



Provided by the author(s) and University of Galway in accordance with publisher policies. Please cite the published version when available.

Title	Microplastics in sewage sludge: effects of treatment
Author(s)	Mahon, A.M.; O'Connell, B.; Healy, Mark G.; O'Connor, I.; Officer, R.; Nash, R.; Morrison, Liam
Publication Date	2016-12-12
Publication Information	Mahon, A. M., O'Connell, B., Healy, M. G., O'Connor, I., Officer, R., Nash, R., & Morrison, L. (2017). Microplastics in Sewage Sludge: Effects of Treatment. <i>Environmental Science & Technology</i> , 51(2), 810-818. doi: 10.1021/acs.est.6b04048
Publisher	American Chemical Society
Link to publisher's version	http://dx.doi.org/10.1021/acs.est.6b04048
Item record	http://hdl.handle.net/10379/6698
DOI	http://dx.doi.org/10.1021/acs.est.6b04048

Downloaded 2024-04-26T20:59:30Z

Some rights reserved. For more information, please see the item record link above.



1 Published as: Mahon, A.M., O'Connell, B., Healy, M.G., O'Connor, I., Officer, R., Nash, R.,
2 Morrison, L. 2017. Microplastics in sewage sludge: effects of treatment. Environmental Science and
3 Technology 51(2): 810 – 818. DOI: [10.1021/acs.est.6b04048](https://doi.org/10.1021/acs.est.6b04048)

4
5

6 **Microplastics in Sewage Sludge: Effects of Treatment**

7 A.M. Mahon ^{a*}, B. O'Connell ^a, M.G. Healy ^b, I. O'Connor ^a, R. Officer ^a, R. Nash ^a, L.
8 Morrison ^c

9 ^aMarine and Freshwater Research Centre (MFRC), Galway-Mayo Institute of Technology, Dublin Road,
10 Galway, Ireland.

11 ^bCivil Engineering, National University of Ireland, Galway, Ireland.

12 ^cEarth and Ocean Sciences, Schools of Natural Sciences and Ryan Institute, National University of Ireland,
13 Galway, Ireland.

14 *Corresponding Author: Anne Marie Mahon: annemarie.mahon@gmit.ie

15

16 **Abstract**

17 Waste Water Treatment Plants (WWTPs) are receptors for the cumulative loading of
18 microplastics (MPs) derived from industry, landfill, domestic waste water and storm water.
19 The partitioning of MPs through the settlement processes of waste water treatment results in
20 the majority becoming entrained in the sewage sludge. This study characterised MPs in
21 sludge samples from seven WWTPs in Ireland, which use anaerobic digestion (AD), thermal
22 drying (TD), or lime stabilisation (LS) treatment processes. Abundances ranged from 4,196
23 to 15,385 particles kg⁻¹ (dry weight). Results of a general linear mixed model (GLMM)
24 showed significantly higher abundances of MPs in smaller size classes in the LS samples,
25 suggesting that the treatment process of LS shear MP particles. In contrast, lower abundances
26 of MPs found in the AD samples suggest that this process may reduce MP abundances.
27 Surface morphologies examined using Scanning Electron Microscopy (SEM) showed
28 characteristics of melting and blistering of TD MPs and shredding and flaking of LS MPs.
29 This study highlights the potential for sewage sludge treatment processes to affect the risk of

30 MP pollution prior to land spreading and may have implications for legislation governing the
31 application of biosolids to agricultural land.

32

33 **Keywords:** Microplastics; sewage sludge; biosolids; anaerobic digestion; lime stabilisation;
34 thermal drying.

35

36 1. Introduction

37 Microplastics (MPs) are synthetic polymers measuring less than 5 mm in diameter and are
38 derived from a wide range of sources including synthetic fibres from clothing,^{1,2} polymer
39 manufacturing and processing industries,³ and personal care products.⁴ They have the
40 potential to adsorb persistent organic contaminants^{5,6} and priority metals⁷⁻¹¹ from the
41 surrounding environment. These may be released upon digestion by biota or through
42 environmental degradation, leading to possible impacts to human health and ecosystems.¹²⁻¹⁴

43 Over the last 10 years, many studies have investigated the distribution^{1,15} and effects¹⁶⁻¹⁹ of
44 MPs within the marine environment. Indeed, MPs have been found in Polar Regions²⁰ and in
45 a range of freshwater environments worldwide.²¹⁻²⁴ Despite this, few studies have sought to
46 determine land-based sources of MPs.²⁵ Wastewater treatment plants (WWTPs) have been
47 identified as receptors of MP pollution and effective in capturing the majority of MPs in the
48 sludge during settlement regimes²⁶, as first found by Habib et al. (1998) when they used
49 synthetic fibers as a proxies for the presence of sewage.²⁷ More than 10 million tonnes of
50 sewage sludge was produced in WWTPs in the European Union (EU) in 2010.²⁷ European
51 Union policy on sustainability and recycling of resources²⁸ favours the recycling of sludge.

52 The introduction of EU legislation such as the Landfill Directive²⁹ and the Renewable Energy
53 Directive³⁰ have diverted sewage sludge from landfill and incineration into use for energy
54 production³¹ and agriculture.³² In some countries, such as Ireland, up to 80% of municipal

55 wastewater sludge is reused in agriculture.^{33,34} Guidelines stipulate that the sludge must
56 undergo some type of treatment (after which it is commonly referred to as ‘biosolids’) prior
57 to land application. This may include lime stabilisation (LS), anaerobic digestion (AD),
58 composting, or thermal drying (TD).³¹ As approximately 99% of MPs are retained in sewage
59 sludge generated in WWTPs,³⁵ there is a possibility that land applied sludge, even having
60 undergone treatment, could be a source of MP pollution.

61

62 The regulations for the use of biosolids in the EU and USA stipulate limit levels for pathogen
63 content, maximum metal and nutrient application rates to land,³⁶ and vector (flies and
64 rodents) attraction reduction (USA only). Restrictions in land application of biosolids vary
65 between the EU and USA. Under US federal legislation, the application of biosolids to
66 agricultural land can occur without restriction in volume or duration, if the contamination
67 level reaches an exceptional quality “EQ”.³⁷ In Europe, sewage sludge is dealt with very
68 differently among member states, and application to land is banned in some countries.³⁸⁻⁴⁰

69

70 As most sewage sludge undergoes treatment prior to land-spreading, the effects of these
71 treatments on MP morphology is important but remains largely unknown, with some
72 evidence of increased abundance of fibres at a smaller size range for LS sludge⁴¹ which is
73 probably due to alkaline hydrolysis.⁴² Therefore, the aim of this study was to investigate the
74 first stage of the MP pathway post-WWTP, and the impacts of different treatments. In
75 particular, it aimed to determine if (1) MPs are present in treated sewage sludge from a range
76 of WWTPs employing AD, TD and LS as treatment techniques, and (2) the type of treatment
77 used (TD, AD, LS) employed at the WWTP impacts on MP abundance and characteristics,
78 including size and surface morphology.

79

80 **2. Methodology**

81

82 2.1 WWTP sludge sample collection and preparation

83 Sewage sludge, having undergone treatment including TD, AD or LS, was collected from

84 seven waste WWTPs with population equivalents (PEs) ranging from 6500 to 2.4 million

85 (Table 1). These WWTPs received waste water from industry, storm water run-off and

86 domestic sources, all of which comprised up to 30% of the influent organic loading

87 (measured as biochemical oxygen demand, BOD) (Table1). Three replicate samples of 30 g

88 were obtained from each WWTP and stored at -20°C prior to sample preparation. The treated

89 sewage sludge had dry matter (DM) contents ranging from 24% (AD) to 87% (TD). Pellets of

90 TD sludge were placed in water for 1 week to induce softening, transferred to a water bath

91 (30°C) for 24 hr, and placed in an “end-over-end” shaker (Parvalux, UK) for 12 hr. This

92 shaking procedure was repeated until the pellets were sufficiently softened without

93 compromising the physical characteristics of the MPs. The samples were subsequently

94 washed through a 250 µm sieve, which resulted in complete degradation of the pelleted

95 clumps prior to elutriation. A proportion of the washed through fraction was retained and

96 passed through 212, 63, and 45 µm sieves for particle size determination or particle size

97 fractionation.

98

99 Anaerobically digested and LS sludge were soaked in filtered tap water to soften and

100 homogenise them, and were also washed through 250, 212, 63 and 45 µm sieves to determine

101 particle size fractions. As the LS sludge had an oily appearance, thought to be derived from

102 the break-down of cellulosic material through alkaline hydrolysis, it was decided that the

103 elutriation and other density separation techniques were unsuitable for extraction of MPs.

104 Instead, 10 g from each replicate sample were examined by passing it directly through a filter
105 (GF/C: Whatman TM, 1.2 µm) using vacuum filtration.

106

107 2.2. Microplastics Extraction

108 2.2.1 Elutriation

109 The principal of elutriation was used as the first step in the separation of MPs from other
110 sample components. Elutriation separates lighter particles from heavier ones through an
111 upward flow of liquid and/or gas, and has been widely used in the separation of biota within
112 sediment samples.⁴² To separate MPs from the sludge samples, an elutriation column, based
113 on the design of Claessens et al.⁴³ was constructed.

114

115 2.2.1.1 Column extraction efficiency estimation

116 To check for efficiency of the column in extracting MP, three sediment samples, each
117 weighing 40 g, were spiked with 50 MP particles of high density polyethylene (HDPE) (three
118 colours) and PVC, and run through the column. The HDPE samples used were shavings of
119 approximately 1.0 (L) × 4.0 (W) × 2.0 mm (B). The PVC particles were of a similar
120 dimension, but were more brittle. Therefore, each particle was marked with a blue marker to
121 ensure that particles were not counted twice upon recovery. The number of particles,
122 separated from the sediment matrix, that exited the column, was enumerated and the
123 percentage efficiency was calculated.

124

125 2.2.2. Zinc chloride (ZnCl₂) extraction

126 The MP extraction was filtered through 250 µm mesh, rinsed into a separatory funnel with 1
127 molar ZnCl₂ solution, and brought to a volume of 300 ml. The funnel was plugged,
128 vigorously shaken for 1 min, and allowed to settle (20 min). The settled material was drained

129 and the remainder of the sample was filtered onto glass fibre filters (GF/C: Whatman TM, 1.2
130 µm). The oily appearance of the LS samples rendered this density separation technique
131 unsuitable for extraction of MP.

132

133 2.3. Characterisation of MPs

134 The filters were examined using stereomicroscopy equipped with a polariser (Olympus
135 SZX10) attachment and a Qimaging[®] RetigaTM 2000R digital camera. Microplastics were
136 identified and enumerated based on several criteria including form, colour and sheen used in
137 previous studies as described by Hidalgo Ruz et al.⁴⁴ The form of a synthetic fibre should not
138 taper at either end, while not having a rigidly straight form. Any polymer will not have
139 cellular structure or other organic structures. Artificial fibre particles also have uniformity of
140 colour and exhibit a sheen once passed through the polarized light. Where ambiguity
141 remained following these observations, the suspected polymer was manipulated with a hot
142 pin by which a melted form indicated a positive result. Microplastics were measured and
143 allotted to the following size categories: 250-400 µm, 400-600 µm, 600-1000 µm, and 1000-
144 4000 µm. Suspected MPs were enumerated and measured, and approximately 10% of MP
145 samples from each filter paper were set aside for polymer identification. Microplastics for
146 which any ambiguity remained as to if it was a polymer, were automatically selected for
147 analyses.

148

149 Attenuated total reflectance (ATR) and Fourier transform infrared spectroscopy (FTIR)
150 (Perkin Elmer, USA, Spectrum TwoTM with Universal ATR Accessory and Thermo
151 Scientific, UK, Nicolet iN10 FTIR microscope with germanium Tip Slide-on-ATR) were
152 used to analyse approximately 10% of MP samples. The spectra were obtained with 3-second
153 data collection (16 scans per sample) over the wave number range 600 – 4000 cm⁻¹ using a

154 liquid nitrogen-cooled MCT-A detector at 8 cm⁻¹ resolution. Microplastic samples extracted
155 from the sludge (and pristine plastics for comparative purposes) were gold-coated (Emitecg
156 K550, Quorum technologies, Ltd., UK) and subjected to variable pressure scanning electron
157 microscopy (SEM) in secondary electron mode using a Hitachi model S2600N (Hitachinaka,
158 Japan). The analyses were performed at accelerating voltages of 10 - 20 kv, an emission
159 current (I_c) of 10 µA, and a working distance of 12 - 24mm.⁴⁴

160

161 2.4 Quality control and contamination prevention

162 Cotton laboratory coats and nitrile gloves were used during the sample preparation and
163 analyses. In addition, synthetic clothing was avoided and samples were covered at all times
164 and working surfaces were cleaned with alcohol prior to use. When analysing filter papers, a
165 blank filter paper was exposed to the open laboratory conditions to assess the possibility of
166 air-borne contamination.

167

168 2.5. Data analyses

169 Statistical analyses were carried out using Minitab 17 (2010) and R.⁴⁵ As data were not
170 normally distributed, non-parametric tests were used to test for differences in MP abundances
171 amongst locations (Mann-Whitney Test). To investigate if there were any possible effects of
172 PE on abundance, a Spearman's rank correlation analysis test was utilised. With the
173 exception of one WWTP, there was only one treatment method employed per site (Table 1),
174 so in-site correlation was not possible. Each site was treated as an independent measurement
175 and plotted using a box plot. A generalised linear mixed effect model (GLMM) was used
176 (Eqn. 1) to investigate the high number of MP particles in the smaller class sizes at WWTPs
177 in which LS was employed.

178

179 Microplastic counts = Treatment Type + Population Equivalent + $\frac{1}{\text{Treatment Plant}}$

180 Eqn. 1

181

182 Where $I/\text{Treatment Plant}$ specifies a random intercept model.

183

184 A separate GLMM for each size class was carried out using a Poisson distribution and a
185 random effect term to account for nesting of replicates within WWTPs to determine which
186 explanatory variable was responsible for larger proportions of smaller MP particles at
187 WWTPs in which LS was employed.

188

189 **3. Results and Discussion**

190

191 3.1 Characterisation of treated sewage sludge

192 The characteristics of the sewage sludge treated using AD, LS and TD had varying physical
193 characteristics. The particle size fractionation (g/kg) of the AD samples was smaller than the
194 LS and TD samples (Table 2), and had a sandy appearance. The AD samples were very dark
195 and heavy with some cellulosic material, whereas the TD samples had a lot of cellulosic
196 material entrained, which was difficult to separate during elutriation and zinc chloride
197 extraction. Although this cellulosic material was distinctive from MP material (in that its
198 fibres tapered at the ends and it was often branched) and therefore easy to disqualify, its
199 presence in the samples greatly increased the time and consumables (filter papers) utilised
200 during the filtration process. High levels of cellulose derived from toilet paper in sewage may
201 merit the inclusion of a digestion process using the cellulase enzyme, as has been previously
202 used for the isolation of MPs in North Sea sediments.⁴⁶

203

204 3.2 Microplastics Extraction

205 3.2.1 Elutriation column extraction efficiency estimation

206 The average extraction efficiency rate of the elutriation column for the spiked sediment
207 samples was 90%, 94% and 91% for the red, blue and black HDPE particles, respectively.
208 The elutriation process was less efficient for the PVC particles, which resulted in an average
209 extraction efficiency of 80%. This is an indication that results of MP abundance in this study
210 may be an underestimation. As the efficiency test was conducted only for fragments at one
211 size only, it may not be representative of efficiency of fibre removal.

212

213 3.3 Characterisation of Microplastics

214 3.3. 1 Microplastics abundance

215 Microplastics extracted from the biosolids ranged from an average of 4,196 to 15,385
216 particles kg⁻¹ (DM) among the seven sites, with significant differences in MP abundances
217 between some sites and within Site 1 (1A, 1B) between AD samples and TD samples (Mann
218 Whitney, $w = 15$, $p = 0.0809$; Figure 1). This is likely to be an underestimation due to losses
219 in column efficiency (approx. 20%) and through the use of a 250 µm sieve from which a
220 proportion of fibres may be lost. The abundances found in this study are in the same order of
221 magnitude to the study by Zubris et al.⁴² who reported between 3,000 and 4,000 particles kg⁻¹

222 ¹. In the current study, a lack of correlation between PE and MP abundance kg⁻¹ (Spearman's
223 rank, $r = -0.308$, $p = 0.458$) implies that these differences may have been due to the variation
224 of input sources (industrial, storm water, landfill etc.). However, as no data exist for the
225 temporal variation of MPs in sewage sludge, there is a possibility that these variations are a
226 result of fluxes in MP input, which could be a result of peak MP emission times in relation to
227 household or industrial activity. The significantly lower abundance of MPs in an
228 anaerobically digested biosolid sample compared to all other sample except Site 3, which was

229 also treated with AD, posits an interesting question over the possible role of AD in the
230 degradation of polymers collected from the same site as sample 1A (taken roughly at the
231 same time). Without pre-treatment samples, there is no evidence to prove that the mesophilic
232 anaerobic digestion (MAD) used at the AD WWTPs in this study, facilitated the breakdown
233 of MPs. Indeed, few studies have examined the breakdown of polymers in anaerobic
234 digesters. However, one pilot study investigated the effect of plastic waste on the functioning
235 of anaerobic digestion and found that digesters from which plastic was removed, produced
236 less gas than those to which plastic was added.⁴⁷ As there is already substantial evidence of
237 microbial breakdown of polymers through the activity of exoenzymes (promoting
238 depolymerisation) and assimilation of smaller articles resulting in mineralisation,^{49 50,51} the
239 role of degradation by microorganisms within the AD systems should be further investigated.

240

241 3.3.2 Morphological categorization and polymer identification of microplastics

242 This study confirmed that MPs are retained in the sewage sludge and are largely composed of
243 fibres, similar to what was found by Talvite et al.⁴⁶ and Magnusson and Norén.³⁵
244 Approximately 75.8% of the MP consisted of fibres, followed by fragments, films, other
245 unidentified particles, and spheres, which accounted for only 0.3% of total MP abundance
246 (Table 3). The greatest proportion of MP fragments was found at the LS WWTPs, with Site 6
247 being the only site to have marginally more fragments than fibres (Table 3; Figure 2).

248 Polymers, identified by FTIR, comprised HDPE, polyethylene (PE) polyester, acrylic,
249 polyethylene terephthalate (PET), polypropylene, and polyamide (Figure 3). Some of these
250 contained minerals. Although waste water derived from households generate high quantities
251 of fibres, principally derived from clothes washing of >1900 fibres per wash¹, other industrial
252 sources of fibres such as the fibre manufacturing industry may also be important contributors.

253

254 3.3.3. Size of micrplastics
255 Using the fitted coefficients from the GLMM, a study hypotheses of no difference between
256 all pairwise combinations of the treatment effects were tested. At small and medium particle
257 sizes, the LS treatment was significantly different from both TD and AD treatments (Figure
258 4; $P < 0.001$; sizes classes A and C; $P < 0.05$ size class B). The larger number of smaller MP
259 particles in LS samples corresponded with the larger proportion of smaller particle sizes
260 determined from the particle size fractionation. As it was not possible to obtain pre-treatment
261 samples, it is not possible to wholly assign the differences in size classes to the treatment
262 processes. However, the elevated numbers at the small size classes for LS samples are in
263 agreement with results reported by Zubris and Richards⁴², where there was some evidence of
264 elevated abundance of MPs at smaller size classes. Further investigations are required to
265 investigate accelerated proliferation of MP pollution through sludge treatment processes.

266
267 3.3.4 Surface morphologies of microplastics
268 Scanning electron micrographs of surface textures of polymers entrained in the treated
269 biosolids had some surface morphologies, which varied among treatment type. An unknown
270 polymer fibre, which was thermally dried, had distinct blistering and fracturing, particularly
271 in the fibre curves (Figure 5: A-C). Additionally, polymer fragments from TD samples,
272 identified as HDPE and PE fragments, showed wrinkling, melding and some fracturing,
273 which was quite distinct from pre-treatment samples (Figure 6: G-I; Figure 7: D-F). Surface
274 morphologies of MPs originating from LS biosolids had a more shredded and flaked
275 appearance for the unknown polymer (Figure 5: D-F) and a HDPE sample (Figure 5: D-F).
276 Anaerobically digested samples of an unknown polymer had deep cleavage, which was
277 distinct from any other observations (Figure 5: G-I).

278

279 **4. Conclusions**

280 Although it was not possible to assign wholly the abundances or size distributions to the
281 treatment processes, results suggest that treatment processes may have an effect. If MPs are
282 altered by treatment, the potential for impact may also be influenced depending accordingly.
283 This could add to the unknown risks associated with MPs in sewage sludge. Regardless of
284 treatment regimes, over time, there may be consequences for the accumulation of MPs in
285 terrestrial, freshwater, or marine ecosystems derived from land-spreading of sewage sludge or
286 biosolids.

287

288 Microplastics entrained in biosolids which are applied to land, may be degraded through
289 photo-degradation and thermo-oxidative degradation^{49,53} exacerbating the problem of land-
290 spread MP pollution. The interaction of MPs with contaminants in the soil, could have major
291 consequences for the absorption and transportation of contamination elsewhere. Surface
292 weathering and the subsequent attachment of organic matter and the resulting negative charge
293 attracts metals including cadmium, lead and zinc.⁵³ Whether agricultural land is a sink or a
294 source of MP pollution remains unclear. Microplastic fibres have been found on land 15
295 years post application, and some evidence of vertical translocation through the soil has also
296 been found.⁴¹ Possible impacts arising from land-applied MPs begin in the terrestrial
297 ecosystem with implications for terrestrial species such as earth worms⁵⁵ and birds feeding on
298 terrestrial ecosystems.⁵⁶ As legislation in the EU and the US generally permit the land
299 application of sewage sludge, there is a strong possibility that large amounts of MPs are
300 emitted to freshwater, where currently little is known about their impacts on species and
301 habitats.⁵⁷ Furthermore, buffer zones around freshwater bodies, which may be stipulated in
302 “codes of good practice”, do not take into account the mechanisms of transportation of MP
303 vertically through the soil or with surface runoff following a precipitation event. While

304 legislation currently takes into account pathogens as well as nutrient and metal concentrations
305 of treated sludge,⁵⁸ it does not consider the presence of MPs within the sludge, and their
306 associated risks. The predicted exponential growth of the plastics industry for the coming
307 years⁵⁹ may be accompanied by a significant increase in MPs in the waste stream. Therefore,
308 vigilant management of cumulative sources of MPs such as sewage sludge or biosolids is
309 necessary. In particular, this study has highlighted the potential for treatment processes to
310 alter the counts of MPs which, in turn, increases the available area for absorption/desorption
311 of organic pollutants.

312

313 A review of sewage sludge treatment processes and their implications for MP pollution
314 should be more thoroughly investigated, with before and after treatment comparisons. In
315 particular, the role of degradation by microorganisms within the AD systems should be
316 further investigated. Knowledge gaps regarding the factors critical for the mobilisation and
317 transport of MPs likely to affect the pathway of land-spread sewage sludge MP pollution
318 should be addressed in order to determine MP flow within the terrestrial system and to
319 freshwater systems. Only when experimental data are acquired, can we estimate exposure
320 and associated risks to the environment from MP pollution.

321

322

323 Supporting Information

324 Detailed description of the dimensions of the elutriation column, accompanied by a
325 photograph and schematic representation. Flow rates and technique used for extraction of
326 MPs using the elutriation column are also included.

327

328

329 Acknowledgements

330 We acknowledge the technical assistance of Mark Deegan in construction of our Elutriation
331 system, Mark Croke and David James from Thermo Fisher Scientific UK for FTIR analyses
332 and the Environmental Protection Agency of Ireland for funding this research.

333

334 **Bibliography**

- 335 (1) Browne, M. A.; Crump, P.; Niven, S. J.; Teuten, E.; Tonkin, A.; Galloway, T.;
336 Thompson, R. Accumulation of Microplastic on Shorelines Worldwide: Sources and Sinks.
337 *Environ. Sci. Technol.* **2011**, *45* (21), 9175–9179.
- 338 (2) Astrom, L. Shedding of synthetic microfibers from textiles. University of Gothenburg,
339 Sweden 2016.
- 340 (3) Lechner, A.; Ramler, D. The discharge of certain amounts of industrial microplastic from
341 a production plant into the River Danube is permitted by the Austrian legislation. *Environ.*
342 *Pollut.* **2015**, *200C*, 159–160.
- 343 (4) Fendall, L. S.; Sewell, M. A. Contributing to marine pollution by washing your face:
344 microplastics in facial cleansers. *Mar. Pollut. Bull.* **2009**, *58* (8), 1225–1228.
- 345 (5) Teuten, E. L.; Saquing, J. M.; Knappe, D. R. U.; Barlaz, M. A.; Jonsson, S.; Björn, A.;
346 Rowland, S. J.; Thompson, R. C.; Galloway, T. S.; Yamashita, R.; et al. Transport and release
347 of chemicals from plastics to the environment and to wildlife. *Philos. Trans. R. Soc. London*
348 *B Biol. Sci.* **2009**, *364* (1526), 2027–2045.
- 349 (6) Engler, R. E. The Complex Interaction between Marine Debris and Toxic Chemicals in
350 the Ocean. *Environ. Sci. Technol.* **2012**, *46* (22), 12302–12315.
- 351 (7) Ashton, K.; Holmes, L.; Turner, A. Association of metals with plastic production pellets
352 in the marine environment. *Mar. Pollut. Bull.* **2010**, *60* (11), 2050–2055.
- 353 (8) Holmes, L. A.; Turner, A.; Thompson, R. C. Adsorption of trace metals to plastic resin
354 pellets in the marine environment. *Environ. Pollut.* **2012**, *160*, 42–48.
- 355 (9) Nakashima, E.; Isobe, A.; Kako, S.; Itai, T.; Takahashi, S. Quantification of Toxic Metals
356 Derived from Macroplastic Litter on Ookushi Beach, Japan. *Environ. Sci. Technol.* **2012**, *46*
357 (18), 10099–10105.
- 358 (10) Rochman, C. M.; Hentschel, B. T.; Teh, S. J. Long-Term Sorption of Metals Is Similar
359 among Plastic Types: Implications for Plastic Debris in Aquatic Environments. *PLoS One*
360 **2014**, *9* (1), e85433.
- 361 (11) Brennecke, D.; Duarte, B.; Paiva, F.; Caçador, I.; Canning-Clode, J. Microplastics as
362 vector for heavy metal contamination from the marine environment. *Estuar. Coast. Shelf Sci.*
363 **2016**, *178*, 189–195.
- 364 (12) Cooper, D. A.; Corcoran, P. L. Effects of mechanical and chemical processes on the
365 degradation of plastic beach debris on the island of Kauai, Hawaii. *Mar. Pollut. Bull.* **2010**,
366 *60* (5), 650–654
- 367 (13) Andrade, A. L. Microplastics in the marine environment. *Mar. Pollut. Bull.* **2011**, *62*
368 (8), 1596–1605.

- 369 (14) Bouwmeester, H.; Hollman, P. C. H.; Peters, R. J. B. Potential Health Impact of
370 Environmentally Released Micro- and Nanoplastics in the Human Food Production Chain:
371 Experiences from Nanotoxicology. *Environ. Sci. Technol.* **2015**, *49* (15), 8932–8947.
372 (15) Cole, M.; Lindeque, P.; Halsband, C.; Galloway, T. S. Microplastics as contaminants in
373 the marine environment: a review. *Mar. Pollut. Bull.* **2011**, *62* (12), 2588–2597.
374 (16) von Moos, N.; Burkhardt-Holm, P.; Köhler, A. Uptake and Effects of Microplastics on
375 Cells and Tissue of the Blue Mussel *Mytilus edulis* L. after an Experimental Exposure.
376 *Environ. Sci. Technol.* **2012**, *46* (20), 11327–11335.
377 (17) Cole, M.; Lindeque, P.; Fileman, E.; Halsband, C.; Goodhead, R.; Moger, J.; Galloway,
378 T. S. Microplastic Ingestion by Zooplankton. *Environ. Sci. Technol.* **2013**, *47* (12), 6646–
379 6655.
380 (18) Remy, F.; Collard, F.; Gilbert, B.; Compère, P.; Eppe, G.; Lepoint, G. When
381 Microplastic Is Not Plastic: The Ingestion of Artificial Cellulose Fibers by Macrofauna
382 Living in Seagrass Macrophytodebris. *Environ. Sci. Technol.* **2015**, *49* (18), 11158–11166.
383 (19) Watts, A. J. R.; Urbina, M. A.; Goodhead, R. M.; Moger, J.; Lewis, C.; Galloway, T. S.
384 Effect of microplastic on the gills of the Shore Crab *Carcinus maenas*. *Environ. Sci. Technol.*
385 **2016**.
386 (20) Lusher, A. L.; Burke, A.; O'Connor, I.; Officer, R. Microplastic pollution in the
387 Northeast Atlantic Ocean: Validated and opportunistic sampling. *Mar. Pollut. Bull.* **2014**, *88*
388 (1), 325–333.
389 (21) Eriksen, M.; Mason, S.; Wilson, S.; Box, C.; Zellers, A.; Edwards, W.; Farley, H.;
390 Amato, S. Microplastic pollution in the surface waters of the Laurentian Great Lakes. *Mar.*
391 *Pollut. Bull.* **2013**, *77* (1–2), 177–182.
392 (22) McCormick, A.; Hoellein, T. J.; Mason, S. A.; Schluempf, J.; Kelly, J. J. Microplastic is an
393 Abundant and Distinct Microbial Habitat in an Urban River. *Environ. Sci. Technol.* **2014**, *48*
394 (20), 11863–11871.
395 (23) Castañeda, R. A.; Avlijas, S.; Simard, M. A.; Ricciardi, A. Microplastic pollution in St.
396 Lawrence River sediments. *Can. J. Fish. Aquat. Sci.* **2014**, *71* (12), 1767–1771.
397 (24) Free, C. M.; Jensen, O. P.; Mason, S. A.; Eriksen, M.; Williamson, N. J.; Boldgiv, B.
398 High-levels of microplastic pollution in a large, remote, mountain lake. *Mar. Pollut. Bull.*
399 **2014**, *85* (1), 156–163.
400 (25) Lechner, A.; Ramler, D. The discharge of certain amounts of industrial microplastic
401 from a production plant into the River Danube is permitted by the Austrian legislation.
402 *Environ. Pollut.* **2015**, *200C*, 159–160.
403 (26) Carr, S. A.; Liu, J.; Tesoro, A. G. Transport and Fate of Microplastic Particles in
404 Wastewater Treatment Plants. *Water Res.* **2016**, *91*, 174–182.
405 (27) Habib, D.; Locke, D. C.; Cannone, L. J. Synthetic Fibers as Indicators of Municipal Sewage Sludge,
406 Sludge Products, and Sewage Treatment Plant Effluents. *Water. Air. Soil Pollut.* **1998**, *103* (1), 1–8.
407 (28) Eurostat (2014) Sewage sludge production and disposal.
408 http://appsso.eurostat.ec.europa.eu/nui/show.do?dataset=env_ww_spd&lang=en
409 (29) Commission of the European Communities (COM) (2014) Towards a circular economy:
410 a zero waste programme for Europe. [http://eur-
411 lex.europa.eu/resource.html?uri=cellar:aa88c66d-4553-11e4-a0cb-
412 01aa75ed71a1.0022.03/DOC_1&format=PDF](http://eur-lex.europa.eu/resource.html?uri=cellar:aa88c66d-4553-11e4-a0cb-01aa75ed71a1.0022.03/DOC_1&format=PDF)
413 (30) EC, 1999. Economic Commission. Council Directive of the 26 April 1999 on landfill
414 waste. (1999/31/EC) Available at: [http://eur-lex.europa.eu/legal-
415 content/EN/TXT/?uri=CELEX%3A31999L0031](http://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX%3A31999L0031)
416 (31) EC, 2009. Directive 2009/28/EC of the European parliament and of the Council of 23
417 April 2009 on the promotion of use of energy from renewable sources and amending
418 subsequently repealing Directives 2001/77/EC and 2003/30/EC.

- 419 (32) Gikas, P. Electrical energy production from biosolids: a comparative study between
420 anaerobic digestion and ultra-high-temperature gasification. *Environ. Technol.* **2014** *35*,
421 2140-2146.
- 422 (33) Healy, M.G., Clarke, R., Peyton, D., Cummins, E., Moynihan, E.L., Martins, A., Beraud,
423 P., Fenton, O. Resource recovery from sludge. p. 139 – 162. In: K. Konstantinos and K.P.
424 Tsagarakis, Eds.) *Sewage treatment plants: economic evaluation of innovative technologies*
425 for energy efficiency. IWA, London, 2015. ISBN: 9781780405018.
- 426 (34) EPA. Urban waste water treatment in 2014.
http://www.epa.ie/pubs/reports/water/wastewater/2014%20waste%20water%20report_web.pdf
- 427 (35) Eurostat 2016. Sewage Sludge production and disposal.
<http://appsso.eurostat.ec.europa.eu/nui/submitViewTableAction.do>.
- 431 (36) Magnusson K., Norén F., 2014. Screening of microplastic particles in and downstream
432 from a wastewater treatment plant. Report to the Swedish Environmental Research Institute:
433 C55.
- 434 (37) Lucid, J. D., Fenton, O., Grant, J., & Healy, M. G. Effect of Rainfall Time Interval on
435 Runoff Losses of Biosolids and Meat and Bone Meal when Applied to a Grassland Soil.
436 *Water Air Soil Pollut.* **2014**, *225* (8), 1-11.
- 437 (38) Harrison, E. Z.; McBride, M. B.; Bouldin, D. R. Land application of sewage sludges: an
438 appraisal of the US regulations. *Int. J. Environ. Pollut.* **1999**, *11* (1), 1–36.
- 439 (39) Environmental, economic and social impacts of the use of sewage sludge on land. Final
440 report- Part I; Overview Report. European Commission Service Service Contract.
441 No.070307/2008/517358/ETU/G4; 2013.
- 442 (40) Environmental, economic and social impacts of the use of sewage sludge on land. Final
443 report- Part I; Report on Options and Impacts. European Commission Service Contract
444 No.070307/2008/517358/ETU/G4; 2013.
- 445 (41) Environmental, economic and social impacts of the use of sewage sludge on land. Final
446 report- Part I; Project Interim Reports. European Commission Service Contract
447 No.070307/2008/517358/ETU/G4; 2013.
- 448 (42) Zubris, K. A. V; Richards, B. K. Synthetic fibers as an indicator of land application of
449 sludge. *Environ. Pollut.* **2005**, *138* (2), 201–211.
- 450 (43) Walling, D. E.; Woodward, J. C. Use of a field-based water elutriation system for
451 monitoring the in situ particle size characteristics of fluvial suspended sediment. *Water Res.*
452 **1993**, *27* (9), 1413–1421.
- 453 (44) Claessens, M.; Van Cauwenbergh, L.; Vandegehuchte, M. B.; Janssen, C. R. New
454 techniques for the detection of microplastics in sediments and field collected organisms. *Mar.*
455 *Pollut. Bull.* **2013**, *70* (1-2), 227–233.
- 456 (45) Morrison, L., Feely, M., Stengel, D. B., Blamey, N., Dockery, P., Sherlock, A. and
457 Timmins, É. Seaweed attachment to bedrock: biophysical evidence for a new geophycology
458 paradigm. *Geobiol.* **2009** *7*: 477-487 (DOI: 10.1111/j.1472-4669.2009.00206.x).
- 459 (46) Lorenz, C. (2014): Detection of microplastics in marine sediments of the German Coast
460 via FT-IR spectroscopy, Master thesis, Universität Rostock.
- 461 (47) Morrison, L.; Feely, M.; Stengel, D. B.; Blamey, N.; Dockery, P.; Sherlock, A.;
462 Timmins, E. Seaweed attachment to bedrock: biophysical evidence for a new geophycology
463 paradigm. *Geobiology* **2009**, *7* (4), 477–487.
- 464 (48) Talvitie, J.; Heinonen, M.; Pääkkönen, J.-P.; Vahtera, E.; Mikola, A.; Setälä, O.; Vahala,
465 R. Do wastewater treatment plants act as a potential point source of microplastics?

- 466 Preliminary study in the coastal Gulf of Finland, Baltic Sea. *Water Sci. Technol.* **2015**, *72* (9),
467 1495–1504.
- 468 (49) Kamal, M.R, Huang, B. *Handbook of Polymer Degradation*, Marcek Dekker, New York,
469 127-168, 1992.
- 470 (50) Shah, A. A.; Hasan, F.; Hameed, A.; Ahmed, S. Biological degradation of plastics: a
471 comprehensive review. *Biotechnol. Adv.* **2008**, *26* (3), 246–265.
- 472 (51) Andrade, A. L.; Hamid, S. H.; Hu, X.; Torikai, A. Effects of increased solar ultraviolet
473 radiation on materials. *J. Photochem. Photobiol. B Biol.* **1998**, *46* (1-3), 96–103.Gu, J.-D.
474 Microbiological deterioration and degradation of synthetic polymeric materials: recent
475 research advances. *Int. Biodeterior. Biodegradation* **2003**, *52* (2), 69–91.
- 476 (52) Yoshida, S.; Hiraga, K.; Takehana, T.; Taniguchi, I.; Yamaji, H.; Maeda, Y.; Toyohara,
477 K.; Miyamoto, K.; Kimura, Y.; Oda, K. A bacterium that degrades and assimilates
478 poly(ethylene terephthalate). *Science* **2016**, *351* (6278), 1196–1199.
- 479 (53) Arthat, T.; Doble, M. Biodegradation of Aliphatic and Aromatic Polycarbonates.
480 *Macromol. Biosci.* **2008**, *8* (1), 14–24.
- 481 (54) Contat-Rodrigo, L. Thermal characterization of the oxo-degradation of polypropylene
482 containing a pro-oxidant/pro-degradant additive. *Polym. Degrad. Stab.* **2013**, *98* (11), 2117–
483 2124.
- 484 (55) Turner, A.; Holmes, L. A. Adsorption of trace metals by microplastic pellets in fresh
485 water. *Environmental Chemistry*. 2015, pp 600–610.
- 486 (56) Peyton, D.P., Healy, M.G., Fleming, G.T.A., Grant, J., Wall, D., Morrison, L.,
487 Cormican, M., Fenton, O. Nutrient, metal and microbial loss in surface runoff following
488 treated sludge and dairy cattle slurry application to an Irish grassland soil. *Sci Total Environ.*
489 **2016**, *541*, 218-229.
- 490 (57) Huerta Lwanga, E.; Gertsen, H.; Gooren, H.; Peters, P.; Salánki, T.; van der Ploeg, M.;
491 Besseling, E.; Koelmans, A. A.; Geissen, V. Microplastics in the Terrestrial Ecosystem:
492 Implications for Lumbricus terrestris (Oligochaeta, Lumbricidae). *Environ. Sci. Technol.*
493 **2016**, *50* (5), 2685–2691.
- 494 (58) Zhao, S.; Zhu, L.; Li, D. Microscopic anthropogenic litter in terrestrial birds from
495 Shanghai, China: Not only plastics but also natural fibers. *Sci. Total Environ.* **2016**, *550*,
496 1110–1115.
- 497 (59) Eerkes-Medrano, D.; Thompson, R. C.; Aldridge, D. C. Microplastics in freshwater
498 systems: a review of the emerging threats, identification of knowledge gaps and prioritisation
499 of research needs. *Water Res.* **2015**, *75*, 63–82.
- 500 (60) Healy, M. G.; Fenton, O.; Forrestal, P. J.; Danaher, M.; Brennan, R. B.; Morrison, L.
501 Metal concentrations in lime stabilised, thermally dried and anaerobically digested sewage
502 sludges. *Waste Manag.* **2016**, *48*, 404–408.
- 503 (61) Plastics Europe. An Analysis of European plastic production, demand and waste data for
504 2011. Brussels, Belguim, 2012.

505

506

507

508

509 Table 1. Characteristics of municipal wastewater treatment sites investigated (adapted from
 510 Healy et al., 2016)

Site	WWTP/ agglomeration size (PEs)	Landfill leachate as % of influent BOD load	Industrial, and domestic septic tank sludge ¹ as % of influent BOD load	Type of treatment
1A	2,362,329	<0.01	<0.01	Thermal drying, anaerobic digestion
1B	284,696	0.3	24	Thermal drying
2	179,000	unknown	30	Anaerobic digestion
3	130,000	unknown	0.008	Thermal drying
4	101,000	2.0	unknown	Lime stabilisation
5	31,788	0.25	unknown	Lime stabilisation
6	25,000	0.7	0	Thermal drying
7	6,500	Unknown	Unknown	Thermal drying

511 ¹ Most recent available figures in all WWTPs (2013)

512
 513
 514
 515
 516
 517
 518
 519
 520
 521
 522
 523
 524

525 Table 2. Particle size fraction (g) of lime stabilised (LS), anaerobically digested (AD) and
 526 thermally dried (TD) samples (40 g).

Size fraction	Treatment type		
	LS	AD	TD
> 212 µm	3.004 ± 0.550	31.753± 0.578	35.503± 0.661
> 63 µm	27.410± 0.840	7.948± 0.7778	3.593± 0.894
> 45 µm	9.400± 1.166	0.327± 0.241	0.930± 0.486
< 45 µm	0.200 ± 0.213	0.000± 0.00	0.000± 0.000

527

528

529

530

531

532

533

534

535

536

537

538

539

540

541

542

543

544

545

546

547

548 Table 3. Breakdown of types of average microplastic abundance kg⁻¹ (dry matter) among
 549 sites.

Site no.	Treatment	Microplastic Types				
		Fibres	Fragments	Films	Spheres	other
1A	TD	9,113	511	255	89	44
1B	AD	2,065	611	67	0	0
2	TD	5,583	588	222	44	67
3	AD	4,007	855	111	33	150
4	TD	13,675	1,143	366	33	178
5	LS	10,778	3,075	122	11	78
6	LS	4,762	5,228	11	0	11
7	TD	3,463	511	167	0	56
Total	-	53,447	12,521	1,321	211	583
%	-	78.5	18.4	1.9	0.3	0.9

550

551

552

553

554

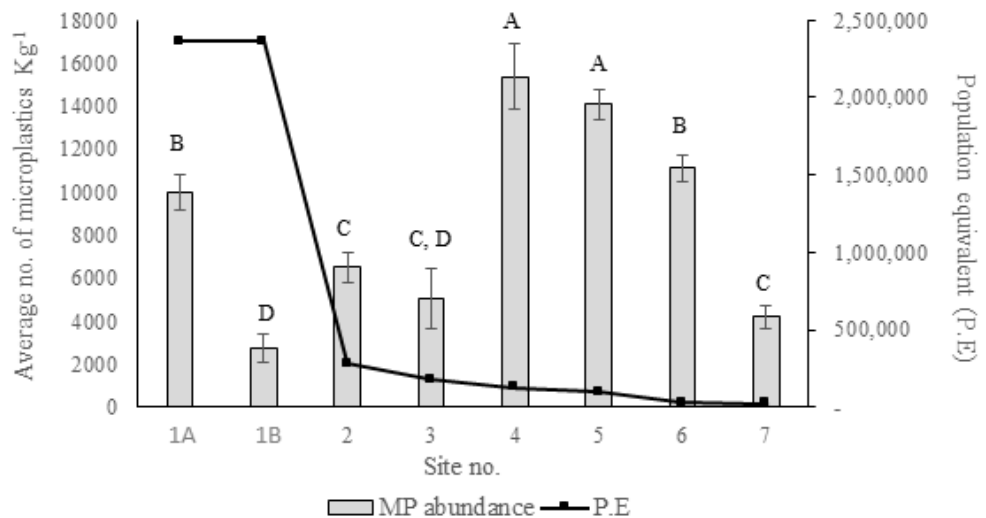
555

556

557

558

559



560

561 Figure 1. Average abundances and corresponding population equivalents of microplastics at 7
 562 sites. Sites sharing the same letter are not significantly different (Mann-Whitney-U test, $p >$
 563 0.005)

564

565

566

567

568

569

570

571

572

573

574

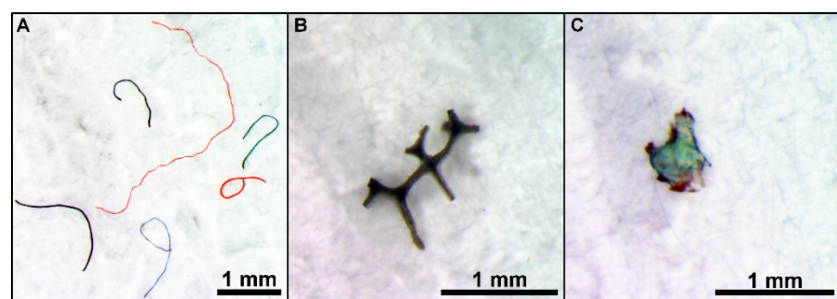
575

576

577

578

579



580

581 Figure 2. Stereomicrograph of mircoplastics fibres (A), other (B) and fragment (C).

582

583

584

585

586

587

588

589

590

591

592

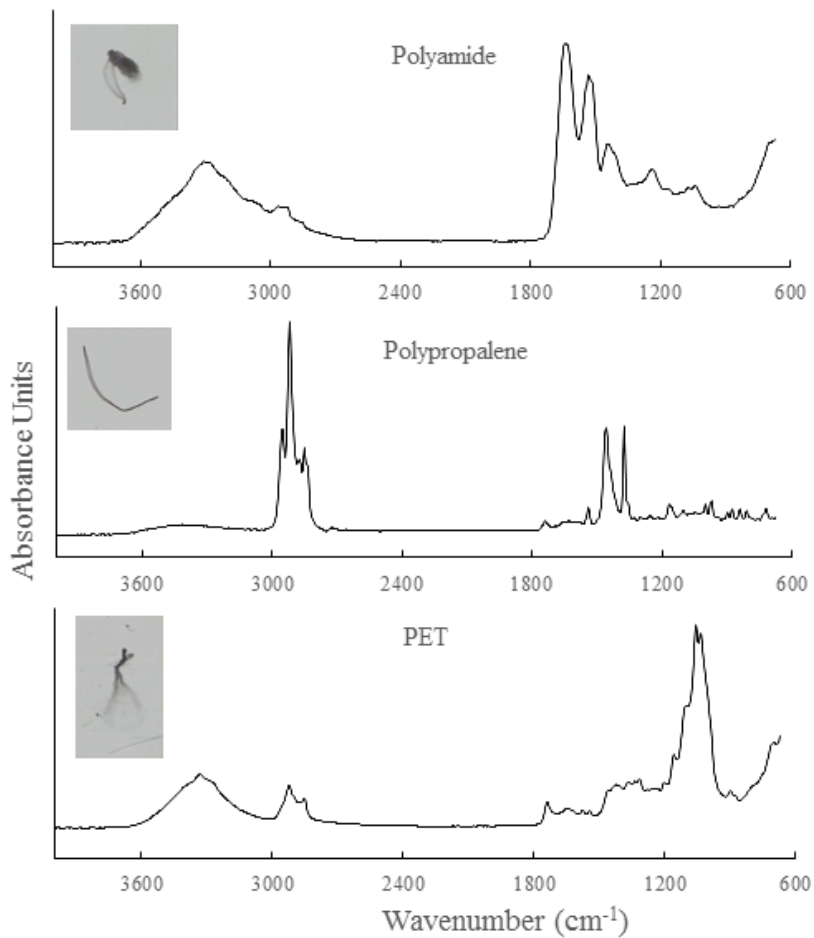
593

594

595

596

597



598

599 Figure 3. Fourier Transform Infrared Spectroscopy (FTIR) spectra within specimen
600 photographs of polyamide, polypropalene and Polyethylene terephthalate (PET).

601

602

603

604

605

606

607

608

609

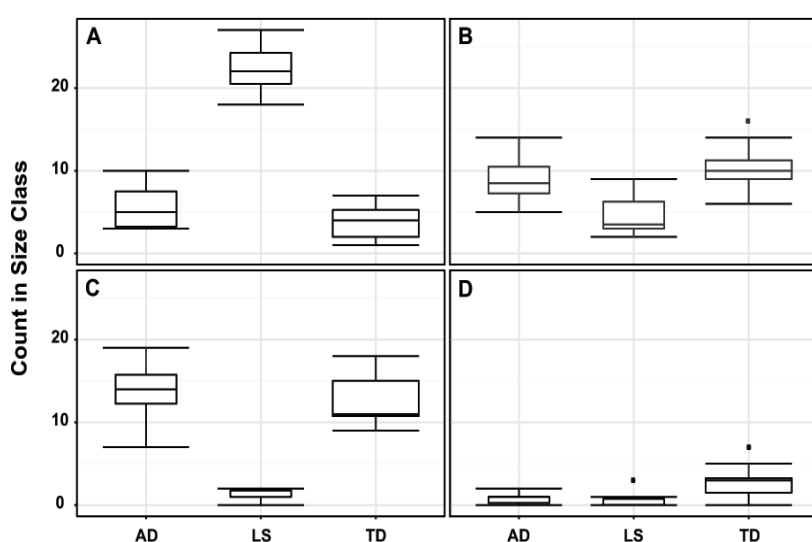
610

611

612

613

614



615

616

617

618

619

620

621

622

623

624

625

Figure 4. Abundance of microplastics in different size classes (**A**: 250-400 μm , **B**: 400-600 μm , **C**: 600-1000 μm , **D**: 1000-4000 μm) as a function of treatment type.

626

627

628

629

630

631

632

633

634

635

636

637

638

639

640

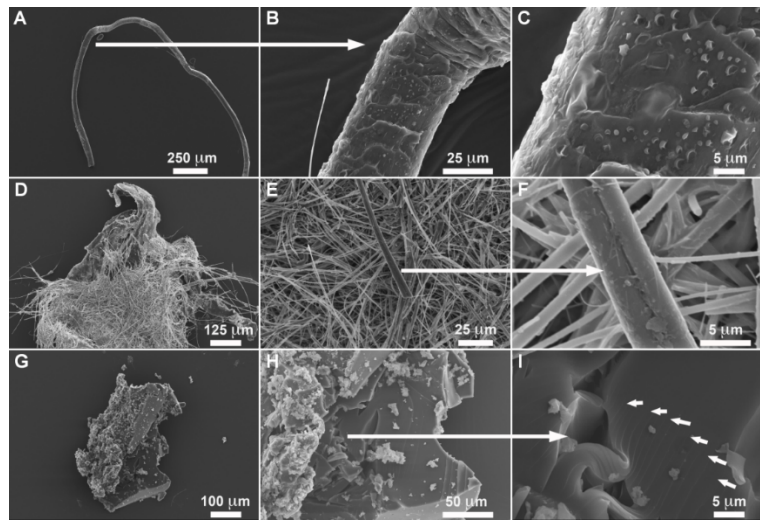
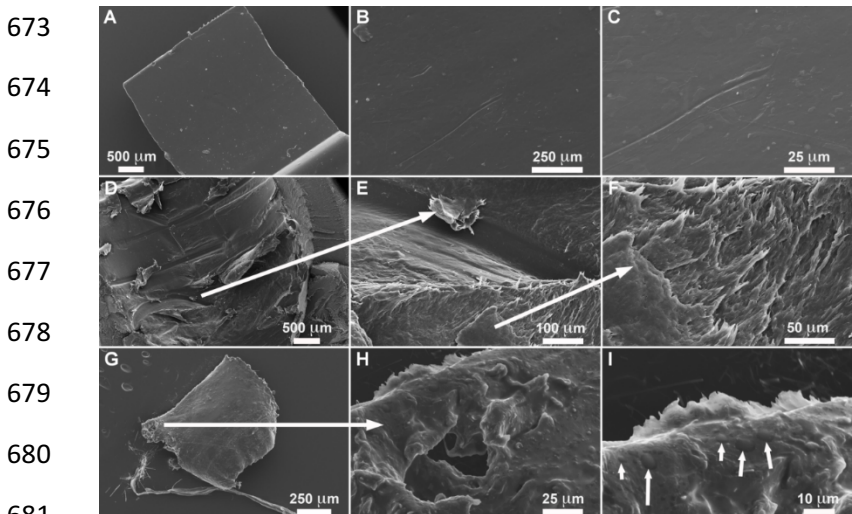


Figure 5. Diversity in morphology and surface texture of microplastics isolated from treated sewage sludge. Scanning electron micrographs of fibrous particle from thermally dried (TD) biosolids (A-C). Multi fibrous particle from lime stabilised (LS) biosolids (D-F). Overview of non-fibrous particle from anaerobically digested (AD) biosolids (G-H). Presence of lamellae or cleavage planes (arrow heads) on microplastic surface (I).

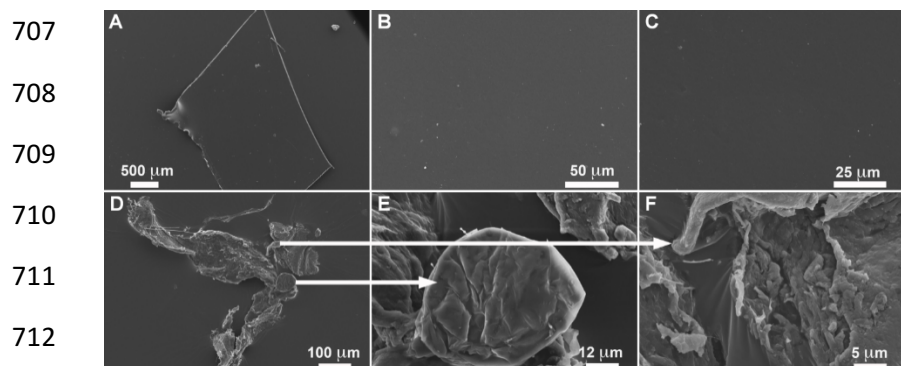
655
656
657
658
659
660
661
662
663
664
665
666
667
668
669
670
671
672



673
674
675
676
677
678
679
680
681
682 Figure 6. Morphological and surface texture comparison between pre-treatment high density
683 polyethylene (HDPE) and HDPE particles isolated from treated sewage sludge. Scanning
684 electron images of pre-treatment HDPE (A-C) showing smooth non-degraded surface.
685 Scanning electron micrographs of HDPE particle from lime stabilised (LS) biosolids (D-F)
686 showing altered and weathered surface texture. Scanning electron micrograph of HDPE
687 particle from thermally dried (TD) biosolids (G-I) with evidence of blistering effect (arrow
688 heads) on polymer surface (I).

689
690
691
692
693
694
695
696
697
698
699
700
701
702
703
704
705

706



714 Figure 7. Morphological and surface texture comparison between pre-treatment polyethylene
715 (PE) and PE particle isolated from sewage sludge. Scanning electron images of pre-treatment
716 PE (A-C) with unaltered surface. Scanning electron micrographs of PE particle from
717 thermally dried (TD) biosolids (D-F) showing wrinkling and fracturing of polymer surface.

718