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1 **Microplastics in sub-surface waters of the Arctic Central Basin**

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11

12 **Abstract**

13 Polar oceans, though remote in location, are not immune to the accumulation of plastic debris. The  
14 present study, investigated for the first time, the abundance, distribution and composition of  
15 microplastics in sub-surface waters of the Arctic Central Basin. Microplastic sampling was carried  
16 out using the bow water system of icebreaker Oden (single depth: 8.5 m) and CTD rosette sampler  
17 (multiple depths: 8 – 4369 m). Potential microplastics were isolated and analyzed using Fourier  
18 Transform Infrared Spectroscopy (FT-IR). Bow water sampling revealed that the median  
19 microplastic abundance in near surface waters of the Polar Mixed Layer (PML) was 0.7 particles  
20 m<sup>-3</sup>. Regarding the vertical distribution of microplastics in the ACB, microplastic abundance  
21 (particles m<sup>-3</sup>) in the different water masses was as follows: Polar Mixed Layer (0 - 375) > Deep  
22 and bottom waters (0 – 104) > Atlantic water (0 – 95) > Halocline i.e. Atlantic or Pacific (0 – 83).

23

24 **Keywords**

25 Microplastic, Marine debris, Arctic Ocean, Sub-surface waters, Pollution, Water column

26

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29

30 **Introduction**

31 The Arctic Ocean, though the smallest in the world, is unique due to its distinct abiotic features  
32 and the highly specialised ecosystem it supports. Key anthropogenic drivers which may put  
33 pressure on this ecosystem include (i) climate change, (ii) harvest and fisheries, (iii) persistent,  
34 bio-accumulative and toxic contaminants, (iv) industrial development, (v) shipping, and (vi)  
35 invasive alien species (CAFF 2017). Plastic contaminants in the world's oceans have emerged as  
36 an issue of global importance due to their ubiquitous distribution, long-range transport potential,  
37 persistence and perhaps most importantly the potential threat they pose to marine organisms  
38 (UNEP 2011). Remote polar oceans such as the Arctic Ocean have not been immune to the entry  
39 of plastics as a combination of long-range transport processes and local anthropogenic activities  
40 have contributed to the plastic debris in these areas.

41

42 Characteristic abiotic features which set the Arctic Ocean apart from other oceanic basins include  
43 (i) a central area of perennial pack ice, (ii) seasonal extremes in solar irradiance, ice and snow  
44 cover, temperature and riverine inflow, and (iii) an upper layer of lower salinity water due to  
45 freshwater input from rivers and seasonal sea-ice melt (CAFF 2013). This unique ecosystem is a  
46 habitat for a vast array of marine organisms, some of which are (i) endemic to the region, (ii)

47 commercially important, (iii) apex predators, (iv) central to the functioning of the ecosystem, and  
48 (v) threatened as evidenced by their inclusion in the IUCN Red List of Threatened Species (CAFF  
49 2013, CAFF 2017).

50

51 Despite its remote location away from major population centres and the low coastal population in  
52 its surrounding shelf areas, both macro and microplastics were detected in the various  
53 environmental compartments of the Arctic Ocean. Between 2002 and 2014, macroplastics were  
54 detected on the seafloor (2500 m depth) of the eastern Fram Strait at the HAUSGARTEN  
55 observatory (Bergmann and Klages 2012; Tekman et al. 2017). Sightings of buoyant macroplastics  
56 were also made during ship and helicopter observation surveys in the Barents Sea and Fram Strait  
57 (Bergmann et al. 2016). A citizen-science study also recently reported the presence of  
58 macroplastics on six beaches of the Svalbard Archipelago (Bergmann et al. 2017a). Arctic sea ice  
59 was reported by Obbard et al. (2014) as having microplastic concentrations (38 – 234 particles m<sup>3</sup>  
60 of ice) several orders of magnitude greater than highly contaminated oceanic waters. Lusher et al.  
61 (2015) first reported on microplastic abundances in surface and sub-surface waters south and  
62 southwest of Svalbard. Amélineau et al. (2016) later reported on microplastic abundance in surface  
63 waters east of Greenland. Regarding Arctic species, microplastics have been detected in the gular  
64 pouches of Little Aulks (*Alle Alle*), (Amélineau et al. 2016), as well as in the stomachs of juvenile  
65 polar cod (*Boreogadus saida*), (Kuhn et al. 2018). Microplastics were also detected in sediments  
66 (collection depths 2340 – 5570 m) from the Fram Strait (Bergmann et al. 2017b). Recently, results  
67 from a circumpolar expedition of the Arctic indicated that concentrations of floating plastic ranged  
68 between 0 – 320 000 items km<sup>-2</sup> in the Greenland and Barents Sea and 0 – 27 000 items km<sup>-2</sup> in  
69 the rest of the Arctic Ocean (Cózar et al. 2017).

70  
71 Plastic contaminants are introduced to the Arctic Ocean due to a combination of (i) long-range  
72 transport processes, e.g. via oceanic currents, biotransport and riverine input, and (ii) local  
73 anthropogenic activities, e.g. shipping. The three oceanic currents which supply the greatest water  
74 volumes to the Arctic Ocean are the (i) West Spitsbergen Current i.e. the polar limb of the North  
75 Atlantic circulation which carries warm water from the North Atlantic Current (9.5 Sverdrup, Sv  
76 =  $10^6\text{m}^3\text{s}^{-1}$ ), (ii) a cold ocean current that enters from the Pacific Ocean via the Bering Strait (1.5  
77 Sv) and, (iii) a branch of the North Atlantic Current, which flows along the Siberian coastline (1.0  
78 Sv), (Zarfl and Matthies 2010). These oceanic currents may also transport plastics to the Arctic  
79 Ocean with the estimated plastic flux to this region ranging between 62 000 to 105 000 tons per  
80 year (Zarfl and Matthies 2010). Models based on a particle-trajectory approach for studying the  
81 fate of marine debris in the open ocean highlighted the northward transport of marine debris to  
82 polar regions and the formation of a sixth so-called garbage patch in the Barents Sea (van Sebille  
83 et al. 2012). Bio-transport is another long-range transport process via which plastics may enter  
84 polar regions. Plastic ingestion was reported in Northern Fulmars (*Fulmaris glacialis*) and Thick-  
85 billed Murres (*Uria lomvia*) in the Arctic (Mallory 2008; Provencher et al. 2012; Trevail et al.  
86 2015). Some studies suggested that the seabirds had ingested plastics during their wintering in the  
87 North Atlantic Ocean and had then transported the contaminants to the Arctic upon migration  
88 (Mallory 2008; Provencher et al. 2012). Riverine discharge from Siberian (Ob, Yenisei and Lena)  
89 and Canadian (Mackenzie) rivers are other potential sources of plastics to the Arctic. Obbard et al.  
90 (2014), however, point out that the contribution of riverine discharge to plastic input in the Arctic  
91 is projected to be low due to the fact that these rivers flow through sparsely populated watersheds.  
92 Local anthropogenic activities are another source of plastics to the Arctic. Increased ship traffic

93 due to shipping and tourism was found to be positively correlated with increased litter densities in  
94 the Fram Strait (Bergmann and Klages 2012; Tekman et al. 2017).

95

96 The intense focus by scientists on the near-surface layer of the ocean for microplastics has been  
97 due in part to the presumption that the majority of particles would be found in this region of the  
98 water column given the inherent densities of individual synthetic polymers. Such a theorization  
99 led to traditional techniques that involved nets, manta trawls as well as the seawater intake of  
100 vessels that sampled only the upper few metres of the water column for microplastics. Yet, several  
101 studies indicated that a mismatch existed between observed and expected plastic concentrations in  
102 surface oceanic waters when estimated plastic production and projected inputs to the oceans were  
103 considered (Cózar et al. 2014; Eriksen et al. 2014). It was therefore proposed that several  
104 mechanisms potentially influenced the vertical distribution of microplastics within the water  
105 column and led to their transport out of surface waters. Some of these mechanisms included (i)  
106 incorporation into marine aggregates (Long et al. 2015), (ii) biofouling (Fazey and Ryan 2016),  
107 (iii) incorporation into faecal matter (Cole et al. 2016) and, (iv) hydrodynamic factors such as wind  
108 (Kukulka et al. 2012). Despite the theorization that surface waters are not the ultimate repository  
109 for plastic debris in the marine environment (Cózar et al. 2014), few studies ventured beyond  
110 traditional near-surface microplastic monitoring to investigate their vertical distribution in the  
111 water column.

112

113 Microplastic pollution in the Arctic Ocean is an issue that warrants attention due to the potential  
114 threats that these contaminants may pose to the inhabitants of this unique ecosystem. A practical  
115 step towards addressing this issue and evaluating the extent of the problem involves assessing the

116 abundance, distribution and composition of microplastics in Arctic waters. Whilst microplastic  
117 monitoring in the marine environment has traditionally focused on surface waters, the reality is  
118 that the vast majority of marine organisms inhabit sub-surface waters. Monitoring microplastics in  
119 sub-surface waters is particularly relevant as it can also provide some insight into the whereabouts  
120 of the ‘missing plastic’ from surface waters. To our knowledge, the present study sought for the  
121 first time (i) to provide a spatial overview of microplastic abundance, distribution and composition  
122 in the Polar Mixed Layer (PML) of the Arctic Central Basin (ACB) and, (ii) to determine whether  
123 microplastics in the ACB were being transported out of surface waters by assessing their vertical  
124 distribution in the water column.

125

## 126 **2.0 Materials and Method**

### 127 **2.1 Study Area**

128 The Arctic Ocean is comprised of a deep central basin surrounded by extensive continental shelves  
129 (CAFF 2013). The bathymetry of the Arctic Ocean is such that the Lomonosov Ridge separates  
130 the central basin into the Canadian (Amerasian) and Eurasian basins with the basins being further  
131 sub-divided by the (i) Gakkel Ridge, into the Amudsen and Nansen basins and, (ii) Alpha Ridge,  
132 into the Makarov and Canada basins (Jakobssen et al. 2004; Rudels 2015, Figure 1). A major  
133 structuring element of the Arctic marine ecosystem is sea ice which floats on the surface layer  
134 impeding surface mixing and influencing freshwater and heat fluxes (CAFF 2013). In the Arctic  
135 Ocean, there is a distinct vertical stratification of the water column giving rise to three major water  
136 layers (i) Polar Surface Water (PSW) which includes the Polar Mixed Layer (PML) and the  
137 halocline, (ii) Atlantic Water and, (iii) deep and bottom waters (Rudels 2015, Figure 1). The PML  
138 (approximately 50 m deep) is the uppermost surface layer of low salinity water formed as a result

139 of sea ice melt and the influx of freshwater from riverine sources (Rudels et al. 1991; CAFF 2013;  
140 Jakobsson et al. 2014). Beneath the PML is a halocline (50 – 250 m), characterised by a strong  
141 salinity increase with depth and comprised of either Pacific waters or Atlantic waters with the  
142 Pacific halocline being deeper than the Atlantic halocline (Rudels et al. 1991; Jakobsson et al.  
143 2004). Below the halocline lies an intermediate water layer comprising of dense saline Atlantic  
144 water. The deep and bottom waters also referred to as Arctic deep water ranges from a depth of  
145 approximately 900 m and extend to the seafloor (Rudels et al. 1991; CAFF 2013).

146

## 147 **2.2 Sample Collection**

### 148 *Underway samples*

149 This study was conducted onboard the Swedish icebreaker Oden during the Arctic Ocean 2016  
150 expedition. The vessel departed Longyearbyen, Svalbard on August 8<sup>th</sup> 2016 and traversed  
151 approximately 4943 nautical miles in the Arctic Ocean until its return on September 19<sup>th</sup> 2016  
152 (Figure 2). Sub-surface oceanic water pumped onboard the vessel via the bow water system was  
153 sampled for microplastics according to Lusher et al. (2014). Sampling was conducted for a period  
154 of approximately 6 weeks (9 August to 16 September 2016). Since each sample constituted the  
155 filtration of approximately 2000 L of water, the total survey effort for this study was approximately  
156 116 000 L of water (58 samples).

157

158 Seawater from a continuous intake located at the keel of the ship (depth 8.5 m) was pumped  
159 onboard the vessel using a rotary positive displacement pump (Universal II Series Pump,  
160 Waukesha Cherry-Burrell) at a flow rate of 85 L/min (at optimal capacity) and transported to the  
161 laboratory via stainless steel pipes. Prior to reaching the laboratory, the seawater passed through a



162 stainless steel primary filter (pore size 2.5 mm) which was beyond the control of the investigator.  
163 The discharge of grey water in relation to the seawater intake was not an issue since the seawater  
164 intake was located towards the front of the vessel whilst grey water was discharged mid-vessel. In  
165 the laboratory, seawater from the vessel's bow water system was allowed to flow through a covered  
166 stainless steel sieve (250  $\mu\text{m}$ ) by means of a connection hose fitted into the wooden sieve cover.  
167 For the duration of the sampling, the stainless steel sieve was supported in a wooden stand. Based  
168 on Lusher et al. (2014), approximately 2000 L of water was filtered for each sample. The length  
169 of time taken for the filtration of the specified volume of water was determined by calculation of  
170 the flow rate of the seawater. A flow meter, attached at a point prior to the entry of the water into  
171 the sieve, was also used to verify the volume of water filtered. Once the specified volume of water  
172 was filtered, the sieve was removed and Milli-Q water was used to wash retained material from  
173 the sieve into a clean container. The collected material was then filtered under vacuum onto glass  
174 microfiber paper (GF/C); Whatman: 47 mm, pore size: 1.2  $\mu\text{m}$ , using a Buchner funnel and a  
175 vacuum flask (Lusher et al. 2014). Each filter paper was then placed into a clean plastic petri dish,  
176 covered and stored in a freezer (-20 °C) until returned to the laboratory. At the start and at the end  
177 of each sample, positioning data were collected. Data for various environmental variables were  
178 obtained from the vessel's (i) thermosalinometer (water temperature, salinity) and (ii) weather  
179 station (wind speed and direction).

180

### 181 *CTD samples*

182 A rosette water sampler containing 24 Niskin bottles coupled to a Sea-Bird SBE 911 conductivity-  
183 temperature-depth (CTD) sensor suite (hereafter referred to as CTD) was used to collect sub-  
184 surface water samples and hydrographic data at 9 sampling locations in the Arctic Ocean. Upon

185 deployment from the vessel, the CTD entered the water and was allowed to descend to the bottom  
186 layer. During the descent, Niskin bottles were open with water flowing through them. It was during  
187 the up-cast that multiple Niskin bottles were closed at specific depths in order to facilitate the  
188 collection of a specified volume of water. A total of 48 water samples were retrieved during the 9  
189 CTD casts to sample for microplastics. At 7 of the CTD casts, 6 water depths were sampled with  
190 48 L of water collected at each depth i.e. 4 Niskin bottles (12 L) per depth. At 2 CTD casts, 3 water  
191 depths were sampled with 21 L of water collected at each depth i.e. 3 Niskin bottles (7 L) per  
192 depth. At a particular sampling location, the overall goal was to collect samples in the near-surface,  
193 mid-water and bottom layers within the water column thereby reflecting the main water masses.  
194 As such, exact sampling depths were determined by the information provided by the salinity and  
195 temperature sensors on each downcast. The deepest samples at each CTD cast were collected at  
196 least 10 m above the seafloor at a given location. Following each CTD cast, Niskin bottle taps  
197 were rinsed with Milli-Q water and a clean hose was attached. Water from bottles closed at the  
198 same depth was passed through the same stainless steel sieve (250  $\mu\text{m}$ ) held in a covered wooden  
199 stand. Once water from all bottles at a specific depth had been filtered, the sieve was removed and  
200 Milli-Q water was used to wash retained material from the sieve into a clean container. The  
201 collected material was filtered under vacuum onto glass microfiber paper (GF/C); Whatman: 47  
202 mm, pore size: 1.2  $\mu\text{m}$ , using a Buchner funnel and a vacuum flask. Each filter paper was then  
203 folded and placed into an aluminium foil packet and stored in a freezer (-20 °C) until returned to  
204 the laboratory.

205

### 206 **2.3 Method Validation and Contamination Prevention**

207 For the underway samples, potential contamination during sample processing was evaluated by (i)  
208 leaving clean plastic petri dishes with filter paper exposed to the air during vacuum filtration, and  
209 (ii) passing an aliquot (250 mL) of Milli-Q water through clean GF/C filter paper under vacuum.  
210 For the CTD samples, potential contamination was assessed by filling a clean Niskin bottle with  
211 Milli-Q water and subjecting it to the exact process a sample underwent. Measures taken to prevent  
212 contamination in the laboratory included (i) wearing lab coats (cotton/polyester blend), cotton  
213 clothing and gloves (nitrile) during sample processing, (ii) placing a wooden cover over the  
214 stainless steel sieve during filtration to prevent airborne contamination, and (iii) washing all  
215 containers used during sample processing with Milli-Q water before reuse.

216

#### 217 **2.4 Laboratory analyses**

218 Filter papers were removed from the freezer, left to dry and then visually examined under a  
219 dissecting microscope (Olympus SZX10) equipped with a polariser and camera (Q Imaging Retiga  
220 2000R). Potential microplastics were isolated and processed (photographed and length  
221 measurements taken) prior to transferring to a clean filter paper in a labelled petri dish (Kanhai et  
222 al. 2017). All potential microplastics were analysed by Fourier transform infrared (FT-IR)  
223 spectroscopy on a Thermo Scientific Nicolet iN10 FT-IR spectrometer. The instrument was  
224 equipped with a potassium bromide (KBr) beamsplitter and an internal mercury cadmium telluride  
225 (MCT) detector which was cooled with liquid nitrogen. Microscope-reflectance sampling was  
226 performed and spectra were recorded as the average of 256 scans in the spectral wave number  
227 range of 4000 - 675  $\text{cm}^{-1}$  at a resolution of 4  $\text{cm}^{-1}$ . Thermo Scientific's OMNIC Picta Version 9  
228 spectroscopy software was used for processing and evaluating all spectra. Prior to analysing each  
229 sample, background scans were performed and sample spectra were automatically corrected. Each

230 sample spectrum was compared with those of known standard polymers in the (i) Hummel  
231 Polymer Sample library, (ii) Polymer Laminate Films library, and (iii) Wizard library. Values of  
232 between 0 and 100 % were produced for each match between sample and reference spectra with  
233 the highest percentage representing the closest match. Particles for which there was uncertainty  
234 regarding the identity of the polymer (specifically fragments and some fibres) were subjected to  
235 further FT-IR spectroscopy on a Bruker Vertex 70 Infrared Spectrometer coupled to a Hyperion  
236 1000 microscope (Kanhai et al. 2017). Samples which produced spectra with a match < 60 % were  
237 automatically rejected while those with a match of > 70 % were automatically accepted. All spectra  
238 with matches > 60 % were individually examined to ensure that there was clear evidence of peaks  
239 from the sample corresponding to known peaks of standard polymers and that instances of the  
240 misidentification of natural and semi-synthetic polymers was reduced (Comneau-Stancu et al.  
241 2017).

242

## 243 **2.5. Statistical analyses**

244 All statistical analyses were performed using R version 3.2.3 (R Core Team 2015). Descriptive  
245 statistics, histograms and box plots were generated and tests of normality (Supplementary Tables  
246 1 and 2) were conducted on all data to determine whether parametric or non-parametric statistical  
247 analyses were appropriate. Correlation analyses were performed between individual  
248 environmental variables and microplastic abundance for both underway and CTD samples. A  
249 generalized additive model (GAM) was developed using the underway data and a generalized  
250 linear mixed model (GLMM) was developed using the CTD data to determine which  
251 environmental variables had an effect on microplastic abundance.

252

## 253 **3. Results**

### 254 **3.1. Quality Control**

255 In conjunction with the collection of samples via the underway system of the vessel, a total of 24  
256 blanks (air contamination-12, method-12) were run (Supplementary Table 3). No synthetic  
257 polymers were found in the method blanks. However, a single synthetic fibre (blue, polyethylene  
258 terephthalate, 0.438 mm) was found in the last air contamination blank. For 6 of the 9 CTD casts,  
259 at least one method blank was run (Supplementary Table 4). Between 0 and 3 synthetic fibres were  
260 found in each of the method blanks. The synthetic fibres that were found included polyethylene  
261 terephthalate (n = 8), polyacrylonitrile (n = 1) and polyvinyl chloride (n = 1).

262

### 263 **3.2. Overview of findings**

#### 264 *Underway samples*

265 A total of 303 particles were isolated from the underway samples and analysed by FT-IR  
266 spectroscopy. Of these, 46 particles were excluded because of uncertainty regarding their identity  
267 (< 60 % match to reference spectra) and in the minority of cases (n = 6) due to their length (< 250  
268  $\mu\text{m}$ ). Of the remaining particles (n = 257), 14 were macro-particles (i.e. > 5 mm in length) and 243  
269 were micro-particles (< 5 mm in length). Of the macro-particles, 11 were confirmed as  
270 macroplastics having the following polymer types: polyethylene terephthalate (4), polyamide  
271 blend (4) and polyacrylonitrile (3). Of the 243 micro-particles, 110 were natural (cellulosic), 16  
272 were semi-synthetic (cellulose-based e.g. rayon) and 117 were synthetic. All further analyses and  
273 discussions focus on the 117 confirmed microplastics.

274

275 The majority (94 %) of microplastics were fibres and 6 % were fragments. In terms of colour, the  
276 most prevalent were blue (49 %) and transparent (25 %) (Figure 3a). Approximately 62 % of the  
277 microplastics occurred in the larger size classes of 1.0 – 2.0 mm and 2.0 – 5.0 mm (Figure 4a).  
278 Microplastic polymer types included polyester (n = 88), blends (n = 11), polyacrylonitrile (n = 8),  
279 polyamide (n = 5) and polyvinyl chloride (n = 5), (Figure 5a). The overall category of ‘polyester’  
280 included both polyethylene terephthalate (PET) and other polyesters while blends included either  
281 polyamide blends or polyester blends.

282

### 283 *CTD samples*

284 A total of 157 particles were isolated from the CTD samples and analysed by FT-IR spectroscopy.  
285 Of these, 14 were excluded for the reasons mentioned above for the underway samples. Of the  
286 remaining particles, 2 were categorised as macroplastics (> 5 mm) and included PET and  
287 polyacrylonitrile. Of the 141 micro-particles (< 5 mm), 39 were natural (cellulosic), 8 were semi-  
288 synthetic (cellulose-based e.g. rayon) and 94 were synthetic. All further analyses and discussions  
289 focus on the 94 confirmed microplastics.

290

291 Overall, the characteristics of the microplastics isolated from CTD samples were similar to those  
292 from the underway samples in that (i) the majority (96 %) of microplastics were fibres and 4 %  
293 were fragments, (ii) the most prevalent colours were blue (46 %) and transparent (22 %) (Figure  
294 3b), (iii) the majority (64 %) of microplastics were in the larger size classes of 1.0 – 2.0 mm and  
295 2.0 – 5.0 mm (Figure 4b), and (iv) microplastic polymer types included polyester (n = 74), blends  
296 (n = 12), polyacrylonitrile (n = 6), polyamide (n = 1) and polyvinyl chloride (n=1), (Figure 5b).

297 The overall category of polyester included both polyethylene terephthalate (PET) and polyester  
298 while blends included only polyamide blends.

299

### 300 **3.2 Microplastic abundance and distribution in the Arctic Ocean**

301 Based on the underway samples (collection depth 8.5 m), microplastic abundance in sub-surface  
302 waters in the Arctic Ocean ranged between 0 – 7.5 particles  $m^{-3}$  with a median of 0.7 particles  $m^{-3}$   
303 (interquartile range 0.4 – 1.0), (Figure 2, Supplementary Table 5). For the majority of the  
304 sampling sites, microplastic abundance ranged between 0 – 1.0 particles  $m^{-3}$ . However, at a few  
305 sites, microplastic abundances were between 2 – 2.5 particles  $m^{-3}$  and at two sites it was at 5 and  
306 7.5 particles  $m^{-3}$  respectively. Based on the CTD samples (collection depths between 8 – 4369 m),  
307 microplastic abundance in sub-surface waters in the Arctic Ocean ranged between 0 – 375 particles  
308  $m^{-3}$  with a median of 20.8 particles  $m^{-3}$  (interquartile range 20.8 – 62.5) (Figures 2 and 5,  
309 Supplementary Table 6). With the exception of CTD cast 4, the CTD casts (1 – 3) nearer the  
310 periphery of the Arctic Central Basin (ACB), i.e. in the Nansen Basin (Yermak Plateau), reflected  
311 a comparatively higher abundance of microplastics in the water column than other CTD casts  
312 within the ACB (Figures 2, 6). It must be noted however that CTD casts 1 – 3 sampled the upper  
313 850 m of the water column and as such would have sampled particles from the Polar Mixed Layer  
314 (PML), Atlantic halocline and Atlantic water (Figures 1, 6). This is in contrast to the other CTD  
315 casts which sampled a much more extensive vertical range throughout the water column by  
316 including deep bottom water in excess of 1000 m depth. Overall, there was no statistically  
317 significant correlation between microplastic abundance and depth (Spearman's rank correlation,  
318  $\rho = 0.06$ ,  $p\text{-value} = 0.7$ ). However, upon examination of individual CTD casts, it is apparent that  
319 microplastic abundance was not uniform at various depths in the water column and that there were

320 certain depths that reflected higher microplastic abundances (Figure 6). Additionally, microplastic  
321 abundance (particles  $m^{-3}$ ) in the different water masses of the ACB was as follows: Polar Mixed  
322 Layer (0 - 375) > Deep and bottom waters (0 – 104) > Atlantic water (0 – 95) > Halocline i.e.  
323 Atlantic or Pacific (0 – 83), (Table 1).

324

### 325 **3.3. Association between environmental variables and microplastic abundance in samples**

326 Correlation analyses were conducted to determine whether there was any association between  
327 environmental variables and microplastic abundance in the samples. For both the underway and  
328 CTD samples, there was no statistically significant correlation between microplastic abundance  
329 and any of the ancillary environmental variables of temperature, salinity, wind direction, wind  
330 speed, depth and density (Supplementary Tables 1 and 2). Specifically, there was no statistically  
331 significant correlation between microplastic abundance at depth (Supplementary Table 2).  
332 However, for the underway samples, there was a statistically significant weak negative correlation  
333 between microplastic abundance and latitude (Spearman's rank correlation,  $\rho = -0.286$ , p-value  
334 = 0.03).

335

336 Using the underway data, a Generalized Additive Model (GAM) was developed to further  
337 determine whether environmental variables influenced microplastic count in the underway  
338 samples. In this model, the response variable was microplastic count (number of microplastics per  
339 sample) and initial explanatory variables included location (latitude, longitude), physico-chemical  
340 properties associated with sub-surface waters (temperature, salinity) and weather data (wind  
341 direction, wind speed). In the model, the Poisson family distribution of error terms was specified  
342 with a log link function since microplastic abundance data were count data. The output of the



343 initial model was examined and based on this non-parametric smoothers were accordingly applied  
344 to the explanatory variables. A scale invariant tensor product smooth (te) was applied to latitude  
345 and longitude while a cubic regression spline (bs = cr) was applied to all other variables except  
346 wind direction to which no smoother was applied (based on initial model plots). Non-significant  
347 explanatory variables (as evidenced by their p-values) were eliminated in a stepwise manner until  
348 a GAM with the lowest Akaike Information Criterion (AIC) score and the fewest explanatory  
349 variables was obtained. The final GAM (R-sq = 0.396) was as shown below:

350

351 Microplastic count ~ te(**latitude, longitude**) + s(**temperature**, br = "cr") + s(**wind speed**, bs =  
352 "cr")

353

354 All of the explanatory variables that were present in the final model (shown in bold) were found  
355 to have a significant influence on microplastic count in water samples from the Arctic Ocean (wind  
356 speed, p-value = 0.0006, latitude, longitude, p-value = 0.0007, temperature, p-value = 0.0483).

357

358 A generalized linear mixed model (GLMM) was developed using the CTD data to determine the  
359 influence of environmental variables on microplastic count in the CTD samples. In this model, the  
360 response variable was microplastic count (number of microplastics per sample) and initial  
361 explanatory variables included location (latitude, longitude), physicochemical properties  
362 associated with sub-surface waters (temperature, salinity), depth at which water was sampled and  
363 ctd number. All the explanatory variables were included in the model as fixed effects, with the  
364 exception of ctd number which was included as a random effect. In the model, the Poisson family  
365 distribution of error terms was specified with a log link function since microplastic abundance data

366 were count data. Based on the preliminary finding that there was a statistically significant  
367 correlation between depth and salinity (Spearman's rank correlation,  $\rho = 0.852$ ,  $p\text{-value} =$   
368  $8.156e-13$ ), temperature and salinity (Spearman's rank correlation,  $\rho = 0.506$ ,  $p\text{-value} = 0.00074$ )  
369 and depth and density (Spearman's rank correlation,  $\rho = 0.973$ ,  $p\text{-value} = < 2.2 e-16$ ),  
370 interactions between these variables (denoted by ':') were included in the initial model. Non-  
371 significant explanatory variables (as evidenced by their  $p$ -values) were eliminated in a stepwise  
372 manner until a model with the lowest Akaike Information Criterion (AIC) score and the fewest  
373 explanatory variables was obtained. The significance of the random effect (ctd number) in the final  
374 model was verified by using analysis of variance (ANOVA) to compare the full final model to a  
375 reduced model (random effect deleted). The model with the random effect was shown to be  
376 significantly different (ANOVA,  $p\text{-value} = 0.0008$ ,  $\Delta\text{AIC} = 9.19$ ) from the model without the  
377 random effect. The model with the lower AIC score (AIC = 190.59) was retained as the final mixed  
378 effects model as shown below:

379

380 Microplastic count ~ **latitude** + **temperature:salinity** + (1|ctd)

381

382 Latitude ( $p\text{-value} = 0.0198$ ) and the physicochemical parameters of temperature and salinity ( $p$ -  
383 value =  $7.46 e-05$ ), as shown in bold, were the explanatory variables that were found to have a  
384 significant influence on microplastic count in the CTD samples.

385

#### 386 **4.0 Discussion**

387 The discovery of microplastics in virtually every environmental phase (sea ice, water, sediments,  
388 biota) of the Arctic and Southern Oceans has revealed that polar oceans, though remote, are not

389 immune to the entry of plastic contaminants to their ecosystems (Bergmann and Klages 2012;  
390 Obbard et al. 2014; Lusher et al. 2015; Amélineau et al. 2016; Bergmann et al. 2016; Bergmann  
391 et al. 2017a; Bergmann et al. 2017b; Cincinelli et al. 2017; Cózar et al. 2017; Isobe et al. 2017;  
392 Tekman et al. 2017; Waller et al. 2017). The present study expands the knowledge base about  
393 plastics in the Arctic by providing evidence for the existence of microplastics in the Polar Mixed  
394 Layer (PML) as well as some insight into the vertical distribution of microplastics in the Arctic  
395 Central Basin (ACB). This region of the Arctic, though of low productivity, has been recognised  
396 as an Ecologically/Biologically Significant Marine Area (EBSA) due to its uniqueness/rarity,  
397 provision of a critical habitat and ability to support specialised biota (CAFF 2017). There is cause  
398 for concern about microplastics in Arctic waters since laboratory studies have shown that these  
399 contaminants may (i) hinder algal photosynthesis/growth (Bhattacharya et al. 2010; Besseling et  
400 al. 2014), (ii) reduce feeding and energy reserves of lugworms (Besseling et al. 2013; Wright et  
401 al. 2013), (iii) reduce filtering activity and decrease lysosomal membrane stability in mussels (Von  
402 Moos et al. 2012; Wegner et al. 2012), (iv) reduce feeding and reproductive output in copepods  
403 (Cole et al. 2015) and, (v) cause liver stress, negatively impact upon cholinergic neurotransmission  
404 and lead to endocrine disruption in fish (Oliveira et al. 2013; Rochman et al. 2013; Rochman et al.  
405 2014). It must be pointed out, however, that some laboratory experiments which reported negative  
406 effects of microplastics on marine organisms used microplastic concentrations of 42 to 10 000  
407 particles/mL or 42 million to 10 billion particles  $m^{-3}$  (Phuong et al. 2016). In context, microplastic  
408 abundance in the ACB as reported by the present study ranged from 0 – 7.5 particles  $m^{-3}$  (based  
409 on underway sampling) and 0 – 375 particles  $m^{-3}$  (based on CTD sampling). Although the  
410 ecological impact of microplastics upon the Arctic ecosystem presently remains unknown, it is  
411 plausible that these contaminants could pose a threat to its inhabitants.

412

413 The present study showed for the first time the pervasiveness of microplastics throughout the water  
414 column of the Arctic Central Basin. Between depths of 8 – 4400 m, microplastic abundance in the  
415 ACB ranged between 0 – 375 particles m<sup>-3</sup> (based on CTD sampling). Such findings provide  
416 evidence that in natural conditions microplastics are being vertically transported out of surface  
417 waters. These findings also give some indication as to the whereabouts of the ‘missing plastic’  
418 from oceanic surface waters (Cózar et al. 2014; Eriksen et al. 2014). Recently, Courtene-Jones et  
419 al. (2017) also reported on microplastic abundance (70.8 particles m<sup>-3</sup>) in deep oceanic waters  
420 (2227 m at the Rockall Trough, North East Atlantic Ocean) and similarly suggested the possibility  
421 of vertical re-distribution of microplastics within the water column. Although it remains unclear  
422 as to which mechanisms are specifically operating in the ACB to influence the vertical transport  
423 of particles, previous studies have provided several possibilities. Specifically, some laboratory  
424 experiments showed that aggregates of algae species (*Chaetoceros neogracile*, *Rhodomonas*  
425 *salina*) were capable of incorporating and concentrating microplastics and that the microplastics  
426 impacted the sinking rates of the aggregates (Long et al. 2015). In the Arctic Ocean, it is certainly  
427 plausible that marine aggregates may be playing a role in the vertical transport of microplastics  
428 due to the existence of phytoplankton in the ACB (CAFF 2017) and the fact that transparent  
429 exopolymer particles (TEPs); which are excreted by algae and are important components of marine  
430 aggregates, have been reported in sub-surface waters as far north as the Fram Strait (Engel et al.  
431 2017). Biofouling is another possibility as field studies have shown that plastic particles exposed  
432 to natural conditions became sufficiently fouled, had their average material density affected  
433 leading them to sink (Fazey and Ryan 2016). The incorporation of microplastics into faecal matter  
434 is another means by which microplastics may be vertically transported out of surface waters given

435 that laboratory experiments have shown that zooplankton may egest microplastics within densely  
436 packed faecal pellets which in natural conditions would sink or in some cases be eaten by other  
437 biota (Cole et al. 2016).

438  
439 Sea ice is an integral component of the Arctic Ocean's ecosystem and as such possibly exerts an  
440 influence on microplastic abundance in sub-surface waters. Sea ice floating on the surface of the  
441 water column in the Arctic Ocean can potentially act as (i) a source of microplastics upon melting,  
442 (ii) a physical barrier to wind and as such reduce vertical mixing of surface waters and, (iii) a  
443 physical barrier to influx of polluted surface waters. Based on the analysis of sub-sections of four  
444 ice cores, sea ice in the Arctic Ocean was reported to contain orders of magnitude more  
445 microplastic than contaminated oceanic waters suggesting that sea ice potentially acts as both a  
446 sink and a source of microplastics (Obbard et al. 2014). Apart from Obbard et al. (2014) no data  
447 exists in the published literature regarding either the spatial or vertical distribution of microplastics  
448 in sea ice from the Arctic Ocean. In the upper water column, the absence of sea ice cover means  
449 that wind stress can generate turbulence and lead to vertical mixing of buoyant plastic debris  
450 (Kukulka et al. 2012). More recently, Cózar et al. (2017) suggested that sea ice can also act as a  
451 physical barrier preventing the surface advance of polluted Atlantic water into the Arctic Ocean.  
452 The present study highlighted that the Polar Mixed Layer (PML) of the ACB reflected the highest  
453 overall microplastic abundance (particles  $m^{-3}$ ): Polar Mixed Layer (0 - 375) > Deep and bottom  
454 waters (0 – 104) > Atlantic water (0 – 95) > Halocline i.e. Atlantic or Pacific (0 – 83). As previously  
455 mentioned, the PML is the uppermost surface layer of low salinity water (approximately 50 m  
456 deep) formed as a result of sea ice melt and the influx of freshwater from riverine sources (Rudels  
457 et al. 1991; CAFF 2013). It is possible that one of the reasons that the highest microplastic

458 abundances were recorded in this layer is due to its proximity to microplastic sources such as  
459 melting sea ice as well as sea-going vessels (especially if they are releasing wastewater to the  
460 environment). Furthermore, in the present study, the highest microplastic abundances were  
461 reported nearer to the periphery of the Arctic Central Basin (ACB), i.e. in waters north of Svalbard.  
462 It is possible that the lack of permanent sea ice cover in this region of the Arctic allows incoming  
463 Atlantic water to have a greater influence on near surface waters thereby resulting in higher  
464 microplastic abundances. A recent circumpolar expedition of the Arctic Ocean similarly reported  
465 that the north eastern Atlantic sector of the Arctic was a hotspot of plastic debris due to the  
466 influence of incoming Atlantic water (Cózar et al. 2017).

467

468 The present study showed that there was a predominance of fibrous microplastics (> 90%) in sub-  
469 surface waters of the ACB. This dominance of fibres in sub-surface waters was similarly reported  
470 in (i) the north east Pacific Ocean (75 %), (ii) the north east Atlantic Ocean (96 %), (iii)  
471 south/southwest of Svalbard (95 %), and (iv) the Atlantic Ocean (96 %) (Desforges et al. 2014;  
472 Lusher et al. 2014; Lusher et al. 2015; Kanhai et al. 2017). Fibrous microplastics in the marine  
473 environment most likely originate from textile materials and fishing gear (Andrady 2017). Studies  
474 have indicated that washing clothes may lead to the release of fibrous materials in the order of >  
475 1900 fibres per wash or as much as 700 000 fibres per 6 kg load of acrylic fabric (Browne et al.  
476 2011; Napper and Thompson 2016). A recent study in the Ross Sea revealed that the highest  
477 concentration of fibrous microplastics (54 %) was found close to the effluent of a sewage treatment  
478 plant at the scientific Mario Zucchelli Station, Antarctica (Cincinelli et al. 2017). Fibrous  
479 microplastics may enter the Arctic Ocean through a combination of long range transport processes  
480 (e.g. via oceanic currents, riverine input) or more in-situ activities such as the release of wastewater

481 from vessels operating in the region. Another mechanism which was recently suggested as being  
482 responsible for plastic fibres in Arctic sea ice was atmospheric transport (Cózar et al. 2017). This  
483 is certainly plausible as there have been reports of the atmospheric fallout of synthetic polymers  
484 in both urban and sub-urban environments in France (Dris et al. 2016). It must be highlighted that  
485 while fibrous microplastics seem to be dominant in certain sub-surface waters, other studies  
486 conducted in surface Arctic waters reported a predominance of filaments (97 %) and fragments  
487 (73 %), (Amélineau et al. 2016; Cózar et al. 2017).

488

489 Analytical techniques such as FT-IR and Raman spectroscopy are a central component of  
490 microplastic studies which aim to provide unambiguous identification of synthetic polymers in  
491 environmental samples. Omission of this critical step is likely to be a key contributor to an  
492 overestimation of microplastic abundances due to the inclusion of non-synthetic polymers in  
493 microplastic counts. In the present study, for example, only a percentage (underway - 48 %, CTD  
494 - 67%) of the particles were confirmed as synthetic polymers with the remainder being a  
495 combination of natural and semi-synthetic polymers. Of the synthetic polymers in the present  
496 study, the most abundant (underway – 74 %, CTD – 78 %) was polyethylene terephthalate (PET).  
497 A member of the polyester family, PET is one of the five major types of commodity plastics  
498 commonly found in the marine environment (Andrady 2011; Andrady 2017). As a thermoplastic,  
499 PET is often used in manufacturing beverage containers and packaging materials and its fibres are  
500 used in clothing. Overall, this finding of a high prevalence of polyesters in sub-surface waters of  
501 the Arctic Central Basin was also corroborated by those of other Arctic studies which reported that  
502 polyester was the most prevalent synthetic polymer in waters south/southwest of Svalbard (15 %),  
503 in waters of the east Greenland Sea (53 %), and in sea ice (21 %), (Obbard et al. 2014; Lusher et

504 al. 2015; Amélineau et al. 2016). Other polymers in sub-surface Arctic waters in this study  
505 included polyacrylonitrile, polyamide and polyvinyl chloride. Of note is the fact that the majority  
506 of synthetic polymers in the present study had densities greater than that of seawater (Andrady  
507 2017). Even though investigators are able to generate information regarding the identity of  
508 polymers in environmental samples, definitive statements cannot be made about the origin of the  
509 plastics. Based on the identity and type of synthetic polymers found in sub-surface Arctic waters,  
510 it is likely that they originated from textiles, fishing gear, beverage containers and packaging  
511 materials (Andrady 2011; Andrady 2017).

512

513 Within the water column, the distribution of microplastics is in a state of flux due to the influence  
514 of multiple factors. The development of models based on simultaneously acquired environmental  
515 and microplastic data is immensely useful in this respect as they can provide some discernment  
516 regarding the variables influencing measures of microplastic abundances in the marine  
517 environment. In the present study, the utilisation of a generalized additive model (GAM) and a  
518 generalized linear mixed effects model (GLMM) was particularly relevant due to the ability of  
519 both models to handle non-normal data and in the case of the GLMM to differentiate between  
520 fixed and random effects. Visual inspection of microplastic abundances in sub-surface waters  
521 (Figure 2) revealed that the highest microplastic abundances were located to some extent towards  
522 the periphery of the Arctic Central Basin (ACB). Bearing this in mind, it was presumed that the  
523 ‘location’ of sampling could have influenced the number of microplastics that were found in the  
524 samples. Cózar et al. (2017) also noted this spatial heterogeneity in the distribution of plastics in  
525 the Arctic. For these reasons, latitude and longitude were included as a proxy of location in the  
526 models. In the GAM, both variables were included using a smoother and therefore it was not



527 assumed that latitude and longitude had a linear effect on microplastic count. Furthermore, during  
528 the period of sampling (August/September 2016), the Arctic Central Basin (ACB) was not  
529 completely covered by sea ice. Since the vessel traversed areas of open water, wind was included  
530 in the model as it could have influenced microplastic abundance at specific sampling sites. Both  
531 models suggested that location, oceanographic (temperature, salinity) and atmospheric variables  
532 (wind) had a significant influence on microplastic counts in samples of water from the Arctic  
533 Ocean. The findings of the present study must be taken in the context of the number of samples  
534 used in the generation of the GAM ( $n = 58$ ). Models that are based on a low number of samples  
535 are weak e.g. standard errors are inflated, etc. However, such models allow us to delve a bit deeper  
536 into the factors influencing measurements of microplastic abundance in the Arctic Ocean. Findings  
537 of the present study were also corroborated by previous studies which indicated that water  
538 temperature, salinity and wind also had a significant effect on microplastic abundance (Lusher et  
539 al. 2014; Lusher et al. 2015; Kanhai et al. 2017).

540

541 Comparative assessments between oceanic basins are critical in providing an indication of the  
542 extent of microplastic pollution in the marine environment. A major challenge, which demands  
543 caution when drawing conclusions from such comparisons, is the lack of standardization of  
544 microplastic sampling methods (depth of collection, mesh size of net/sieve, etc). Microplastic  
545 abundances in the present study were not normally distributed and therefore the median was  
546 reported as it is the most relevant measure of central tendency for such data. However, in order to  
547 enable comparability with other published studies, which generally did not report median  
548 microplastic abundances nor made statements about the normality of their data, the mean was also  
549 reported in Supplementary Table 7. In the present study, sub-surface waters (depth 8.5 m) in the

550 Arctic Central Basin (ACB), sampled via the bow water system, had a mean microplastic  
551 abundance of  $0.97 \pm 1.20$  particles  $m^{-3}$ . In comparison to other studies that employed similar  
552 methods (i.e. the underway system of vessels) to sample sub-surface waters, microplastic  
553 abundance in the ACB was lower than values reported (i) in the north eastern Pacific Ocean ( $279$   
554  $\pm 178$  particles  $m^{-3}$ ), (ii) in the North Atlantic Ocean ( $13 - 501$  particles  $m^{-3}$ ), (iii) off Svalbard  
555 ( $2.68 \pm 2.95$  particles  $m^{-3}$ ), (iv) in the north east Atlantic Ocean ( $2.46 \pm 2.43$  particles  $m^{-3}$ ), and (v)  
556 in the Atlantic Ocean ( $1.15 \pm 1.45$  particles  $m^{-3}$ ), (Desforges et al. 2014; Lusher et al. 2014; Enders  
557 et al. 2015; Lusher et al. 2015; Kanhai et al. 2017, Supplementary Table 7). The only oceanic basin  
558 for which there were reports of lower microplastic abundances in sub-surface waters was the Ross  
559 Sea ( $0.17 \pm 0.34$  particles  $m^{-3}$ ), (Cincinelli et al. 2017; Supplementary Table 7). Although the  
560 methods used to sample sub-surface waters for microplastics in the above mentioned studies were  
561 similar, the fact remains that the variation of several factors e.g. mesh size of sieve ( $1 - 300 \mu m$ ),  
562 sampling depth ( $3 - 11$  m), etc., amongst the studies could have impacted the reported microplastic  
563 abundances. In terms of assessing the vertical distribution of microplastics in the marine  
564 environment, Bagaev et al. (2017) was the only other published study which utilised a similar  
565 sampling method (Niskin bottles) in the Baltic Sea. Being cognisant of the fact that no  
566 confirmatory analytical techniques or blanks were used by Bagaev et al. (2017), microplastic  
567 abundance at multiple depths in sub-surface waters of the Arctic Ocean (mean:  $46 \pm 62$  particles  
568  $m^{-3}$ ; range:  $0 - 375$  particles  $m^{-3}$ ; depths sampled:  $8 - 4400$  m) was lower than reported for the  
569 Baltic Sea (mean:  $310 \pm 520$  particles  $m^{-3}$ ;  $70 - 2600$  particles  $m^{-3}$ ; depths sampled:  $1 - 218$  m).  
570 Similar to the findings of the present study whereby the highest microplastic abundances were  
571 found in the uppermost water layer i.e. the PML, Bagaev et al. (2017) reported that near- surface  
572 and near-bottom water layers in the Baltic Sea had higher fibre concentrations than intermediate

573 layers and that this was possibly due to greater turbulence and density stratification in those layers.  
574 Of note is the fact that microplastic abundance in deep waters of the ACB (0 – 104 particles m<sup>-3</sup>,  
575 depths 1000 – 4400 m, sieve 250 µm) was similar to those reported for deep waters at the Rockall  
576 Trough, North East Atlantic Ocean (70.8 particles m<sup>-3</sup>, depth 2227 m, sieve 80 µm), (Courteney-  
577 Jones et al. 2017). Overall, it must be acknowledged that an underestimation of microplastic  
578 abundance in the Arctic Central Basin (ACB) could have occurred in the present study as the mesh  
579 size of the sieve was only 250 µm leading to an exclusion of smaller sized particles. Nevertheless,  
580 the Arctic's remote geographic location away from major population centres, its low population  
581 in its surrounding continental shelves and relatively low in-situ anthropogenic activities (e.g.  
582 shipping) are all factors which may explain the lower microplastic abundances in sub-surface  
583 waters within the Arctic Central Basin (ACB). From an oceanographic perspective, the reduced  
584 contribution of Atlantic water in its upper water layers due to the dominance of the polar mixed  
585 layer is another possible explanation for lower microplastic abundances in the Arctic in  
586 comparison to other oceanic basins. Presumably more polluted, Atlantic water which originates  
587 from the more densely populated southern latitudes has its surface advance into the Arctic Ocean  
588 hindered due to freshwater released from melting ice and other physical barriers such as the sea  
589 ice itself and the Novaya Zemlya islands (Cózar et al. 2017). However, a plausible future scenario  
590 for the Arctic in the context of a changing climate is that microplastic abundance in near-surface  
591 layers of the Arctic Ocean may increase upon melting of contaminated sea ice and opening up of  
592 shipping lanes due to a decrease in sea ice extent (Obbard et al. 2014; Cózar et al. 2017).

593

594 Of interest is the fact that the present study managed to sample microplastics in sub-surface waters  
595 at approximately 8.5 m depth by two independent methods i.e. by the bow water system of the

596 vessel (underway sampling) and the rosette water sampler (CTD sampling). CTD samples (n = 9)  
597 retrieved from an average depth of 8.5 m indicated that microplastic abundance in the ACB ranged  
598 between 0 – 148 particles m<sup>-3</sup>, with a median of 20.8 particles m<sup>-3</sup>. By comparison, samples  
599 collected via the underway system (n = 58) at 8.5 m indicated that microplastic abundance in the  
600 ACB ranged between 0 – 7.5 particles m<sup>-3</sup>, with a median of 0.7 particles m<sup>-3</sup>. Although both  
601 methods sampled water at an average depth of 8.5 m, calculated microplastic abundances from  
602 both methods are not directly comparable due to the differences associated with the methods.  
603 Whereas underway sampling involved filtration of a greater volume of water (approximately 2000  
604 L) over a longer distance and a longer sampling time (> 2 hours), the CTD sampling involved the  
605 collection and subsequent filtration of a smaller volume of water (21 L or 48 L) at a single location  
606 in a shorter period of time (minutes). The advantage of using the underway system is that  
607 microplastic abundances over a larger spatial area can be quantified whilst the vessel is in transit.  
608 By comparison, CTD sampling facilitates the quantification of microplastic abundance at specific  
609 locations making it less likely to mask contamination hotspots. However, some of the major  
610 limitations associated with CTD microplastic sampling are (i) the vessel must stop at sampling  
611 stations to collect samples, (ii) deployment and retrieval of the rosette water sampler is time  
612 consuming and, (iii) only small volumes of water can be collected in comparison to the underway  
613 sampling. The limitation of filtering smaller volumes of water is twofold in that there can be (i)  
614 false negatives whereby microplastics are not sampled despite being present in the environment  
615 or, (ii) microplastics are found in the samples but scaling up to relevant units (particles m<sup>-3</sup>) has a  
616 greater effect on microplastic abundances.

617

618 One of the major challenges that investigators face when quantifying marine microplastic  
619 abundance is sample contamination. In addition to employing strict measures to control  
620 contamination during sampling and processing, it is important that checks are carried out to  
621 quantify potential contamination of samples. For underway samples in the present study, although  
622 method blanks were free of contamination by synthetic particles, a single synthetic fibre was found  
623 in one air contamination check. In context, there were between 0 – 15 synthetic particles in each  
624 underway sample, with an average of 2 synthetic particles per sample. For CTD samples in the  
625 present study, between 0 – 3 synthetic fibres were found in the method blanks. In context, between  
626 0 – 18 synthetic particles were found per CTD sample (21 L or 48 L), with an average of 2 particles  
627 per sample. In both cases, if contamination were an issue, its contribution to the reported  
628 microplastic abundances in the present study would be substantial. However, the possibility of  
629 airborne contamination in the actual underway samples is projected to be low since (i) 92 % of the  
630 air contamination checks (11 of the 12 petri dishes) were free of synthetic particles and, (ii) air  
631 contamination checks had maximum exposure to the atmosphere while actual samples had  
632 minimal exposure. With respect to the CTD samples, it is proposed that synthetic fibres in the  
633 method blanks may have been introduced into the Niskin bottle during the transfer of Milli-Q water  
634 or could have been present from the previous CTD cast and remained in the bottle due to  
635 insufficient rinsing with Milli-Q water prior to the blank. This should not have been an issue for  
636 the actual samples since Niskin bottles were rinsed during the downcast and were closed within  
637 the water column thus preventing the possibility of airborne contamination.

638

## 639 **5.0 Conclusion**

640 The present study demonstrated the pervasiveness of microplastics in sub-surface waters of the  
641 Arctic Central Basin (ACB). Two independent sampling techniques led to the discovery of  
642 microplastics in near surface waters of the Polar Mixed Layer (PML) i.e. at a single depth of 8.5  
643 m as well as throughout the water column i.e. at multiple depths (8 – 4369 m) of the ACB. Such  
644 findings confirm that microplastics are entering the central Arctic Ocean, that they are being  
645 vertically transported out of surface waters and that the water column is one of the reservoirs of  
646 microplastics in this region. Presently, however, uncertainty exists regarding the actual  
647 mechanisms responsible for the vertical transport of microplastics in the Arctic Ocean. Although  
648 there was a predominance of fibrous microplastics, the majority of which were polyester, the exact  
649 sources of microplastics to the Arctic Ocean remain unknown as they could have been introduced  
650 to the ecosystem via long range transport processes or originated from more local sources. The  
651 fact that the highest microplastic abundances were recorded in the PML nearer to the periphery of  
652 the ACB suggests the influence of location-specific factors e.g. absence of sea ice, proximity to  
653 microplastic sources, wind, etc. Knowledge about microplastic abundance, distribution and  
654 composition in the Arctic Ocean is vital as it provides (i) quantitative data on the concentrations  
655 and types of microplastics that polar organisms are exposed to, (ii) a sound starting point for  
656 investigating the potential threat that microplastics pose to the Arctic ecosystem and, (iii) insight  
657 into the whereabouts of the ‘missing plastic’ from oceanic surface waters.

658

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