

Alma Mater Studiorum Università di Bologna Archivio istituzionale della ricerca

Microbial colonization of different microplastic types and biotransformation of sorbed PCBs by a marine anaerobic bacterial community

This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

Published Version:

Rosato A., Barone M., Negroni A., Brigidi P., Fava F., Xu P., et al. (2020). Microbial colonization of different microplastic types and biotransformation of sorbed PCBs by a marine anaerobic bacterial community. SCIENCE OF THE TOTAL ENVIRONMENT, 705, 1-10 [10.1016/j.scitotenv.2019.135790].

Availability: This version is available at: https://hdl.handle.net/11585/731894 since: 2020-02-24

Published:

DOI: http://doi.org/10.1016/j.scitotenv.2019.135790

Terms of use:

Some rights reserved. The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. For all terms of use and more information see the publisher's website.

This item was downloaded from IRIS Università di Bologna (https://cris.unibo.it/). When citing, please refer to the published version.

(Article begins on next page)

MICROBIAL COLONIZATION OF DIFFERENT MICROPLASTIC TYPES AND BIOTRANSFORMATION OF SORBED PCBs BY A MARINE ANAEROBIC BACTERIAL COMMUNITY

Antonella Rosato^a, Monica Barone^b, Andrea Negroni^a, Patrizia Brigidi^b, Fabio Fava^a, Ping Xu^c, Marco Candela^b, Giulio Zanaroli^a*

^aDept. of Civil, Chemical, Environmental and Materials Engineering (DICAM), *Alma Mater Studiorum* University of Bologna, Via Terracini 28, 40131 Bologna, Italy

^bDept. of Pharmacy and Biotechnology (FaBit), *Alma Mater Studiorum* University of Bologna,

Via Belmeloro 6, 40126 Bologna, Italy

^cSchool of Life Science & Biotechnology, Shanghai Jiao Tong University, 800 Dong Chuan Road, Shanghai, 200240, People's Republic of China

*corresponding author (giulio.zanaroli@unibo.it; +39 0512090924)

Keywords

Microplastics; biofilm; microbial community; marine sediment; polychlorinated biphenyls; reductive dehalogenation



Highlights

- Marine bacterial biofilm composition changes on different MP types.
- MP-associated bacterial communities differ from the surrounding sediment ones.
- MP-sorbed PCBs do not significantly affect the MP-biofilm composition.
- PCBs sorbed on MPs can be rapidly dehalogenated by anaerobic marine biofilms.

1 Abstract

We investigated the colonization dynamics of different microplastic (MP) pellets, namely, polyethylene (PE), polyethylene terephthalate (PET), polystyrene (PS), polypropylene (PP) and polyvinyl chloride (PVC), either pristine or contaminated with polychlorinated biphenyls (PCBs), by an organohalide respiring marine microbial community and its biotransformation activity towards PCBs sorbed on MPs, in anaerobic laboratory microcosms of a marine sediment.

8 All MPs were rapidly colonized by the microbial community within 2 weeks of incubation, 9 when approximately 10¹⁰ 16S rRNA gene copies cm⁻² were detected on PVC, 10⁹ copies cm⁻² on PE, and 10⁸ copies cm⁻² on PET, PP and PS. A greater biofilm growth on PVC pellets than 10 11 other MPs was confirmed by quantification of the reducing sugars of the EPS and biofilm 12 staining with crystal violet. Illumina sequencing of the 16S rRNA genes and Principal 13 Coordinate Analysis (PCoA) revealed that the biofilm community on MPs significantly differed 14 from the sediment community, being enriched of chemoorganotrophic fermenting species, and was significantly affected by the type of polymer. The presence of sorbed PCBs did not 15 significantly affect the overall community composition, and mainly resulted in the enrichment 16 17 of Dehalococcoidia, i.e., of the organohalide respiring members of the community.

18 Reductive dechlorination of PCBs sorbed to MPs was observed after two weeks of incubation, 19 when the average number of chlorines per biphenyl molecule was reduced from 5.2 to 4.8 -20 4.3, and was faster (35.2 ± 1.9 to 61.2 ± 5.8 µmoles of Cl removed kg_{MP}-1 week-1) than that of 21 sediment-sorbed ones (33.9 ± 9.1 µmoles of Cl removed kg_{sediment}-1 week-1), which started only 22 after 10 weeks of incubation. These data suggest that microbial colonization of contaminated 23 MPs might change the composition of sorbed PCB mixtures and therefore the toxicity 24 associated to PCB-polluted MPs.

29 1 Introduction

30 Plastics use and production has enormously increased since 1950, reaching around 335 31 million tonnes in 2016, with 60 million tonnes generated in Europe alone (PlasticsEurope, 32 2018). The most common polymers are high-density and low-density polyethylene (PE), 33 polyethylene terephthalate (PET), polystyrene (PS), polypropylene (PP) and polyvinyl 34 chloride (PVC), which together represent approximately 90% of the global plastic production 35 (Andrady and Neal, 2009). A huge amount of plastic materials ends up in the marine 36 environment, becoming an ever-increasing problem due to the toxicity, persistence and 37 universal presence of such debris. The estimation of plastic released in the oceans varies from 38 4.8 to 12.7 million tons annually (Haward, 2018), with size ranging from meters to 39 micrometers (Ryan et al., 2009). The term microplastics (MPs) was introduced within the last 40 decade to describe small particles of plastic, commonly defined as < 5mm in diameter (Frias 41 and Nash, 2019). Once discharged into the marine environment, plastic litter undergoes 42 different processes, such as weathering, fragmentation and fouling (Cole et al., 2011). In 43 particular, biomass accumulation on MPs due to bio(fouling) can lead to an increase of their 44 density and thus their sinking (Morét-Ferguson et al., 2010; Chubarenko et al, 2016; Miao et 45 al., 2019a). The anaerobic sediment has been indicated as a possible long-term sink for MPs 46 (Andrady, 2011; Van Cauwenberghe et al., 2015). Microplastic contamination in marine sediments has been reported by several authors (Thompson et al., 2004; McDermid and 47 48 McMullen, 2004; Claessens et al., 2011), with some of the highest concentrations of MPs found 49 in sediments located in Nova Scotia and Arctic Ocean (up to 8000 and 6595 MPs/kg of 50 sediment, respectively) (Mathalon and Hill, 2014; Bergmann et al., 2017).

51 The first evidence of microbial colonization of plastic fragments dates back to the early 70s 52 (Carpenter et al., 1972). Recent studies reported that MPs are readily colonized by 53 environmental microbial communities in few hours or days of incubation in seawater

54 (Ogonowski et al., 2018) and coastal marine sediments (Harrison et al., 2014). The bacterial 55 assemblages colonizing MPs in freshwater and marine environments have been reported to 56 significantly differ in taxonomic composition and structure from those present the in the surrounding water and/or sediment (Miao et al., 2019b; Frère et al., 2018; Ogonowski et al., 57 58 2018; Rummel et al., 2017; De Tender et al., 2015). Microplastic-associated microbial 59 communities were also reported to have a lower alpha diversity (richness, evenness, and 60 diversity) than those associated to natural substrates, indicating a remarkable differentiation 61 between microbial communities and a substrate-type-coupled species sorting (Miao et al., 62 2019b; Ogonowski et al., 2018). MPs can thus be considered a distinct ecological habitat for 63 diverse microbial communities, the "plastisphere" (Zettler et al., 2013), potentially 64 characterized by distinct microbial and ecological functions (Miao et al., 2019b; Arias-Andres 65 et al., 2018). Less conclusive information has been reported on the selective colonization of MPs of different materials, possibly as the consequence of the potential confounding temporal 66 67 and environmental variability associated to the dynamic exposure conditions typically occurring in situ (Ogonowski et al., 2018). Recently, Li et al. (2019) reported that 68 69 environmental factors, such as the salinity and the nutrients (total nitrogen and total 70 phosphorus), affected the growth rate of biofilms on five types of plastics debris (polyvinyl 71 chloride, polypropylene, polyethylene, polystyrene, and polyurethane) in the Haihe Estuary. 72 The same authors also showed that some genera in the bacterial communities exhibit 73 selectivity for the different polymer types (Li et al., 2019).

Given their small size, MPs may be ingested and accumulated by a wide range of marine organisms, causing direct effects such as physical damage in their intestinal tract, or in other tissues or organs (Van Cauwenberghe et al., 2015). However, the uptake of MPs might also have indirect impacts, since they can absorb and concentrate persistent organic pollutant (POPs) by partition and then transfer such contaminants to the marine food web through

ingestion (Andrady, 2011; Wang et al., 2018). Polychlorinated biphenyls (PCBs), polycyclic
aromatic hydrocarbons (PAHs), dichloro-diphenyl-trichloroethane (DDTs), polybrominated
diphenyl ethers (PBDEs), alkylphenols and bisphenol A (BPA), are the main contaminants that
have been found on plastics debris in the marine environment, at concentrations ranging from
0.001 to 10 mg/kg (Hirai et al., 2011).

84 Despite their production ban in the 1980s, PCBs are still common contaminants in marine 85 sediments and, as reported recently, on marine plastic pellets, where they have been detected 86 at concentrations up to 7.5 mg/kg (Taniguchi et al., 2016). Due to their high lipophilicity, PCBs 87 are accumulated through the food chain up to humans, where they act as endocrine disrupters 88 and possible carcinogens. Besseling et al. (2013) showed that an exposure of lugworm 89 (Arenicola marina) to low concentrations of polystyrene MPs mixed with PCB-contaminated sediment enhanced the bioaccumulation of PCBs. Similar results were obtained by Koelmans 90 91 et al. (2013) in their model analysis with A. marina, although their experiments showed that 92 the role of plastic in the bioaccumulation of POPs may not be considered a significant hazard. 93 Therefore, the release of MPs in the marine environment can have contrasting effects on the 94 PCBs bioaccumulation in marine organisms, which can depend on different causes, such as the 95 experimental conditions or the type of PCB congeners, and further investigations in this 96 regard are necessary (Ziccardi et al., 2016). The toxicity of PCBs varies with the position and 97 number of chlorine atoms on the biphenyl ring, being the coplanar congeners more toxic 98 (Tanabe et al., 1987; Hashmi et al., 2017). Under anoxic conditions, PCBs can undergo 99 reductive dechlorination by organohalide respiring microorganisms occurring in marine 100 sediments, typically belonging to the Dehalococcoidia class of the Phylum Chloroflexi 101 (Zanaroli et al., 2015; Nuzzo et al., 2017). This process replaces chlorine atoms mainly in the 102 meta and para positions with hydrogen, thus converting highly chlorinated congeners into 103 less chlorinated products that are often less toxic, less prone to bioaccumulation (less

104 hydrophobic) and more amenable to degradation by indigenous aerobic bacteria (Fava and 105 Agathos, 2006). If taking place on MP-sorbed PCBs, this microbial process might change the 106 composition of PCBs, and thus the toxicity and bioavailability of the sorbed PCB mixture for 107 consumers ingesting MPs. The effect of microbial colonization of plastics debris on the 108 environmental fate and the biodegradation/biotransformation of the pollutants absorbed on 109 the MPs surface has not been adequately evaluated yet. Microbial colonization of low-density 110 PE pellets incubated in marine and river sediments under aerobic conditions has been shown 111 to increase the biotransformation of MP-sorbed DDTs and PAHs, but not of PCBs (Wu et al., 112 2017), suggesting that it might affect the toxicity associated with polluted microplastics. 113 However, to the best of our knowledge, no studies investigated the influence of microbial 114 colonization of MPs on MP-sorbed pollutants under anoxic conditions typically present in 115 sediments.

The aim of this study was to investigate the colonization dynamics on different types of MPs (low-density PE, PET, PS, PP and PVC) by an anaerobic marine bacterial community containing organohalide respiring bacteria, and the effect of the microbial biofilm on the biotransformation of sorbed PCBs. The process was studied in anaerobic slurry microcosms consisting of marine sediment suspended in seawater, i.e. under biogeochemical conditions mimicking those occurring in situ and controlled exposure conditions.

122

123 2 Materials and Methods

124 2.1 Microplastics and their contamination

Five types of MPs were selected for this study: low density polyethylene pristine pellets (PE), crystalline poly(ethylene terephthalate) pristine pellets (PET), general purpose polystyrene pristine pellets (PS), homo-polypropylene pristine pellets (PP), and poly(vinyl chloride) soft pristine pellets (PVC). The size of the plastic particles ranged from approximately 2.5 to 3 mm. 129 All MPs were sterilized in 70% ethanol for 15 min under shaking, followed by rinsing 3 times 130 with sterilized deionized water, before their use. MPs were contaminated in the laboratory 131 with a commercial mixture of PCBs (Aroclor 1254, UltraScientific, Bologna, Italy) at the final 132 concentration of 30 mg_{PCBs}/kg_{MPs}. This concentration, which is approximately the same order 133 of magnitude of the highest PCBs concentration reported on marine plastic debris (7.5 mg/kg; 134 Taniguchi et al., 2016), was selected in order to obtain final concentrations of single PCB 135 congeners (or co-eluting congeners) of the mixture in the range 20 µg/kgMPs - 4.8 mg/kgMPs, 136 and thus to better assess the microbial dechlorination processes of the spiked PCB mixture. 137 The contamination protocol was adapted from Beckingham and Ghosh (2017); synthetic marine water (1.5 L) was spiked with 0.6 mL of Aroclor 1254 stock solution (20000 mg/L in 138 139 acetone) under shaking (acetone:water 0.04%), 400 g of MPs were added immediately after 140 and incubated under shaking (180 rpm) for 10 days. MPs were recovered by sieving and dried

141 at room temperature under sterile conditions. The actual amount of PCBs sorbed on MPs was 142 indirectly estimated by measuring the residual mass of PCBs in the water phase via solid 143 phase extraction with polydimethylsiloxane (PDMS) fibers. Five-cm PDMS fibers (outer 144 diameter 558.8 µm, inner diameter 486 µm, thickness annulus 35.4 µm, fiber volume 0.597 145 μ L/cm) were incubated in the water phase under mixing (150 rpm) at room temperature for 146 40 days. PCBs were then eluted for 16 h in 0.1 mL hexane and analysed as described in section 147 2.3. The water concentration of PCBs was calculated from their concentration in the PDMS 148 fiber using the fiber-water coefficient (K_{PDMS-W}). The latter was calculated, for each PCB 149 congener, from the octanol-water partition coefficient (K_{ow}) using the equation $log K_{PDMS-W}$ = 150 $0.725 log K_{ow} + 0.479$ (Thomas et al., 2014). Octanol-water partition coefficients for all PCB 151 congeners were obtained from Hawker and Connell (1988). An almost negligible mass of PCBs 152 was detected in the water phase, corresponding to 0.71%, 0.15%, 0.16%, 0.01% and 0.01% (in case of PET, PP, PS, PE and PVC, respectively) of the total mass of PCBs initially added forMPs contamination. The residual mass of PCBs was assumed to be sorbed on MPs.

155

156 2.2 Microcosms set up, incubation and sampling

157 Sacrificial anaerobic slurry microcosms (200 mL total volume) consisting of sediment (20% 158 w/v) and seawater collected from Piallassa Baiona (Ravenna, Italy) were set up according the 159 procedure described in Nuzzo et al. (2017). Microcosms were autoclave-sterilized for 1 h on 160 three consecutive days, inoculated at 5% (v/v) with a PCB-dechlorinating microbial culture 161 enriched previously from marine sediments (Nuzzo et al., 2017) and supplemented with 5 g of 162 MPs, corresponding to a number of MPs ranging from approximately 175 to approximately 163 300, depending on the density and size of the MP type. This resulted in a final concentration 164 ranging from approximately 4375 to approximately 7500 MPs/kg of sediment, which is 165 comparable with the concentrations of microplastics reported in the most MP-impacted 166 marine sediments (Bergmann et al., 2017; Mathalon and Hill, 2014). For each type of MPs, the 167 following microcosms sets were prepared: i) supplemented with MPs contaminated by PCBs; 168 ii) supplemented with pristine MPs (not contaminated). A sterile control set was set up for 169 each MP type by supplementing sterile, not inoculated slurry microcosms with MPs 170 contaminated by PCBs. In addition, a MP-free biologically active control was set up by spiking 171 Aroclor 1254 PCBs in the sediment and inoculating the same marine culture.

The microcosms were incubated at 20°C in the dark under static conditions for 28 weeks. After 2, 5, 10, 14, 19 and 28 weeks of incubation, 3 microcosms for each MP type were sacrificed and MPs were recovered through sieving on sterilized 0.5 mm sieves for the analysis of biofilm microbial communities (except from sterile controls) and of the sorbed PCBs (where spiked).

178 2.3 PCB extraction and analysis

179 PCBs were batch extracted from 1 g of MPs (i.e., approximately 35 to 60 MPs, depending on 180 the MP type) with 4 mL of hexane overnight (30°C, mixing at 150 rpm) and sonication (Hong 181 et al., 2017). PCBs in the organic extracts were analyzed with a 6890N gas-chromatograph 182 equipped with a 63Ni electron capture detector (µECD) and a 6890 series-automatic sampler 183 (Agilent Technologies) using a $30m \times 0.25mm$ HP-5 capillary column (Agilent Technologies) 184 under the conditions described elsewhere (Fava et al., 2003). Calibration curves were 185 obtained and verified monthly using standard mixtures of Aroclor 1254 and Aroclor 1242 (0.5 186 to 30 mg/L concentration range).

187

188 2.4 Quantification of the bacterial biofilm

Prior to each analysis, the MPs were separated from the sediment by sieving (mesh 0.5 mm), rinsed three times with sterile water, in order to remove not-attached cells, and air dried under sterile conditions. Three different approaches were used for the quantification of the biofilm growth on MPs: i) metagenomic DNA extraction followed by qPCR of bacterial 16S rRNA genes; ii) cell/biofilm staining with crystal violet; iii) quantification of the reducing sugars after alkaline hydrolysis of the EPS polysaccharides (Costerton et al., 1995).

195 Metagenomic DNA was extracted from a mixture of replicate samples (0.5 g of MPs, i.e., 196 approximately 18-30 MPs, depending on the MP type) with the UltraClean Soil DNA kit 197 (MoBio Laboratories, Carlsbad, CA, USA) following the procedure described by the provider, 198 with an additional enzymatic cell lysis step before mechanical cell lysis. In particular, MPs 199 were incubated in the bead solution (provided with the kit) in the presence of Lysozyme (11.9) 200 µL of a 100 mg mL⁻¹ solution) at 37 °C on a rotary shaker for 30 min, and then of Proteinase K 201 (3 µL of a 20 mg mL-1 solution) at 37 °C on a rotary shaker for 45 min, prior to addition of 202 SDS and mechanical cell lysis (bead beating on vortex at maximum speed for 10 min). Total

203 DNA was quantified using Qubit® dsDNA HS Assay Kit with a Qubit 3.0 fluorimeter, following 204 the manufacturer's specifications. The 16S rRNA genes were quantified via qPCR with primers 205 905f (5'-AAACTCAAAGGAATTGACGG-3') and 1044r (5'-GACARCCATGCASCACCTG-3') using 206 the reactions conditions described in Nuzzo et al. (2017). 7-point standard curves were 207 included in each plate using *E. coli* 16S rRNA gene. Samples and standards were analyzed in 208 triplicate reactions and 16S rRNA gene copy numbers per cm² of MPs were finally calculated.

209 A colorimetric biofilm quantification protocol using crystal violet was adapted from Burton et 210 al. (2007) and Zanaroli et al. (2011). MPs (0.5 g, i.e., approximately 18-30 MPs, depending on 211 the MP type) were stained 20 minutes with 2 mL of crystal violet solution (0.05% w/v). MPs 212 were then rinsed with sterile distilled water and air-dried for 15 min. The crystal violet on 213 MP-associated biofilm was solubilized by adding 2 mL of 96% (v/v) ethanol and the 214 absorbance of the destaining solution was spectrophotometrically measured at 570 nm 215 versus a blank solution obtained with the same procedure using ethanol-sterilized MPs (see 216 above).

Reducing sugars occurring in the EPS polysaccharides of the biofilm were measured via the colorimetric method described by Bailey (1988), using 3,5 dinitrosalicylic acid, potassium tartrate and NaOH. MPs (0.5 g, i.e., approximately 18-30 MPs, depending on the MP type) were incubated in a boiling water bath for 15 minutes. After cooling in ice, the absorbance of the supernatant was measured at 540 nm versus a blank solution obtained with the same procedure using ethanol-sterilized MPs (see above).

To identify statistically significant differences between MP types, multiple pair-wise
comparisons were performed with the Tukey test (P < 0.05).

225

226 2.5 Analysis of the biofilm community via PCR-DGGE of the 16S rRNA genes

227 PCR-DGGE analysis of 16S bacterial rRNA genes was used to investigate changes in richness 228 (Rr) and community structure (Co) of the bacterial biofilm on different MPs over time. The 229 V3-V5 variable regions of the bacterial 16S rRNA genes were PCR amplified with primers GC-230 357f, containing a 40 bp GC-clamp (5'-CGC CCG CCG CGC CCC GCG CCC GGC CCG CCC CCG 231 CCC C CC TAC GGG AGG CAG CAG-3') and 907r (5'-CCG TCA ATT CCT TTG AGT TT-3') (Sass et 232 al., 2001). The PCR program consisted of an initial denaturation at 95 °C for 5 min, 30 cycles of repeated denaturation at 95 °C for 30 s, annealing at 55 °C for 30 s, and elongation at 72 °C for 233 234 1 min, followed by a final elongation at 72 °C for 10 min. The PCR reaction (50 µL) consisted 235 of mixtures containing 1× colorless GoTaq[®] Flexi buffer (Promega Corporation, Italy), 1.5 236 mM MgCl₂, 0.2 mM each dNTP, 0.4 µM each primer, 1 U of GoTaq® G2 Flexi DNA Polymerase 237 (Promega Corporation, Italy), and of 4 µL template DNA.

The amplicons were resolved with a D-Code Universal Mutation Detection System (Bio-Rad, Milan, Italy) on a 7% (w/v) polyacrylamide gel (acrylamide-N,N'-methylenebisacrylamide, 37:1) containing a denaturing gradient from 40% (top) to 60% (bottom) denaturant (100% denaturant: 7M urea, 40% v/v formamide) as described in Nuzzo et al. (2017). DGGE image analysis was used to calculate the community richness (Rr) and community organization (Co) indexes (Nuzzo et al., 2017).

244

245 2.6 Analysis of the biofilm community via Illumina sequencing of 16S rRNA genes and 246 statistical analysis

DNA extracted from biofilms of selected samples was subjected to sequencing in order to investigate the bacterial community composition. The selection of samples was based on the observed colonization dynamics in terms of i) amount of bacterial biofilm formed and, ii) changes of the community richness and organization over time, detected via PCR-DGGE analysis (section 3.1), as well as on the observed reductive dechlorination of MP-sorbed PCBs
(section 3.3). V1-V3 hypervariable regions of 16S rRNA were PCR amplified and sequenced on
Illumina Miseq with 300 bp paired-end approach (BMR genomics, Padova, Italy).

254 The 16S rRNA gene raw sequences were processed using the open source software pipeline 255 Quantitative Insights Into Microbial Ecology 2 (QIIME 2) version 2017.12 (http://qiime2.org). 256 SILVA was used as reference database to classify the representative sequences from our 257 dataset, and Jaccard distance matrices as beta-diversity measures. The Principal Coordinate 258 Analysis (PCoA) plot was created in R (version 3.5.1) through community ecology vegan 259 package. The significance of separation among study groups was determined by permutation 260 test with pseudo-*F* ratios using the function adonis of vegan package. Significant differences in 261 the relative abundance of Dehalococcoidia between MPs groups - pristine and PCB-262 contaminated - were evaluated with the Wilcoxon test.

263

264 **3 Results**

265 3.1 Colonization dynamics of different MPs

The colonization dynamics was evaluated during a 28-weeks experiment using different approaches: i) quantification of the Bacterial 16S rRNA genes via qPCR, as an approximation of cell density on the MPs surface (16S rRNA gene copies per cm² of MPs), ii) cell/biofilm staining based on crystal violet, which targets both the bacterial cells and the extracellular polymeric substances (EPS) they produce when growing attached to surfaces, and iii) quantification of reducing sugars occurring in the polysaccharide fraction of EPS.

The qPCR analysis revealed that all MPs were rapidly colonized by the bacterial community within the first 2 weeks of incubation, without any remarkable difference between the PCBcontaminated MPs and the pristine (not-contaminated) MPs (Figure 1A,D). The highest cell density was found on PVC pellets, which reached approximately 10¹⁰ 16S rRNA gene copies 276 per cm² of MPs, followed by PE (approximately 10⁹ 16S rRNA copies cm⁻²), and then PET, PP 277 and PS (approximately 10⁸ 16S rRNA copies cm⁻²). Since the surface area of MPs was 278 estimated using the geometry of each plastic pellet, i.e., not considering the irregularities of 279 the shapes and the porosities, the calculated superficial cell density may be an 280 underestimation of the actual microbial colonization. No remarkable cell density changes 281 were observed along the rest of the incubation, indicating that cells attachment and growth 282 were not taking place and/or that were balanced by cells death and detachment from the surface (McDougald et al., 2012). 283

284 Staining with crystal violet (Figure 1B,E) and quantification of the reducing sugars (Figure 285 1C,F) revealed a biofilm increase up to weeks 10-15 on PE, PS, PP and PET, and up to the end 286 of incubation (week 28) on PVC, without remarkable differences in the presence and absence 287 of the sorbed pollutants. The biofilm increase over time may indicate the occurrence of a 288 biofilm maturation phase, taking place after a rapid cells adhesion and growth. The biofilm 289 amount then changed less markedly, showing slight decreases followed by small increases 290 that may indicate the succession of detachment and regrowth events involving portions of the 291 biofilm, which are typical of these dynamic structures (McDougald et al., 2012). The greatest 292 biomass concentration, approximately one order of magnitude higher than on the other MP 293 types, was detected on PVC MPs also according to crystal violet and quantification of reducing 294 sugars and was significantly higher than that on other MP types (P<0.05, Tukey test) 295 according to all measured parameters.

A preliminary investigation of the evolution of the biofilm community established on the different MPs was carried out through PCR-DGGE analysis of the 16S rRNA genes. PCR-DGGE profiles (Figure S1) were mainly used to evaluate if major changes were taking place over time in the community developed on each MP type, rather than for an in-depth assessment of the community diversity and structure or to compare the biofilm communities on the

301 different MP types. The following parameters were considered: i) community richness (Rr), 302 i.e., the number of species, and ii) community organization (Co), which indicates if the 303 individuals of the population are homogeneously distributed between the species (even 304 community) or if individuals belonging to few species are predominant over those belonging 305 to all others species (uneven community).

306 The Rr and Co varied markedly between the different type of MPs, but not between the MPs 307 contaminated by PCBs and those not contaminated (Figure S2). In particular, the biofilm 308 communities on PET, PS and PP did not change remarkably in Rr and Co over time, and were 309 characterized by an average Rr of approximately 25 and 18, for the pristine and PCB-310 contaminated MPs, respectively. The Co resulted to be approximately 25 or lower for all this 311 plastic samples (PET, PS and PP, contaminated and pristine), which indicates a high degree of 312 community evenness. In contrast, a remarkable change of the biofilm community appeared to 313 occur on PE during the first 10 weeks of incubation, where a noteworthy reduction of Rr and a 314 substantial increase of Co where observed. This indicates that a fraction of the biofilm 315 microbial community on the PE surface was able to become dominant over time, causing a 316 loss of biodiversity. Conversely, an increase of Rr and limited fluctuations of the Co were 317 observed on PVC over time. This indicates that an increase of biodiversity took place without 318 affecting the functional organization of the community.

319

320 3.2 Composition of the biofilm community on different MPs

The composition of the biofilm communities was then investigated more in detail through Illumina sequencing of the 16S rRNA genes. Samples were selected based on the observed colonization dynamics in terms of i) amount of bacterial biofilm formed and, ii) changes of the community richness and organization over time (section 3.1), as well as on the observed reductive dechlorination of MP-sorbed PCBs (section 3.3). In particular, the biofilm 326 community composition was investigated for all samples after 2 weeks of incubation, since at 327 this sampling point all MP types were colonized by the maximum cells concentration, which then remained almost constant (Figure 1A,D), and a remarkable reductive dechlorination of 328 329 sorbed PCBs was observed on all MP types (section 3.3, Figure 5); in addition, the final 330 sampling point (28 weeks of incubation) was analysed only for PE and PVC pellets, since 331 substantial changes in Rr and Co were observed between 2 and 28 weeks of incubation only 332 on these materials (Figure S2). The PCB-dechlorinating marine culture inoculated in the 333 microcosms was also analysed to investigate if adhesion and biofilm formation on MPs 334 involves specific members of the community or the whole population. In addition, the 335 sediment present in the microcosms supplemented with PCB-contaminated MPs was analysed 336 to evaluate any differences between the microbial communities adhering to the MPs and 337 those associated to the sediment particles.

At the phylum level (Figure 2), the microbial inoculum mainly consisted of Proteobacteria 338 339 (58%), Chloroflexi (30%) and, at lower extent, Firmicutes (6%). After 2 weeks of incubation, 340 Firmicutes became dominant in the biofilm communities on all MP types, having a relative 341 abundance that ranged from 88% (pristine PS) to 49% (PCB-contaminated PVC). Conversely, 342 Proteobacteria represented a minor fraction of the biofilm communities, ranging from 1 to 343 27%, and the fraction of Chloroflexi decreased on all plastic pellets (0.3-13%,), except for PVC 344 contaminated by PCBs (45%). Plastic biofilm communities also included, at quite high relative 345 abundances, the phyla Lentisphaerae (0-26%), Deinococcus-Thermus (0-7%), Actinobacteria 346 (0-13%) and Acidobacteria (0-9%), which were not detected in the inoculum or represented 347 less than 2% of the original microbial community.

Relevant changes in composition were detected after 28 weeks of incubation on PE and PVC,when Chloroflexi became more abundant and Firmicutes decreased.

The bacterial communities associated to the sediment surrounding the different PCBcontaminated MPs were very similar to each other and mainly consisted of bacteria belonging to the phyla Firmicutes (30-43%), Chloroflexi (20-28%), Proteobacteria (12-22%), and to less extent, Acidobacteria (6-9%), Synergistetes (4-8%), Lentisphaerae (3-7%), and Actinobacteria (2-3%). These bacterial communities therefore remarkably differed from the microbial inoculum and those colonizing the MPs.

356 At lower taxonomic levels (Table S1), the original marine microbial community was 357 dominated by bacteria of the genera *Sulfurovum* (20.4%) and *Sulfurimonas* (15.5%), which are 358 both typically sulphur oxidizing, nitrate-reducing bacteria, Dethiosulfatibacter (5.3%), a 359 sulphur and thiosulfate-reducing genus, *Magnetovibrio* (3.6%), which includes members 360 capable of anaerobic thiosulfate oxidation using nitrous oxide, Celeribacter and another 361 member (unidentified genus) of the Rhodobacteraceae family (12.0%) and Dehalobium 362 (17.7%), an organohalide respiring Dehalococcoidia, along with other non-dehalorespiring 363 members of the Chloroflexi phylum (unidentified genus, Anaerolineacea family, 12.8%). Most 364 of these genera were not detected in the biofilm communities grown on pristine MPs, or they 365 were at much lower abundances. After 2 weeks of incubation, the biofilm present on PS and 366 PVC was dominated by a Firmicutes bacterium of the Peptostreptococcaceae family 367 (unidentified genus), which represented 42.4% and 37.6% of the community, respectively. 368 This bacterium was also present at high relative abundance on PE (19.4%) and, at lower 369 abundances, on PET (9.4%) and PP (8.5%) after 2 weeks of incubation. The microbial 370 community on PE and PP after 2 weeks of incubation was mainly dominated by a different 371 Firmicutes bacterium belonging to the SRB2 family of Thermoanaerobacterales (unidentified 372 genus), which represented approximately 29% of the biofilm community on both MPs types. 373 The same bacterium was also detected at high abundance on PET (17.6%) and at much lower 374 abundance on PS (5.6%), while was not detected on PVC. A member of the Clostridiaceae 1

375 family (unidentified genus) was also dominant or present at relatively high abundances on 376 PET (20.5%), PP (21.0%) and PS (16.4%), whereas the community on PVC included at high 377 abundances a member of Clostridiales (22.5%), Celeribacter (15.0%) and a member of 378 Dehalocccoidia (10.0%). Other community members that enriched on pristine MPs after 2 379 weeks of incubation were an uncultured Coriabacteriaceae (unidentified genus) in PVC (12.5%), PE (6.6%), PET (5.3%) and PP (4.3%), a member of Deinococci in PP (6.8%), PE 380 381 (5.9%) and PS (5.3%), Sedimentibacter in PE (10.2%), PET (7.3%), PP (6.3%) and PS (3.1%), 382 and *Clostridium senso strictu 1* only in PET (7.9%) and PS (5.4%).

After 28 weeks of incubation, the SRB2 family of *Thermoanaerobacterales* and *Sedimentibacter* were lost by the biofilm community on PE, where *Peptostreptococcaceae* further enriched (40.3%) along with *Anaeroplasma* (6.5%) and *Anarolinaceae* (17.9%). A remarkable change was also observed in the biofilm composition of PVC after 28 weeks of incubation, when the uncultured *Coriobacteriaceae*, Dehalococcoidia, Clostridiales and *Celeribacter* drastically reduced, and uncultured *Anaerolinaceae* remarkably enriched (35.4%).

The presence of sorbed PCBs had very limited effects on the overall composition of the biofilm on MPs, resulting mainly in the enrichment of the class Dehalococcoidia, i.e., the organohalide respiring Chloroflexi members of the community (Figure 3). This behavior appears even more marked in the case of PVC, where the increase of Dehalococcoidia in the biofilm community in the presence of PCBs reaches the statistics significance (P-value = 0.05, Wilcoxon test).

The PCoA analysis based on Jaccard similarity index was then used to assess the overall OTUlevel compositional differences in the microbial communities adhering to the different plastic types (PE, PET, PS, PP, PVC) and sediment. The data show a clear separation of microbial community profiles by MPs type and between MPs and the surrounding sediment (P-value <

0.03, permutation test with pseudo-*F* ratio) (Figure 4A). Permutation test with pseudo-*F* ratio
of the significance of pairwise separation between each MP type and MPs and the sediment is
reported in Table S2. The PCoA analysis also demonstrated that the communities on PCBcontaminated and pristine MPs did not significantly differ between each other (P-value = 0.4)
(Figure 4B). The sorption of PCBs on MPs therefore did not significantly change the taxonomic
composition of the overall microbial community adhering to them.

406

423

407 3.3 PCB dechlorination

408 No dechlorination of MP-sorbed PCBs was observed in sterile controls during the 28 weeks of 409 incubation. On the contrary, a significant reductive dechlorination of PCBs associated to all 410 different types of MPs was observed in the microcosms inoculated with the PCB-411 dechlorinating culture after only two weeks of incubation, when the average number of 412 chlorines per biphenyl molecule was reduced from 5.2 to 4.8 on PP, 4.7 on PE, 4.6 on PET, 4.4 413 on PS and 4.3 on PVC (Figure 5), corresponding to dechlorination rates of 35.2 ± 1.9 (PP), 44.6 \pm 2.6 (PE), 56.8 \pm 6.2 (PET), 57.8 \pm 5.4 (PS) and 61.2 \pm 5.8 (PVC) µmoles of Cl removed kg_{MP}⁻¹ 414 415 week⁻¹. The reductive dechlorination of sorbed PCBs then proceeded more slowly, leading to 416 an average number of chlorines per biphenyl molecules after 28 weeks of incubation in the 417 range 4.4-3.9. Remarkably, the biotrasformation of PCBs sorbed on MPs was faster than that 418 of PCBs sorbed on sediment, which started only after 10 weeks of incubation and proceeded 419 with a dechlorination rate of 33.9 \pm 9.1 µmoles of Cl removed kg_{sediment}⁻¹ week⁻¹ until week 28, 420 when the average number of chlorines per biphenyl molecule was reduced to 4.0 (Figure 5). 421 In general, highly similar dechlorination patterns were observed for PCBs sorbed on different MP types, as indicated by the depletion of the same highly-chlorinated congeners and the 422

424 chlorinated congeners and the lower accumulation of the low-chlorinated ones detected on

accumulation of the same low-chlorinated ones (Figure 6). The lower reduction of the highly-

425 PE and PP pellets at the end of incubation was in accordance with the less extensive 426 dechlorination process occurred on these MPs. The dechlorination patterns of PCBs sorbed on 427 MPs was similar to that observed on PCBs sorbed on sediment (without MPs), with a main 428 difference: the accumulation in the sediment of the co-eluting hexa-/penta-chlorinated 429 congeners 234-35, 235-34 and 2356-24, which did not accumulate on the MPs, and the 430 concomitant accumulation of the tri-chlorinated congeners 25-3 and 24-3 on MPs and not in 431 the sediment. This indicates that a meta and para dechlorination of 235-34 and 234-35 to 25-432 3 and 24-3, respectively, occurred only on MPs.

433

434 **4 Discussion**

435 Microbial colonization of MPs in the marine environment may have several implications on 436 their fate and sedimentation behavior, the fate of MP-sorbed pollutants and their toxicity and 437 bioavailability to marine organisms (Cole et al., 2011; Wang et al., 2018; Mohamed Nor and 438 Koelmans, 2019). While the colonization of plastic debris by marine microorganisms have 439 been recently reported in seawater (Dussud et al., 2018; Frère et al., 2018; Xu et al., 2019), 440 very limited information is available on plastic colonization in marine sediments (Harrison et 441 al., 2014), which are the ultimate sink, as well as an entry point in the food chain through 442 benthic organisms, of plastic debris and hydrophobic pollutants in the marine environment 443 (Kaiser et al., 2017). Very limited information is available also on the fate of pollutants sorbed 444 on plastics and, in particular, on the potential role of microbial biofilms in the 445 biodegradation/biotransformation of plastic-sorbed pollutants (Wu et al., 2017). In this study, 446 the colonization dynamics of microplastic (MP) pellets of different materials, namely, PE, PET, 447 PS, PP and PVC, either pristine or contaminated with polychlorinated biphenyls (PCBs), was 448 investigated in laboratory microcosms of an anoxic marine sediment and seawater collected 449 from the same site, i.e., under laboratory biogeochemical conditions mimicking those

450 occurring in situ and controlled exposure conditions. The sediment was inoculated with a 451 marine anaerobic microbial community previously selected for its ability to dehalogenate 452 PCBs (Nuzzo et al., 2017), in order to better assess the potential role of microbial reductive 453 dehalogenation processes on the fate of MP-sorbed PCBs. The use of a well-defined source 454 community was also made to better evaluate the possible effects of the MP material and of 455 MP-sorbed pollutants on the surface colonization dynamics and on structure and composition 456 of the biofilm (Ogonowski et al. 2018).

All MPs were rapidly colonized during the first 2 weeks of incubation, up to a cell density range of 1.0×10^8 to 1.0×10^{10} cells/cm². Harrison et al. (2014) reported a comparable, rapid colonization of low-density PE by bacteria of coastal marine sediments incubated in Petri dishes, that reached a density of 1.0×10^6 to 1.0×10^9 16S rRNA genes DNA per mm² of PE within 7 days. Our work confirms that MPs are a good anthropogenic substrate for colonization by marine sediment microbial communities also under strictly anoxic conditions, which are typically present in situ few millimeters below the sediment surface.

464 Remarkable differences were observed in terms of colonization of different MP types. In 465 particular, biofilm formation was remarkably higher on PVC pellets than on other MP types, 466 both in terms of cell density reached and amount of EPS produced. Such a higher biofilm 467 growth on PVC is likely the consequence of a higher availability of energy and carbon sources 468 that might have been released from the MPs. Many organic additives are commonly 469 supplemented to plastic materials in order to improve their proprieties, such as tensile 470 strength, flexibility and durability (Teuten et al., 2009). In particular, PVC is among the 471 plastics containing the highest amounts of additives, mainly plasticizers and stabilizers, that may reach up to 50% w/w of the plastic material. Phthalates (alkyl/aryl esters of 1,2-472 benzenedicarboxylic acid), as well as low molecular weight, easily biodegradable citrate, 473 474 adipate and hexanoate esters, are very commonly used plasticizers in PVC (Markarian, 2007;

Babinsky, 2006). The microbial degradation of phthalates to methane and carbon dioxide has
been also reported under anaerobic conditions (Chang et al., 2005; Liang et al., 2008). All
these additives can thus sustain the growth of anaerobic microbes/microbial communities
and could have therefore favored a more extensive biofilm growth on PVC pellets.

All MP types selected specific fractions of the inoculated microbial community. The dominant members of the latter were mainly associated with sulphur cycling, a common process taking place in marine sediments (Wasmund et al., 2017), as well as with organohalide respiration, which was a specific feature of the marine microbial community inoculated in the microcosms (Nuzzo et al., 2017). Most of these members were not detected in any of the biofilm communities grown on MPs, which were dominated by taxa mainly including chemoorganotrophic, fermenting bacteria.

486 Remarkable differences were also observed in terms of composition between the microbial 487 communities associated to the different MP types and the surrounding sediment. Recent 488 studies have demonstrated that the bacterial assemblages that develop on the surface of 489 plastic material in seawater or freshwater systems can significantly differ from those 490 developing on non-plastic substrates, such as wood, cobblestone, cellulose and glass, as well 491 as from the sediment and water bacterial communities (Miao et al., 2019b, Ogonowski et al., 492 2018; Frère et al., 2018; De Tender et al., 2015). Some of these studies have also reported that 493 the microbial communities developed on PE and PP were quite similar to each other, and 494 more distinct to those associated with PS pellets (Ogonowski et al., 2018; Frère et al., 2018). 495 The differentiation of biofilm communities on different MP types occurred in this study may 496 be attributed to many factors. For example, Ogonowski et al. (2018) recently reported a 497 significant correlation between the substrate hydrophobicity and bacterial composition on 498 different plastic and non-plastic (glass, cellulose) materials. Since the hydrophobicity of the 499 materials tested in this study typically changes in a relatively narrow range (contact angle 80500 100; Ogonowski et al., 2018; Gotoh et al., 2011; McGinty and Brittain, 2008), the difference in 501 biofilm composition could be more probably related to other features, such as the presence of 502 different organic and inorganic additives and the different rates they are released from the 503 plastic material. Indeed, the additives released from different MPs may either promote the 504 growth of some microbial species able to use them as carbon and energy source, or limit the 505 growth and survival of some other species that are sensitive to these chemicals. However, the 506 very high variety of additives used in different plastics, as well as the use of different additives 507 for the same plastic type by different producers, make impossible to identify a typical 508 additives composition for each plastic and thus to relate the specific compositions of the 509 biofilm communities observed in this study to the release of specific chemicals. Interestingly, 510 Klaeger et al. (2019) recently reported that the release of biodegradable residual monomers 511 and oligomers from polymers can lead to a remarkable overestimation of the plastic materials 512 degradation, in particular during the first 19 days of incubation. Indeed, styrene and vinyl 513 chloride monomers have been widely reported to leach from PS and PVC pellets, respectively 514 (Pilevar et al., 2019; Fayad et al., 1997) and are known to be susceptible, as well as other 515 hydrocarbons, to anaerobic biodegradation (Varjani, 2017). This suggests that the leaching of 516 residual monomers and/or oligomers from the different plastic types may have also 517 contributed to the differentiation of biofilm compositions on the different MPs we observed 518 especially during the first 2 weeks of incubation. This is in agreement also with the selective 519 enrichment of Dehalococcoidia (i.e., of the organohalide respiring Chloroflexi members of the 520 inoculated community) in the biofilm promoted by the sorbed PCBs on all MPs, without any 521 significant change of the overall community.

522 The PCBs sorbed on the different MP types were rapidly converted into less chlorinated 523 congeners by the anaerobic marine biofilms. Indeed, the rate of PCB dechlorination we 524 observed may be much higher than that potentially taking place in situ, given the extremely

525 low concentration of organohalide respiring bacteria typically present in environmental 526 samples and the specialized enriched community with marked organohalide-respiring activity we used in our study under controlled conditions. However, to our knowledge, this is the first 527 528 report describing the potential occurrence of reductive dechlorination processes towards 529 PCBs sorbed on MPs. Very few studies investigated the biotransformation/biodegradation of 530 pollutants sorbed on MPs; Wu et al. (2017) reported for example that colonization of low-531 density PE pellets increased the biotransformation of dichloro-diphenyl-trichloroethanes 532 (DDTs) and polycyclic aromatic hydrocarbons (PAHs) in marine and river sediments 533 incubated aerobically, although had not significant effect on PCBs. This claims the need of 534 further studies on the biotransformation of pollutants sorbed on MPs under different 535 environmental conditions and by different microbial communities.

536 Moreover, our study shows that PCBs sorbed on MPs underwent reductive dechlorination 537 much more rapidly than those sorbed on the sediment. While Mato et al. (2001) showed that 538 sorption of organic compounds, such as PCBs and DDE, is approximately two orders of 539 magnitude higher in PP pellets than in marine sediments, probably due to the plastic 540 hydrophobic surfaces, our results suggest that PCBs are more bioavailable to the 541 dechlorinating marine microbes when sorbed on MPs, or that the colonization of MPs might 542 increase the bioavailability of MP-sorbed PCBs, e.g., through biosurfactants production, better 543 than of the sediment-sorbed ones. PCB dechlorination was more extensive on PVC, PS and 544 PET, followed by PE and PP, which might indicate a slight different degree of PCB 545 bioavailability on different MP materials. The most abundant highly-chlorinated PCB 546 congeners were extensively converted into less chlorinated products, that are commonly less 547 toxic and less hydrophobic, thus potentially more bioavailable and less prone to 548 bioaccumulation. The toxicity of contaminated MPs is potentially due to the combination of 549 several factors, e.g., the intrinsic toxicity of the sorbed pollutants, their bioavailability and

capability to bioaccumulate; the observed change in congener composition of the PCB mixture
associated to MPs might thus have remarkable effects on the overall toxicity associated to
polluted MPs, that deserves more investigations.

553

554 **5 Conclusions**

555 The colonization dynamics of different MP pellet types (PE, PET, PS, PP, and PVC) by an 556 anaerobic marine microbial community was investigated for the first time in anoxic marine 557 sediments, which represent the main sink for MPs in the marine environment. This study 558 showed that microbial colonization took place very rapidly on all MPs and that the biofilm 559 composition differed significantly between the five plastic types and from the surrounding 560 sediment community. The colonization of MPs is thus a selective process that may depend on 561 several factors, such as the different properties of polymer surface and the presence of 562 different additives and/or different residual monomers/oligomers that may be released from 563 the plastics and promote or limit selectively the growth of distinct bacterial species. However, 564 the factors driving the enrichment of different biofilm communities on the MPs may be 565 multiple and diverse and need further investigation. This work also showed that PCBs sorbed 566 on different MP types can be dehalogeated much faster than those sorbed on sediments, 567 suggesting that they are highly bioavailable. In addition, their susceptibility to be 568 bioconverted to even more bioavailable, although less toxic, low-chlorinated PCBs indicate 569 that microbial biofilms growing on contaminated MPs, by changing the congener composition 570 of the sorbed PCB mixture, might remarkably affect the toxicity of contaminated MPs in the 571 marine environment, as well as the uptake and bioaccumulation of MP-associated PCBs by 572 marine organisms. Further investigation should be carried out in order to assess ecotoxicity 573 changes associated to the biotransformation of PCBs sorbed to MPs particles.

574

575 Acknowledgements

576 This research was supported through the Joint Programming Initiative Healthy and 577 Productive Seas and Oceans (JPI- Oceans) PLASTOX project (grant agreement No 696324) by 578 the Italian Ministry of Education, University and Research (MIUR, Project Grant 6962 579 31/03/2015).

We thank Dr. Stefania Piarulli, Dr. Joanne Wong and Prof. Laura Airoldi (University of
Bologna) for assistance in sampling of sediment samples from Piallassa Baiona (Ravenna,
Italy).

583

584 **References**

- Andrady, A.L., 2011. Microplastics in the marine environment. Marine pollution bulletin,
 62(8), 1596-1605.
- Andrady, A.L., Neal, M.A., 2009. Applications and societal benefits of plastics. Philosophical
 Transactions of the Royal Society B: Biological Sciences, 364(1526), 1977-1984.
- Arias-Andres, M., Kettner, M.T., Miki, T., Grossart, H.-P., 2018. Microplastics: New substrates
 for heterotrophic activity contribute to altering organic matter cycles in aquatic
 ecosystems. Science of the Total Environment, 635, 1152–1159.
- Babinsky, R., 2006. PVC additives: a global review. Plastics, Additives and Compounding,
 8(1), 38-40.
- Bailey, M.J., 1988. A note on the use of dinitrosalicylic acid for determining the products of
 enzymatic reactions. Applied Microbiology and Biotechnology, 29(5), 494-496.
- Beckingham, B., Ghosh, U., 2017. Differential bioavailability of polychlorinated biphenyls
 associated with environmental particles: Microplastic in comparison to wood, coal and
 biochar. Environmental Pollution, 220, 150-158.

- Bergmann, M., Wirzberger, V., Krumpen, T., Lorenz, C., Primpke, S., Tekman, M. B., Gerdts,
 G., 2017. High quantities of microplastic in Arctic deep-sea sediments from the
 HAUSGARTEN observatory. Environmental science & technology, 51(19), 11000-11010.
- Besseling, E., Wegner, A., Foekema, E.M., Van Den Heuvel-Greve, M.J., Koelmans, A.A., 2013.
 Effects of microplastic on fitness and PCB bioaccumulation by the lugworm Arenicola
 marina (L.). Environmental science and technology, 47(1), 593-600.
- Burton, E., Yakandawala, N., LoVetri, K., Madhyastha, M.S., 2007. A microplate
 spectrofluorometric assay for bacterial biofilms. Journal of industrial microbiology and
 biotechnology, 34(1), 1-4.
- Carpenter, E.J., Anderson, S.J., Harvey, G.R., Miklas, H.P., Peck, B.B., 1972. Polystyrene
 spherules in coastal waters. Science, 178(4062), 749-750.
- Chang, B.V., Liao, C.S., Yuan, S.Y., 2005. Anaerobic degradation of diethyl phthalate, di-nbutyl phthalate, and di-(2-ethylhexyl) phthalate from river sediment in Taiwan.
 Chemosphere, 58(11), 1601-1607.
- Chubarenko, I., Bagaev, A., Zobkov, M., Esiukova, E., 2016. On some physical and dynamical
 properties of microplastic particles in marine environment. Marine Pollution Bulletin,
 108(1-2), 105-112.
- Claessens, M., De Meester, S., Van Landuyt, L., De Clerck, K., Janssen, C.R., 2011. Occurrence
 and distribution of microplastics in marine sediments along the Belgian coast. Marine
 Pollution Bulletin, 62(10), 2199-2204.
- Cole, M., Lindeque, P., Halsband, C., Galloway, T.S., 2011. Microplastics as contaminants in
 the marine environment: a review. Marine pollution bulletin, 62(12), 2588-2597.
- Costerton, J.W., Lewandowski, Z., Caldwell, D.E., Korber, D.R., Lappin-Scott, H.M., 1995.
- 622 Microbial biofilms. Annual review of microbiology, 49(1), 711-745.

- De Tender, C.A., Devriese, L.I., Haegeman, A., Maes, S., Ruttink, T., Dawyndt, P., 2015.
 Bacterial community profiling of plastic litter in the Belgian part of the North Sea.
 Environmental science and technology, 49(16), 9629-9638.
- Dussud, C., Hudec, C., George, M., Fabre, P., Higgs, P., Bruzaud, S., Delort, A.M., Eyheraguibel,
 B., Meistertzheim, A.L., Jacquin, J., Cheng, J., Callac N., Odobel, C., Rabouille, S., Ghiglione, J.F.,
 2018. Colonization of non-biodegradable and biodegradable plastics by marine
 microorganisms. Frontiers in microbiology, 9, 1571.
- Fava, F., Agathos, S.N., 2006. Uncertainty and research needs in the area of the biological
 restoration of contaminated sediments. In Assessment and remediation of contaminated
 sediments (pp. 239-246). Springer, Dordrecht.
- Fava, F., Zanaroli, G., Young, L.Y., 2003. Microbial reductive dechlorination of pre-existing
 PCBs and spiked 2, 3, 4, 5, 6-pentachlorobiphenyl in anaerobic slurries of a contaminated
 sediment of Venice Lagoon (Italy). FEMS microbiology ecology, 44(3), 309-318.
- Fayad, N.M., Sheikheldin, S.Y., Al Malack, M.H., El Mubarak, A.H., Khaja, N., 1997.
 Migration of vinyl chloride monomer (VCM) and additives into PVC bottled drinking water.
 Journal of Environmental Science and Health Part A, 32(4), 1065-1083.
- Frère, L., Maignien, L., Chalopin, M., Huvet, A., Rinnert, E., Morrison, H., Kerninon, S.,
 Cassone, A.-L, Lambert, C., Reveillaud, J., Paul-Pont, I., 2018. Microplastic bacterial
 communities in the Bay of Brest: Influence of polymer type and size. Environmental
 pollution, 242, 614-625.
- Frias, J.P.G.L., Nash, R., 2019. Microplastics: Finding a consensus on the definition. Marine
 pollution bulletin, 138, 145-147.
- Gotoh, K., Yasukawa, A., Kobayashi, Y., 2011. Wettability characteristics of poly(ethylene
 terephthalate) films treated by atmospheric pressure plasma and ultraviolet excimer light.
 Polymer Journal, 43, 545–551.

- Harrison, J.P., Schratzberger, M., Sapp, M., Osborn, A.M., 2014. Rapid bacterial colonization
 of low-density polyethylene microplastics in coastal sediment microcosms. BMC
 microbiology, 14(1), 232.
- Hashmi, M.Z., Zhang, J., Li, B., Su, X., Tariq, M., Ahmad, N., Malik, R.N., Ullah, K., Chen, C., Shen,
 C., 2017. Effects of structurally different noncoplanar and coplanar PCBs on HELF cell
 proliferation, cell cycle, and potential molecular mechanisms. Environmental toxicology,
 32(4), 1183-1190.
- Haward, M., 2018. Plastic pollution of the world's seas and oceans as a contemporary
 challenge in ocean governance. Nature communications, 9(1), 667.
- Hawker, D.W., Connell, D.W., 1988. Octanol-water partition coefficients of polychlorinated
 biphenyls congeners. Environmental Science & Technology, 22, 382-387.
- Hirai, H., Takada, H., Ogata, Y., Yamashita, R., Mizukawa, K., Saha, M., Kwan, C., Moore, C.,
 Gray, H., Laursen, D., Zettler, E.R., Farrington, J.W., Reddy, C.M., Peacock, E.E., Ward, M.W.,
 2011. Organic micropollutants in marine plastics debris from the open ocean and remote
 and urban beaches. Marine Pollution Bulletin, 62(8), 1683-1692.
- Hong, S.H., Shim, W.J., Hong, L., 2017. Methods of analysing chemicals associated with
 microplastics: a review. Analytical Methods, 9(9), 1361-1368.
- Kaiser, D., Kowalski, N., Waniek, J.J., 2017. Effects of biofouling on the sinking behavior of
 microplastics. Environmental Research Letters, 12(12), 124003.
- Klaeger, F., Tagg, A.S., Otto, S., Bienmüller, M., Sartorius, I., Labrenz, M., 2019. Residual
 monomer content affects the interpretation of plastic degradation. Scientific reports, 9(1),
 2120.
- Koelmans, A.A., Besseling, E., Wegner, A., Foekema, E.M., 2013. Plastic as a carrier of POPs
 to aquatic organisms: a model analysis. Environmental science and technology, 47(14),
 7812-7820.

673	• Li, W., Zhang, Y., Wu, N., Zhao, Z., Xu, W.A., Ma, Y., Niu, Z., 2019. Colonization Characteristics
674	of Bacterial Communities on Plastic Debris Influenced by Environmental Factors and
675	Polymer Types in the Haihe Estuary of Bohai Bay, China. Environmental science $\&$
676	technology, 53(18), 10763-10773.

- Liang, D.W., Zhang, T., Fang, H.H., He, J., 2008. Phthalates biodegradation in the
 environment. Applied microbiology and Biotechnology, 80(2), 183.
- Markarian, J., 2007. PVC additives–What lies ahead? Plastics, Additives and Compounding,
 9(6), 22-25.
- Mathalon, A., Hill, P., 2014. Microplastic fibers in the intertidal ecosystem surrounding
 Halifax Harbor, Nova Scotia. Marine pollution bulletin, 81(1), 69-79.
- Mato, Y., Isobe, T., Takada, H., Kanehiro, H., Ohtake, C., Kaminuma, T., 2001. Plastic resin
 pellets as a transport medium for toxic chemicals in the marine environment.
 Environmental science and technology, 35(2), 318-324.
- McDermid, K.J., McMullen, T.L., 2004. Quantitative analysis of small-plastic debris on
 beaches in the Hawaiian archipelago. Marine pollution bulletin, 48(7-8), 790-794.
- McDougald, D., Rice, S.A., Barraud, N., Steinberg, P.D., Kjelleberg, S., 2012. Should we stay or
 should we go: mechanisms and ecological consequences for biofilm dispersal. Nature
 Reviews Microbiology, 10, 39–50.
- McGinty, K.M., Brittain, W.J., 2008. Hydrophilic surface modification of poly (vinyl chloride)
 film and tubing using physisorbed free radical grafting technique. Polymer, 49(20), 4350 4357.
- Miao, L., Hou, J., You, G., Liu, Z., Liu, S., Li, T., Mo, Y., Guo, S., Qu, H., 2019a. Acute effects of
 nanoplastics and microplastics on periphytic biofilms depending on particle size,
 concentration and surface modification. Environmental Pollution, 255, 113300.

- Miao, L., Wang, P., Hou, J., Yao, Y., Liu, Z., Liu, S., Li, T., 2019b. Distinct community structure
 and microbial functions of biofilms colonizing microplastics. Science of the Total
 Environment, 650, 2395-2402.
- Mohamed Nor, N.H., Koelmans, A.A., 2019. Transfer of PCBs from microplastics under
 simulated gut fluid conditions is biphasic and reversible. Environmental science and
 technology, 53(4), 1874-1883.
- Morét-Ferguson, S., Law, K.L., Proskurowski, G., Murphy, E.K., Peacock, E.E., Reddy, C.M.,
 2010. The size, mass, and composition of plastic debris in the western North Atlantic
 Ocean. Marine Pollution Bulletin, 60(10), 1873-1878.
- Nuzzo, A., Hosseinkhani, B., Boon, N., Zanaroli, G., Fava, F., 2017. Impact of bio-palladium nanoparticles (bio-Pd NPs) on the activity and structure of a marine microbial community.
 Environmental Pollution, 220, 1068-1078.
- Ogonowski, M., Motiei, A., Ininbergs, K., Hell, E., Gerdes, Z., Udekwu, K.I., Bacsik, Z.,
 Gorokhova, E., 2018. Evidence for selective bacterial community structuring on
 microplastics. Environmental microbiology, 20(8), 2796-2808.
- Pilevar, Z., Bahrami, A., Beikzadeh, S., Hosseini, H., Jafari, S.M., 2019. Migration of styrene monomer from polystyrene packaging materials into foods: characterization and safety evaluation. Trends in Food Science and Technology.
- PlasticsEurope, 2018. Plastics the Facts 2017. An Analysis of European Plastics
 Production, Demand and Waste Data. Brussels, Belgium.
- Rummel, C.D., Jahnke, A., Gorokhova, E., Kühnel, D., Schmitt-Jansen, M., 2017. Impacts of
 biofilm formation on the fate and potential effects of microplastic in the aquatic
 environment. Environmental Science and Technology Letters, 4(7), 258-267.

Ryan, P.G., Moore, C.J., van Franeker, J.A., Moloney, C.L., 2009. Monitoring the abundance of
 plastic debris in the marine environment. Philosophical Transactions of the Royal Society
 B: Biological Sciences, 364(1526), 1999-2012.

Sass, A.M., Sass, H., Coolen, M.J., Cypionka, H., Overmann, J., 2001. Microbial communities in
 the chemocline of a hypersaline deep-sea basin (Urania basin, Mediterranean Sea). Appl.
 Environ. Microbiol., 67(12), 5392-5402.

- Tanabe, S., Kannan, N., Subramanian, A., Watanabe, S., Tatsukawa, R., 1987. Highly toxic
 coplanar PCBs: occurrence, source, persistency and toxic implications to wildlife and
 humans. Environmental Pollution, 47(2), 147-163.
- Taniguchi, S., Colabuono, F.I., Dias, P.S., Oliveira, R., Fisner, M., Turra, A., Izar, G.M., Abessa,
 D.M.S., Saha, M., Hosoda, J., Yamashita, R., Takada, H., Lourenço, R.A., Magalhães, C.A.,
 Bícego, M.C., Montone, R.C., 2016. Spatial variability in persistent organic pollutants and
 polycyclic aromatic hydrocarbons found in beach-stranded pellets along the coast of the
 state of São Paulo, southeastern Brazil. Marine pollution bulletin, 106(1-2), 87-94.
- 734 Teuten, E.L., Saquing, J.M., Knappe, D.R., Barlaz, M.A., Jonsson, S., Björn, A., Rowland, S.J., 735 Thompson, R.C., Galloway, T.S., Yamashita, R., Ochi, D., Watanuki, Y., Moore, C., Viet, P.H., 736 Tana, T.S., Prudente, M., Boonyatumanond, R., Zakaria, M.P., Akkhavong, K., Ogata, Y., Hirai, 737 H., Iwasa, S., Mizukawa, K., Hagino, Y., Imamura, A., Saha, M., Takada H., 2009. Transport 738 and release of chemicals from plastics to the environment and to wildlife. Philosophical 739 Transactions of the Royal Society of London B: Biological Sciences, 364(1526), 2027-2045. Thomas, C., Lamperta, D., Reible, D., 2014. Remedy performance monitoring at 740 741 contaminated sediment sites using profiling solid phase microextraction (SPME) 742 polydimethylsiloxane (PDMS) fibers. Environmental science. Processes & Impacts, 16, 445-
- 743 452.

744	•	Thompson, R.C., Olsen, Y., Mitchell, R.P., Davis, A., Rowland, S.J., John, A.W., McGonigle, D.,
745		Russell, A.E., 2004. Lost at sea: where is all the plastic? Science, 304(5672), 838-838.
746	•	Van Cauwenberghe, L., Devriese, L., Galgani, F., Robbens, J., Janssen, C.R., 2015.
747		Microplastics in sediments: a review of techniques, occurrence and effects. Marine
748		environmental research, 111, 5-17.
749	•	Varjani, S.J., 2017. Microbial degradation of petroleum hydrocarbons. Bioresource
750		Technology, 223, 277-286.
751	•	Wang, F., Wong, C.S., Chen, D., Lu, X., Wang, F., Zeng, E.Y., 2018. Interaction of toxic
752		chemicals with microplastics: a critical review. Water research, 139, 208-219.
753	•	Wasmund, K., Mußmann, M., Loy, A., 2017. The life sulfuric: microbial ecology of sulfur
754		cycling in marine sediments. Environmental Microbiology Reports, 9(4), 323-344.
755	•	Wu, C.C., Bao, L.J., Liu, L.Y., Shi, L., Tao, S., Zeng, E.Y., 2017. Impact of polymer colonization
756		on the fate of organic contaminants in sediment. Environmental science and technology,
757		51(18), 10555-10561.
758	•	Xu, X., Wang, S., Gao, F., Li, J., Zheng, L., Sun, C., He, C. Wang, Z. Qu, L., 2019. Marine
759		microplastic-associated bacterial community succession in response to geography,
760		exposure time, and plastic type in China's coastal seawaters. Marine Pollution Bulletin, 145,
761		278-286.
762	•	Zanaroli, G., Negroni, A., Calisti, C., Ruzzi, M., Fava, F., 2011. Selection of commercial
763		hydrolytic enzymes with potential antifouling activity in marine environments. Enzyme
764		and microbial technology, 49(6), 574-579.
765	•	Zanaroli, G., Negroni, A., Häggblom, M.M., Fava, F., 2015. Microbial dehalogenation of
766		organohalides in marine and estuarine environments. Current Opinion in Biotechnology,

767 33, 287-295.

768	• Zettler, E.R., Mincer, T.J., Amaral-Zettler, L.A., 2013. Life in the "Plastisphere": Microbial
769	Communities on Plastic Marine Debris. Environmental Science and Technology, 47(13),
770	7137-7146.

Ziccardi, L.M., Edgington, A., Hentz, K., Kulacki, K.J., Kane Driscoll, S., 2016. Microplastics as
 vectors for bioaccumulation of hydrophobic organic chemicals in the marine environment:
 A state - of - the - science review. Environmental toxicology and chemistry, 35(7), 1667 1676.

775 Figures

Figure 1. Colonization dynamics of PCB-contaminated (A-C) and pristine (D-F) MPs determined via qPCR of 16S rRNA genes (A, D), reducing sugars in the EPS (B, E), and biofilm staining with crystal violet (C, F). Values are the mean of triplicate cultures ± standard deviation.

Figure 2. Composition at the Phylum level of the inoculated microbial community, the biofilm grown on all PCB-contaminated and pristine MPs after 2 weeks of incubation, the biofilm grown on PCB-contaminated and pristine PE and PVC after 28 weeks of incubation and the microbial community associated to the sediment particles surrounding the PCB-contaminated MPs after 2 weeks of incubation.

Figure 3. Boxplots showing the relative abundance (%) of Dehalococcoidia in pristine and
PCB-contaminated MPs. *: P-value = 0.05; Wilcoxon test.

Figure 4. PCoA plots of bacterial communities in the inoculum, in the sediment and associated
to MPs (A) and of bacterial communities associated to PCB-contaminated and to pristine MPs
(B).

Figure 5. Reductive dechlorination of PCBs sorbed on the sediment (biotic control microcosms) and on the different MP types (biologically active microcosms and sterile controls). Values (average number of chlorines/biphenyl molecule) are the mean of triplicate cultures ± standard deviation.

Figure 6. PCB congeners and their concentrations on MPs (A) and in the sediment (B) at the
beginning and the end of incubation (28 weeks). Value are the mean of triplicate cultures ±
standard deviation.

797 Supplementary information

Table S1. Composition at the genus level of the inoculated microbial community, of thebiofilm grown on all MP types after 2 weeks of incubation and after 28 weeks of incubation on

- 800 PE and PVC MPs, and of the microbial community associated to the sediment particles.
- 801 **Table S2.** Results of permutation test with pseudo-F ratio statistics applied to ordination
- analysis based on Jaccard similarity index (related to Figure 4A).
- 803 Figure S1. PCR-DGGE gels of the 16S rRNA genes of the inoculated marine microbial culture
- 804 (A) and the biofilm microbial community grown on pristine MPs (A) and on PCB-805 contaminated-MPs (B).
- Figure S2. Evolution of the biofilm community richness (Rr) and organization (Co) over time
 on PCB-contaminated (A) and pristine (B) MPs, based on PCR-DGGE analysis of 16S rRNA
 genes (Figure S1).





uncultured Bacteria Thermotogae Tenericutes Synergistetes Proteobacteria Planctomycetes ■ Lentisphaerae Firmicutes Elusimicrobia Deinococcus-Thermus Cyanobacteria Chloroflexi Bacteroidetes Atribacteria Armatimonadetes Actinobacteria Acidobacteria

Figure 3 Click here to download Figure: Figure 3.pdf

Dehalococcoidia



Relative abundance (%)

Figure 4 Click here to download Figure: Figure Jacard

4.2%

MDS2 -

В



MDS1 – 9.5% MDS1 – 9.5%

MDS1 – 9.5%



Incubation Time (weeks)



PCB congeners

Figure S1 Click here to download Supplementary material for on-line publication only: Figure S1.pdf Figure S2 Click here to download Supplementary material for on-line publication only: Figure S2.pdf
 Table S1

 Click here to download Supplementary material for on-line publication only: Table S1.xlsx

Table S2Click here to download Supplementary material for on-line publication only: Table S2.xlsx