average 81%) and very homogeneously
17 213 distributed in Barents Sea surface sediments (Fig. 4). This shows that the bulk grain size measurements of
18 214 Barents Sea sediments are strongly modulated by their carbonate content.

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25 215 Since CaCO_3 in Barents Sea surface sediments is assumed to be mainly of marine origin, higher CaCO_3
26 216 content indicates higher primary productivity, which could be expected to result in higher organic matter fluxes
27 217 towards the seafloor. But the CaCO_3 content in Barents Sea surface sediments shows an opposite pattern to the
28 218 OC distribution, i.e., low OC content in the south-western part coincides with high CaCO_3 content, and vice
29 219 versa in the north-eastern part (Fig. 3) [56]. A possible reason could be OC dilution through higher CaCO_3
30 220 contents in the south-western area of the Barents Sea. However, a calculation of OC contents on a CaCO_3 basis
31 221 (see supplementary Fig. S2) does not indicate a strong dilution effect of OC through inorganic carbon. Moreover,
32 222 in the very productive Storfjord trough south of Svalbard (Station B7, B9-B11), both OC and CaCO_3 show
33 223 relatively high concentrations. Steinsund et al. [60] attributed differences in the CaCO_3 content to carbonate
34 224 dissolution in the north-eastern Barents Sea caused by dense, cold, saline and CO_2 -rich bottom water produced
35 225 by sea ice formation. However, while this may explain the lower carbonate content north of the polar front, it
36 226 cannot explain the described regional differences in the OC content, since OC is not susceptible to dissolution
37 227 by CO_2 -rich waters. Moreover, dense cold bottom water currents produced by sea ice brine formation also occur
38 228 in areas where CaCO_3 concentrations are high, for example in the Storfjord trough (Station B7, B9-B11) [61-63].
39 229 Hebbeln et al. [64] showed that the carbonate content in the surface sediments of the Polar North Atlantic reflect
40 230 the influx of temperate Atlantic waters into the Nordic Seas, where the highest carbonate content follows the
41 231 main axis of the Norwegian Current and decreases with lower water temperature northwards and to the west.
42 232 Moreover, sea ice cover reconstruction based on a sediment core from the south-western Barents Sea showed
43 233 that seasonal sea ice cover during the early Holocene was accompanied by lower carbonate content and a clear
44 234 increase in the total sedimentary organic carbon concentrations [65]. These findings indicate that low carbonate

content in the north-east Barents Sea is likely related to cold Arctic [39] water masses, with lower carbonate production, while higher CaCO_3 content in the south-western Barents Sea sediments are probably related to the warmer Atlantic water inflow (Fig. 3). Hence, we suggest that the opposite distribution pattern of OC and CaCO_3 in the seasonally sea ice-covered north-western Barents Sea and the ice-free southern area (Fig. 3) could be related to differences in primary productivity and vertical OM flux rates. Wassmann et al. [66 and references therein] showed that the main phytoplankton bloom development occurs in May/June in the southern Barents Sea and is relatively predictable. The spring bloom in the northern Barents Sea, however, depends on the sea ice conditions which are highly variable, and the bloom develops more rapidly than in the southern Barents Sea. It follows that while predators are well-adapted to the spring bloom in the southern Barents Sea, the rapid and unpredictable development of the spring bloom in the marginal ice zone typically decouples phytoplankton development from zooplankton grazing [39]. Thus, despite the ice cover, OC pelagic-benthic fluxes are probably higher in the northern Barents Sea due to lower OM consumption in the water column. Additionally, the export of ice algae (diatoms) might substantially contribute to high OM export fluxes in the marginal ice zone [67]. Beyond OM export quantity, high pelagic consumption and recycling also reduces the quality of vertically exported OM, while low to moderate pelagic consumption allows OM of higher quality to reach the seafloor [68]. In accordance with investigations of the pelagic-benthic coupling and related OM fluxes from the water column to the seabed in the Arctic and Northeast Atlantic [9, 66] we suggest that the increased sedimentary OC contents in the northern Barents Sea (Fig. 3) are related to higher rates of OC delivery to the seafloor. This trend in OM export appears to be matched by similar trends in the benthic macro- and megafauna. A clear and consistent south-north distribution pattern of benthic organisms with generally more taxa, higher biomass and higher abundance in the northern Barents Sea implies increased OM fluxes, which support the benthic ecosystem [40]. If we use the environmental setting of the southern ice free Barents Sea as an analogue for a future ice free northern Barents Sea, these findings imply that with ongoing climate change, the northern Barents Sea may transform from a cold and stratified Arctic to a southern Barents Sea-like warm and well-mixed Atlantic-dominated climate regime [6]. This change may lead to a shift from the current “sea ice algae–benthos” ecosystem to a “phytoplankton–zooplankton” dominated ecosystem [9]. Since our findings indicate a link between marine productivity and the geochemical composition of Barents Sea surface sediments, ongoing sea ice reduction and the associated alteration of pelagic primary productivity are expected to cause accompanied shifts in the Barents Sea surface sediment composition. Compared to the modern situation, the northern Barents Sea surface sediments might contain higher contents of CaCO_3 and less OC, which could result in reduced OC burial rates in the future.

Preservation of organic matter promoted by iron in Barents Sea surface sediments

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4 267 To evaluate the preservation of OC in the seasonally ice covered northern Barents Sea and the ice-free
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6 268 southern area, we determined the amount of organic carbon associated with reactive iron phases by applying a
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8 269 citrate–dithionite iron reduction method [29]. In the following, we will discuss the sources of total and reactive
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10 270 iron in Barents Sea surface sediments. Thereafter, we evaluate the spatial distribution pattern of OC bound to
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12 271 iron and show that the fraction of total organic carbon bound to reactive iron phases is not related to sea ice
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14 272 cover.

15 273 In accordance with the previously published spatial distribution pattern of iron in surface sediments from
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17 274 the southern Barents Sea, our results show that the bulk iron contents in Barents Sea surface sediments are
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19 275 highest to the eastern side of the Svalbard archipelago (stations B14-B18) (Fig. 5 and 6 B; Knies et al. [56]). Values
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21 276 decrease towards the south with intermediate concentrations south of Svalbard (station B9-B13), and lowest
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23 277 values in the south-western Barents Sea (stations B1-B6). Higher iron contents in northern Barents Sea sediments
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25 278 are probably related to bedrock erosion by glaciers on Svalbard [69-71], deposition from sea ice [72, 73] and
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27 279 erosion of Barents Sea Mesozoic bedrock [71, 74]. Our results show that the reactive iron (Fe_R) abundance is
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29 280 strongly related to the sedimentary bulk iron content ($r = 0.94$, $n=22$, supplementary Fig. S3). Thus, the Fe_R
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31 281 contents and the relative contributions of dithionite-extractable reactive iron oxides show a south-north gradient
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33 282 as well (Fig. 6 C and D). The reactive iron fraction of the total iron content (fFe_R) in samples from the south-
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35 283 western stations B1-B13 is on average 16.2%, whereas fFe_R contents in samples north of the Polar Front (B14-18)
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37 284 are on average 27.9%. Thus, as sediment samples from seasonally sea ice covered stations contain the highest
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39 285 OC content and show highest fFe_R contribution (Fig. 6 A and D) we would expect them to have a high potential
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41 286 to bind OC to iron oxides as well. Indeed, we find that the amount of OC bound to iron ($OC-Fe_R$) is on average
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43 287 about three times higher in the northern Barents Sea compared to the south-western area (Fig. 5 E). The strong
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45 288 relationship between Fe_R and $OC-Fe_R$ is in accordance with Ma et al. [34] who investigated literature data of
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47 289 $OC-Fe_R$ and suggest that $OC-Fe_R$ contents in marine surface sediments are highly dependent on OC and Fe_R
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49 290 availability. Moreover, our data show no clear spatial relation between sea ice cover and $OC-Fe_R$ content.
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51 291 Stations B6, B7 and B11 were affected by winter sea ice at least for the past 40 years (Fig. 1) [2]. But compared to
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53 292 B11, $OC-Fe_R$ concentrations at B6 and B7 are very low. B13 is not affected by sea ice but $OC-Fe_R$ concentrations
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55 293 are high (Fig. 6 E). This implies that sea ice cover has no direct impact on the preservation of OC through Fe_R
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57 294 sorption.

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59 295 In contrast to $OC-Fe_R$, the spatial distribution of the OC fraction of the total sedimentary OC pool bound to
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296 Fe_R ($fOC-Fe_R$) (Fig. 6 F) shows no relationship to either TOC or Fe_R contents and, therefore, does not show a
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298 spatial south-north gradient. Also, an association to sea ice cover, proximity to land, grain size distribution or
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sediment composition were not identified either. In fact, the fraction of OC bound to Fe_R in the southern Barents Sea is very similar to that in the northern Barents Sea region (Fig. 6F), even though sample locations are very different in terms of their environmental settings, sediment sources, OC and Fe_R contents (see discussion above). Thus, a relatively high fraction of OC can be bound to Fe_R even if absolute Fe_R contents are relatively low. This suggests that the amount of OC bound to reactive iron is not dependent on the total amount of Fe_R available, but that other factors such as the organic matter type and composition as well as redox processes play an important role. This assumption is in accordance with findings from the Eurasian Arctic Shelf. Salvadó et al. [30] showed that the composition of the OC associated with the Fe phases changes with the OM source (i.e., marine versus terrigenous), and that in Arctic shelf areas dominated by marine OM, $f\text{OC-Fe}_R$ can be lower than in areas dominated by remobilized terrigenous OC, e.g. from thawing permafrost. Also Zhao et al. [31] found that in estuarine sediments in southern China, Fe_R was largely associated with terrigenous OC. Moreover, the association between OC and Fe_R is formed mainly through co-precipitation/chelation and/or adsorption [29, 33, 75]. Coprecipitation has a higher sorption capacity of OC and occurs when upward diffusing pore water Fe^{2+} is oxidized at the redox interface in the presence of dissolved OC. Thus, it has been proposed that Fe redox processes are “ultimately the overarching determinant” of $f\text{OC-Fe}_R$ in marine sediments [34]. Even though most observations suggest that the oxygen penetration depth in the Barents Sea is >1 cm [49, 58] and that the first centimetre of Barents Sea surface sediments is affected and homogenised through physical and/or biological mixing [52, 57], the redox interface might still reach into the first centimetre, e.g. due to high Fe^{2+} upward fluxes or seasonal changes of the oxygen penetration depth through primary productivity variability. At seven stations (B3, B13-B18) we analysed the OC bound to iron in 0.5 cm depth intervals. The results show no significant differences between the TOC, Fe and Fe_R contents in the 0-0.5 cm and 0.5-1 cm sections (supplementary Fig. S4), confirming that the first centimetre is well mixed. Compared to the 0.5-1 cm section, $f\text{Fe}_R$, OC-Fe_R and $f\text{OC-Fe}_R$ contents are in general slightly higher in the first half centimetre. This implies that the effect of redox processes (Fe^{2+} upward fluxes) on the $f\text{OC-Fe}_R$ content in the first centimetre of Barents Sea sediments is minor.

Besides the investigation of natural samples, recent experimental laboratory studies on the composition of Fe_R -associated organic matter revealed that varying $\text{OC}_F:\text{Fe}_R$ molar ratios are related to the binding mechanism of OC with Fe_R phases: adsorption results in lower $\text{OC}_F:\text{Fe}_R$ ratios (≤ 1), while co-precipitation yields ratios between 6 and 10 [76]. In turn, the impact of adsorption and co-precipitation on organic matter loadings ultimately depends on the organic matter composition and redox processes [33, 75]. In Barents Sea surface sediments, $\text{OC}_F:\text{Fe}_R$ molar ratios vary between 0.9 and 3.8 (average = 1.8) and are in the range for sediments overlain by oxic bottom waters [29] (supplementary Tab. S6 and Fig. S5). The majority of $\text{OC}_F:\text{Fe}_R$ values show only small variations between about 1-2; only stations B1, B2 and B11 show relatively high values of 2.9, 3.3 and 3.8, respectively. This might indicate that besides the large differences in the biogeochemical characteristics of

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4 331 the Barents Sea shelf regimes, the composition of OC bound to Fe_R is relatively similar, maybe due to generally
5 332 low contributions of terrigenous OM at all investigated locations [77]. However, $OC_F:Fe_R$ values of stations B3
6 333 and B14-B18 show average values of 1.6 and 1.9 for the upper and lower half centimetre, respectively
7 334 (supplementary Tab. S6). This indicates that the effect of coprecipitation is either very small or that factors other
8 335 than the binding mechanisms of OC to Fe oxides, such as mineralogy or Fe-oxide reactivity influence the
9 336 $OC_F:Fe_R$ ratio. Moreover, competitive sorption by arsenic (As) or phosphorus species onto Fe oxide surfaces, can
10 337 influence the $OC_F:Fe_R$ ratio. For example, As contents in our Barents Sea samples are strongly related to Fe_R
11 338 contents ($r = 0.9$, $n = 15$) but show a weak correlation with $fOC-Fe_R$ ($r = 0.5$, $n = 15$), hence it is likely that surface
12 339 sorption sites on Fe oxides can be “blocked” by As and thus are unavailable for OC binding. To further evaluate
13 340 differences in the $OC_F:Fe_R$ ratios in natural sediments from the Barents Sea and globally, we need to develop a
14 341 better understanding of the composition and type of the organic matter bound to iron oxides and the timing of
15 342 when this bonding occurs.

343 Implications and Conclusion

26 344 Strong regional differences in the surface sediment composition between the northern, seasonally sea ice-
27 345 covered and the southern, ice-free region of the western Barents Sea reveal that $CaCO_3$ content shows an
28 346 opposite pattern to the OC distribution, i.e., low OC content in the south-western part coincide with high $CaCO_3$
29 347 content, and vice versa in the north-eastern part. We propose that this is likely related to the modern ecosystem
30 348 structure with higher primary productivity but lower vertical organic carbon flux rates in the southern than in
31 349 the northern Barents Sea. Low $CaCO_3$ content in the north-east Barents Sea might be related to cold Arctic water
32 350 masses, with lower carbonate production, while higher $CaCO_3$ content in the south-western Barents Sea
33 351 sediments is probably related to the warmer Atlantic water inflow.

34 352 Arctic warming will result in higher water temperatures, increased river run-off and reduced sea ice cover.
35 353 Thus, the northern Barents Sea may transform from a cold and stratified Arctic to a southern Barents Sea-like
36 354 warm and well-mixed Atlantic-dominated climate regime. This enormous environmental change will certainly
37 355 induce substantial marine ecosystem changes. More extensive open water conditions and enhanced nutrient
38 356 inputs through rivers are expected to enhance primary productivity. However, less sea ice cover in the northern
39 357 Barents Sea may also lead to a shift of the typical “sea-ice algae–benthos” ecosystem to a “phytoplankton–
40 358 zooplankton” dominated ecosystem. The proposed link between marine productivity and the geochemical
41 359 composition of Barents Sea surface sediments implies that ongoing “Atlantification” of the Barents Sea will
42 360 affect the Barents Sea surface sediment composition and that compared to the modern situation the northern
43 361 Barents Sea surface sediments might contain higher contents of $CaCO_3$ and less OC in the future. Thus, a rise in

primary productivity may lead to higher atmospheric CO₂ uptake but higher carbon turnover rates/remineralisation in the water column may decrease vertical OC fluxes in the northern Barents Sea.

To better constrain the controls on, and efficiency of, carbon burial in the Arctic shelf seas, we analysed the fraction of organic carbon bound to dithionite-extractable iron phases (fOC-Fe_R). Consistent with the global estimate by Lalonde et al. [29] 21% of the total organic carbon is on average associated to iron in Barents Sea surface sediments. We found that a relatively high fraction of OC can be bound to reactive iron even if absolute reactive iron contents are relatively low. Moreover, our findings indicate that the amount of OC bound to reactive iron is not dependent on the total amount of reactive iron available, but that the organic matter type and composition seem to be important factors in natural sediments. Furthermore, the spatial distribution of the organic carbon bound to iron seems to be unrelated to sea ice cover, Atlantic water inflow proximity to land, grain size distribution or sediment composition. Future Arctic warming might therefore neither enhance nor decrease carbon burial through the adsorption to iron oxides.

Additional Information

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Data Accessibility

The datasets supporting this article have been uploaded as part of the supplementary material.

Competing Interests

We declare we have no competing interests.

Author contributions

J.C.F. was the lead author and wrote the manuscript. J.C.F., M.A.S., A.T. and C.M. conducted fieldwork/sampling together and compiled datasets. J.C.F., M.A.S., A.F., I.M., G.D.A., R.H. and J.P. carried all the required analytical work and J.K. provided organic and inorganic elemental data. All authors contributed early ideas, revised the initial manuscript and provided a lively discussion.

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References

- 1 Meier WN, Hovelsrud GK, van Oort BEH, Key JR, Kovacs KM, Michel C, et al. Arctic sea ice in transformation: A review of recent observed changes and impacts on biology and human activity. *Reviews of Geophysics*. 2014;52(3):185-217.
- 2 Fetterer F, Knowles K, Meier WN, Savoie M, Windnagel AK. *Sea Ice Index, Version 3* Boulder, Colorado USA: NSIDC: National Snow and Ice Data Center; 2017 [
- 3 Post E, Bhatt US, Bitz CM, Brodie JF, Fulton TL, Hebblewhite M, et al. Ecological consequences of sea-ice decline. *Science*. 2013;341(6145):519-24.
- 4 Smedsrud LH, Esau I, Ingvaldsen RB, Eldevik T, Haugan PM, Li C, et al. The Role of the Barents Sea in the Arctic Climate System. *Reviews of Geophysics*. 2013;51(3):415-49.
- 5 Loeng H. Features of the physical oceanographic conditions of the Barents Sea. *Polar Research*. 1991;10(1):5-18.
- 6 Lind S, Ingvaldsen RB, Furevik T. Arctic warming hotspot in the northern Barents Sea linked to declining sea-ice import. *Nature Climate Change*. 2018;8(7):634-9.
- 7 Polyakov IV, Pnyushkov AV, Alkire MB, Ashik IM, Baumann TM, Carmack EC, et al. Greater role for Atlantic inflows on sea-ice loss in the Eurasian Basin of the Arctic Ocean. *Science*. 2017;356(6335):285-91.
- 8 Barton BI, Lenn Y-D, Lique C. Observed Atlantification of the Barents Sea Causes the Polar Front to Limit the Expansion of Winter Sea Ice. *Journal of Physical Oceanography*. 2018;48(8):1849-66.
- 9 Piepenburg D. Recent research on Arctic benthos: common notions need to be revised. *Polar Biology*. 2005;28(10):733-55.
- 10 Dalpadado P, Arrigo KR, Hjøllø SS, Rey F, Ingvaldsen RB, Sperfeld E, et al. Productivity in the barents sea--response to recent climate variability. *PLoS One*. 2014;9(5):e95273.
- 11 Wassmann P. Arctic marine ecosystems in an era of rapid climate change. *Progress in Oceanography*. 2011;90(1-4):1-17.
- 12 Wassmann P, Carroll J, Bellerby RGJ. Carbon flux and ecosystem feedback in the northern Barents Sea in an era of climate change: An introduction. *Deep Sea Research Part II: Topical Studies in Oceanography*. 2008;55(20-21):2143-53.
- 13 Arrigo KR, van Dijken GL. Secular trends in Arctic Ocean net primary production. *Journal of Geophysical Research*. 2011;116(C9).
- 14 Arrigo KR, van Dijken G, Pabi S. Impact of a shrinking Arctic ice cover on marine primary production. *Geophysical Research Letters*. 2008;35(19).
- 15 Haug T, Bogstad B, Chierici M, Gjørseter H, Hallfredsson EH, Høines ÅS, et al. Future harvest of living resources in the Arctic Ocean north of the Nordic and Barents Seas: A review of possibilities and constraints. *Fisheries Research*. 2017;188:38-57.
- 16 Stein R, MacDonald RW. *The Organic Carbon Cycle in the Arctic Ocean*. Berlin Heidelberg: Springer; 2004.
- 17 Berner RA. The long-term carbon cycle, fossil fuels and atmospheric composition. *Nature*. 2003;426(6964):323-6.
- 18 Müller PJ, Suess E. Productivity, sedimentation rate, and sedimentary organic matter in the oceans - I. Organic carbon preservation. *Deep Sea Research Part a Oceanographic Research Papers*. 1979;26(12):1347-62.
- 19 Ingall ED, Vancappellen P. Relation between Sedimentation-Rate and Burial of Organic Phosphorus and Organic-Carbon in Marine-Sediments. *Geochimica Et Cosmochimica Acta*. 1990;54(2):373-86.
- 20 Canfield DE. Factors influencing organic carbon preservation in marine sediments. *Chem Geol*. 1994;114:315-29.
- 21 Pedersen T, Calvert SE. Anoxia vs. Productivity: What Controls the Formation of Organic-Carbon-Rich Sediments and Sedimentary Rocks? *AAPG Bulletin*. 1990;74.

- 1
2 22 Hartnett HE, Keil RG, Hedges JI, Devol AH. Influence of oxygen exposure time on organic carbon
3 preservation in continental margin sediments. *Nature*. 1998;391(6667):572-4.
4 23 Burdige DJ. Preservation of organic matter in marine sediments: controls, mechanisms, and an imbalance in
5 sediment organic carbon budgets? *Chem Rev*. 2007;107(2):467-85.
6 24 Hatcher PG, Spiker EC, Szeverenyi NM, Maciel GE. Selective Preservation and Origin of Petroleum-
7 Forming Aquatic Kerogen. *Nature*. 1983;305(5934):498-501.
8 25 Hedges JI, Keil RG. Sedimentary organic matter preservation: an assessment and speculative synthesis.
9 *Marine Chemistry*. 1995;49(2):81-115.
10 26 Mayer LM. Relationships between Mineral Surfaces and Organic-Carbon Concentrations in Soils and
11 Sediments. *Chemical Geology*. 1994;114(3-4):347-63.
12 27 Hemingway JD, Rothman DH, Grant KE, Rosengard SZ, Eglinton TI, Derry LA, et al. Mineral protection
13 regulates long-term global preservation of natural organic carbon. *Nature*. 2019;570(7760):228-31.
14 28 Berner RA. Sedimentary Pyrite Formation. *Am J Sci*. 1970;268(1):1-&.
15 29 Lalonde K, Mucci A, Ouellet A, Gelinás Y. Preservation of organic matter in sediments promoted by iron.
16 *Nature*. 2012;483(7388):198-200.
17 30 Salvadó JA, Tesi T, Andersson A, Ingri J, Dudarev OV, Semiletov IP, et al. Organic carbon remobilized from
18 thawing permafrost is resequenced by reactive iron on the Eurasian Arctic Shelf. *Geophysical Research*
19 *Letters*. 2015;42(19):8122-30.
20 31 Zhao B, Yao P, Bianchi TS, Shields MR, Cui XQ, Zhang XW, et al. The Role of Reactive Iron in the
21 Preservation of Terrestrial Organic Carbon in Estuarine Sediments. *Journal of Geophysical Research:*
22 *Biogeosciences*. 2018;123(12):3556-69.
23 32 Barber A, Brandes J, Leri A, Lalonde K, Balind K, Wirick S, et al. Preservation of organic matter in marine
24 sediments by inner-sphere interactions with reactive iron. *Sci Rep*. 2017;7(1):366.
25 33 Shields MR, Bianchi TS, Gélinas Y, Allison MA, Twilley RR. Enhanced terrestrial carbon preservation
26 promoted by reactive iron in deltaic sediments. *Geophysical Research Letters*. 2016;43(3):1149-57.
27 34 Ma W-W, Zhu M-X, Yang G-P, Li T. Iron geochemistry and organic carbon preservation by iron
28 (oxyhydr)oxides in surface sediments of the East China Sea and the south Yellow Sea. *Journal of Marine*
29 *Systems*. 2018;178:62-74.
30 35 Linkhorst A, Dittmar T, Waska H. Molecular Fractionation of Dissolved Organic Matter in a Shallow
31 Subterranean Estuary: The Role of the Iron Curtain. *Environ Sci Technol*. 2017;51(3):1312-20.
32 36 Sirois M, Couturier M, Barber A, Gélinas Y, Chaillou G. Interactions between iron and organic carbon in a
33 sandy beach subterranean estuary. *Marine Chemistry*. 2018;202:86-96.
34 37 Wang D, Zhu MX, Yang GP, Ma WW. Reactive Iron and Iron-Bound Organic Carbon in Surface Sediments
35 of the River-Dominated Bohai Sea (China) Versus the Southern Yellow Sea. *Journal of Geophysical Research:*
36 *Biogeosciences*. 2019;124(1):79-98.
37 38 Carmack E, Barber D, Christensen J, Macdonald R, Rudels B, Sakshaug E. Climate variability and physical
38 forcing of the food webs and the carbon budget on panarctic shelves. *Progress in Oceanography*. 2006;71(2-
39 4):145-81.
40 39 Wassmann P, Reigstad M, Haug T, Rudels B, Carroll ML, Hop H, et al. Food webs and carbon flux in the
41 Barents Sea. *Progress in Oceanography*. 2006;71(2-4):232-87.
42 40 Jørgensen LL, Ljubin P, Skjoldal HR, Ingvaldsen RB, Anisimova N, Manushin I. Distribution of benthic
43 megafauna in the Barents Sea: baseline for an ecosystem approach to management. *ICES Journal of Marine*
44 *Science*. 2015;72(2):595-613.
45 41 Loeng H, Ozhigin V, Adlandsvik B. Water fluxes through the Barents Sea. *Ices Journal of Marine Science*.
46 1997;54(3):310-7.
47 42 Jakobsen T, Ozhigin VK. The Barents Sea - ecosystem, resources, management. Half a century of Russian -
48 Norwegian cooperation: Tapir Akademisk Forlag; 2011.
49 43 Sakshaug E. Primary and Secondary Production in the Arctic Seas. In: Stein R, MacDonald RW, editors. *The*
50 *Organic Carbon Cycle in the Arctic Ocean*. Berlin, Heidelberg: Springer Berlin Heidelberg; 2004. p. 57-81.
51 44 Eriksen E, Skjoldal HR, Gjørseter H, Primicerio R. Spatial and temporal changes in the Barents Sea pelagic
52 compartment during the recent warming. *Progress in Oceanography*. 2017;151:206-26.
53
54
55
56
57
58
59
60

- 1
2 45 Harris CL, Plueddemann AJ, Gawarkiewicz GG. Water mass distribution and polar front structure in the
3 western Barents Sea. *Journal of Geophysical Research: Oceans*. 1998;103(C2):2905-17.
- 4 46 Drinkwater KF. The influence of climate variability and change on the ecosystems of the Barents Sea and
5 adjacent waters: Review and synthesis of recent studies from the NESSAS Project. *Progress in Oceanography*.
6 2011;90(1-4):47-61.
- 7 47 Carroll J, Zaborska A, Papucci C, Schirone A, Carroll ML, Pempkowiak J. Accumulation of organic carbon
8 in western Barents Sea sediments. *Deep Sea Research Part II: Topical Studies in Oceanography*. 2008;55(20-
9 21):2361-71.
- 10 48 Nickel M, Vandieken V, Brüchert V, Jørgensen BB. Microbial Mn(IV) and Fe(III) reduction in northern
11 Barents Sea sediments under different conditions of ice cover and organic carbon deposition. *Deep Sea*
12 *Research Part II: Topical Studies in Oceanography*. 2008;55(20-21):2390-8.
- 13 49 Vandieken V, Nickel M, Jørgensen BB. Carbon mineralization in Arctic sediments northeast of Svalbard:
14 Mn(IV) and Fe(III) reduction as principal anaerobic respiratory pathways. *Mar Ecol Prog Ser*. 2006;322:15-27.
- 15 50 Pathirana I, Knies J, Felix M, Mann U. Towards an improved organic carbon budget for the western Barents
16 Sea shelf. *Climate of the Past*. 2014;10(2):569-87.
- 17 51 Tamelander T, Reigstad M, Hop H, Carroll ML, Wassmann P. Pelagic and sympagic contribution of organic
18 matter to zooplankton and vertical export in the Barents Sea marginal ice zone. *Deep Sea Research Part II:*
19 *Topical Studies in Oceanography*. 2008;55(20-21):2330-9.
- 20 52 Zaborska A, Carroll J, Papucci C, Torricelli L, Carroll ML, Walkusz-Miotk J, et al. Recent sediment
21 accumulation rates for the Western margin of the Barents Sea. *Deep Sea Research Part II: Topical Studies in*
22 *Oceanography*. 2008;55(20-21):2352-60.
- 23 53 Morata N, Renaud PE. Sedimentary pigments in the western Barents Sea: A reflection of pelagic–benthic
24 coupling? *Deep Sea Research Part II: Topical Studies in Oceanography*. 2008;55(20-21):2381-9.
- 25 54 Belt ST, Cabedo-Sanz P, Smik L, Navarro-Rodriguez A, Berben SMP, Knies J, et al. Identification of paleo
26 Arctic winter sea ice limits and the marginal ice zone: Optimised biomarker-based reconstructions of late
27 Quaternary Arctic sea ice. *Earth and Planetary Science Letters*. 2015;431:127-39.
- 28 55 Knies J, Martinez P. Organic matter sedimentation in the western Barents Sea region: Terrestrial and marine
29 contribution based on isotopic composition and organic nitrogen content. *Norwegian Journal of Geology*.
30 2009;89(1-2):79-89.
- 31 56 Knies J, Jensen HKB, Finne TE, Lepland A, Sæther OM. Sediment composition and heavy metal distribution
32 in Barents Sea surface samples: Results from Institute of Marine Research 2003 and 2004 cruises. *Geological*
33 *Survey of Norway*; 2006. Report No.: 2006.067.
- 34 57 Maiti K, Carroll J, Benitez-Nelson CR. Sedimentation and particle dynamics in the seasonal ice zone of the
35 Barents Sea. *Journal of Marine Systems*. 2010;79(1-2):185-98.
- 36 58 Hulth S, Blackburn TH, Hall POJ. Arctic sediments (Svalbard): consumption and microdistribution of
37 oxygen. *Marine Chemistry*. 1994;46:293-316.
- 38 59 Stein R, Grobe H, Wahsner M. Organic-Carbon, Carbonate, and Clay Mineral Distributions in Eastern
39 Central Arctic-Ocean Surface Sediments. *Marine Geology*. 1994;119(3-4):269-85.
- 40 60 Steinsund PI, Hald M. Recent Calcium-Carbonate Dissolution in the Barents Sea - Paleoceanographic
41 Applications. *Marine Geology*. 1994;117(1-4):303-16.
- 42 61 Knies J, Pathirana I, Cabedo-Sanz P, Banica A, Fabian K, Rasmussen TL, et al. Sea-ice dynamics in an Arctic
43 coastal polynya during the past 6500 years. *arktos*. 2016;3(1).
- 44 62 Schauer U. The release of brine-enriched shelf water from Storfjord into the Norwegian Sea. *Journal of*
45 *Geophysical Research*. 1995;100(C8).
- 46 63 Honjo S, Manganini SJ, Wefer G. Annual Particle-Flux and a Winter Outburst of Sedimentation in the
47 Northern Norwegian Sea. *Deep-Sea Res*. 1988;35(8):1223-34.
- 48 64 Hebbeln D, Henrich R, Baumann KH. Paleoceanography of the last interglacial/glacial cycle in the Polar
49 North Atlantic. *Quaternary Science Reviews*. 1998;17(1-3):125-53.
- 50 65 Berben SMP, Husum K, Cabedo-Sanz P, Belt ST. Holocene sub-centennial evolution of Atlantic water
51 inflow and sea ice distribution in the western Barents Sea. *Climate of the Past*. 2014;10(1):181-98.

- 66 Wassmann P, Peinert R, Smetacek V. Patterns of Production and Sedimentation in the Boreal and Polar Northeast Atlantic. *Polar Research*. 1991;10(1):209-28.
- 67 Boetius A, Albrecht S, Bakker K, Bienhold C, Felden J, Fernandez-Mendez M, et al. Export of algal biomass from the melting Arctic sea ice. *Science*. 2013;339(6126):1430-2.
- 68 Reigstad M, Wexels Riser C, Wassmann P, Ratkova T. Vertical export of particulate organic carbon: Attenuation, composition and loss rates in the northern Barents Sea. *Deep Sea Research Part II: Topical Studies in Oceanography*. 2008;55(20-21):2308-19.
- 69 Rasmussen TL, Thomsen E. Pink marine sediments reveal rapid ice melt and Arctic meltwater discharge during Dansgaard–Oeschger warmings. *Nature Communications*. 2013;4(1).
- 70 Flink AE, Noormets R, Fransner O, Hogan KA, ÓRegan M, Jakobsson M. Past ice flow in Wahlenbergfjorden and its implications for late Quaternary ice sheet dynamics in northeastern Svalbard. *Quaternary Science Reviews*. 2017;163:162-79.
- 71 Vogt C, Knies J. Sediment pathways in the western Barents Sea inferred from clay mineral assemblages in surface sediments. *Norwegian Journal of Geology*. 2009;89(1-2):41-55.
- 72 Lannuzel D, Vancoppenolle M, van der Merwe P, de Jong J, Meiners KM, Grotti M, et al. Iron in sea ice: Review and new insights. *Elementa-Sci Anthrop*. 2016;4.
- 73 Elverhoi A, Pfirman SL, Solheim A, Larssen BB. Glaciomarine Sedimentation in Epicontinental Seas Exemplified by the Northern Barents Sea. *Marine Geology*. 1989;85(2-4):225-50.
- 74 Bjørlykke K, Bue B, Elverhoi A. Quaternary sediments in the northwestern part of the Barents Sea and their relation to the underlying Mesozoic bedrock. *Sedimentology*. 1978;25(2):227-46.
- 75 Chen C, Dynes JJ, Wang J, Sparks DL. Properties of Fe-organic matter associations via coprecipitation versus adsorption. *Environ Sci Technol*. 2014;48(23):13751-9.
- 76 Wagai R, Mayer LM. Sorptive stabilization of organic matter in soils by hydrous iron oxides. *Geochimica et Cosmochimica Acta*. 2007;71(1):25-35.
- 77 Stevenson MA, Abbott GD. Exploring the composition of macromolecular organic matter in Arctic Ocean sediments under a changing sea ice gradient. *Journal of Analytical and Applied Pyrolysis*. 2019;140:102-11.

Figure captions

Figure 1: Map of the western Barents Sea and sampling locations (red dots). The northern Barents Sea is seasonally ice-covered and winter maximum and median sea ice coverage over the past forty years [2] are shown as white area and blue line, respectively. The boundary between the relatively warm northward flowing North Atlantic Current and the southward flowing cold Arctic currents forms the oceanographic Polar Front (yellow line).

Figure 2: Published linear sedimentation rates (LSR) in the Barents Sea. Data and references are provided in supplementary table S1.

Figure 3: Spatial distribution of CaCO₃ (left) and total organic carbon (right) in Barents Sea surface sediments. For further legend details see Fig. 1.

Figure 4: Grain size distribution in Barents Sea surface sediments in a) decarbonated and b) bulk sediment samples.

Figure 5: Spatial distribution of iron in Barents Sea surface sediments. Data from this study and Knies et al. [56].

Figure 6: Distribution of A) TOC, B) bulk Fe, C) reactive iron, D) reactive iron fraction of total iron ($f\text{Fe}_R$), E) organic carbon bound to reactive iron (OC- Fe_R) and F) the organic carbon fraction of total organic carbon bound to reactive iron ($f\text{OC-Fe}_R$) in Barents Sea surface sediments (0-1 cm). Circles mark stations which are

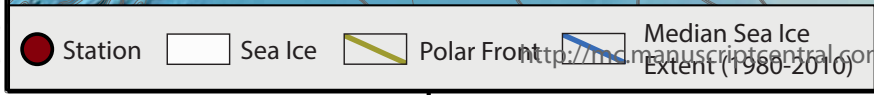
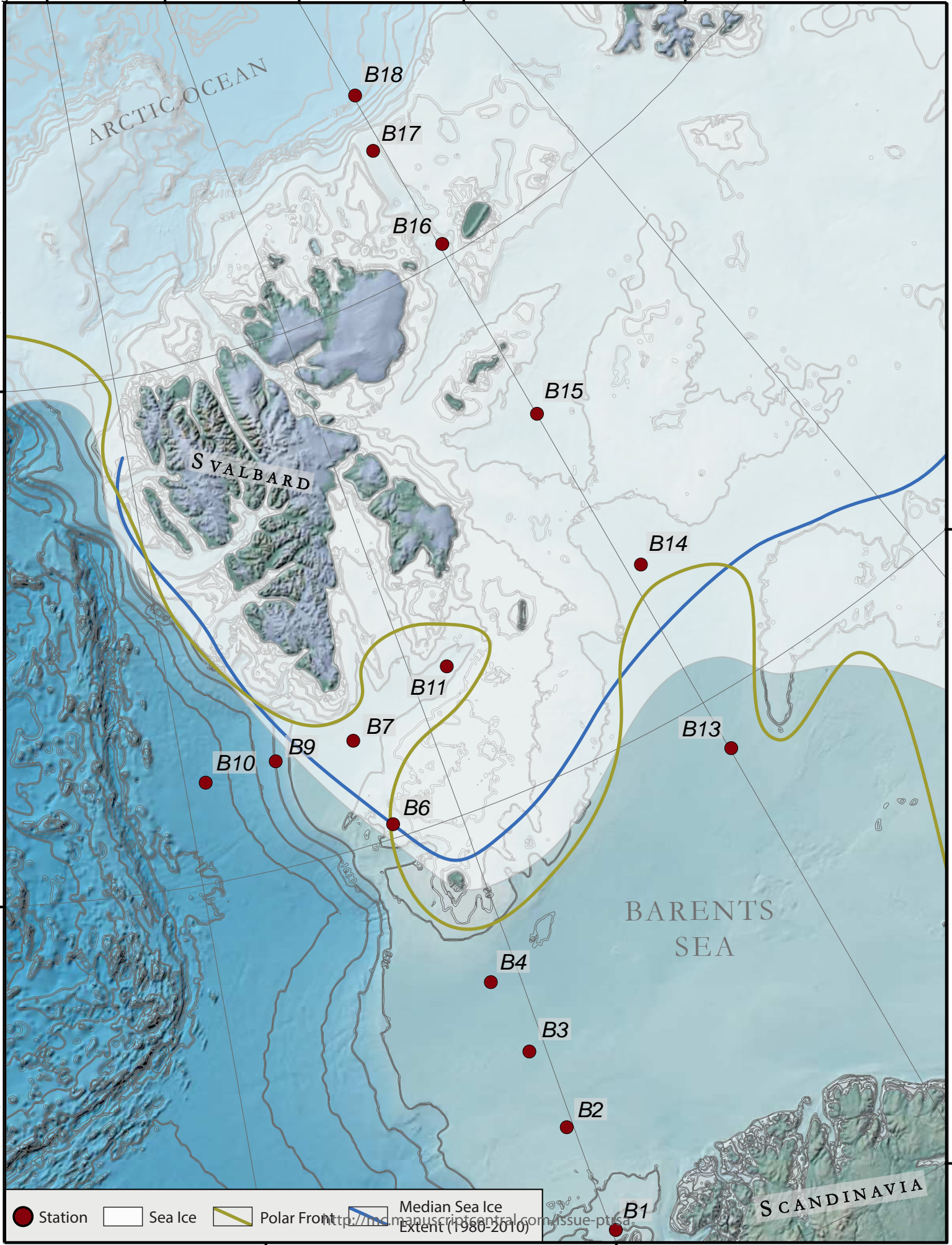
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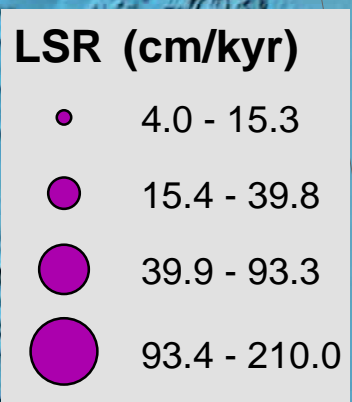
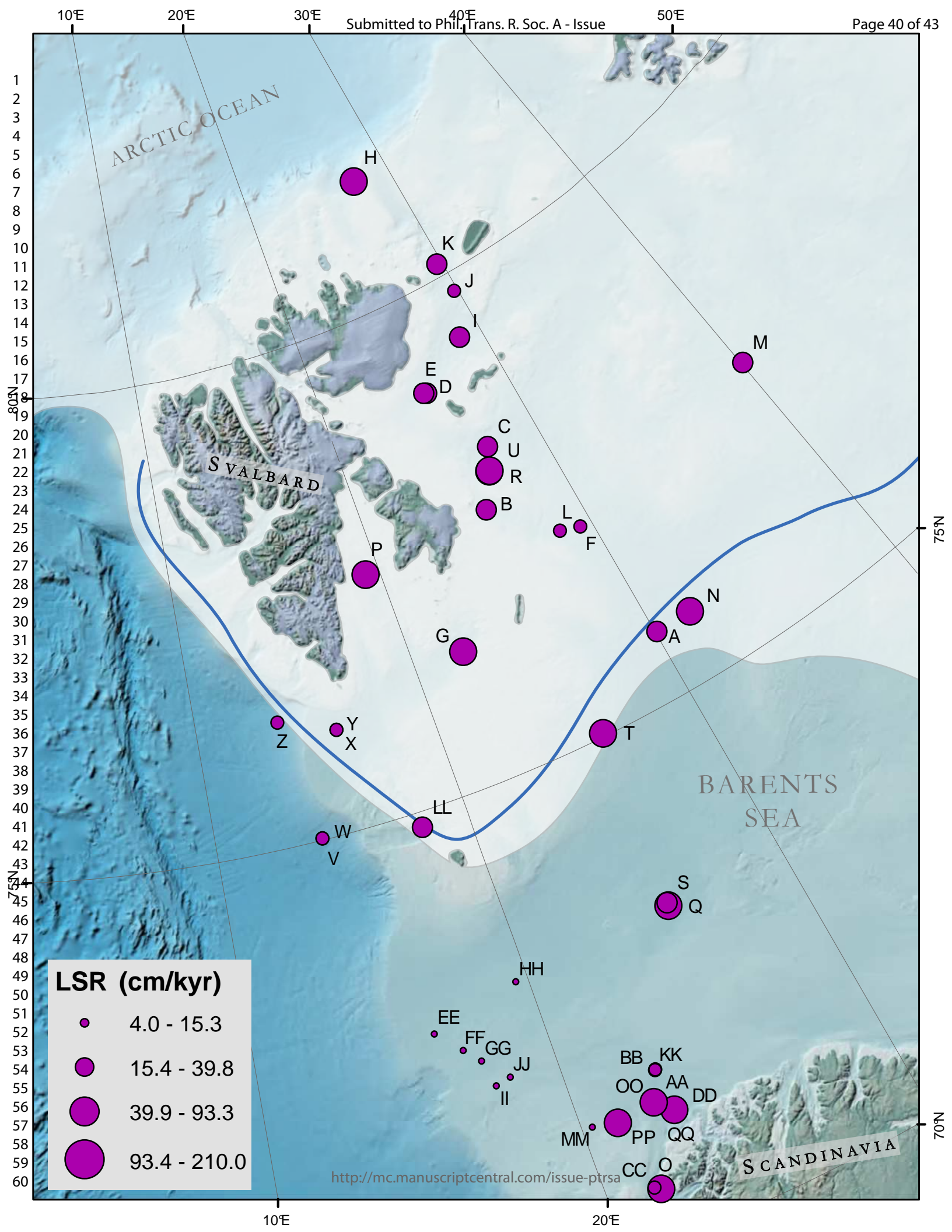
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2 seasonally sea ice covered and crosses are stations which are ice free during winter. Station locations (B1-B18)
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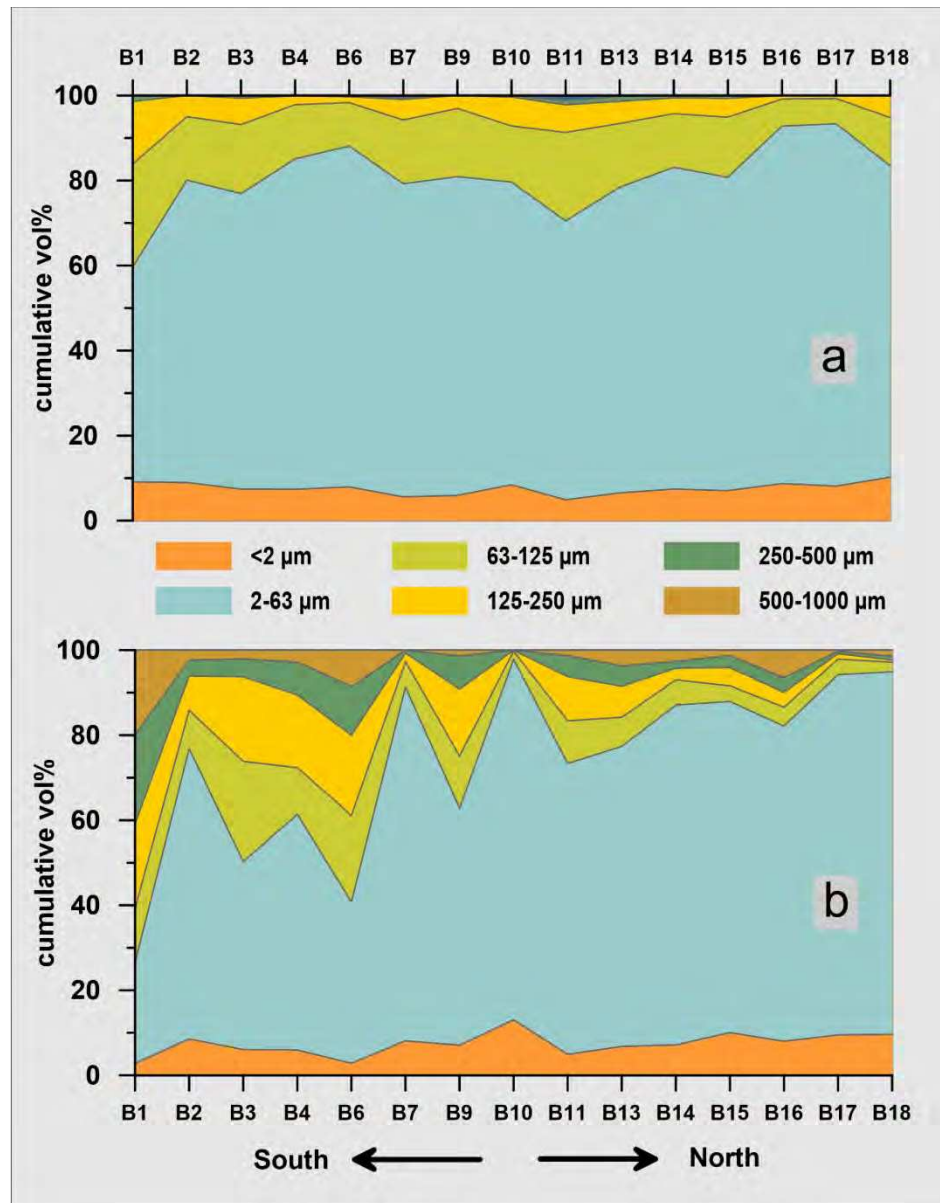
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17 **Supplementary material**

18 Supporting information associated with this article (figure S1 to S5 and table S1 to S7) can be found in the
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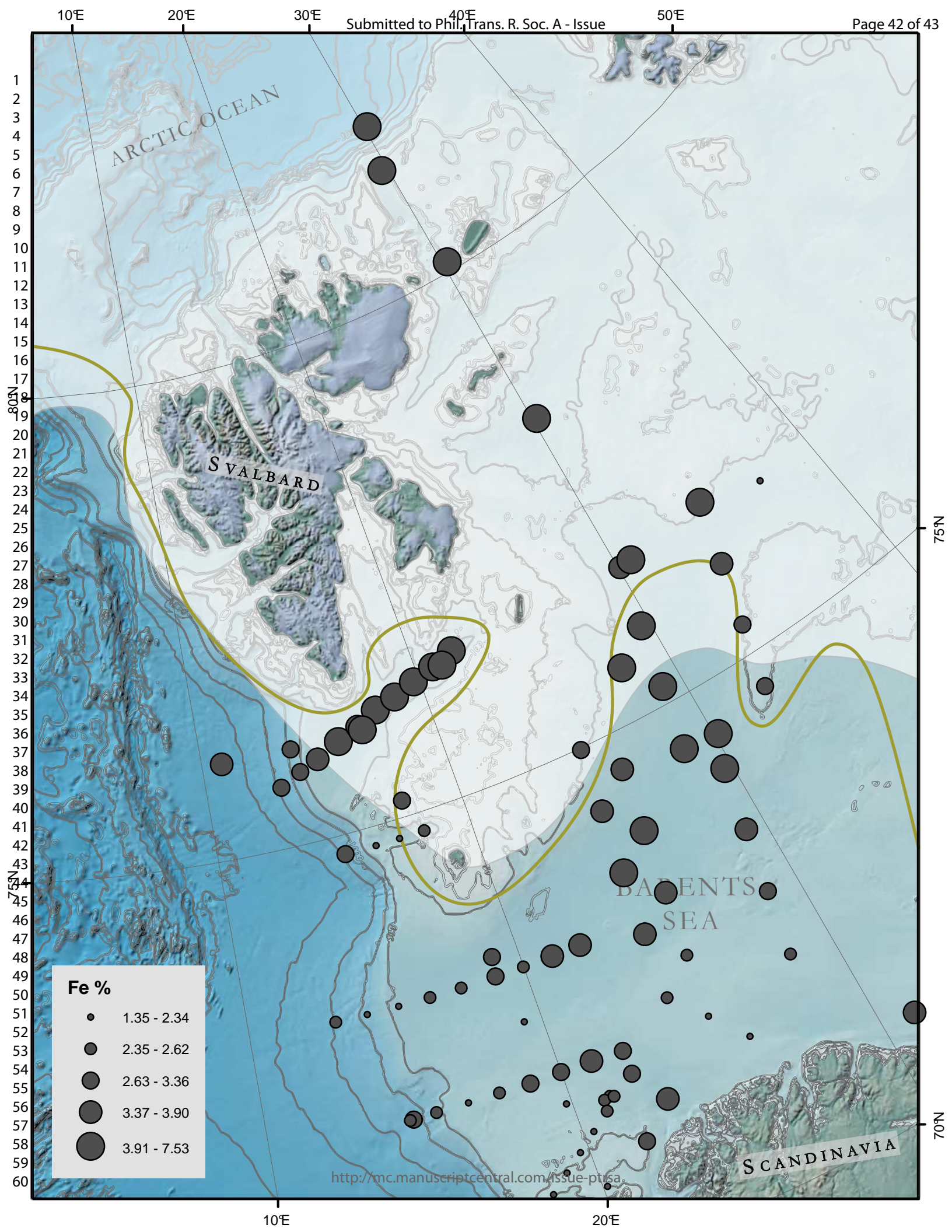


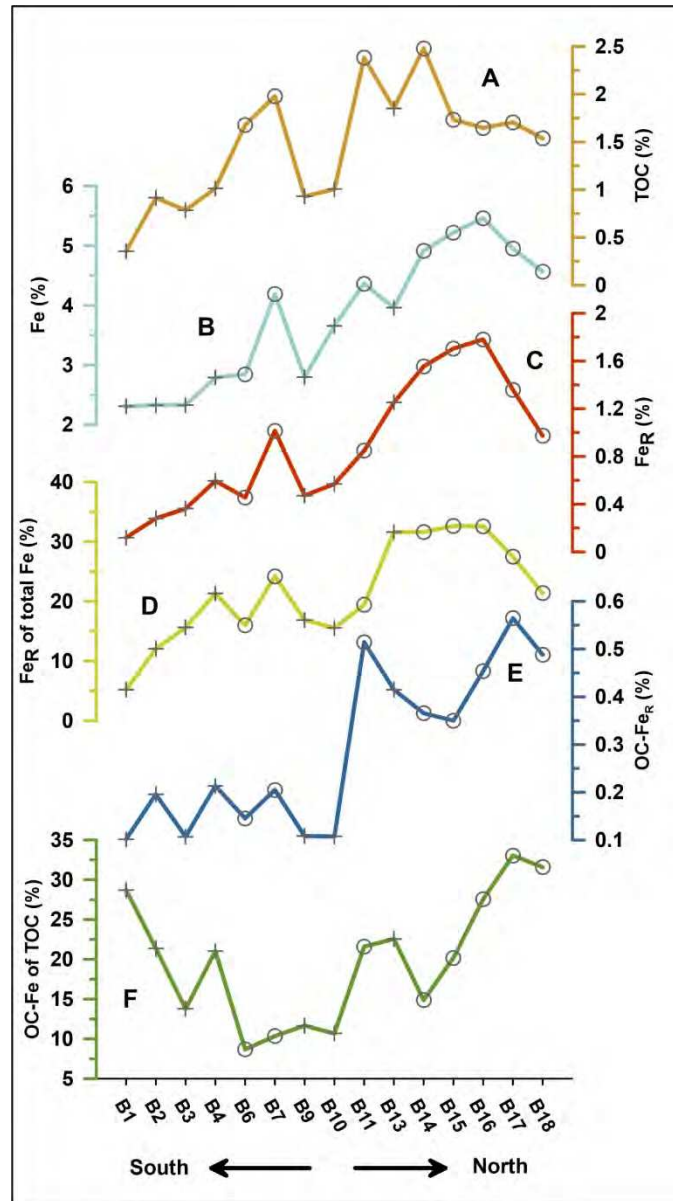




Grain size distribution in Barents Sea surface sediments in a) decarbonated and b) bulk sediment samples.

124x158mm (600 x 600 DPI)





Distribution of A) TOC, B) bulk Fe, C) reactive iron, D) reactive iron fraction of total iron (fFeR), E) organic carbon bound to reactive iron (OC-FeR) and F) the organic carbon fraction of total organic carbon bound to reactive iron (fOC-FeR) in Barents Sea surface sediments (0-1 cm). Circles mark stations which are seasonally sea ice covered and crosses are stations which are ice free during winter. Station locations (B1-B18) and ice coverage is shown in Fig. 1.

113x201mm (600 x 600 DPI)