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1 **Distribution of ^{210}Pb and ^{210}Po in the Arctic water column during the 2007 sea-ice**
2 **minimum: particle export in the ice-covered basins**

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19 **ABSTRACT**

20 ^{210}Pb and ^{210}Po are naturally occurring radionuclides that are commonly used as a proxy
21 for particle and carbon export. In this study, the distribution of the $^{210}\text{Po}/^{210}\text{Pb}$ pair was
22 investigated in the water column of the Barents, Kara and Laptev Seas and the Nansen,
23 Amundsen and Makarov Basins in order to understand the particle dynamics in the Arctic
24 Ocean during the 2007 sea-ice minimum (August-September). Minimum activities of
25 total ^{210}Pb and ^{210}Po were found in the upper and lower haloclines (approx. 60-130 m),
26 which are partly attributed to particle scavenging over the shelves, boundary current
27 transport and subsequent advection of the water with low ^{210}Pb and ^{210}Po activities into
28 the central Arctic. Widespread and substantial (>50%) deficits of ^{210}Po with respect to
29 ^{210}Pb were detected from surface waters to 200 m on the shelves, but also in the basins.
30 This was particularly important in the Makarov Basin where, despite very low
31 chlorophyll-a levels, estimates of annual new primary production were three times higher

32 than in the Eurasian Basin. In the Nansen, Amundsen and Makarov Basins, estimates of
33 annual new primary production correlated with the deficits of ^{210}Po in the upper 200 m
34 of the water column, suggesting that in situ production and subsequent export of biogenic
35 material were the mechanisms that controlled the removal of ^{210}Po in the central Arctic.
36 Unlike ^{210}Po , ^{234}Th deficits measured during the same expedition were found to be very
37 small and not significant below 25 m in the basins (Cai et al., 2010), which indicates,
38 given the shorter half-life of ^{234}Th , that particle export fluxes in the central Arctic would
39 have been higher before July-August in 2007 than later in the season.

40 Keywords:

41 Particle export, annual new primary production, scavenging, $^{210}\text{Po}/^{210}\text{Pb}$, Arctic Ocean,
42 2007 sea-ice minimum

43 Highlights:

44 Largest dataset of ^{210}Pb and ^{210}Po in the Arctic Ocean to date
45 Minimum ^{210}Pb and ^{210}Po activities in the halocline reflect scavenging on the shelf
46 Potential use of the $^{210}\text{Po}/^{210}\text{Pb}$ proxy as an indicator of annual new primary production
47 Substantial deficits of ^{210}Po in surface and subsurface waters of the Makarov Basin
48 Higher particle export in the central Arctic before July-August in 2007

49 **1. Introduction**

50 The Arctic Ocean is undergoing rapid changes in response to global warming, including
51 a rapid sea-ice retreat (e.g. Stroeve et al., 2012). During the past 10 years, the Arctic sea-
52 ice extent has experienced two record minima in September 2007 and 2012, related to
53 anomalously high temperatures and southerly winds (Comiso et al., 2008), or intense
54 cyclone events (Parkinson and Comiso, 2013). Sea ice has also thinned and lost volume
55 (Kwok et al., 2009; Laxon et al., 2013; Renner et al., 2014). The freshwater storage in the
56 upper Arctic Ocean has concurrently increased, primarily due to a reduction in the
57 average salinity in the layer between the surface and the lower halocline and, to a lesser
58 extent, due to the thickening of that layer (Haine et al., 2015; Rabe et al., 2014, 2011).
59 These changes have been accompanied by an increase in upper ocean stratification

60 (Korhonen et al., 2013). This scenario impacts primary production (e.g. Ardyna et al.,
61 2014; Arrigo and van Dijken, 2015), marine ecosystems (Wassmann et al., 2011) and
62 pelagic-benthic coupling (Wassmann and Reigstad, 2011), but there are many
63 uncertainties due to the limitations in obtaining in situ data. Therefore, impacts on
64 biogeochemical cycles in the Arctic remain poorly understood, including the transport of
65 particles to deep waters and its implications for carbon export (Reid et al., 2009; Tremblay
66 et al., 2015; Wassmann, 2011).

67 The present understanding is that particle and carbon export from the upper water column
68 to depth is widely heterogeneous in the Arctic, being substantially higher on the shelves
69 than in the basins in consistency with the distribution of primary production (Findlay et
70 al., 2015; Honjo et al., 2010; Randelhoff and Guthrie, 2016; Wassmann et al., 2004). In
71 the late summer of 2007, the time of the present study, deficits of ^{234}Th with respect to its
72 parent, ^{238}U , in the upper 100 m of the water column were basically restricted to the Arctic
73 shelves and continental margins (Cai et al., 2010); in the deep basins ($>80^\circ\text{N}$), very small
74 ^{234}Th deficits were measured in the upper 25 m and $^{234}\text{Th}/^{238}\text{U}$ ratios were
75 indistinguishable from 1 at deeper depths, indicating very small particle export in the
76 central Arctic. This low export is in line with previous studies conducted in the central
77 Arctic during the summer season (see compilation of ^{234}Th flux data in Roca-Martí et al.,
78 2016), although there are exceptions, for instance, in the Canada Basin (e.g. Baskaran et
79 al., 2003; Ma et al., 2005).

80 The natural radionuclides ^{210}Pb (half-life, $T_{1/2} = 22.3$ years) and ^{210}Po ($T_{1/2} = 138.4$ days)
81 have been widely used as particle tracers in the marine environment (e.g. Bacon et al.,
82 1988, 1976; Cochran and Masqué, 2003), but their application in the Arctic is scarce
83 (Moore and Smith, 1986; Roca-Martí et al., 2016; Smith et al., 2003; Smith and Ellis,
84 1995). Both have a strong affinity for particle surfaces, but ^{210}Po is also incorporated into
85 the cytoplasm of bacteria and phytoplankton (Cherrier et al., 1995; Fisher et al., 1983)
86 and is preferentially assimilated by zooplankton with respect to ^{210}Pb (Stewart and Fisher,
87 2003). When sinking of biogenic particles occurs, the different biogeochemical
88 behaviours of ^{210}Pb and ^{210}Po create a deficit of the latter in the ocean surface that can be
89 used to estimate particulate organic carbon (POC) sinking fluxes (see review by Verdeny
90 et al., 2009). The $^{210}\text{Po}/^{210}\text{Pb}$ pair has, potentially, some advantages over the most
91 commonly used proxy for POC export, the $^{234}\text{Th}/^{238}\text{U}$ pair, when studying export on a

92 seasonal scale: ^{210}Po has a stronger preference for POC than ^{234}Th (Friedrich and Rutgers
93 van der Loeff, 2002) and, with a mean life of 200 days, ^{210}Po integrates a time scale of
94 several months, while ^{234}Th (mean life of 35 days) misses events that had occurred more
95 than one month before sampling (Stewart et al., 2007a; Verdeny et al., 2009).

96 Here, we examine the distribution of ^{210}Pb and ^{210}Po (total and particulate) on three
97 shelves (Barents, Kara and Laptev) and in three deep basins (Nansen, Amundsen and
98 Makarov) of the Arctic Ocean in order to investigate the processes governing the
99 dynamics of particles and the particle export around the time of the sea-ice minimum in
100 summer 2007. This is combined with information on physical characteristics of the study
101 areas, estimates of annual new primary production and the export production derived
102 from ^{234}Th data from the same expedition (Cai et al., 2010).

103 **2. Materials and methods**

104 **2.1. Study area**

105 Seawater samples were collected during the ARK-XXII/2 expedition that took place from
106 28 July to 7 October in 2007 along the shelves of the Barents, Kara and Laptev Seas and
107 the Nansen, Amundsen and Makarov Basins (R/V Polarstern, Schauer, 2008). Stations
108 have been classified into three categories according to the water depth (Table 1, shelf:
109 <350 m; slope: 350-1050 m; basin: >1050 m), and divided into five sections (S1-S5, see
110 Figure 1), similarly to Cai et al. (2010).

111 **2.2. ^{210}Pb and ^{210}Po**

112 Pb-210 and ^{210}Po activities were measured in the dissolved and particulate fractions in
113 the upper ~500 m of the water column, while only total activities were determined for
114 deeper samples (Table S1). Surface seawater (10 m) was sampled at 50 stations (Table 1)
115 using the ship seawater intake. Vertical profiles were collected at 17 stations (Table 1)
116 using Niskin bottles attached to a conductivity-temperature-depth (CTD) rosette sampler.
117 The sample volumes were about 20 L (dissolved and total) and 40 to 160 L (particulate).
118 Samples were filtered through 1 μm pore-size Nuclepore filters (142 mm, Whatman) to
119 separate the dissolved and particulate fractions. After filtration, the filters were dried at
120 room temperature and stored for later processing at the Alfred Wegener Institute (AWI).
121 Samples for the analyses of the total and dissolved fractions were acidified with 20 mL

122 concentrated HNO₃ immediately after collection and filtration, respectively, and spiked
123 with known amounts of ²⁰⁸Po (T_{1/2} = 2.9 years) and stable Pb. After addition of FeCl₃ as
124 a carrier and vigorous stirring, samples were allowed to equilibrate for about 24 h. Pb and
125 Po were then co-precipitated with Fe(OH)₃ by adjusting the pH to ~8.5 with ammonium
126 hydroxide solution. After a few hours, supernatants were removed carefully by
127 decantation and samples were centrifuged. Precipitates were transferred to plastic bottles
128 and stored until further analyses at AWI.

129 Pb-210 and ²¹⁰Po were determined following the method of Fleer and Bacon (1984),
130 based on the procedure of Flynn (1968). Filters (i.e. particulate fraction) were spiked with
131 ²⁰⁸Po and stable Pb and digested using a microwave with a mixture of HNO₃, HF and
132 H₂O₂ (10, 0.3 and 2 mL, respectively). Pb and Po isotopes were then co-precipitated with
133 Fe(OH)₃ as described for water samples. Iron precipitates were dissolved in 0.5 M HCl
134 and ascorbic acid was added to the solutions to reduce Fe³⁺ to Fe²⁺ before placing the
135 silver discs and hence allow the auto-deposition of Po (80°C, >4 hours). The silver discs
136 were then counted for Po by alpha spectrometry using silicon surface barrier alpha
137 detectors (EG&G Ortec, USA) for about seven days to achieve counting statistics <5%.
138 Samples were re-spiked with ²⁰⁹Po (T_{1/2} = 125 years) and stored for 15 to 26 months in 8
139 M HNO₃ for later determination of ²¹⁰Pb via ²¹⁰Po ingrowth. Samples were plated for Po
140 as described above and counted once more by alpha spectrometry. Pb-210 and ²¹⁰Po
141 activities at sampling time were calculated applying appropriate ingrowth and decay
142 corrections (Fleer and Bacon, 1984). The chemical yield of stable Pb was determined by
143 inductively coupled plasma-optical emission spectrometry (ICP-OES, Thermo Fisher
144 Scientific, USA). The activity uncertainties, which account for counting, detector
145 background, spike activities and sample volume, were on average 5% for ²¹⁰Pb and 9%
146 for ²¹⁰Po for the dissolved and total fractions, and 5% for ²¹⁰Pb and 15% for ²¹⁰Po for the
147 particulate fraction. The greater uncertainties of ²¹⁰Po are due to the time elapsed between
148 sampling and the first Po plating (range: 55-117 days; median: 78 days).

149 The AWI laboratory participated in the ²¹⁰Pb and ²¹⁰Po inter-calibration exercise
150 organized by GEOTRACES in 2008-2009 (Church et al., 2012). This study
151 recommended a minimum sample activity of 0.1 dpm for both radionuclides. In this work,
152 the reported activities of ²¹⁰Pb and ²¹⁰Po are always higher than 0.1 dpm for the dissolved
153 and total fractions, while the particulate activities of both radionuclides are below this

154 threshold in about 10% of the samples. The complete ^{210}Pb and ^{210}Po dataset can be found
155 in Table S1 (Supplemental Material) and is available at PANGAEA (Friedrich, 2011).

156 **2.3. CTD observations and nutrients**

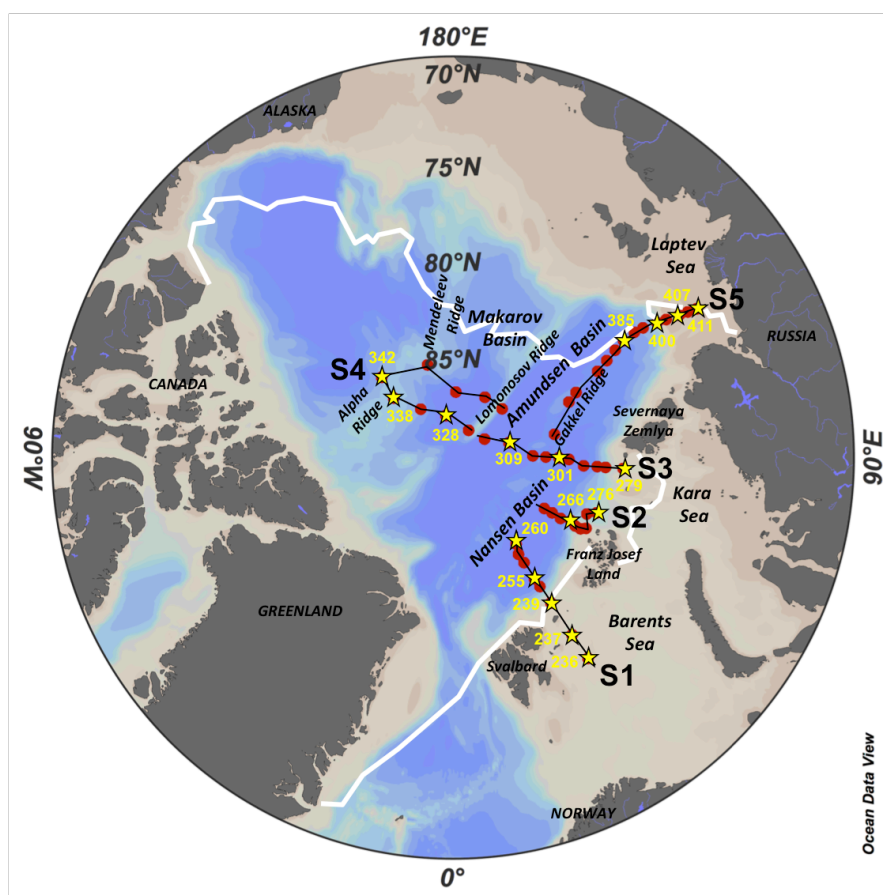
157 Profiles of temperature and salinity in the water column were obtained using a CTD
158 system with a Carousel Water Sampler (Sea-Bird Electronics Inc., USA). Salinity was
159 calibrated using discrete samples from the rosette, and processed with an on-board
160 salinometer. Discrete nutrient water samples from the rosette were processed using a
161 Technicon TRAACS 800 continuous flow auto-analyser and standards previously
162 prepared on land. Details of the measurements and processing are described in Schauer
163 (2008). The data are published in Laan et al. (2008), Wisotzki and Bakker (2008) and
164 Schauer and Wisotzki (2010).

165 **2.4. Annual new primary production**

166 New primary production during the productive season after the Arctic winter (from April
167 to sampling time) was calculated by using estimates of the winter mixed layer depth and
168 the seasonal nutrient uptake based on measurements made during the ARK-XXII/2
169 expedition, following Rudels et al. (1996) and Korhonen et al. (2013). The method
170 interpolates data as in Reiniger and Ross (1968) and is described in detail in Boetius et
171 al. (2013) and Fernández-Méndez et al. (2015). The winter mixed layer depth was
172 estimated by the temperature minimum below the summer mixed layer (August-
173 September 2007), which is assumed to be a remnant of the previous winter convection
174 and homogenization (Rudels et al., 1996). Briefly, in the summer season the net ice melt
175 and warming due to solar radiation both make the surface layer less dense. Because the
176 freshening and warming do not reach below the shallow summer mixed layer, the
177 temperature minimum below this layer and the summer halocline is very likely a remnant
178 of the previous winter mixed layer. Then, the total inventory of nutrients used up since
179 the previous winter is estimated by integrating the difference between the nutrient
180 concentration profiles and the nutrient concentrations at the winter mixed layer depth
181 measured in late summer in the vertical from the surface to the winter mixed layer depth.
182 The annual total inorganic nitrogen (nitrite+nitrate), phosphate and silicate uptake was
183 then converted to carbon units using the Redfield-Brzezinski ratio 106C:16N:15Si:1P
184 (Brzezinski, 1985; Redfield et al., 1963), as done by Boetius et al. (2013), Ulfso et al.

185 (2014) and Fernández-Méndez et al. (2015). The elemental stoichiometry 106C:16N was
186 confirmed for the central Arctic by analysis of suspended particulate organic matter
187 collected during several research programmes (n = 255; Frigstad et al., 2014).

188 Annual new primary production has been only estimated at those stations located in the
189 basins (where ^{210}Pb and ^{210}Po vertical profiles were taken; Table 2). This method assumes
190 that lateral inputs of nutrients from rivers and shelves to surface and subsurface waters in
191 the deep central Arctic ($>78^\circ\text{N}$) have a limited impact on the annual new primary
192 production estimates because the horizontal distance between the shelves and the basin
193 stations is too large to be covered from the winter mixing to the time of sampling with
194 published rates of transport (Ekwurzel et al., 2001; Rutgers van der Loeff et al., 2018).
195 This method does not take into account nitrification, phosphorus remineralization and
196 silica dissolution occurring at or above the winter mixed layer depth during the season.
197 Ulfsbo et al. (2014) found good agreement when comparing this method (using nitrogen
198 and phosphate) to three alternative approaches using concurrent observations during late
199 summer.



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201 **Figure 1:** Location of the stations sampled for surface waters (red dots) and vertical profiles (yellow stars
202 with station numbers) during the ARK-XXII/2 cruise (July-October 2007). The study area is divided into
203 five sections: S1, stations 236-260; S2, stations 261-276; S3, stations 279-312; S4, stations 320-363; S5,
204 stations 371-411. The contour white line represents the minimum sea-ice extent in September 2007
205 (<http://www.meereisportal.de>).

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Table 1: Coordinates, sampling date and water column depth (down to the seafloor) of the stations sampled for ^{210}Pb and ^{210}Po analyses during the ARK-XXII/2 expedition. It is indicated whether the stations were only sampled for surface seawater (10 m) or depth profiles were collected (n refers to the number of samples, see Table S1 for further details). Stations have been classified into sections (S1-S5) and areas (location and shelf/slope/basin, see text for further details).

Section	Station	Long. (°E)	Lat. (°N)	Date (2007)	Samples collected	Water depth (m)	Area
S1	236	33.98	77.50	31 Jul.	Depth profile (n = 7)	196	Barents Shelf
	237	33.97	79.00	31 Jul.	Depth profile (n = 7)	272	Barents Shelf
	239	34.00	80.99	1 Aug.	Depth profile (n = 7)	224	Barents Shelf
	249	33.98	82.00	2 Aug.	Surface seawater	2281	Nansen Basin
	255	33.89	82.52	4 Aug.	Depth profile (n = 11)	3135	Nansen Basin
	257	34.05	83.50	5 Aug.	Surface seawater	3958	Nansen Basin
	258	34.00	84.00	6 Aug.	Surface seawater	4055	Nansen Basin
	260	36.08	84.51	8 Aug.	Depth profile (n = 10)	4054	Nansen Basin
S2	261	60.92	84.64	11 Aug.	Surface seawater	3846	Nansen Basin
	263	60.96	84.17	11 Aug.	Surface seawater	3713	Nansen Basin
	264	60.43	83.65	12 Aug.	Surface seawater	3512	Nansen Basin
	266	61.81	83.12	14 Aug.	Depth profile (n = 13)	3011	Nansen Basin
	268	60.81	82.81	14 Aug.	Surface seawater	1609	Nansen Basin
	271	60.80	82.50	15 Aug.	Surface seawater	327	Barents Shelf
	272	61.99	82.25	15 Aug.	Surface seawater	231	Barents Shelf
	274	67.10	82.52	16 Aug.	Surface seawater	1176	Nansen Basin
	276	68.95	82.09	17 Aug.	Depth profile (n = 8)	680	Kara Slope
S3	279	86.23	81.24	19 Aug.	Depth profile (n = 7)	336	Kara Shelf
	285	86.34	82.14	20 Aug.	Surface seawater	724	Kara Slope
	290	86.44	82.58	21 Aug.	Surface seawater	2071	Nansen Basin
	295	86.30	83.27	22 Aug.	Surface seawater	3357	Nansen Basin
	299	89.06	84.05	23 Aug.	Surface seawater	3694	Nansen Basin
	301	89.76	84.56	24 Aug.	Depth profile (n = 14)	3758	Nansen Basin
	303	90.23	85.25	25 Aug.	Surface seawater	3985	Nansen Basin
	306	91.18	85.92	26 Aug.	Surface seawater	4019	Gakkel Ridge (basin)

	309	104.98	87.04	28 Aug.	Depth profile (n = 14)	4449	Amundsen Basin
	312	120.15	88.12	29 Aug.	Surface seawater	3009	Amundsen Basin
S4	320	150.33	88.41	31 Aug.	Surface seawater	1952	Makarov Basin
	328	-170.33	87.82	2 Sept.	Depth profile (n = 13)	3992	Makarov Basin
	333	-146.39	87.03	4 Sept.	Surface seawater	3285	Makarov Basin
	338	-134.96	85.69	6 Sept.	Depth profile (n = 11)	1570	Makarov Basin
	342	-138.30	84.50	7 Sept.	Depth profile (n = 12)	2289	Makarov Basin
	349	-164.55	85.07	9 Sept.	Surface seawater	1996	Makarov Basin
	352	177.54	86.64	10 Sept.	Surface seawater	4005	Makarov Basin
	358	151.96	86.51	11 Sept.	Surface seawater	1459	Makarov Basin
	363	134.92	86.47	13 Sept.	Surface seawater	3991	Amundsen Basin
S5	371	102.73	84.66	16 Sept.	Surface seawater	4271	Gakkel Ridge (basin)
	377	115.55	83.41	18 Sept.	Surface seawater	4301	Gakkel Ridge (basin)
	379	117.85	82.86	18 Sept.	Surface seawater	4413	Gakkel Ridge (basin)
	382	120.72	81.36	19 Sept.	Surface seawater	5343	Gakkel Ridge (basin)
	383	122.21	80.66	19 Sept.	Surface seawater	3902	Gakkel Ridge (basin)
	384	123.46	80.00	20 Sept.	Surface seawater	3653	Gakkel Ridge (basin)
	385	124.36	79.35	20 Sept.	Depth profile (n = 13)	3525	Gakkel Ridge (basin)
	387	124.61	78.64	21 Sept.	Surface seawater	2865	Gakkel Ridge (basin)
	391	124.24	78.13	21 Sept.	Surface seawater	2435	Gakkel Ridge (basin)
	400	123.42	77.37	22 Sept.	Depth profile (n = 11)	1049	Laptev Slope
	404	122.87	76.90	23 Sept.	Surface seawater	94	Laptev Shelf
	407	122.13	76.18	23 Sept.	Depth profile (n = 2)	75	Laptev Shelf
	409	121.77	75.71	23 Sept.	Surface seawater	65	Laptev Shelf
	411	121.36	75.20	24 Sept.	Depth profile (n = 3)	48	Laptev Shelf

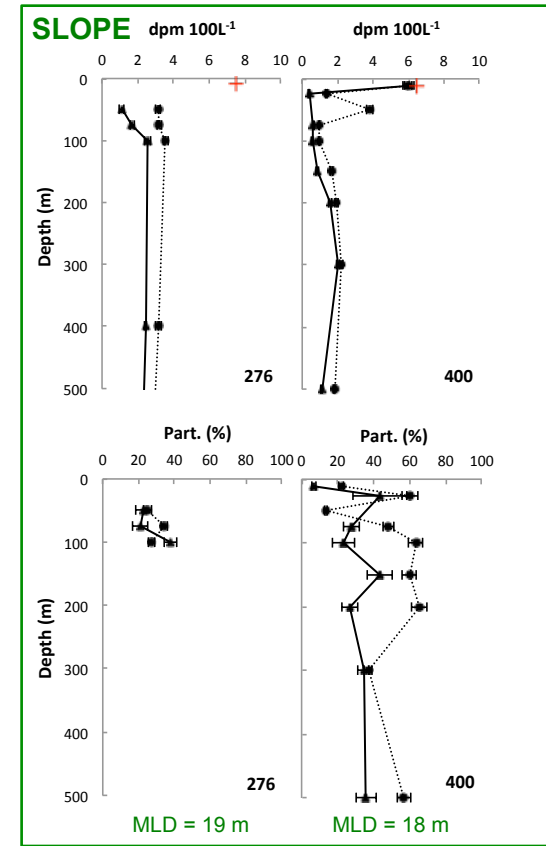
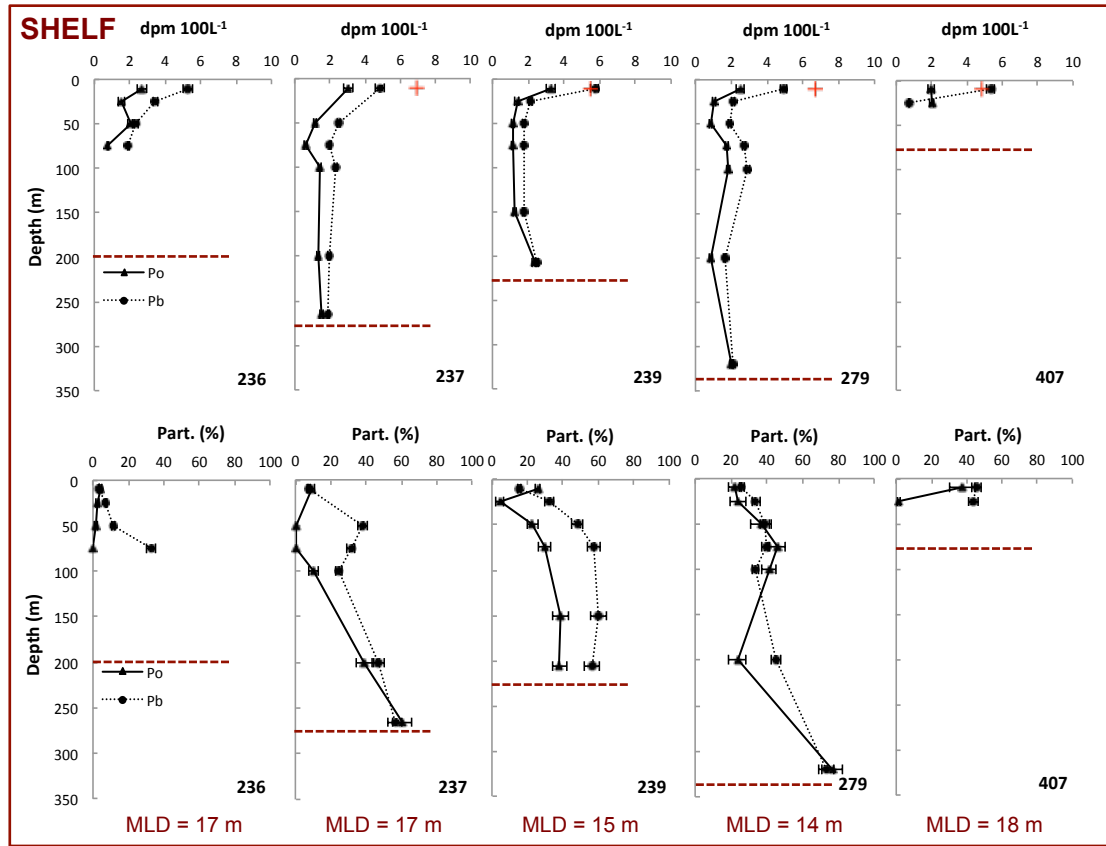
210 **3. Results and discussion**

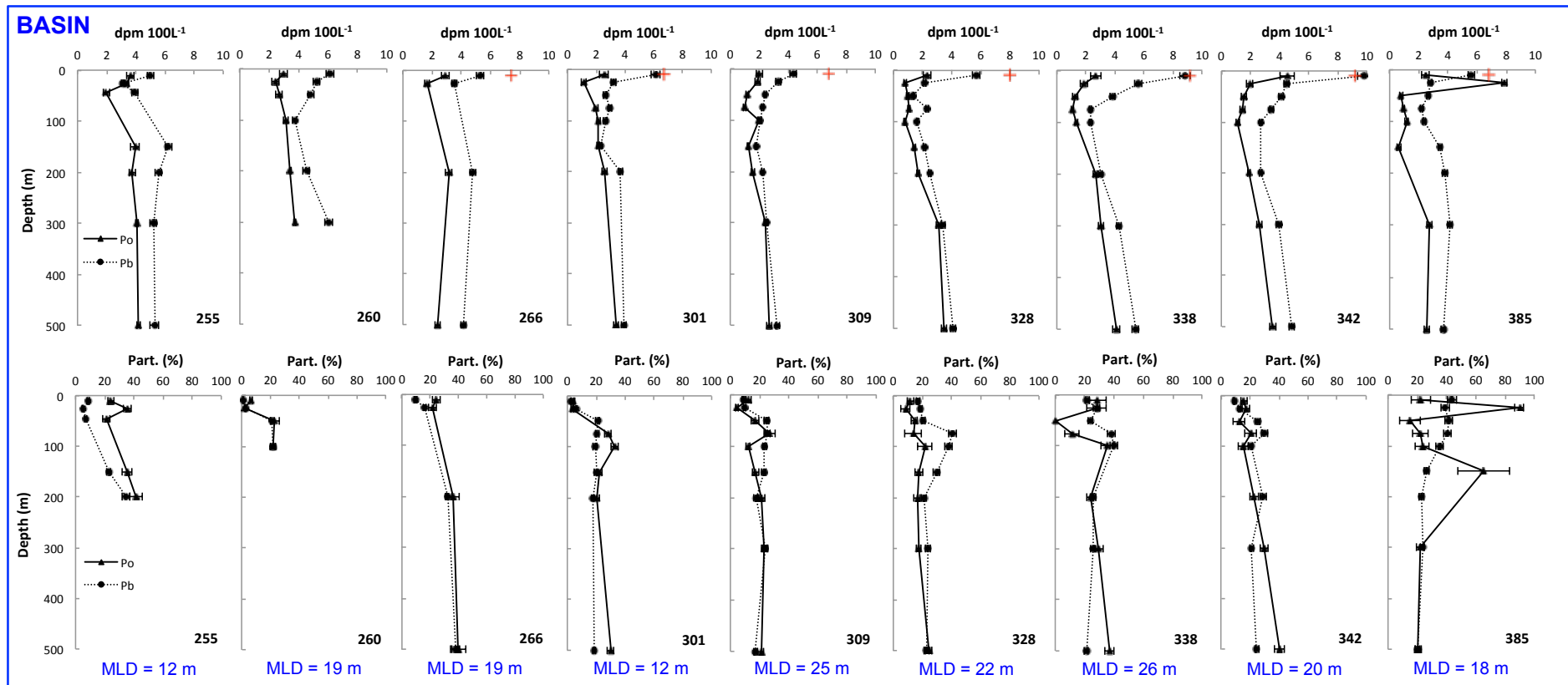
211 First, the activities of both radionuclides and the $^{210}\text{Po}/^{210}\text{Pb}$ ratios in surface waters and
212 vertical profiles are discussed in relation to the area and water masses, and compared to
213 literature data (Section 3.1). Second, the annual new primary production estimates
214 obtained for the basin stations are presented (Section 3.2). Last, total ^{210}Po deficits are
215 discussed in parallel with the annual new primary production estimates and the origin of
216 freshwater in the upper water column (Bauch et al., 2011), and compared to ^{234}Th -derived
217 particle export estimates (Cai et al., 2010) (Section 3.3).

218 Characteristics of the study area, including hydrography (see Figure S1), sea-ice
219 conditions and nutrient regime can be found in Supplemental Material.

220 **3.1. ^{210}Pb and ^{210}Po in seawater**

221 The activities of ^{210}Pb and ^{210}Po and the $^{210}\text{Po}/^{210}\text{Pb}$ ratios in surface waters (10 m) and
222 vertical profiles are presented in Figures 2-5 and in Supplemental Material. Vertical
223 profiles were taken down to 25 to 320 m (shelf), 650 to 1015 m (slope), and 1000 to 4365
224 m (basin).





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Figure 2: Vertical activity profiles for total ^{210}Po (solid line) and ^{210}Pb (dotted line) and relative contribution of the particulate activities of both radionuclides for the upper 500 m of the water column in the shelf (<350 m), slope (350-1050 m) and basin (>1050 m) environments. The red cross indicates the total activity of ^{226}Ra in surface waters (Rutgers van der Loeff et al., 2012) for comparison with ^{210}Pb activity (see Section 3.1.1). Notice the different scale on the y-axis between the shelf and the slope/basin profiles. The horizontal dashed line in the shelf panel indicates the bottom depth. The mixed layer depth (MLD) for each profile is given at the bottom of the panels. Station 411 is not shown, since the total and particulate activities of ^{210}Pb and ^{210}Po could only be determined at one investigated depth (Table S1, Supplemental Material).

232 3.1.1. Total ^{210}Pb and ^{210}Po activities and $^{210}\text{Po}/^{210}\text{Pb}$ ratios

233 Total activities of ^{210}Pb and ^{210}Po in the mixed layer (including data from surface stations
234 and vertical profiles) were, on average, 5.8 ± 1.6 dpm 100L^{-1} ($n = 39$) and 2.6 ± 0.9 dpm
235 100L^{-1} ($n = 38$), respectively. The average ^{210}Pb and ^{210}Po activities found below the
236 mixed layer depth in the vertical profiles (3.5 ± 1.7 dpm 100L^{-1} for ^{210}Pb , $n = 123$; and
237 2.6 ± 1.7 dpm 100L^{-1} for ^{210}Po , $n = 120$), show that the mixed layer was enriched in ^{210}Pb
238 with respect to underlying waters.

239 The ocean surface usually presents higher activities of ^{210}Pb with respect to its parent,
240 ^{226}Ra , due to the atmospheric deposition of ^{210}Pb produced from the decay of ^{222}Rn ,
241 representing the major input of ^{210}Pb into the ocean surface (e.g. Nozaki et al., 1980).
242 Considering surface ^{226}Ra data from the same expedition (range: 3.3 to 9.5 dpm 100L^{-1} ;
243 Rutgers van der Loeff et al., 2012) and assuming an atmospheric depositional flux of
244 ^{210}Pb of 0.06 dpm $\text{cm}^{-2} \text{y}^{-1}$ in the Arctic (Huh et al., 1997) the mean residence time of
245 ^{210}Pb in surface waters is 2.0 ± 0.8 years (range: 1.1 to 4.2 years), without significant
246 differences between the shelf and basin environments (Wilcoxon test, $p > 0.05$). This
247 residence time falls in the upper range of previous estimates from diverse oceanic regions
248 including Arctic locations (0.1 to 2.5 years; see references in Masqué et al., 2002 and
249 Smith et al., 2003). Yet, this estimate is very sensitive to the value chosen for the
250 atmospheric flux and does not take into account the role of the sea-ice cover, which
251 intercepts and accumulates a fraction of the atmospheric fluxes during sea-ice transit,
252 releasing them where and when melting occurs (Cámara-Mor et al., 2011; Chen et al.,
253 2012; Masqué et al., 2007; Roberts et al., 1997).

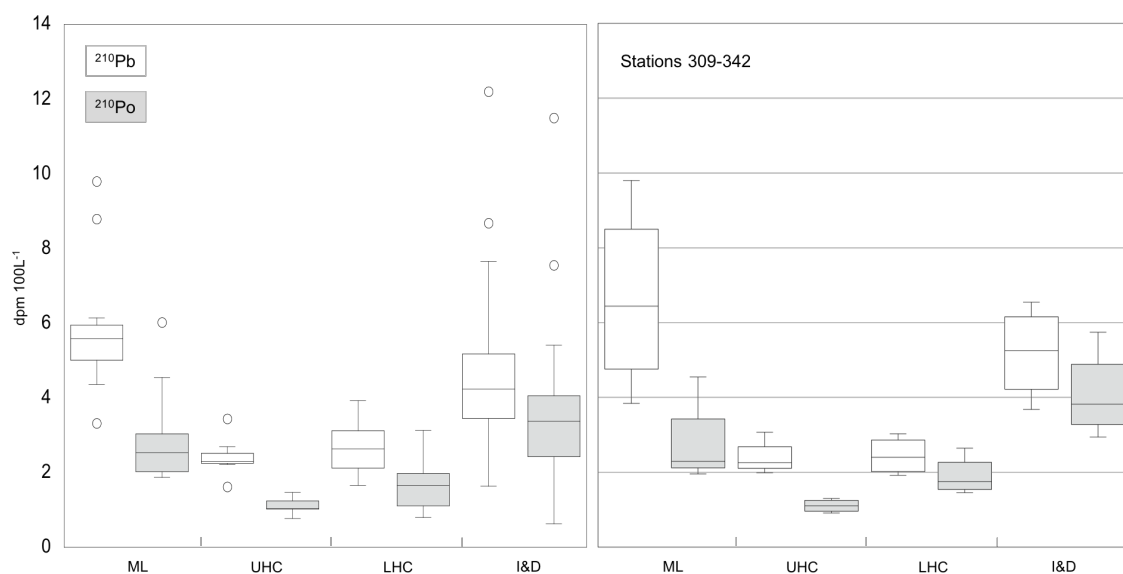
254 Whereas in surface waters of ice-free areas of the world ocean $^{210}\text{Pb}/^{226}\text{Ra}$ ratios are
255 usually larger than 1 as a result of atmospheric deposition, in our study these ratios were
256 >1.0 only at some stations (Table S1, Supplemental Material and Figure 2), where
257 significant amounts of sea-ice meltwater were found (2-8%; Bauch et al., 2011). These
258 stations were located near the ice edge on the Barents and Laptev shelves (239, 407 and
259 409), and under the sea-ice cover in the Nansen and Makarov Basins (303 and 342).
260 Indeed, considering all stations, there was a positive relationship between the fraction of
261 sea-ice meltwater (Bauch et al., 2011) and $^{210}\text{Pb}/^{226}\text{Ra}$ ratios in surface waters ($p < 0.01$;
262 Spearman correlation coefficient, $\rho = 0.63-0.67$ for the two approaches used by Bauch et

263 al., 2011, n = 20-22; not shown). This suggests a role of sea ice in regulating the amount
264 of atmospherically-derived ^{210}Pb in surface waters, preventing its input where sea ice is
265 present and enhancing ^{210}Pb activities when it melts through the release of accumulated
266 ^{210}Pb in sea ice and/or the direct input of atmospheric ^{210}Pb to seawater. Here, we cannot
267 quantify the effect of sea-ice melt on driving the enrichment of ^{210}Pb in surface waters
268 without considering the concentration of ^{210}Pb in sea ice and its removal by scavenging
269 once released into the ocean. In the case of ^{210}Po , its atmospheric flux to the ocean surface
270 usually accounts for only about 10% of that of ^{210}Pb (Lambert et al., 1982; Baskaran,
271 2011). However, in the Arctic, the inputs of ^{210}Po to the ocean surface due to sea-ice melt
272 could lead to greater $^{210}\text{Po}/^{210}\text{Pb}$ ratios resulting from the ingrowth of ^{210}Po from ^{210}Pb
273 decay in sea ice (range: 0.4-1.0 in sea-ice cores; Masqué et al., 2007; Roca-Martí et al.,
274 2016). Then, a preferential removal of ^{210}Po over ^{210}Pb by particle export in surface waters
275 (Nozaki et al., 1997) may explain the non-significant enrichment of ^{210}Po in the mixed
276 layer.

277 In the upper halocline (stations 309, 328, 338 and 342), total ^{210}Pb and ^{210}Po activities
278 were on average 2.4 ± 0.6 dpm 100L^{-1} and 1.1 ± 0.2 dpm 100L^{-1} (n = 7), respectively,
279 similarly to the lower halocline, where the activities averaged 2.7 ± 0.7 dpm 100L^{-1} and
280 1.6 ± 0.7 dpm 100L^{-1} (n = 17), respectively. These values are low in comparison with
281 overlying and intermediate/deep waters (Figure 3). Low total activities of ^{210}Pb and ^{210}Po
282 in the upper halocline were firstly reported for the Canadian Expedition to Study the
283 Alpha Ridge (CESAR; Moore and Smith, 1986), and later for the Makarov and Canada
284 Basins for both ^{210}Pb and ^{210}Po (Smith et al., 2003) or only for ^{210}Pb (Hu et al., 2014;
285 Lepore et al., 2009). This observation was explained by particle scavenging of ^{210}Pb and
286 ^{210}Po at the sediment-water interface over the Chukchi and Beaufort shelves, where the
287 upper halocline is formed, and subsequent advective transport into the interior Arctic (e.g.
288 Rutgers van der Loeff et al., 2012).

289 Over the Alpha Ridge, the mean ^{210}Pb and ^{210}Po activities in the upper halocline ($2.7 \pm$
290 0.5 dpm 100L^{-1} and 1.2 ± 0.2 dpm 100L^{-1} , respectively, n = 4, stations 338 and 342) are
291 found to be significantly higher than at the CESAR site (0.75 ± 0.10 dpm 100L^{-1} and 0.52
292 ± 0.05 dpm 100L^{-1} , respectively; Moore and Smith, 1986; see location in Figure S2). This
293 difference probably derives from temporal variability in the Pacific Water pathways,

294 which are partly controlled by atmospheric circulation patterns (Morison et al., 2012;
 295 Steele et al., 2004; Timmermans et al., 2014).



296

297 **Figure 3:** Box plots for total ^{210}Pb and ^{210}Po activities in the mixed layer (ML: surface to MLD), upper
 298 halocline (UHC), lower halocline (LHC) and intermediate and deep waters (I&D) obtained from all the
 299 vertical profiles (left panel) or only those from stations 309, 328, 338 and 342 (right panel) that showed all
 300 water layers (ML, UHC, LHC and I&D). The bottom and top of the boxes mark the 25th and 75th
 301 percentiles, respectively, and the middle line represents the median (50th percentile). The lines extending
 302 from the bottom and top of the boxes mark the minimum and maximum values. Outliers are displayed as
 303 empty circles. Number of samples considered: ML $n = 17$; UHC $n = 7$; LHC $n = 17$; I&D $n = 64$ (^{210}Po) or
 304 65 (^{210}Pb) (left panel); and ML $n = 6$; UHC $n = 7$; LHC $n = 5$; I&D $n = 21$ (right panel).

305 The lower halocline, in contrast to the upper halocline, is a common feature in the Arctic
 306 Ocean, with origin in the Barents Sea or the Nansen Basin (Rudels, 2009). Smith et al.
 307 (2003) reported lower ^{210}Pb activities in lower halocline waters formed in the Barents Sea
 308 compared to those formed by haline convection in the Nansen Basin. This difference was
 309 ascribed to enhanced particle scavenging in the productive and particle-rich Barents Sea,
 310 with respect to the Nansen Basin, and removal during boundary current transport into the
 311 central Arctic.

312 In summer 2007, ^{210}Pb activities in the lower halocline were higher at the stations located
 313 north of the Barents Sea and Franz Josef Land (3.5 ± 0.3 dpm 100L^{-1} , stations 255, 260,
 314 266 and 276) than at stations located further east in the Eurasian sector and the Makarov
 315 Basin (2.3 ± 0.5 dpm 100L^{-1} , stations 279, 301, 309, 328, 338, 342, 385 and 400) by a
 316 factor of 1.5 (Wilcoxon test, $p < 0.01$). Temperature and salinity profiles indicate a
 317 potential shelf influence in the lower halocline from station 300 onwards. Yet, we cannot
 318 confirm this without data from previous years, since the advective time scales of the lower

319 halocline from its interaction with the shelves to the central Arctic may be more than a
320 few years (mean of 9.6 ± 4.6 years in Ekwurzel et al., 2001). On the other hand, Laptev
321 shelf waters had an influence on the continental slope driven by the high polynya activity
322 in the Laptev Sea in April 2007 (Bauch et al., 2010). Brine-enriched waters in the surface
323 regime were detected out of the shelf, which could have weakened stratification and
324 enhanced winter convection into the range of the lower halocline. Indeed, a winter mixed
325 layer deeper than 100 m was found on the continental slope at station 400 and
326 surroundings, indicating an influence from the Laptev shelf. Moreover, the halocline
327 around stations 255-276 is generally not influenced by the Barents Sea (Rudels, 2009;
328 Rudels et al., 2004, 1996). Therefore, all these observations suggest that the lower
329 halocline at stations 279-400 had been more affected by shelf processes than at stations
330 255-276. Thus, enhanced particle scavenging during formation and/or transport of the
331 lower halocline to the Makarov and eastern Eurasian Basins would explain the lower
332 activities of ^{210}Pb observed.

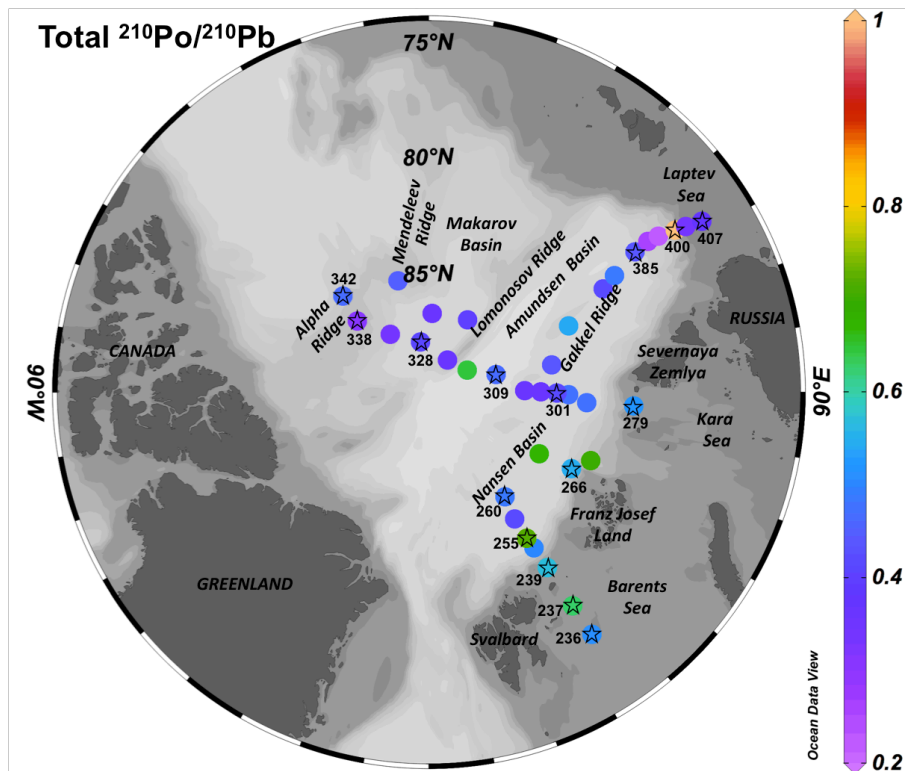
333 In intermediate and deep waters, total activities of ^{210}Pb and ^{210}Po increased to 4.3 ± 1.7
334 dpm 100L^{-1} ($n = 65$) and 3.4 ± 1.6 dpm 100L^{-1} ($n = 64$), respectively. These activities, in
335 the case of ^{210}Pb , were mostly lower than those found in the mixed layer, but, in the case
336 of ^{210}Po , they were generally higher than in overlying waters (Figure 3).

337 Over the Alpha Ridge, good agreement was found between the activities measured at
338 stations 338 and 342 at 1000 m, which averaged 5.0 ± 1.1 dpm 100L^{-1} for ^{210}Pb and 4.06
339 ± 0.09 dpm 100L^{-1} for ^{210}Po , and those found at the CESAR site at the same depth ($3.9 \pm$
340 0.2 dpm 100L^{-1} and 4.3 ± 0.2 dpm 100L^{-1} , respectively; Moore and Smith, 1986). In the
341 Eurasian Basin, the ^{210}Pb and ^{210}Po activities measured in the present study at 300 m were,
342 on average, 5.6 ± 0.6 dpm 100L^{-1} and 3.9 ± 0.3 dpm 100L^{-1} , respectively, in the Nansen
343 Basin (stations 255 and 260), and 2.49 ± 0.11 dpm 100L^{-1} and 2.43 ± 0.12 dpm 100L^{-1} ,
344 respectively, in the Amundsen Basin (station 309). These activities are also comparable
345 to those measured in the same basins and water depth in summer 2012 (Roca-Martí et al.,
346 2016). Based on the inspection of temperature-salinity properties in summer 2007 (not
347 shown), Atlantic Water seems to originate in the Barents Sea inflow branch at station 309,
348 the Fram Strait branch at station 255 or a mixture of the two at station 260. Therefore,
349 besides removal during transit, enhanced particle scavenging in the Barents Sea could
350 also help explain the lower activities of both radionuclides in the Amundsen Basin with

351 respect to those measured at stations affected by the Fram Strait branch, similar to the
352 interpretation by Smith et al. (2003) for the lower halocline.

353 Regarding the total $^{210}\text{Po}/^{210}\text{Pb}$ ratios, significant deficits of ^{210}Po (i.e. ratio <0.90 ,
354 considering uncertainties) were observed throughout most of the water column in the
355 study area. The most substantial deficits of ^{210}Po were found in surface and subsurface
356 waters spanning depths from 10 to 200 m (Figures 2 and 4), covering the mixed layer and
357 the halocline(s): total $^{210}\text{Po}/^{210}\text{Pb}$ ratios averaged 0.46 ± 0.19 in the mixed layer ($n = 38$),
358 0.46 ± 0.14 in the upper halocline ($n = 7$) and 0.6 ± 0.3 in the lower halocline ($n = 17$).
359 Po-210 deficits in the upper water column are usually attributed to biological particle
360 production and subsequent scavenging and export, which will be discussed in Section 3.3.

361 In the upper 25 m, we find good agreement between the mean total $^{210}\text{Po}/^{210}\text{Pb}$ ratio in
362 summer 2007 (0.5 ± 0.2 , $n = 14$; including all profiles except stations 237, 276 and 411
363 for lack of data) and that measured in the Eurasian Basin in summer 2012 (0.66 ± 0.19 , n
364 $= 7$; Roca-Martí et al., 2016). These results are also consistent with those obtained at
365 single depths (10-20 m) from the Chukchi Sea to the Mendeleev Ridge in summer 1994
366 (0.6 ± 0.3 , $n = 6$; Smith et al., 2003). Total $^{210}\text{Po}/^{210}\text{Pb}$ ratios of 0.50 to 0.60 correspond
367 to a ^{210}Po residence time of 7 to 10 months based on a steady-state balance and the
368 assumption of negligible atmospheric flux of ^{210}Po (Nozaki et al., 1998). Although the
369 rather low biological productivity of the Arctic ice-covered waters, this estimate
370 compares well with the residence time obtained for the mixed layer in the Atlantic, Pacific
371 and Indian Oceans (~ 7 months; Bacon et al., 1976; Cochran et al., 1983; Nozaki et al.,
372 1976; Shannon et al., 1970). Similarly, the mean total $^{210}\text{Po}/^{210}\text{Pb}$ ratio in the upper 200
373 m was 0.6 ± 0.3 ($n = 12$; including all profiles except stations 260, 266, 276, 407 and 411
374 for lack of data). This result is lower than that found in summer 2012 (0.9 ± 0.6 , $n = 7$;
375 Roca-Martí et al., 2016) and at the CESAR site in spring 1983 (0.96 ± 0.04 , $n = 1$; Moore
376 and Smith, 1986).



377

378 **Figure 4:** Total $^{210}\text{Po}/^{210}\text{Pb}$ activity ratios in surface waters (10 m). Stars indicate the stations sampled for
 379 vertical profiles (station numbers shown).

380 Apart from the ^{210}Po deficits found in the upper ocean, they were also detected in the
 381 mesopelagic and bathypelagic zones. Total $^{210}\text{Po}/^{210}\text{Pb}$ ratios averaged 0.8 ± 0.5 in
 382 intermediate and deep waters ($n = 64$), where equilibrium between both radionuclides
 383 was only approached in one-third of the instances (ratio >0.90 , Table S1, Supplemental
 384 Material). The magnitude of these ^{210}Po deficits at depth could indicate that they do not
 385 solely reflect the settling of particles to the deep ocean. Considering the previous
 386 literature, mean total $^{210}\text{Po}/^{210}\text{Pb}$ ratios were near equilibrium in the Eurasian Basin (1.0
 387 ± 0.5 , $n = 14$, depth range: 300-400 m; Roca-Martí et al., 2016) and at the CESAR site
 388 (0.9 ± 0.2 , $n = 3$, depth range: 800-1200 m; Moore and Smith, 1986), but were
 389 significantly far from equilibrium during the Arctic Ocean Section (0.6 ± 0.4 , $n = 17$,
 390 depth range: 300-4200 m; Smith et al., 2003) that went from the Chukchi Sea to the Fram
 391 Strait (Figure S2). A deficiency of ^{210}Po in the ocean interior has also been observed in
 392 other oceanic regions (e.g. Kim and Church, 2001; Nozaki et al., 1997, 1990; Sarin et al.,
 393 1994; Thomson and Turekian, 1976), but its origin is still unclear. Hence, while this study
 394 cannot elucidate the reasons for these deficits, the interpretation of the $^{210}\text{Po}/^{210}\text{Pb}$ pair at
 395 depth remains uncertain until further investigations are made.

396

397 **3.1.2. Particulate $^{210}\text{Po}/^{210}\text{Pb}$ ratios**

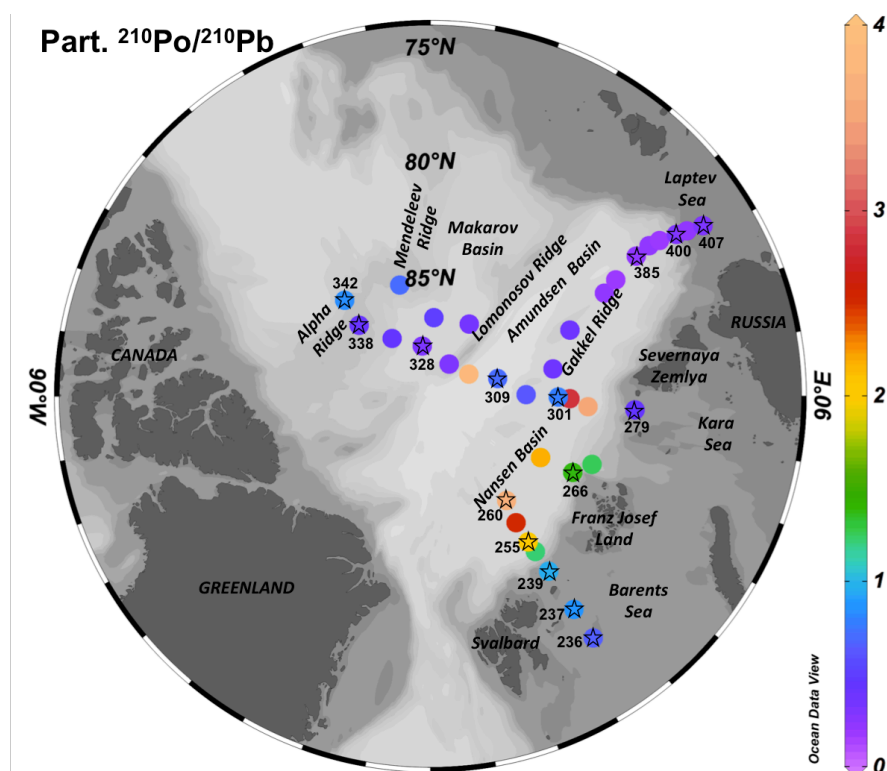
398 Particulate ^{210}Pb and ^{210}Po activities can be found in the Supplemental Material.

399 Particulate $^{210}\text{Po}/^{210}\text{Pb}$ ratios averaged 0.5 ± 0.5 in the mixed layer ($n = 38$), 0.30 ± 0.18
400 in the upper halocline ($n = 7$), 0.5 ± 0.3 in the lower halocline ($n = 19$), and 0.8 ± 0.5 in
401 intermediate waters down to 500 m ($n = 28$). Only 15% of the samples showed ^{210}Po
402 enrichment (i.e. particulate $^{210}\text{Po}/^{210}\text{Pb}$ ratio >1.10), which were mainly located in the
403 Nansen Basin (up to 4-8 at 10-25 m; Figure 5) around the area where large deficits of
404 ^{210}Po were found (see Section 3.3). The two studies that have previously reported
405 particulate ^{210}Pb and ^{210}Po activities in the Arctic Ocean have also presented $^{210}\text{Po}/^{210}\text{Pb}$
406 ratios lower than 1 for particles $\geq 0.45 \mu\text{m}$ (Moore and Smith, 1986) and $\geq 53 \mu\text{m}$ (Roca-
407 Martí et al., 2016) at depths from 25 to 800 m over the Alpha Ridge and the Eurasian
408 Basin. The overall low ratios obtained seem to be inconsistent with the preferential
409 removal of ^{210}Po by particle export shown by the widespread deficiency of ^{210}Po in the
410 upper ocean in summer 2007 (see Sections 3.1.1 and 3.3).

411 The particles we analysed ($\geq 1 \mu\text{m}$) were obtained from 40-160 L of seawater collected
412 with Niskin bottles, which would basically represent the suspended (or small) particle
413 fraction (Bishop et al., 2012). Indeed, the quantification of ^{210}Pb and ^{210}Po in large
414 particles ($>53\text{-}70 \mu\text{m}$), those that contribute most to particle fluxes (see references in Lam
415 and Marchal, 2015), requires filtration of large volumes, particularly in the central Arctic
416 (several hundreds of liters may not be enough, Roca-Martí et al., 2016). Therefore, we
417 believe that the particulate activities presented here correspond to small, suspended
418 particles rather than rare, large, sinking particles. Yet, these particle pools could have a
419 similar composition by aggregation and disaggregation. Previous studies have shown that
420 the degree of this particle exchange can vary geographically and seasonally in relation to
421 the dominant particle type and plankton community structure (Abramson et al., 2010;
422 Lam and Marchal, 2015). While we do not know to what extent the collected particles
423 would represent the large particles in late summer, particulate $^{210}\text{Po}/^{210}\text{Pb}$ ratios <1
424 indicate that these particles would not represent the sinking particle pool that created the
425 ^{210}Po deficiency in seawater. Hence, these observations suggest that: (i) large particles in
426 the late summer are not well represented in this study, and/or (ii) ^{210}Po export mostly
427 occurred earlier in the season and particle composition changed during spring/summer.

428 This latter possibility is consistent with the conclusions drawn by Roca-Martí et al.
429 (2016).

430 Particle types that could be found during the Arctic late summer that could help explain
431 the measured particulate $^{210}\text{Po}/^{210}\text{Pb}$ ratios <1 include: sea-ice drafted particles enriched
432 in ^{210}Pb via atmospheric input ($^{210}\text{Po}/^{210}\text{Pb}$ ratios <1 in sea-ice sediments; Roca-Martí et
433 al., 2016), material remineralized by chemical and biological processes (Stewart et al.,
434 2007b), faecal pellets (Rodríguez y Baena et al., 2007; Stewart et al., 2005), picoplankton
435 aggregates (Stewart et al., 2010) and substrates rich in transparent exopolymer particles
436 (Quigley et al., 2002). Also, low particulate $^{210}\text{Po}/^{210}\text{Pb}$ ratios (≤ 1) in shelf waters could
437 be due to the presence of terrestrial and riverine particles (Nozaki et al., 1998; Tateda et
438 al., 2003). Sampling of size-fractionated particles, ideally at different times during the
439 growing season, coupled with particle composition analyses should be carried out in
440 future works to infer what controls the particulate $^{210}\text{Po}/^{210}\text{Pb}$ ratios in the Arctic and,
441 specifically, those lower than 1.



442

443 **Figure 5:** Particulate $^{210}\text{Po}/^{210}\text{Pb}$ activity ratios in surface waters (10 m). Stars indicate the stations sampled
444 for vertical profiles (station numbers shown).

445

3.2. Annual new primary production

446 The annual new primary production estimates in the Nansen, Amundsen and Makarov
447 Basins are presented in Table 2. The nitrogen consumed above the winter mixed layer
448 depth during the productive season was on average $90 \pm 40 \text{ mmol m}^{-2}$ in the Eurasian
449 sector (stations 255, 260, 266, 301, 309 and 385), whereas it was $250 \pm 90 \text{ mmol m}^{-2}$ in
450 the Canadian sector (stations 328, 338 and 342). Annual nitrogen-derived primary
451 production estimates were similar across the Nansen and Amundsen Basins, with an
452 average of $7 \pm 3 \text{ g C m}^{-2} \text{ yr}^{-1}$, although it was about four times lower north of Franz Josef
453 Land (station 266). In the Makarov Basin, annual new primary production was always
454 higher than in the Eurasian Basin, averaging $20 \pm 7 \text{ g C m}^{-2} \text{ yr}^{-1}$. Phosphate-derived
455 estimates were similar to those derived from nitrogen (<6% difference) at most of the
456 stations. Ulfsbo et al. (2014) also found higher estimates in the Makarov Basin,
457 particularly over the Mendeleev Ridge, when compared to the other basins. Globally, the
458 annual new primary production was on average $12 \text{ g C m}^{-2} \text{ yr}^{-1}$ considering all the
459 nitrogen- and phosphate-derived estimates, which is similar to reported estimates of about
460 $9 \text{ g C m}^{-2} \text{ yr}^{-1}$ including values from the Nansen, Amundsen, Makarov and Canada Basins
461 during years 2011 and 2012 (Fernández-Méndez et al., 2015; Ulfsbo et al., 2014).

462 Silicate-derived new primary production estimates were on average $2.3 \pm 1.9 \text{ g C m}^{-2} \text{ yr}^{-1}$
463 in the Eurasian sector, whereas they averaged $32 \pm 17 \text{ g C m}^{-2} \text{ yr}^{-1}$ in the Makarov Basin.
464 Comparison between silicate-derived new primary production with the other two
465 estimates gives an indication of the contribution of diatoms to new production. Diatom
466 production estimates varied widely across the study area, amounting between 10 to 70%
467 of annual new primary production in the Eurasian sector and about 100% in the Makarov
468 Basin. This suggests that diatom production had an important role in the Canadian sector
469 of the study area during the productive season in 2007. Yet, we note that the higher
470 silicate-derived new primary production estimates obtained in the Makarov Basin
471 compared to those from nitrogen and phosphate indicate that the former estimates may
472 be overestimated. This bias can be due to the advected silicate maximum in the upper
473 halocline and uncertainties in determining silicate concentrations in the winter mixed
474 layer, together with the variability in N/Si ratios reported for Arctic diatoms (Fernández-
475 Méndez et al., 2015). Therefore, silicate-derived new primary production estimates in the
476 Makarov Basin should be taken with caution.

477 **Table 2:** Nitrite+nitrate, phosphate and silicate drawdown above the winter mixed layer depth (WMLD)
478 and annual new primary production estimated using the Redfield-Brzezinski ratio (106C:16N:15Si:1P).

479

Station	WMLD (m)	Nutrient deficits (mmol m ⁻²)			Annual new production (g C m ⁻² yr ⁻¹)		
		Nitrite+nitrate	Phosphate	Silicate	Nitrite+nitrate	Phosphate	Silicate
255	38	86	5.0	52	6.8	6.4	4.4
260 ^a	81	88	4.9	17	7.0	6.2	1.5
266	22	26	1.6	11	2.0	2.0	0.91
301	60	120	7.3	15	9.8	9.3	1.3
309	59	130	8.4	59	10	11	5.0
328	54	190	16	210	15	21	18
338	60	210	13	310	17	17	27
342	67	350	30	590	28	38	50
385 ^a	50	88	5.9	10	7.0	7.6	0.86

480

^a Different casts were taken for nutrient deficits and WMLD determination.

481

3.3. Particle export in the Arctic as revealed by ²¹⁰Po deficits

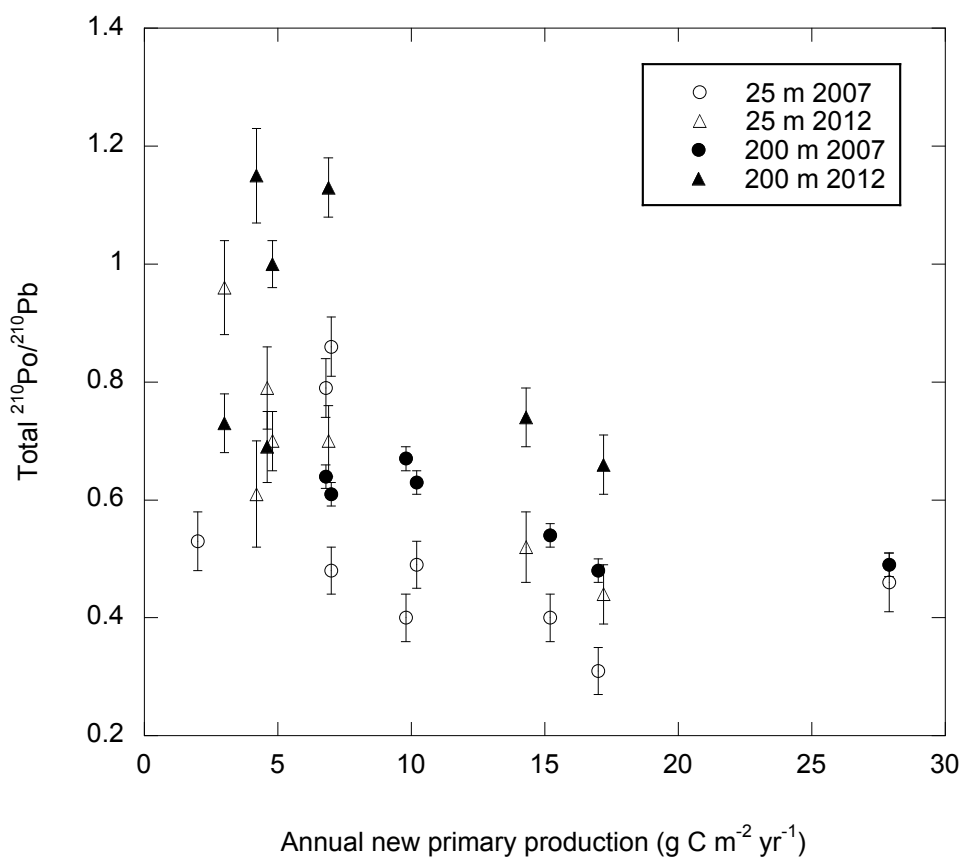
482 On the Arctic shelves visited in late summer 2007, Cai et al. (2010) found ²³⁴Th deficits
 483 in the upper 100 m of the water column in association with enhanced Chl-a
 484 concentrations, indicating that in situ production and export of biogenic particles were
 485 the main mechanism for ²³⁴Th removal. Nevertheless, we did not observe significant
 486 correlations between Chl-a concentrations and ²¹⁰Po deficits (not shown). This may not
 487 be surprising given that Chl-a concentration is a snapshot of the sampling time (i.e. late
 488 summer), whereas ²¹⁰Po integrates a time scale of several months, covering, in this case,
 489 from the onset of the growing season, which occurs as early as March (Wassmann and
 490 Reigstad, 2011). Satellite images of the ice-free area in the Barents Sea revealed five-fold
 491 higher Chl-a concentrations in May-June with respect to those in July-August in 2007,
 492 showing that the sampling was conducted in a post-bloom situation (Klunder et al., 2012).
 493 Indeed, the concentrations of nutrients in the mixed layer were low during the expedition
 494 (see details in Supplemental Material).

495 Seasonal estimates of primary production can be helpful to obtain a wider perspective on
 496 production of organic matter than that at the time of sampling. However, the method used
 497 here to estimate annual new primary production does not allow obtaining reliable results
 498 in the shelf environment, where the advective input of nutrients could significantly bias
 499 the estimates. Therefore, we cannot provide evidence of the relation between ²¹⁰Po
 500 deficits and annual new primary production (and subsequent export) in the Barents, Kara
 501 and Laptev Seas.

502 As described in Section 3.1.1, ²¹⁰Po deficits were not restricted to the shelf environment,
 503 but they were also pronounced in the central Arctic, without showing any gradient from

504 the shelves to the central Arctic (Figures 2 and 4). During the expedition, the basins were
505 characterized by generally low Chl-a levels and very low particle export, as reported by
506 Cai et al. (2010) using ^{234}Th . This apparent discrepancy between ^{234}Th and ^{210}Po may be
507 explained by significant export fluxes that occurred more than one month before
508 sampling, which would be recorded by ^{210}Po but missed by ^{234}Th . In this line, Rutgers
509 van der Loeff et al. (2012) argued that the ^{234}Th -based scavenging rate could not explain
510 the distribution of ^{228}Th in the central Arctic in summer 2007 due to seasonal variations
511 in scavenging and the different half-lives of ^{234}Th and ^{228}Th . In the following discussion,
512 we explore whether the results of ^{210}Po and ^{234}Th indicate a seasonal variation in particle
513 export by comparing the deficits of ^{210}Po with the estimates of annual new primary
514 production obtained from the Nansen, Amundsen and Makarov Basins.

515 Although the number of data points is small, we find a negative relationship between
516 annual new primary production estimates based on nitrogen consumption and total
517 $^{210}\text{Po}/^{210}\text{Pb}$ ratios in the upper 25 and 200 m of the basin stations ($p < 0.05$ for each depth;
518 25 m: Spearman correlation coefficient, $\rho = -0.72$, $n = 9$; 200 m: $\rho = -0.79$, $n = 7$; Figure
519 6). This suggests that greater deficits of ^{210}Po in the upper water column of the Arctic
520 were related to higher in situ new production and subsequent export of biogenic material
521 to depth. This would be analogous to the situation reported for the Eurasian Basin in 2012,
522 based on $^{210}\text{Po}/^{210}\text{Pb}$ and nutrient sampling also conducted during the late summer (from
523 11 August to 22 September; Roca-Martí et al., 2016). Indeed, if we combine the results
524 from both studies, the significance of the observed relationship increases ($p < 0.01$ for
525 each depth; 25 m: Spearman correlation coefficient, $\rho = -0.73$, $n = 16$; 200 m: $\rho = -0.68$,
526 $n = 14$; Figure 6). This shows the potential use of the $^{210}\text{Po}/^{210}\text{Pb}$ proxy as an indicator of
527 annual new primary production in the central Arctic, although further investigations will
528 be necessary.



529

530 **Figure 6** Total ²¹⁰Po/²¹⁰Pb activity ratios in the upper 25 m (white) and 200 m (black) vs nitrogen-derived
 531 new primary production estimates that encompass the Arctic growing season in summers 2007 (circles; this
 532 study) and 2012 (triangles; Roca-Martí et al., 2016).

533 Total ²¹⁰Po/²¹⁰Pb ratios were lower for waters in the salinity range from about 27 to 34,
 534 covering the mixed layer and the upper halocline (basically confined in the Canadian
 535 sector), than for higher salinities and, hence, deeper waters. The lower salinity waters
 536 were influenced by Pacific Water or freshwater inputs from river runoff and local
 537 precipitation (jointly referred to as river water) and net sea-ice melting (see details in
 538 Bauch et al., 2011; Roeske et al., 2012; Rutgers van der Loeff et al., 2012). The
 539 contribution of these water sources to surface and subsurface waters is discussed below
 540 in relation to the total ²¹⁰Po/²¹⁰Pb ratios measured in each area in order to investigate the
 541 processes that drove the ²¹⁰Po deficits in 2007.

542 In the Makarov Basin, ²¹⁰Po/²¹⁰Pb ratios were especially low with values generally <0.50
 543 for waters extending down to the upper halocline. Over the Alpha Ridge, sea-ice
 544 meltwater amounted to 1 to 2% of surface waters at stations 338 and 342 (Bauch et al.,
 545 2011; Rutgers van der Loeff et al., 2012). Roca-Martí et al. (2016) showed that a complete
 546 melting of sea ice in the central Arctic would change the total ²¹⁰Po/²¹⁰Pb ratios by less

547 than 10% in surface waters. Therefore, the substantial deficits of ^{210}Po observed would
548 not be explained by the input of sea-ice meltwater depleted in ^{210}Po with respect to ^{210}Pb .
549 On the other hand, the upper 100 m of stations 328, 338 and 342 were composed of 5 to
550 15% of river water and a large fraction of Pacific Water (up to >80%; Bauch et al., 2011).
551 Seawater with salinities of ~30 to 32 (part of the Polar Mixed Layer) found in the central
552 Arctic, especially near the Lomonosov Ridge (stations 320 and 328), originated from the
553 bottom of the Laptev Sea (Bauch et al., 2011), whereas waters with salinities of 32.5 to
554 33.3 (part of the upper halocline) found over the Alpha and Mendeleev Ridges came from
555 the Chukchi Sea (Roeske et al., 2012). These waters from the Laptev and Chukchi Seas
556 carried river and Pacific-derived water, respectively, and were transported to the central
557 Arctic by the Transpolar Drift. Thus, a hypothesis would be that the deficiency of ^{210}Po
558 observed in the upper water column of the Makarov Basin was created in the shelf regime
559 as a consequence of biological activity and particle export. This possibility would be
560 consistent with a transit time of less than three months, comparing the most substantial
561 deficits of ^{210}Po that occurred near the continental margin to the deficits measured over
562 the Alpha Ridge ($^{210}\text{Po}/^{210}\text{Pb}$ ratios of ~0.2 vs. ~0.4, respectively). However, it was
563 estimated that the shelf waters encountered at the surface of the Makarov Basin and the
564 Lomonosov Ridge in summer 2007 travelled for at least seven months (Rutgers van der
565 Loeff et al., 2018).

566 Another possibility could be that ^{210}Po deficits in the Makarov Basin originated in situ.
567 During the productive season in 2007, the average annual new primary production in the
568 Makarov Basin was higher than in the Eurasian Basin by a factor of 3, and diatoms
569 appeared to dominate the phytoplankton community (see Section 3.2). Consistent with
570 this, Middag et al. (2009) measured minimum concentrations of aluminium in the upper
571 300 m of the Makarov Basin during the same expedition, which were related to biological
572 uptake favoured by silica inputs from the upper halocline. Particularly, at stations 338
573 and 342, nitrate was depleted from the surface to 50 m depth. The mixed layer depth at
574 these stations decreased from 60-67 m in winter to 20-26 m at the sampling time and,
575 therefore, the uptake of nitrate had to take place early in the productive season when the
576 seasonal mixed layer was deep enough. This would most likely have occurred before July,
577 since from May surface warming and sea-ice melt establish stratification and, thus, limit
578 the mixing of nutrients within the winter mixed layer (Kawaguchi et al., 2012; Korhonen
579 et al., 2013). Therefore, the ^{210}Po deficiency found in the Makarov Basin likely originated

580 in situ by means of enhanced new primary production and subsequent export. Over the
581 Alpha Ridge, in particular, this might have been related to ice-algae export resulting from
582 sea-ice melt, as observed in the Eurasian Basin during the record sea-ice minimum in
583 2012. In that year, a widespread deposition of sea-ice diatom aggregates was observed on
584 the seafloor (>3000 m, Boetius et al., 2013) and the largest deposits were found together
585 with the strongest depletion of ^{210}Po in seawater (Roca-Martí et al., 2016).

586 In the Barents Sea and Nansen Basin, the major deficits of ^{210}Po (total $^{210}\text{Po}/^{210}\text{Pb}$ ratios
587 ≤ 0.50) were found in cold and relatively low salinity waters in the upper 75 m of section
588 1 (stations 236, 237, 249, 255, 257 and 260). Most of this section had a significant fraction
589 of sea-ice meltwater in the upper 50 to 100 m ($\sim 2\%$; Bauch et al., 2011). Substantial
590 depletions of ^{234}Th , ^{226}Ra , Ba and dissolved Fe were concurrent with the ice edge (81-
591 82°N), together with particularly high phytoplankton biomass, suggesting the occurrence
592 of an ice-edge bloom (Cai et al., 2010; Klunder et al., 2012; Roeske et al., 2012; Rutgers
593 van der Loeff et al., 2012). As observed for ^{210}Po , ^{234}Th was depleted in the upper 100 m
594 south of the ice edge (Cai et al., 2010), where Chl-a levels were found to be very high
595 two months prior to the sampling ($\sim 5 \text{ mg m}^{-3}$; Klunder et al., 2012). In the nutrient-rich
596 Barents Sea, the association between freshwater from sea-ice melt and ^{210}Po deficits may
597 be explained by enhanced primary production when melting occurs. This would improve
598 light conditions for phytoplankton growth and subsequent particle settling, in accordance
599 with previous studies of particle fluxes conducted in the area (Coppola et al., 2002;
600 Lalande et al., 2008; Wassmann et al., 2004; Wiedmann et al., 2014).

601 In the Laptev section, river water had an important presence in the entire water column
602 of the Laptev Sea and in the upper 25 m offshore (Bauch et al., 2011). The inflow of
603 waters from the Lena River supplies significant amounts of nutrients to the Laptev Sea
604 (e.g. Le Fouest et al., 2013), which may have nourished the high phytoplankton biomass
605 observed on the shelf (Cai et al., 2010). Moreover, a large reduction of the sea-ice cover
606 was observed in the Laptev sector in summer 2007 compared to earlier years (Comiso et
607 al., 2008), and sea-ice meltwater in surface waters was detected from the shelf to station
608 385 (Bauch et al., 2011). This sea-ice decline extended the growing season by 50 to 80
609 days and boosted primary production (Arrigo et al., 2008). The maximum inventory of
610 Chl-a was measured at station 407 (Cai et al., 2010), where a substantial deficit of ^{210}Po
611 was found in surface waters overlying an excess at 25 m (total $^{210}\text{Po}/^{210}\text{Pb}$ ratios of 0.37

612 and 2.8, respectively). This suggests export driven by biogenic particles in surface waters
613 and remineralization or particle disaggregation below, in agreement with the conclusions
614 drawn from Ba, ^{226}Ra and dissolved Fe in the Laptev Sea (Klunder et al., 2012; Roeske
615 et al., 2012; Rutgers van der Loeff et al., 2012). The ^{210}Po deficits found north of the
616 Laptev Sea (stations 371-400) could be related to biogenic fluxes that occurred prior to
617 sampling, as they are significant in this area from June to August (Fahl and Nöthig, 2007;
618 Lalande et al., 2009b). Indeed, sediment traps deployed on the Laptev slope revealed
619 about two-fold higher annual POC export in 2006-2007 relative to 2005-2006 (Lalande
620 et al., 2009a). This was mainly attributed to an increase in POC export during and
621 following sea-ice melt in 2007, with maximum fluxes in July.

622 **4. Conclusions**

623 We investigated the distribution of ^{210}Pb and ^{210}Po in the water column of various seas
624 (Barents, Kara and Laptev) and basins (Nansen, Amundsen and Makarov) of the Arctic
625 Ocean in summer 2007. Total activities of ^{210}Pb and ^{210}Po were minimum in the upper
626 and lower haloclines, especially in the Makarov and eastern Eurasian Basins, at
627 approximately 60-130 m. This is partly ascribed to particle scavenging on the shelves
628 where these water masses are formed, boundary current transport and subsequent
629 transport into the central Arctic.

630 During the sea-ice minimum in 2007, widespread deficits of ^{210}Po in the upper water
631 column were observed all over the Arctic, both on the shelves and in the basins. In the
632 shelf areas of the Barents and Laptev Seas, ^{210}Po deficits were related to elevated
633 phytoplankton biomass and particle export. These deficits were usually associated with
634 sea-ice meltwater and riverine water inputs, which may improve light and/or nutrient
635 conditions for photosynthesis. In the basins, estimates of annual new primary production
636 were negatively correlated to total $^{210}\text{Po}/^{210}\text{Pb}$ ratios in the upper 200 m of the water
637 column, suggesting that in situ production and subsequent export controlled the removal
638 of ^{210}Po also in this environment. This shows the potential use of the $^{210}\text{Po}/^{210}\text{Pb}$ proxy as
639 an indicator of annual new primary production in the central Arctic, although more data
640 will be necessary. The ^{210}Po deficits were most substantial (total $^{210}\text{Po}/^{210}\text{Pb}$ ratios <0.50)
641 in the mixed layer and the upper halocline of the Makarov Basin, where estimates of
642 annual new primary production were found to be particularly high.

643 Unlike ^{210}Po , ^{234}Th deficits were very small and not significant below 25 m over the
644 basins in August-September 2007 (Cai et al., 2010). Given the shorter half-life of ^{234}Th ,
645 this indicates that particle export fluxes in the Arctic basins would have been higher
646 before July-August than later in the summer, in line with the conclusions drawn by Roca-
647 Martí et al. (2016) during the record sea-ice minimum in 2012.

648 Deficits of total ^{210}Po in intermediate and deep waters (approx. 200-3000 m) were
649 frequently observed, as well as particulate $^{210}\text{Po}/^{210}\text{Pb}$ ratios below 1, although the reasons
650 are not clear. Sampling of size-fractionated particles in various periods of the growing
651 season together with analyses of particle composition shall be helpful to better understand
652 the biogeochemical behaviour of these radionuclides.

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