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Accumulation and distribution of microplastics in coastal sediments from the Inner Oslofjord, Norway --Manuscript Draft--

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HIGHLIGHTS

- Microplastics were detected in marine sediments of the western Inner Oslofjord.
- The distribution of microplastics in sediments was statistically analysed.
- Considerations were made on the release of microplastics from a WWTP.
- Microplastics can be redistributed in the fjord by shallow and deep currents.
- The occurrence of microplastics in sediments could affect benthic fauna.

Accumulation and distribution of microplastics in coastal sediments from the Inner Oslofjord, Norway

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Abstract

Microplastic presence in benthic marine systems is a widely discussed topic. The influence of the natural matrix on microplastic distribution within the sedimentary matrix is often overlooked. Marine sediments from the western Inner Oslofjord, Norway, were investigated for temporal trends, with a particular focus on the relationship between sediment grain-sizes and microplastic distribution. Density separation, optical microscopy and chemical validation were used to categorize microplastics. Microplastic concentrations ranged from 0.02 to 1.71 MPs g⁻¹ dry weight (dw). Fibres were the most common (76%), followed by fragments and films (18%, 6%). Common polymers were polyesters (50%), polypropylene (18%), polymethylmethacrylate (9%), rayon and viscose (5%) and elastane (4%). Microplastics appear to accumulate preferentially according to their morphology and polymer type in certain sediment grain-sizes. Microplastics inputs to the Oslofjord appear to derive from a wastewater treatment plant in the vicinity. Although, the redistribution of microplastics within the fjord needs further investigation.

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1. Introduction

Plastic products have a fundamental role in everyday life, with many applications from packaging and textiles, to building, automotive, and within the medical field (PlasticsEurope, 2019). Versatility and low production costs have fuelled the global demand of plastic items, reaching an annual production of 368 million tons in 2018 to which Europe contributes with 57.9 million tons (16% of the global production; PlasticsEurope, 2020). The increasing production rates have been linked to the growing concern of plastics escaping waste handling and entering the environment (Borrelle et al., 2020). Plastics can be moved between terrestrial and aquatic reservoirs, with marine environments acting as a recipient of discarded plastic items, both from land and sea (Van Sebille et al., 2012; Martin et al., 2017; Law, 2017).

33 Plastic debris can be found in the marine environment in a wide variety of sizes, ranging from meters to mi-
34 crometres (Barnes et al., 2009). Microplastics are described as a “*heterogeneous mixture of differently shaped*
35 *materials referred to as fragments, fibres/filaments, beads/spheres, films/sheets and pellets*” (EFSA, 2016;
36 Lusher et al., 2017a). In the current paper we defined them in the 1 μm to the 1000 μm size range as suggested
37 by Hartmann et al. (2019). Microplastics can enter the environment as “primary” and “secondary” microplastics
38 (Andrady, 2011; Cole et al., 2011; Hidalgo-Ruz et al., 2012). Whilst secondary microplastics result from frag-
39 mentation and weathering of larger plastic items, primary microplastics result in the direct release of small
40 particles such as pellets, powders, and fibres (Sundt et al., 2014; Lebreton and Andrady, 2019). Primary mi-
41 croplastics, specifically designed to be small in their application, such as microbeads incorporated into per-
42 sonal care products that can be carried with wastewaters via sewers and released to aquatic systems (Napper
43 and Thompson, 2016; De Falco et al., 2019). Most treatment schemes employ an initial screening of influent
44 to eliminate macrodebris and settling to remove dense particles and grit. During this first step it has been
45 estimated that the 78% of microplastics can be removed, whilst sequential steps can often remove the remain-
46 ing 20% (Hale et al., 2020). Microplastics density, size and shape, highly influence retaining capacities of
47 wastewater treatment plants (WWTPs). Similarly, the filters employed at WWTPs are not specifically designed
48 to retain microplastic fibres which may escape the filtering process and can be released within effluent. Re-
49 search has suggested that effluent may contribute to 35% of the world ocean’s microplastic burden (Boucher
50 and Friot, 2017; Salvador Cesa et al., 2017; Long et al., 2019).

51 The fate of microplastics in the marine environment is influenced by several complex biotic or abiotic processes
52 (Van Sebille et al., 2020). Firstly, the physical characteristics of plastic polymers - including their density - play
53 a fundamental role in the vertical distribution of microplastics in the water and in benthic habitats (Murray and
54 Cowie, 2011). Whilst low-density microplastics tend to be buoyant, plastics with a density that exceeds that of
55 seawater ($> 1.02 \text{ g cm}^{-3}$) will readily sink (Van Cauwenberghe et al., 2015). Additionally, concurrent processes
56 such as additives leaching out of microplastics, biofouling and incorporation within marine aggregates can
57 cause changes in density of floating microplastics, which may finally sink to the seabed (Cózar et al., 2014;
58 Martin et al., 2017). Marine sediments have been identified as a possible final destination for microplastics in
59 the marine environment (Woodall et al., 2014). Here bottom currents, sediment depth, sedimentation rate,
60 biota redistribution, submarine physiography, distance from the shoreline and human activities (Shen et al.,
61 2019; Harris, 2020; Canals et al., 2021), may further influence their distribution and accumulation.

62 As a general trend observed for coastal surface waters, microplastics in sediments appear to positively corre-
63 late with nearby population density (Van Cauwenberghe et al., 2015). In Norwegian coastal areas most of the
64 population as well as the main industrial and tourism centres are concentrated close to fjords (Gomiero et al.,
65 2019). In particular, fjords have the greatest sediment trapping efficiency of all coastal sedimentary environ-
66 ments (Smith et al., 2015). This is enhanced by the transition from freshwater to saltwater, where sediments
67 suspended in the fresh river water mix with saltwater and give rise to the so-called estuarine turbidity maximum
68 (Burchard et al., 2018). As an example, average numbers of microplastics identified in Norwegian fjords ranged
69 from 190 to 77,000 particles kg^{-1} and exhibit a median of ~ 7000 particles kg^{-1} (Noren, 2007; Kazmiruk et al.,
70 2018; Black et al., 2018; Haave et al., 2019; Singdahl-Larsen, 2019). This median value is an order of magni-
71 tude greater than that for shallow coastal environments or other

72 estuarine environments (Harris, 2019). Although the contamination rate of fjords sediments by microplastics
73 is substantial, the related research is patchy (Noren, 2007; Black et al., 2018; Kazmiruk et al., 2018; Haave et
74 al., 2019; Olsen et al., 2020). Considering currently available published literature, half of the research on mi-
75 croplastics distribution in fjord sediments was conducted on the Norwegian coast near Bergen (Haave et al.,
76 2019) and in the Bunnefjord, eastern Inner Oslofjord (Olsen et al., 2020). In the studies mentioned above, the
77 distribution and the behaviour of microplastics considering sediment's characteristics is generally overlooked.
78 There is the need to understand how microplastics distribute within the sediment column to have a general
79 overview of the possible areas of accumulation and how the natural matrix could influence microplastic occur-
80 rence (Martin et al., 2021).

81 Therefore, the aims of this study were to: i) optimize the approach to sampling microplastics in the Inner
82 Oslofjord sediments, by quantifying the distribution of microplastics in sediments within and between sites; ii)
83 understand sources and paths of microplastics and how these particles tend to accumulate in specific areas;
84 and, iii) explain how microplastics abundance varies concerning age of sediments and grain-sizes.

2. Materials and Methods

2.1. Study area

87 The Oslofjord is located on the south-eastern Norwegian coastline and it stretches 100 km northwards from
88 the Inner Skagerrak, from which it is separated by a sill at about 120 meters of water depth (Staalstrøm, 2005).
89 The whole fjord system has a surface area of approximately 1644 km² (Staalstrøm and Ghaffari, 2015) and it
90 is further divided into the Inner and Outer Oslofjord by the 12 km long Drøbak Sound, which develops with a
91 shallow sill (19.5 m water depth) at its northern end (Staalstrøm et al., 2012). The Inner Oslofjord, is a fjord
92 system consisting of two main basins, the Vestfjord on the west side and the Bunnefjord on the east side, with
93 a water depth exceeding 150 m in both basins and with an average depth of 49 m (Staalstrøm, 2005). The two
94 inner basins are divided by the Nesodden Peninsula and by two sills at 55 m water depth extending from the
95 northern tip of the Nesodden to the mainland (Lepland et al., 2010).

96 The seafloor is characterized by numerous small basins, shoals and ridges that extend in the NE-SW di-
97 rection, following fault patterns (Solheim and Grøne, 1983; Lepland et al., 2010). In the Inner Oslofjord area,
98 the sediment accumulation rate (SAR) may vary considerably from zero in areas of bottom erosion, to 2.6
99 mm/year or higher (Lepland et al., 2010). The uppermost part of the seabed's sedimentary sequence is typi-
100 cally very loose and rich in organic matter. Anoxic conditions have developed at the bottom of several deep
101 basins because of restricted deep-water renewals, and eutrophication resulting from a high supply of municipal
102 waste, organic material, pollutants and nutrients (Lepland et al., 2010). Deep water is renewed by the denser
103 water from the Outer fjord/Inner Skagerrak during the winter and early spring (October/November—April). The
104 main driving force is northerly winds over the Oslofjord/Inner Skagerrak area and the strength and duration of
105 northerly winds determine the amount of renewed water (Gade, 1968). Furthermore, the fine grain-size and
106 fluffy surface character of sediments in this depositional basin indicate weak bottom currents and limited bot-
107 tom transport. The main currents near the bottom have a north-south direction and originate from flow-topog-
108 raphy interaction (Staalstrøm and Ghaffari, 2015). In general, the water masses in the fjord are a mixture of

109 Skagerrak water and locally formed waters affected by rivers, where the latter constitute the principal supply
110 of water into the region (Baalsrud and Magnusson, 2002).

141 2.2. Sample collection

112 Surface sediments were sampled on the 7th November 2019 from three stations aboard the R/V Trygve Braarud
113 (University of Oslo, UiO). They were collected along a transect in the deep-water channel that runs from the
114 Vestfjordens Avløpssekskap wastewater treatment plant (VEAS WWTP) to the Inner Oslofjord inlet in proximity
115 of Håøya island (Figure 1). The locations of sediment sampling were decided according to prevailing tidally
116 driven water currents, suitable depth (~100 m water depth; Table 1), seabed topography and to avoid areas
117 subject to trawling (Green et al., 2020). Due to safety distance from VEAS diffuser (discharge point) and suit-
118 able depth conditions, two stations were placed northeast of the diffuser (ST3, 0 km from the diffuser) and
119 southeast of the diffuser (ST2, 0.4 km from the diffuser). The third station was located at Gråøyrenna (ST1,
120 11.2 km away from the diffuser).

121 Surface sediments were collected by a double Gemini corer (10 cm inner diameter) and were sampled using
122 a steel slicer which was rinsed between samples with seawater, and wooden spatulas, discarded after each
123 sample slice. The water-sediment interface was preserved in the cores. Both corer tubes were subsampled to
124 provide duplicate samples from each depth at each sediment station. Sediments were sliced into 1 cm layers
125 from 0 to 5 cm sediment column depth, resulting in a total of 10 samples for each station. Once sliced, samples
126 were placed into cleaned glass jars, rinsed with Reverse Osmosis-water (RO-water) in the laboratory, previ-
127 ously to fieldwork, and were stored at +10°C in a fridge until processing the following day. During fieldwork, 30
128 procedural blanks were collected. These consisted of sample jars which were kept open for the duration of the
129 core slicing. They were closed and stored in the same way as the samples.

130 2.3. Sample preparation and analysis

131 Wet sediments stored in glass jars were oven dried at 40°C. Dry sediment was subsampled in 10 g replicates
132 and moved into 50 ml Falcon tubes. A procedural blank was included between 5 samples to test background
133 laboratory contamination. During this phase of the work, procedural blanks consisted of empty 50 ml Falcon
134 tubes which were previously rinsed using RO-water following NIVA laboratory protocols.

135 To extract microplastics from sediments, a high-density Sodium Iodide (NaI) solution (1.8 g cm⁻³) was applied
136 because it is capable of isolating the majority of common polymer types from sediment matrices (Hurley et al.,
137 2018). Two extractions were performed per sediment sample (and corresponding field and procedural blanks).
138 Prefiltered-NaI solution was added to the Falcon tubes with the dry sediment. Similarly, the procedural blanks
139 (empty Falcon tubes) were filled with the high-density solution. The samples and blanks were shaken to mix
140 for 30 seconds, after which they were left to stand for at least one day, until turbidity dissolved. Sediments and
141 blanks were filtered from the solution with a vacuum filtering system (Büchner), samples were passed through
142 a 47 mm Ø Whatmann GF glass fibre filter (pore size: 1.6 µm). When clay clusters occurred, they were crushed
143 with a steel spoon between the first and the second extraction.

144 After sample preparation, a total of 170 glass fibre filters were analysed, 30 of which were field blanks and 10
145 were processing blanks. For the visual identification, a stereomicroscope (NIKON S'MZ745T') with a zoom
146 magnification of 7.5x and a long working distance of 115 mm was used. A magnification up to 50x was used
147 with a detection limit of ~30 μm . The microscope was equipped with a camera (Infinity 1, Lumenera) and
148 pictures were processed with 'Infinity Analyze' software. During this step, potential microplastics were counted
149 and sorted into fragments, films and fibres as suggested by Lusher et al. (2020). The longest (L) and shortest
150 (S) axes of particles were measured, and the colour of particles was recorded.

151 Lastly, chemical validation of potential microplastics was performed using single-point measurements with
152 Attenuated Total Reflectance – $\mu\text{FT-IR}$ ('Perkin Elmer Spotlight 400 $\mu\text{FT-IR}$ Microscope'). The ATR is a rapid
153 surface analysis method that can be used for particles down to a diameter of 2 mm (Renner et al., 2016). To
154 improve the quality of the spectra generated, particles were prepared for analysis using a diamond compres-
155 sion cell (DCC) accessory. Particles were carefully transferred from filter papers to the DCC with use of extra
156 fine micro forceps. The DCC was used to compress particles to a thin, homogeneous thickness. The DCC was
157 then loaded onto the $\mu\text{FT-IR}$ microscope stage for analysis. Measurements were obtained in transmission
158 mode at 4 cm^{-1} spectral resolution for the range 4,000 to 600 cm^{-1} . Spectra were produced from a composite
159 of 3 scans. Background measurements were taken before each batch of particle was analysed (Bråte et al.,
160 2020). Once spectra were obtained, they were compared to library spectra. This included the commercially
161 available libraries: PerkinElmer ATR Polymers library, STJapan Polymers ATR library; the BASEMAN library
162 (Pimpke et al., 2018); and, in-house curated libraries (including reference polymers, different textile materials,
163 and potential sources of laboratory contamination). Library search matches were compared against the tested
164 particle. The basic principle of library searching is to calculate a numeric score which ranges from 0 to 1 and
165 describes the difference of two spectra. In the case of two identical spectra, the score is equal to 1, whilst if
166 the two spectra differ from each other, the score decreases (Mecozzi et al., 2016). All spectra and matches
167 were manually inspected to confirm the match. Matches between the polymer and reference spectra which
168 exceeded 0.7 were automatically accepted, in accordance with current international procedures (e.g. Pimpke
169 et al., 2018; Knutsen et al., 2020; Olsen et al., 2020; Lusher et al., 2020). Microplastics with a match between
170 0.6 and 0.7 were subjected to further visual examination of spectra characteristics before being accepted or
171 rejected (Lusher et al., 2015; Comnea-Stancu et al. 2017; Kim et al., 2018). This additional analysis was
172 needed because values which fell >0.6 and <0.7 were often a consequence of bio-fouled particles or very thick
173 fragments (i.e. with a thickness $>100 \mu\text{m}$). A total of 45 (16% of the total confirmed) low-scoring spectra (>0.6)
174 were included based on visual inspection of the spectra.

175 All the particles isolated from the samples and the blanks were analysed for chemical characterization, except
176 those that were lost during processing ($n= 20$, 5.6%). This approach exceeds the recommendation for reporting
177 under European Union's Marine Strategy Framework Directive (MSFD) (Gago et al., 2016). Lost particles were
178 included in the final counts if their colour and form clearly fit within the classification of "Anthropogenic". This
179 follows the protocol of Lusher et al. (2020) when using visual to define microplastics. Transparent, light yellow
180 and black fibres were excluded from the data set to avoid any false positive as these are more likely to be of
181 natural origin.

182 After chemical validation the number of microplastics per sample was corrected to remove non-plastic materi-
183 als. Particles were removed if they produced spectra of natural particles, such as merino wool (n= 35), silk (n=
184 12), non-modified cellulose (e.g. identified as tissue paper) (n= 1). Rayon and viscose were considered micro-
185 plastics because of the origin of those fibres could be linked with the release of acetate fibres from cigarette
186 butts, sanitary products such as feminine tampons or garments (Wright et al., 2015). Field blanks did not
187 present evidence of procedural contamination during sampling, whilst only two fibres were observed in the
188 procedural blanks: these were identified as silk and merino wool. No sample correction was applied as these
189 materials were not considered to be plastic.

190 2.4. Contamination control

191 Strict controls were followed during sample collection and processing in the laboratory to eliminate post-dep-
192 ositional contamination. Prior to fieldwork, all sample glass containers were pre-washed thoroughly with RO-
193 water (filter Millipak® Express 20' for 0.22 μm particulate) before use. During sample collection, nitrile gloves
194 (NBR) and security coveralls were used, glass and metal equipment was used whenever possible.

195 Laboratory analyses were conducted in the "Mikroplast lab" at NIVA in Oslo, a dedicated laboratory for micro-
196 plastics analyses. Processing was performed in a fume hood/laminar flow whilst, microscopy work was per-
197 formed outside of the laminar flow bench. However, to minimize the contamination from the laboratory atmos-
198 phere, the samples on filter papers were enclosed in petri dishes while they were analysed under the micro-
199 scope. The upper lid was removed just only when a potential microplastics needed to be manipulated, or while
200 taking pictures.

201 Reagents were vacuum filtered through 'Whatmann glass microfiber filters GF/A' ($\text{\O} = 47$ mm, pore size: 1.6
202 μm) immediately prior to use. All equipment was cleaned with RO-water and the use of plastic laboratory
203 equipment was kept to a minimum. In the laboratory gloves are mandatory as personal protective equipment
204 (PPE; NIVA security protocols), NBR blue gloves were used. This polymer is resistant to oils and acids and
205 due to its colour is easily recognizable during optical identification. Any particles identified to match NBR pol-
206 ymer and morphology combination were excluded from further analysis.

207 2.5. Sediments analyses and sediment accumulation rate

208 Grain size analyses were performed at the Earth Science Department, University of Pisa. Sediments were
209 investigated with the particle size analyser: the 'CAMSIZER x2 (RETSCH technology, particle characteriza-
210 tion)'. The particle size analyser is based on the principle of dynamic image analysis which provides precise
211 particle size and shape information of sediment grains. Ultrabright LED stroboscopic light sources and two
212 high-resolution cameras achieve a frame rate than 300 images per second which are evaluated in real time
213 by the CAMSIZER x2 software. For the analysis of sediments, the X-Flow module was used. Thanks to this
214 feature, liquid suspensions of sediments can be investigated. An ultrasonic bath and a strong centrifugal pump
215 ensured efficient dispersion of particles contained in the suspension. With this analysis mode, particles within
216 a dimensional range of 0.8 mm and 1000 μm can be detected (CAMSIZER x2, user's manual).

217 Prior to grain-size analysis, sediments were stored in 50 ml Falcon tubes. Demineralized water was added to
218 each Falcon tube 48 hours before grain-size analysis in order to dilute the sediment suspension. The solution
219 was picked up with a 5 ml pipette and few drops were released in the dispersion bath of the particle size
220 analyser. From here, the solution traversed a close loop to the flow cell where the camera system captured
221 particle images, for which an upper limit of 30,000 images per cycle was set.

222 Sediment dating was calculated from two previous works, to assess the degree of microplastic burial within
223 the sediment column after deposition. In the work of Dolven et al. (2012), four cores were collected and dated,
224 three in the Bunnefjord (Ep1, Cp3, B18x) and one in the Vestfjord (Cj3; Figure 1). The samples were collected
225 in February and April 2009 and the sites were chosen considering the highest possible sedimentation rate and
226 the least disturbed locations. The samples were sent to the Gamma Dating Centre in Denmark and were
227 analysed for ^{210}Pb -, ^{226}Ra - and ^{137}Cs - activity via gamma spectrometry carried out on a Canberra Ultraslow-
228 background Ge-detector. The other work considered is the Decelles' (2019) master thesis. The core V-60-A18
229 was sampled in May 2018 using a double Gemini corer (Figure 1). The core was sent to the Environmental
230 Radioactivity Research Centre at the University of Liverpool for the radiometric dating. The samples were
231 analysed based upon direct gamma assay of ^{210}Pb , ^{226}Ra and ^{137}Cs radionuclides. From the data of these two
232 works, the mean sediment accumulation rate (SAR) for the Vestfjord was calculated.

233 2.6. *Multivariate analysis*

234 Past4.03 software was used to perform Canonical Correspondence Analysis (CCA, Legendre and Legendre,
235 1998). This type of multivariate analysis was used to examine linear relationships between the dataset and
236 environmental variables. The occurrence of different grain-sizes (clay, silt, sand) was considered as environ-
237 mental variable and was compared to the occurrence of fibres, films, fragments and the type of polymers
238 (polyesters, polypropylene, polymethylmethacrylates, etc.).

239 3. Results

240 3.1. *Plastic particles distribution*

241 After chemical confirmation, a total of 310 microplastics were found in all environmental samples. Considering
242 the full sum of plastic particles within each core, the majority of microplastics were found at ST2 (62%), followed
243 by ST3 (33%) and ST1 (5%). A total of 191 particles were identified at ST2 which represented a concentration
244 of 1.04 microplastics per gram of sediment dry weight (MPs g^{-1} dw). A total of 102 particles were found at ST3,
245 equating to 0.52 MP g^{-1} dw. The lowest concentration was found at ST1. Only 17 microplastics were found,
246 equating to 0.08 MPs g^{-1} dw (Figure 2).

247 Between sites, there appeared to be some differences between the number of microplastics at different depths
248 within each core. For example, there was a difference in abundance of microplastics when the sediment layers
249 (1 cm slices) were investigated at ST2.

250 The layer 4 – 5 cm appeared to be the most influenced by the presence of microplastics, 1.7 MPs g^{-1} dw,
251 followed by the superficial layer (0 – 1 cm) with 1.12 MPs g^{-1} dw. The layers 1 – 2 cm and 2 – 3 cm showed a

252 similar distribution with 0.94 and 0.91 MPs g⁻¹ dw respectively. Whereas, section 3 – 4 cm (0.53 MPs g⁻¹ dw)
253 had the lowest concentration of microplastics. The distribution of microplastics at ST3 decreases towards the
254 top of the core. The highest concentration was observed in 4 – 5 cm (0.80 MPs g⁻¹ dw) whereas the lowest
255 concentration was observed at 1 – 2 cm (0.34 MPs g⁻¹ dw). The remaining layers 0 – 1 cm, 2 – 3 cm, 3 – 4 cm
256 presented 0.46, 0.50 and 0.52 MPs g⁻¹ dw respectively.

257 The sediment layers with the highest microplastic concentrations at ST1 were 4 – 5 cm and 3 – 4 cm, with
258 0.17 and 0.10 MPs g⁻¹ dw each. The lowest microplastic concentration was found in layer 2 – 3 cm (0.02 MPs
259 g⁻¹ dw). This was followed by layer 0 – 1 cm (0.06 MPs g⁻¹ dw) and layer 1 – 2 cm (0.07 MPs g⁻¹ dw).

260 3.2. *Microplastics dimension and morphology*

261 The plastic particles found in Oslofjord's sediments had a wide range of dimensions. The smallest particle
262 detected with the optical microscope had the longest axis dimension of 35 μm, whereas the longest was 8754
263 μm long. Considering the longest dimension of particles (L), two main dimensional classes were identified:
264 microplastics (30 μm < L < 1000 μm) and mesoplastics (1000 μm < L < 10,000 μm). Microplastics were further
265 divided in subclasses. The size classes of 30 – 200 μm, 200 – 400 μm and 400 – 600 μm showed a similar
266 distribution, comprising of between 25% and 21% of the overall particle count. Abundances decreased in the
267 subclasses 600 – 800 μm (16%) and 800 – 1000 μm (14%). The subclass of 1000 – 2000 μm contained the
268 majority of the observed particles, with an abundance of the 54%. Largest particles were gathered in the 2000
269 – 3000 μm subclass and accounted for the 21%.

270 Fibres were the most common shape found, they accounted for 76% of all particles, followed by fragments
271 (18%) and films (6%). Fibres were pervasive throughout cores, whilst the distribution of fragments and films
272 were patchier.

273 Fibres accounted for the 59% of the particles at ST1, followed by films (24%) and fragments (18%) Fibres
274 were found in every layer and were most abundant in 1 – 2 cm (n= 3) and 4 – 5 cm (n= 4). Fragments were
275 only present in the layers 0 – 1 cm (n= 1), and in 4 – 5 cm (n= 2). Films were found in deeper layers: 3 – 4 cm
276 (n= 3) and 4 – 5 cm (n= 1).

277 Similarly, fibres accounted for 78% of the particles found in core ST2, whilst fragments and films accounted
278 the remaining 19% and 3% respectively. Generally, fibres were abundant in every layer, although they de-
279 creased in layer 3 – 4 cm. Fibres and fragments were more abundant in layer 4 – 5 cm where 38 fibres and
280 13 fragments were counted. Films occurred in layers 1 – 2 (n= 2), 2 – 3 (n= 2) and 4 – 5 cm (n= 1).

281 Fibres were found in all layers at ST3, but were more abundant in layer 4 – 5 cm (n= 25). No other particle
282 morphologies were observed in the first layer (0 – 1 cm) whereas in layer 1 – 2 cm three fragments were
283 present in addition to fibres. The abundance of fragments decreased from layer 2 – 3 cm (n= 6) to 4 – 5 cm
284 (n= 3), whereas the distribution of films accounted for one particle for each of these last layers.

285 Considering all stations and all intervals, the most frequent colours observed were transparent (37%), blue
286 (21%), black (13%), green (10%), red and yellow (7% each). Few particles were pink (3%), orange (1%) and
287 brown (< 1%).

288 3.3. Chemical characterisation

289 A total of 359 potential microplastics were initially counted, of this number a total 285 (79%) particles were
290 chemically confirmed as microplastics. Due to sample loss, 21 particles were just visually confirmed and not
291 excluded from plastics counts, 4 particles were confirmed as plastics but could not be assigned to a polymer
292 (Table 2). Lost and undetermined particles accounted together for the 7%. In total across every station, the
293 most abundant polymer was polyesters (PES) (50%, 154 particles) followed up by polypropylene (PP) (18%,
294 56 particles), acrylates and polymethylmethacrylates (A+PMMA) (9%, 26 particles). These three polymers
295 represented more than 75% of all particles (Figure 3).

296 3.4 Sediments analyses

297 Sediment grain-size distribution was similar between samples in the Oslofjord. Generally, cores were domi-
298 nated by both silt (from very fine to very coarse) and sand (from very fine to medium) (Friedman and Sanders,
299 1978). A low portion of clay was detected (1 – 4%). The range between coarse silt and fine sand was the most
300 common, with some exceptions. The silt/sand ratio was favourable for the sand grain-size in 9 layers of the 15
301 total layers analysed between stations (ST1, ST2, ST3). The end members (clay and coarse sand) were found
302 only in few layers. Overall, ST1 and ST3 were characterised by relatively coarser sediments, whilst the abun-
303 dances of finer sediments were slightly higher for ST2.

304 3.5. Age of sediments

305 An average SAR of 0.296 ± 0.02 cm/year was calculated for the Oslofjord with varying accumulation rates
306 found throughout the fjord. The maximum of 0.42 and 0.36 cm/year were observed at station Ep1 and Cj3
307 respectively. The station Cp3 and V-60-A18 showed a similar pattern of approximately 0.20 cm/year (Figure
308 4).

309 The age of sediments for the ST1 layers was calculated considering the 0.29 cm/year average accumulation
310 rate of the Oslofjord. The year 2019 was set as the upper limit of the 0 – 1 cm layer, thus 2015 was identified
311 as the lower limit. Starting from here, the lower limits of the core bottom layer (4 – 5 cm) were individuated at
312 2005 for the upper boundary and 2002 for the lowest one.

313 Considering the estimated SAR for station V-60-A18 (0.2 cm/year), the age of sediment layers was calculated
314 for both ST2 and ST3 due to the proximity of the three stations. As for ST1, 2019 was considered as the upper
315 boundary of layer 0 – 1 cm, then 2014 was identified as the lower limit. Lastly, 1994 was individuated as the
316 lowest boundary of the core bottom layer (4 – 5 cm) with 1999 as the upper limit of this last layer.

317 3.6. Multivariate analysis

318 From CCA was observed that plastic particles appeared to accumulate preferentially according to their mor-
319 phology (Figure 5). Fibres and fragments were more influenced by sandy sediments, whilst films accumulated
320 mostly in silty sediments. Polymers too showed a preferential accumulation according to sediments. Such as
321 PES, low-density polyester (LD-PE), expanded polystyrene and polystyrene (EPS+PS) and polyvinylchloride
322 (PVC) which settled in finer grain-sizes (silt), like A+PMMA, elastane mix (E), rayon and viscose (R+VI), poly-
323 amide (PA) and polycarbonate (PC) but with a stronger correlation with clay. PP is the only polymer that oc-
324 curred preferentially in sand, whilst VI accumulated in fine sediments but did not show a preference between
325 silt or clay.

326 4. Discussions

327 4.1. Visual identification quality

328 Of the 359 potential microplastics identified by visual microscope assessment, 285 particles (79%) were con-
329 firmed as microplastics using FT-IR analyses. Whilst 21 particles (6%) could not be chemically confirmed as
330 they were lost prior to chemical inspection.

331 The protocol presented in Lusher et al. (2020) allowed the identification of potential microplastics under an
332 optical microscope. It addresses the definition of particles' morphology, considering their optical properties,
333 whilst defining their physical behaviour manipulating them with forceps during the analysis. The protocol has
334 been deemed efficient, especially considering fragments (59 potential, 50 chemically confirmed, 85% correct)
335 and fibres (255 potential, 217 chemically confirmed, 85% correct). Bright and anthropogenic colours (red, blue,
336 green, yellow, etc.) appeared to be reliable markers of a particles' polymeric origin, even considering very
337 small particles – e.g. the smallest fragment found (35 μm) was identified thanks to its greenish colour (con-
338 firmed with FT-IR as PP). Transparent fragments and fibres were visually identified and were the two most
339 common combinations (37%) among the other morphotypes characterised by different colours. However, no
340 transparent fragment $<250 \mu\text{m}$ were found. It is possible that smaller transparent fragments may have been
341 overlook, and thus underestimated.

342 Some issues were encountered for films when applying visual identification: of 28 potential plastic films, 18 of
343 these were confirmed with FT-IR analysis. Discarded films were all fragmented, yellowish in colour, easily
344 foldable and, when folded, did not broke. These characteristics can be easily shared with both anthropogenic
345 films and organic matter (e.g. algae). For these latter, FT-IR spectra were characteristic of material with a
346 cellulose origin. The best match reference spectra were those of silk and wool, although it is not possible to
347 differentiate these from natural cellulose, such as algae. This misinterpretation could be either related to inex-
348 perience of the user or operation error, meaning that sufficient training and or experience is required for mi-
349 croplastic identification in environmental samples. This supports conclusions by several authors that visual
350 identification alone (without the use of confirmation techniques) cannot be used for the identification of smaller
351 microplastics ($<1 \text{ mm}$) (Isobe et al., 2019; Lusher et al., 2020; AMAP 2021). Lastly, FT-IR confirmation anal-
352 yses allowed users to not overestimate microplastic abundances, as seen here with films.

353 4.2. Abundances

354 The abundances and the distribution of microplastics between stations in the Oslofjord varied widely even
355 when considering stations in proximity (i.e., 0.37 km between ST2 and ST3). Microplastics were found at every
356 station and within every core. The sampling site downstream and close to the VEAS WWTP diffuser (ST2)
357 was the most polluted location identified in this study, where a total of 104 particles were identified in 100 g of
358 sediment (0 – 5 cm, normalized values). This value was not reflected at other sites within the same quantity of
359 sediment ($n_{ST3}= 52$, $n_{ST1}= 8$; normalized values). Interestingly, there was a difference in microplastic numbers
360 between ST2 and ST3 even with the close proximity. The abundance of plastic particles at ST2 was doubled
361 that of ST3 One of the principal sources of microplastics in this area is believed to be the VEAS WWTP system.
362 A diffuser releases treated water into the Oslofjord via an outlet tunnel and five long distribution pipes in prox-
363 imity to the stations downstream (ST2) and upstream (ST3). This outlet spreads the treated effluent from a
364 depth of 20 metres (<https://www.veas.nu/produkter/vann>). The distribution of outlet waters was assessed be-
365 fore building and the operation of the system during a three-day dye drop experiment (Bjerkeng et al., 1978).
366 The flow from VEAS showed that the dye released in the Oslofjord moves generally toward the southeast,
367 carried by tidally driven surface waters (Staalstrøm and Røed, 2016), with varying concentrations considering
368 the distance from the discharge point. The finding of the dye drop experiment appeared to be consistent with
369 the quantity of microplastics found at ST2 which is located more southerly than the VEAS diffuser. ST3 would
370 be expected to have a higher occurrence of microplastics because it is located close to the diffuser. However,
371 the results of this study showed a reverse distribution considering ST2 and ST3. Considering that the micro-
372 plastic composition was mostly dominated by fibres and by polymers that are less dense than seawater, it
373 could be expected that microplastics did not readily sink to the seafloor but tended to be buoyant in the water
374 column for a period of time. The average SAR for these two sites was calculated to be of 0.2 cm/year but these
375 estimates can be valid for a mineral grain, such as spherical quartz, which can have a density of 2.65 g cm^{-3}
376 (Harris, 2020). The polymer with the highest density observed during analysis was polystyrene with a density
377 of 1.05 g cm^{-3} in the non-expanded form. Thus, the SAR for microplastics is expected to be even lower than
378 that for sediment particles, even when microplastics are weathered or biofouled.

379 Additionally, when considering the seafloor geomorphology of the area, it is more likely that microplastics
380 accumulate in proximity of ST2 than of ST3. Indeed, whilst the first site is located in an almost flat area with a
381 bathymetric high in its southernmost part, ST3 is located in proximity of a threshold where the basin gets
382 deeper below the sampling station. It is predictable that the major burden of microplastics which pollutes that
383 area will probably be found in the bottom part of this enclosed area.

384 The deepest site sampled in the Oslofjord was ST1 ($113.5 \pm 0.38 \text{ m}$) which also contained the lowest number
385 of microplastics ($n= 17$, 0 – 5 cm). Interestingly, this site is influenced by the action of both superficial and
386 bottom currents which cause major changes of water masses between the Outer and the Inner Oslofjord.
387 Between September and November, the deep water entering the Inner Oslofjord from the Outer fjord, has a
388 velocity of 0.25 m/s at a water depth between 111 m and 116 m with the strongest current in the north-south
389 direction (Staalstrøm and Ghaffari, 2015). This current splits into two main northward branches, due to the
390 presence of several islands in the Oslofjord inlet. Plus, the presence of the Drøbak Sound further reduces the
391 strength of this bottom current. The current that flows above ST1, has a flux diffusivity rate of $\sim 2 \text{ cm}^2/\text{s}$ at a
392 depth between 90 and 125 m (Staalstrøm et al., 2012), resulting in an area more influenced by bottom sedi-
393 ment transport. Thus, it is unlikely for high numbers of microplastics to accumulate in this area. Instead, surface

394 currents show a maximum strength outward from the Inner Oslofjord (Albretsen et al., 2004). Collignon et al.
395 (2012) observed that strong currents could increase the mixing and the vertical redistribution of microplastics
396 in the upper and mid layers of the water column. Thus, the strong overflowing current could even transport
397 slowly sinking microplastics that will accumulate on the seafloor when increasing in density or when flow con-
398 ditions allow settling. It is thus uncertain if the main supply of microplastics at ST1 comes from the Outer
399 Oslofjord via bottom currents and turbulent flows, or if even sub-superficial currents contribute to the transport
400 and deposition of microplastics. As only one sample was analysed in this area, further research is required to
401 better understand the pattern of distribution of microplastics and elucidate the hypothesis.

402 4.3. Vertical distribution and plastic particles characteristics

403 Following the estimation of the SAR, microplastics were calculated to accumulate in the Oslofjord sediments
404 over the last 26 years (Table 2). Sediments at ST1 were 8 years younger than those of ST2 and ST3. Indeed,
405 the bottom age for core ST1 was calculated to be 2002, whereas 1994 was calculated for both ST2 and ST3.
406 The layer 4 – 5 cm of ST1 settled in correspondence of the transition between layer 3 – 4 cm and 2 – 3 cm of
407 ST2 and ST3. From the analysis of microplastics abundance between cores, this correlation seems to be
408 confirmed because the relative distribution of fibres, films and fragments follow the same distribution in these
409 layers. A general trend can be observed between sites, especially considering ST1 and ST3 where the two
410 cores show a general decreasing accumulation of microplastics from 2002 to 2014/2015. This is in agreement
411 with the study performed in the eastern Inner Oslofjord (Singdahl-Larsen, 2019). Additionally, the lowest quan-
412 tity of microplastics deposited for both ST1 and ST3 was for the year 2009. This trend is reversed at ST2, as
413 the quantity of microplastics increased from 2002 to 2014 suggesting a higher accumulation capability for
414 microplastics in the ST2 area.

415 The present study shows that settling of microplastics is not sufficient to explain their distribution in benthic
416 habitats. Benthic organisms were not investigated here although their influence on the distribution of micro-
417 plastics cannot be ruled out. It is well known that bioturbation has a key role in the vertical and horizontal
418 redistribution of microplastics in sediments (Graham and Thompson, 2009; Taylor et al., 2016; Courtene-
419 Jones et al., 2017; Näkki et al., 2017; Bour et al., 2018). The intensity of bioturbation is dependent on species
420 composition due to the difference on specific characteristics such as typical burrowing depth and habitat of
421 certain species (Josefson et al., 2012). The small size of microplastics makes them available for interactions
422 with marine biota in many other ways including feeding mode and thus ingestion/egestion at different trophic
423 levels (Wright et al., 2013). Existing data indicate that microplastics are widely distributed in digestive tracts of
424 a range of organisms living in marine benthic habitats (Gonçalves et al. 2019; Iannilli et al., 2019; Sfriso et al.,
425 2020). As an example, in a recent study of Bråte et al. (2020) for the Nordic Council of Ministers, several Nordic
426 environments were analysed. Many stations were sampled in the Baltic Sea, in Denmark, on the Norwegian
427 Coast, in Faroe Islands, Iceland and Greenland. Between all these locations, it was demonstrated that the
428 species inhabiting the Inner Oslofjord are the most polluted ones between those analysed (*Limelicola balthica*,
429 *Abra nitida* and *Thyasira* spp., *Mytilus* spp.). An average of 61 microplastics per individual was reported. In a
430 similar study, Bour et al. (2018) highlighted that the ingestion rate and occurrence rate of microplastics in
431 deposit-feeders (*Ennucula tenuis*, *Ophiura alida*, *Brissopsis lyfera*, *Hediste diversicolor*) was equal that of
432 predators (*Hippoglossoides platessoides*, *Enchelyopus cimbrius*, *Trisopterus esmarki*), but higher than that for

433 filter-feeding species (*Amphiura filiformis*, *Sabella pavonina*, *Crangon allmanni*). Authors themselves sug-
434 gested to improve the investigation of the distribution of microplastics in water and sediments of the Oslofjord
435 to evaluate how biological or environmental factors could explain their results. Based on this information, the
436 benthic fauna from the sampled locations in the Oslofjord may play a role in the redistribution of microplastics
437 and future research should investigate benthic fauna in the same locations.

438 Considering the morphological and distributional characteristics of microplastics, it appeared that fibres were
439 pervasive throughout the cores, whereas films and fragments tended to accumulate in deeper layers. Moreo-
440 ver, due to the proximity of inhabited areas, it was expected that beads and pellets would be found in sediment
441 samples, although none were observed. It is possible that the efficiency of the WWTPs that operate in the
442 Oslofjord may have contributed to this observation. Both VEAS (western Inner Oslofjord) and Bekkelaget
443 (BRA, eastern Inner Oslofjord) WWTPs have tertiary treatment facilities meaning that waters undergo both
444 chemical and biological treatment steps. For instance, differently from secondary treatments the biological step
445 in these tertiary systems employs nitrogen removal from wastewaters, whilst in primary systems waters just
446 passes through fine screens (~350 μm in Tomasjord WWTP, Tromsø). Additionally, the VEAS system has a
447 separate treatment for excess stormwater which employ mechanical and chemical cleaning processes (Lusher
448 et al., 2017b). Research has shown that sludge produced from these two of the WWTPs operating in the
449 Oslofjord region (VEAS and Bekkelaget) were capable of retaining a total of 965,535,470 particles/ m^3 per day,
450 mostly composed by beads and fragments. The average particle size of fragments identified in sludge samples
451 was 414 μm for Bekkelaget sludge and 312 μm for VEAS sludge.

452 Microplastics analysed in the present study from the Oslofjord had an average dimension of 256 μm . Thus, it
453 is possible that smaller plastic fragments are not retained by WWTP filtering systems and can be easily re-
454 leased into the marine environment from diffusers (Browne et al., 2011; Carr et al., 2016). As an example,
455 setting the lower microplastics dimensional limit at 400 μm , 81% of the microplastics identified in this study
456 may not be retained in WWTPs and if we consider 300 μm as the lower limit for the longest microplastic's axis,
457 65% of microplastics will be released from both diffusers.

458 Plastic fibres constitute the largest burden of microplastics in Oslofjord sediments. Although larger debris is
459 removed in WWTPs, filters are not specifically designed to retain them (Zubris and Richards, 2005). As stated
460 above, the WWTPs that operate in the Oslofjord, especially Bekkelaget system, have a higher retaining ca-
461 pacity for beads ($n_V= 23$, $n_B= 93$) and fragments ($n_V= 10$, $n_B= 80$) than for fibres ($n_V= 11$, $n_B= 35$; Lusher et al.,
462 2017b). Some experiments demonstrated that a single garment could produce >1900 fibres per wash (Browne
463 et al., 2011), the direct consequence is that a large portion of microplastic fibres which were found in Oslofjord
464 sediments may have derived WWTPs which showed low capacity to retain fibres.

465 Many polymer types were identified in the Oslofjord, with fibres mostly composed by PES (68%) and PP (33%),
466 which was more common for fragments (36%) and films (33%). Generally, PP and PES are very versatile
467 polymers, together they constitute the most demanded resin types globally. Both are used for the clothing
468 production, but PP is also used for food packaging, hinged caps, microwave containers, pipes, etc. For PP,
469 the 2018 annual demand accounted for the 19.3%. PES differentiates for the low- and high- to medium-density
470 forms, it is generally employed for reusable bags, trays and containers, agricultural film, food packaging, etc.

471 in its lighter composition (17.5% 2018 global demand). Whilst the heavier form is used to produce toys, milk
472 bottles, shampoo bottles, pipes, houseware, etc. (12.2% 2018 global demand; PlasticsEurope, 2019). A major
473 part of microplastics analysed in Oslofjord samples looked weathered (51%) and biofouled (28%), indicating
474 a high residence time in seawater before they were able to reach the benthic domain. Lastly, strong correla-
475 tions have been observed from CCA between polymers and sediment grain-size, such as the relationship
476 between PES and fine sediments, PP with sand grain-sizes and clay with PMMA and E.

477 **5. Conclusions**

478 Microplastics have been found in surface sediments (0 – 5 cm) within every sample analysed in this study from
479 the Inner Oslofjord. Considering the sum of all the stations (ST1, ST2, ST3), the total concentration of micro-
480 plastics was about 0.55 MP g⁻¹ dw. The stations upstream (ST3) and downstream (ST2) of the VEAS WWTP
481 diffuser were the most polluted stations, whilst the furthest station outward of the fjord (ST1) showed a low
482 occurrence of microplastics, probably due to the high hydrodynamics of the location in which this sample was
483 collected. Sites ST1 and ST3 showed that the concentration of microplastics decreased from 2002 and
484 2014/2015, indicating a reduction in release of microplastics or an improvement in WWTPs systems' retaining
485 capacity. Reversely, at ST2 the amount of microplastics increased, likely indicating an area susceptible for
486 microplastic accumulation.

487 Besides sinking of microplastics driven by increasing of density, it was supposed that the main transport agents
488 in the locations analysed are shallow currents which redistribute microplastics between locations. For ST1,
489 further analyses are needed to understand the main direction from which microplastics reach this area and
490 thus if the main input is from the Inner or the Outer Oslofjord. From the correlation between microplastics
491 characteristics and composition of sediments found in every sample analysed in this study, it was observed
492 that there are some peculiarities in the deposition of microplastics. For example, fibres and fragments will
493 preferentially accumulate in relatively coarse sediments such as sand from very fine to coarse, whilst films
494 accumulate preferentially in fine sediments like silt. Acrylates, polyesters, elastane, rayon and viscose, poly-
495 amides and polystyrenes will accumulate preferentially in finer sediments such as clay and silt, while polypro-
496 pylene was the only polymer that occurred in sandy sediments in our study.

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Stations	ST1	ST2	ST3
North	6619347.25	6629374.13	6629751.75
East	586924.94	585438.94	585432.41
Depth (m)	113.5	98.5	100

Table 1: Coordinates (in UTM Zone 32N) and depths of sampled stations in the Inner Oslofjord.

	ST1	ST2	ST3	Total
Potential MPs	22	202	135	359
Confirmed MPs	17	191	102	310
Lost	2	11	8	21

Table 2: Quantities of microplastics confirmed across all sampling locations.

ST1 layers (cm)	year	SAR (cm/year) = 0.297	ST2-ST3 layers (cm)	year	SAR (cm/year) = 0.2
0 - 1	2015		0 - 1	2014	
1 - 2	2012		1 - 2	2009	
2 - 3	2009		2 - 3	2004	
3 - 4	2005		3 - 4	1999	
4 - 5	2002		4 - 5	1994	

Table 3: Age of sediments along the layers of ST1, ST2 and ST3 cores.

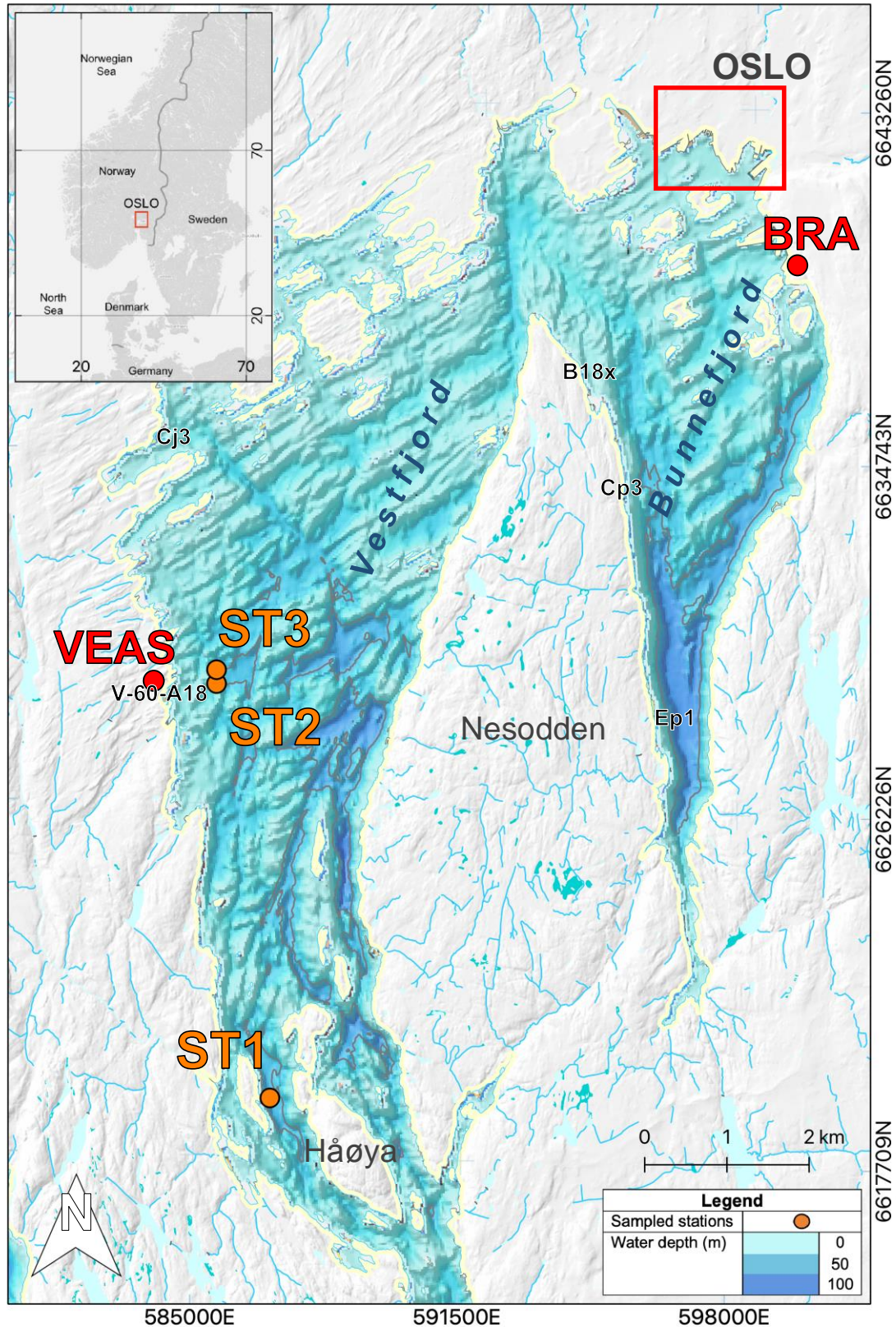


Figure 1: Depth-coloured shaded-relief image of the Inner Oslofjord and sampled stations. Data are derived from the high-resolution bathymetric dataset collected by the Geological Survey of Norway (water depth 0 – 100 m). Red dots indicate the location of the two main WWTPs operating in the Inner Oslofjord, VEAS (Vestfjordens Avløpsselskap) and BRA (Bekkelaget). Dark dots indicate the locations of the dated cores: Cj3, B18x, Cp3, Ep1 (Dolven et al., 2012) and V-60A18 (Decelles, 2019).

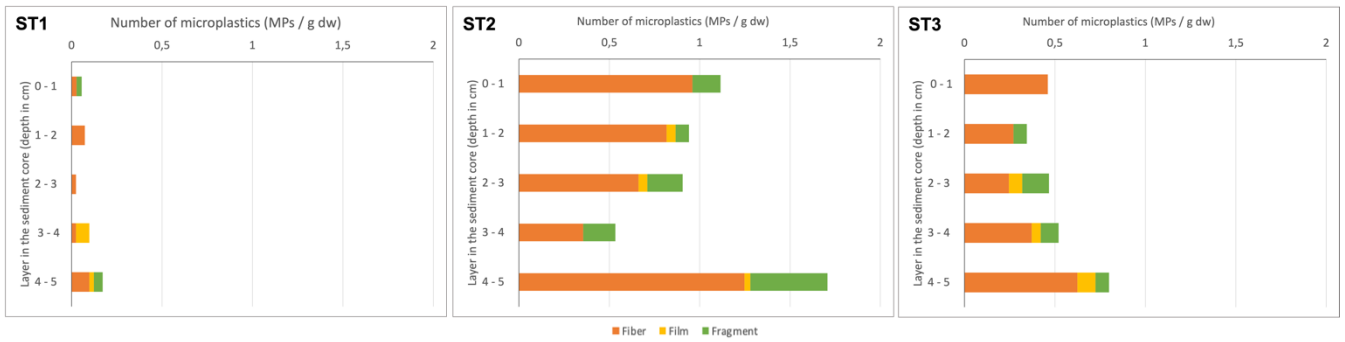


Figure 2: Distribution of plastic particles in Oslofjord samples and layers of each core. Morphologies (fibres, films and fragments) plus microplastics abundances are normalised to MPs / g d.w.

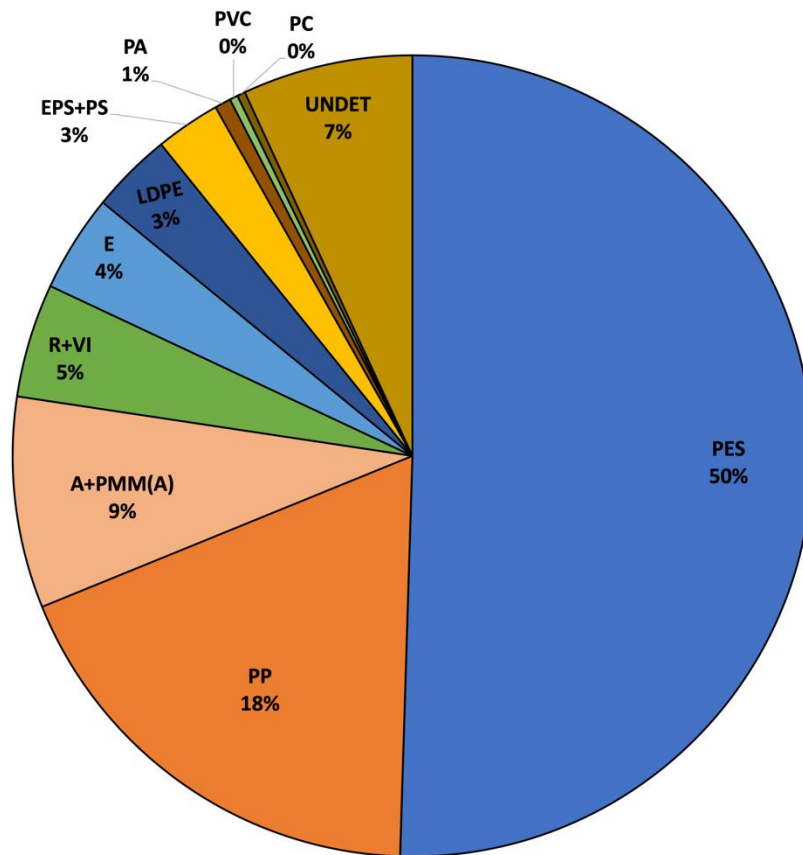


Figure 3: Polymers detected in all Oslofjord samples (ST1, ST2, ST3). Abundances are expressed in percentages.

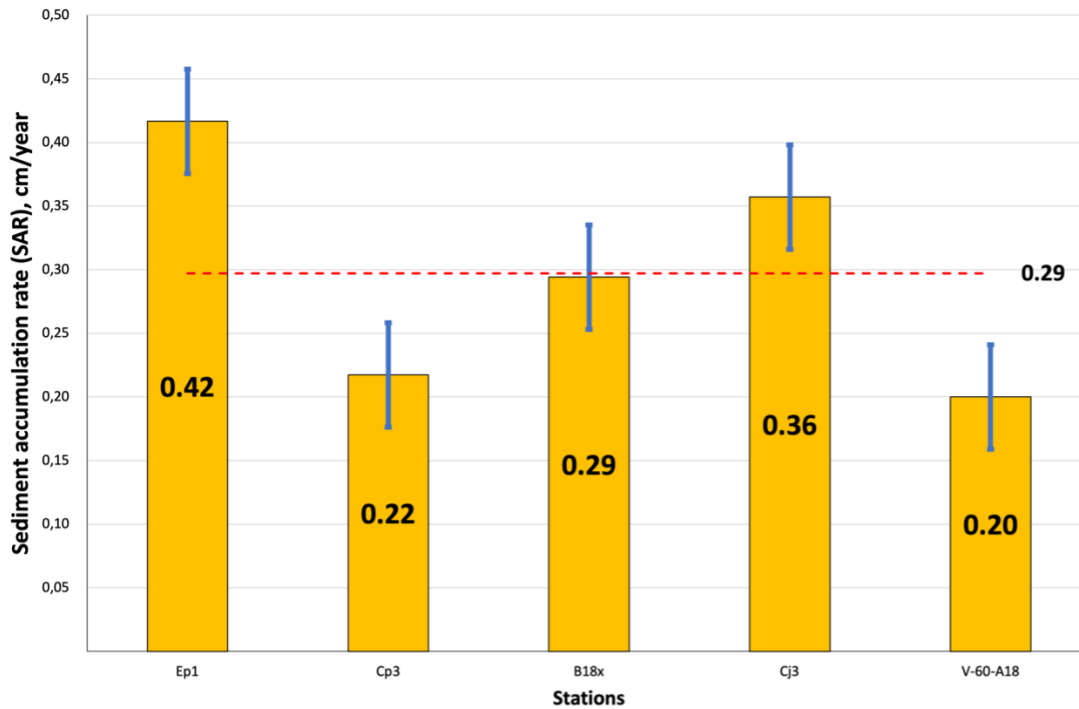


Figure 2: Sediment accumulation rate (SAR) for the dated locations (EP1, Cp3, B18x, Cj3, V-60-A18) in the Inner Oslofjord. The red dotted line indicates the average accumulation rate calculated for the whole fjord.

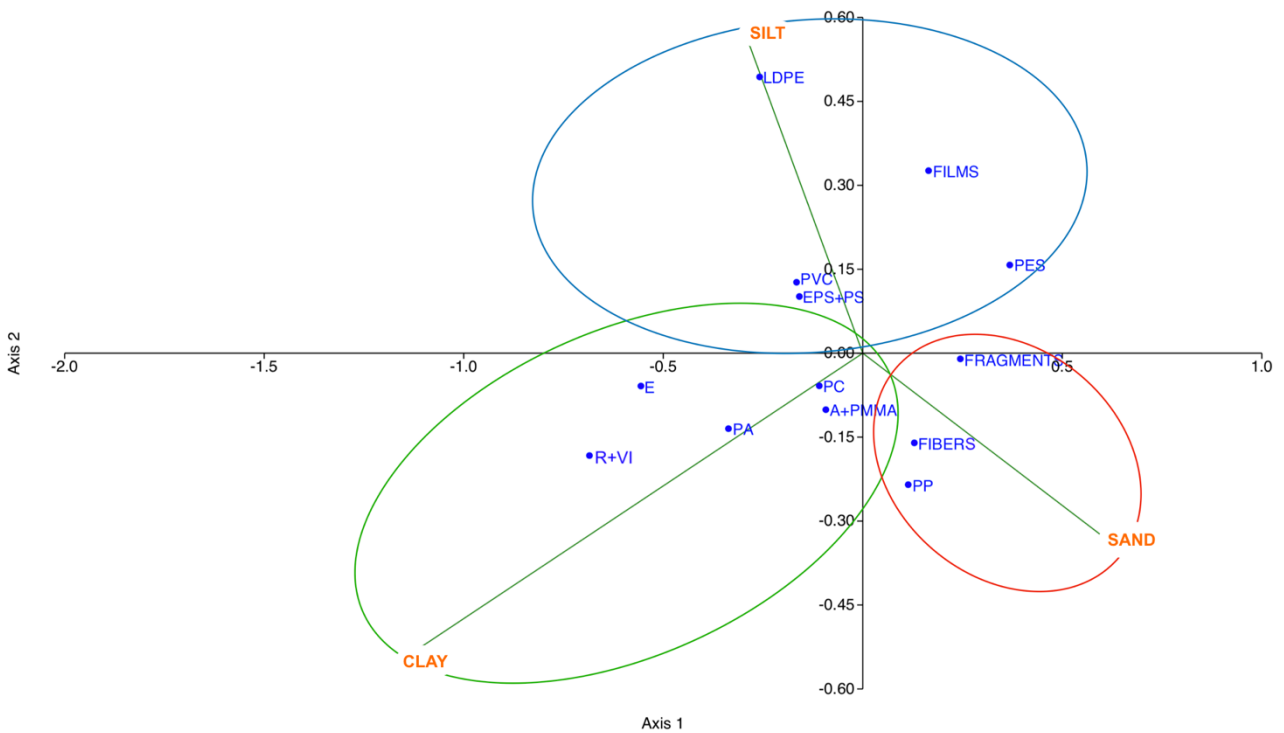
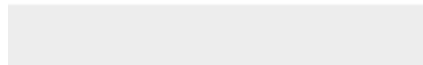


Figure 3: CCA in which grain-sizes are included as environmental variables and are compared to plastic particles characteristics (morphology and polymeric composition). In the red circle are gathered those particles influenced by sandy sediments, in the blue circle particles influenced by silty sediments and in the green circle particles that show a better correlation with the clay grain-size.



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CRedit author statement

Laura Bronzo: Investigation, Data curation, Formal analysis; Writing - Original draft preparation. **Amy L. Lusher:** Methodology, Data curation, Funding acquisition, Reviewing and Editing. **Merete Schøyen:** Investigation. **Caterina Morigi:** Supervision, Reviewing and Editing.

Declaration of interests

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

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: