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| 3 | Mobilisation kinetics of Br, Cd, Cr, Hg, Pb and Sb in |
| 4 | microplastics exposed to simulated, dietary-adapted |
| 5 | digestive conditions of seabirds |
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39 Abstract

| 40 | Samples of beached plastics and historical and contemporary consumer plastics |
|----|---|
| 41 | containing hazardous elements derived from reaction residues or functional additives |
| 42 | have been micronised and subject to extraction conditions representative of the |
| 43 | digestive environment of seabirds. Mobilisation of Br, Cd, Cr, Hg, Pb and Sb into |
| 44 | NaCl solution, an avian physiologically-based extraction test (PBET) and a dietary- |
| 45 | adapted PBET (DA-PBET) incorporating fish oil as part of the avian diet was |
| 46 | monitored by ICP-MS over a 168-h period. Kinetic data were subsequently fitted |
| 47 | using pseudo-first-order and parabolic diffusion models in order to derive rate |
| 48 | constants for the release of hazardous elements during avian digestion of |
| 49 | microplastics. Rate constants were variable and dependent on the nature and origin of |
| 50 | plastic, type of residue or additive, extractant solution employed and model applied. |
| 51 | Resulting estimates of bioaccessibility, defined as the equilibrium or maximum |
| 52 | concentration of an element mobilised over the time course relative to its total |
| 53 | concentration, were variable but considerable in many cases. Specifically, maximum |
| 54 | values of about 65% of Cd and 100% of Pb were observed in consumer |
| 55 | polycarbonate-acrylonitrile butadiene styrene exposed to the avian PBET and beached |
| 56 | polyurethane exposed to the DA-PBET, respectively. The potential health risks of |
| 57 | hazardous elements in microplastics are addressed and criteria for classification based |
| 58 | on the European Toy Safety Directive migration (mobilisation) limits are proposed. |
| 59 | |
| 60 | |
| 61 | Keywords: microplastics; additives; metals; mobilisation kinetics; avian PBET; |
| 62 | bioaccessibility |
| 62 | |

64 **1. Introduction**

65 Although the sources, distribution and physical impacts of microplastic litter have 66 received considerable attention over the past few decades (Ng et al., 2006; Hall et al., 67 2015; Lin et al., 2016), less well studied are the nature, occurrence, mobility and fate 68 of chemical residues and additives (Kwon et al., 2017; Luo et al., 2019). Residues 69 may remain as reactants or catalysts from the manufacturing process of certain 70 plastics or may be more widely distributed amongst plastics through the recycling and 71 blending of end-of-use materials (Turner, 2018a). Additives are deliberately 72 formulated into plastics, either physically or chemically, for specific functions that 73 include flame retardancy, colour, fastness, opacity, lubrication, strength, heat 74 resistance and stabilisation (Pritchard, 1997). While most contemporary additives are 75 regarded as safe, many historical additives are now restricted or inhibited on health 76 and environmental grounds. For example, the Restriction of Hazardous Substances 77 (RoHS) Directive provides limit values of Cd, Cr(VI), Hg, Pb and certain brominated 78 flame retardants in new or recycled electrical and electronic plastics (European 79 Parliament and Council, 2003; 2011), while the Toy Safety Directive specifies 80 migration limits for various metals and metalloids, including Cd, Cr(VI), Hg, Pb and 81 Sb, from plastic toys into a fluid mimicking a child's stomach (European Parliament 82 and Council of the EU, 2009). Despite these restrictions, however, potentially harmful 83 residues and additives remain in products in circulation and are particularly common 84 in marine litter where a heterogeneous assortment of plastics of variable sources and 85 ages are encountered (Massos and Turner, 2017; Shaw and Turner, 2019). 86

While most residues and additives are either designed, or at least considered, to
remain in the polymeric matrix, aging and weathering in the environment facilitates

89 their gradual mobilisation (Hansen et al., 2013; Nakashami et al., 2016). Significant in 90 this respect is the propensity of hazardous additives, such as those listed in the RoHS, 91 to be released into the digestive tract of animals that inadvertently or incidentally 92 ingest plastics because mobilisation may result in their entry into the systemic 93 circulation and subsequent accumulation. The release of small but significant 94 quantities of Cd and Sb (typically < 1% of corresponding total values) from various 95 micronised plastics into near-neutral surfactant- and protein-rich fluids simulating the 96 digestive conditions of deposit-feeding invertebrates over a six-hour time period has 97 recently been demonstrated by Martin and Turner (2019) and James and Turner 98 (2020). In an earlier study, Turner (2018b) showed greater release (ranging from <99 1% to > 20%) of various hazardous elements from polyolefins, polyvinyl chloride and 100 expanded plastics over a more extended timeframe into an acidic solution that mimics 101 the digestive chemistry of a seabird. Using a similar approach Tanaka et al. (2015) 102 demonstrated mobilisation of trace quantities of the polybrominated diphenyl ether 103 flame retardant, deca-BDE, impregnated in polyethylene. However, mobilisation was 104 increased 50-fold when oils that simulate the presence of food were added to the 105 acidified extractant.

106

In the present study, we hypothesize that the mobilisation of a variety of hazardous elements present in a wider variety of weathered and unweathered plastics are impacted by the presence of relatively hydrophobic dietary components in the avian digestive environment. Accordingly, we compare the kinetics of element mobilisation from micronised plastics (microplastics) in a standard physiologically-based extraction test (PBET) with those in a dietary-adapted-PBET (DA-PBET) in which fish oil is added. In the absence of any guidelines or limit values for environmental

- 114 plastics, measures of bioaccessibility arising from the experiments are related to
- 115 available migration limits as defined by the Toy Safety Directive in order to evaluate

116 the potential health implications of the different samples.

117

118 **2. Materials and methods**

119 2.1. Microplastic sample characteristics and preparation

120 Nine samples of plastic that had been archived from previous studies (e.g. Turner, 121 2018c; Turner and Solman, 2016) were selected for the present experiments. The 122 origin and characteristics of the samples, shown in Table 1, indicate five different 123 types of polymer, and five primary or secondary beached plastics that have been 124 exposed to the environment and four historical or contemporary consumer plastics 125 that have not undergone such exposure. Also shown in Table 1 are the concentrations 126 of elements that are hazardous according to the RoHS Directive and determined by X-127 ray fluorescence spectrometry; namely, Br (a proxy for brominated flame retardants), 128 Cr (as an upper bound indicator of Cr(VI)), Cd, Hg, and Pb. Although Sb is not in the 129 current iteration of the RoHS, it is also included because it is used as a flame retardant 130 synergist with brominated compounds (note its association with Br) and is regulated 131 by the Toy Safety Directive. Significantly, all samples are non-compliant or 132 potentially non-compliant (depending on the form of Br and speciation of Cr) with respect to the 1000 µg kg⁻¹ or 100 µg kg⁻¹ (Cd only) limit values specified by the 133 134 RoHS Directive (European Parliament and Council, 2011). 135 136 About 1 g of each plastic sample was formulated to "microplastic" of less than 1 mm

137 in at least two dimensions using a stainless steel grater. Microplastics were stored in

138 individual polyethylene specimen bags and in the dark pending use in the

139 experiments.

140

| 141 | Table 1: Characteristics of the microplastics used in the study. Polymer types were |
|-----|---|
| 142 | identified by Fourier-Transform infrared spectrometry and elemental concentrations |
| 143 | (in $\mu g g^{-1}$ and where $nd = not$ detected) were determined by X-ray fluorescence |
| 144 | spectrometry according to methods outlined elsewhere (Massos and Turner, 2017). |

| sample | type | description | Br | Cd | Cr | Hg | Pb | Sb |
|--------|---|--|--------|------|------|-----|--------|------|
| 1 | polypropylene | boot stud remover - black | nd | 766 | 1180 | nd | 9160 | 109 |
| 2 | polyethylene | child's shape sorter toy - yellow | nd | 6880 | 26.0 | nd | nd | nd |
| 3 | polyvinyl chloride | washing machine drainage hose - grey | 73.7 | nd | nd | nd | 22,900 | nd |
| 4 | polycarbonate + acrylonitrile butadiene styrene | jewellery beads - black, painted red | 15,100 | 34.3 | nd | nd | 123 | 8960 |
| 5 | polyethylene | water treatment medium (beached) - black | 3320 | nd | nd | nd | 22.4 | 1970 |
| 6 | glass-reinforced polyurethane | foam fragment (beached) - brown | 50.2 | nd | 116 | nd | 4860 | nd |
| 7 | polyethylene | unidentified fragment (beached) - green | 1260 | 35.9 | 4970 | nd | 121 | 674 |
| 8 | polyethylene | shot gun cartridge (beached) - red | 6.3 | 1780 | nd | nd | nd | nd |
| 9 | polyethylene | unidentified fragment (beached) - red | nd | 969 | 85.5 | 480 | 21.9 | nd |

146

145

147 2.2 Extraction reagents

Extractions consisted of a 0.1 M solution of NaCl, a simulated avian physiologically-148 149 based extraction test (PBET) and a dietary adapted PBET (DA-PBET). Sodium 150 chloride solution, simulating the pre-digestive conditions in the oesophagus and crop 151 of seabirds, was prepared by dissolving 5.844 g of Aristar NaCl (VWR Chemicals 152 BDH) in 1 L of high purity Elga LabWater (18.2 MΩ.cm resistivity). The standard 153 avian PBET was based on methods outlined elsewhere (Turner, 2018b) and modelled 154 on the chemistry of the proventriculus-gizzard of the northern fulmar, Fulmarus 155 glacialis, a procelliform known to ingest substantial quantities of microplastics 156 (Avery-Gomm et al., 2012) and an indicator species of plastic pollution according to 157 the Oslo and Paris Convention for the Protection of the Marine Environment of the 158 North-East Atlantic (OSPAR, 2008). Here, 10 g of pepsin (lyophilised powder from 159 porcine gastric mucosa; Sigma-Aldrich) were dissolved in 1 L of 0.1 M NaCl solution 160 whose pH was adjusted to 2.5 by the dropwise addition of 1 M HCl (prepared from

161 Fisher Scientific TraceMetal grade concentrated HCl). The DA-PBET was designed 162 to simulate digestive conditions that, additionally, include oils arising from the diet. We used oil from menhaden, fish of the genera Brevoortia and Ethmidium that are 163 consumed by a diverse range of predators and that represent an important food source 164 165 for many marine birds (Buchheister et al., 2017). Standard refined menhaden oil, 166 comprising ~ 20-35 % omega-3 fatty acids as triglycerides and of density 0.93 g ml⁻¹, 167 was purchased from Sigma-Aldrich. 168 169 2.3. Extraction procedure 170 Extractions were performed on 100 mg of each micronised sample in a series of

171 screw-capped polypropylene centrifuge tubes using 50 ml NaCl solution, 50 ml PBET

solution, and 40 ml PBET solution plus 10 ml menhaden oil (DA-PBET). The

173 contents of the tubes were incubated under continuous lateral agitation in a water bath

174 (Clifton, Nickel Electro Ltd, Weston-super-Mare, UK) set at 100 rpm and 40 °C. At

time intervals of approximately 0.5, 1, 3, 6, 24, 48, 96 and 168 h, 4 ml aliquots of

176 NaCl or PBET solution were pipetted from each tube, taking care not to abstract any

177 oil from the DAPBET, and filtered through 0.45 µm Whatman membrane filters

178 (Sigma-Aldrich) with the aid of a Terumo syringe. Filtrates were transferred to

179 individual Sterilin tubes to which 80 µl aliquots of 2% HNO₃ (prepared from Fisher

180 Scientific TraceMetal grade concentrated HNO₃) were added before the contents were

181 stored at room temperature and in the dark. Controls were performed likewise for

182 each extractant but in the absence of micronised microplastics.

183

In some acidified extracts from the PBET and DA-PBET a precipitate was observed
to form on storage. Here, extracts were centrifuged at 3000 rpm for 10 min using an

MSE Super Minor centrifuge (Heathfeld, UK), with 1 ml aliquots of supernatant
subsequently diluted to 4 ml 2% HNO₃ in new Sterilin tubes.

188

189 2.4. Extract analysis

190 Sample extracts were analysed in triplicate for Br, Cd, Cr, Hg, Pb and Sb by collision-

191 cell inductively coupled plasma-mass spectrometry (ICP-MS) using a Thermo

192 Scientific iCAP RQ ICP-MS (Thermo Elemental, Winsford, UK) with a concentric

193 glass nebuliser and conical spray chamber. Radio frequency power was 1550 W and

194 coolant, auxiliary, nebuliser and collision cell gas flow rates were set at 14 L Ar min⁻¹,

195 0.8 L Ar min⁻¹, 1.05 L Ar min⁻¹ and 5 ml He min⁻¹, respectively. Extracts were

analysed in triplicate, and data were obtained over a dwell time of 10 ms with 50

197 sweeps per reading. The instrument was calibrated using four mixed standards (in the

198 range 2 to 20 µg L⁻¹) and one blank prepared from LabKings and SCP Science

199 standard solutions in 0.1 M NaCl. For quality assurance purposes, a certified reference

200 drinking water (EP-L, SCP Science) was analysed in triplicate during each session

and a standard was analysed after every ten samples as a check for instrumental drift.

202 Limits of detection after normalisation to dry mass of microplastic ranged from about

203 0.003 μ g g⁻¹ for Cd, Cr and Pb in the DA-PBET to 0.44 μ g g⁻¹ for Br in NaCl solution

and precision (as relative standard deviation) among replicate analyses was usually
between 3 and 10%.

206

207 2.5. Timed data fitting

Data arising from the timed experiments were fitted with two diffusion models that are based on those outlined by Ruby et al. (1992). The linearised solution to the first model is as follows:

212
$$\ln(C_e - C) = \ln C_e - k_1 t + \ln C_0$$

| 213 | |
|-----|--|
| 214 | where C and C_e represent the elemental concentrations mobilised from the |
| 215 | microplastic on a weight basis at time t and at equilibrium, respectively, k_1 is a |
| 216 | combined, forward and reverse pseudo-first-order rate constant of units h^{-1} , and C_0 is |
| 217 | the weight-normalised concentration mobilised at $t = 0$. The latter is effectively a |
| 218 | constant that accounts for the very rapid, initial mobilisation that is often observed to |
| 219 | occur in such experiments but that cannot otherwise be modelled by a standard |
| 220 | diffusion-controlled reaction (Turner, 2018b). The value of k_1 was obtained from the |
| 221 | gradient arising from linear regression analysis of $\ln(C_e - C)$ - $\ln C_e$ versus <i>t</i> , using |
| 222 | Microsoft Excel Office 365, with the value of C_0 derived from the intercept of the |
| 223 | linear regression, a, as follows: |
| 224 | |
| 225 | $C_0 = C_{\rm e} \left(1 - e^a \right) \tag{1b}$ |
| 226 | |
| 227 | The linearised solution to the second, parabolic model of mobilisation is as follows: |
| 228 | |
| 229 | $C = k_2 t^{1/2} + C_0 \tag{2}$ |
| 230 | |
| 231 | where k_2 is a parabolic diffusion rate constant of units $\mu g [g(h)^{1/2}]^{-1}$ and, as above, C_0 |
| 232 | is the weight-normalised concentration mobilised at $t = 0$. Values of k_2 and C_0 were |
| 233 | obtained directly from the gradient and intercept, respectively, arising from linear |
| 234 | regression analysis of C versus $t^{1/2}$ using Microsoft Excel. |
| 235 | |
| | |

(1a)

237 **3. Results and Discussion**

238 3.1. Elemental mobilisation among the different microplastics and extractants 239 The concentrations of potentially hazardous elements detected in the microplastic 240 extracts on a mass basis and corrected for corresponding control concentrations, C (in $\mu g g^{-1}$), are shown as a function of time in Figures 1 to 6. Note that analytical error 241 242 bars were often smaller than the symbol size and are not shown for clarity. There is a 243 net increase in concentration for all elements (where detected) and all extractants over 244 the time course, and in most cases concentrations either increase continuously over 245 time or exhibit an initial, rapid increase followed by an approach to apparent 246 equilibrium. For a given element, however, differences in the precise timed profiles 247 are evident among the different samples and between the different extractant 248 solutions.

249

250 Mobilisation of Br (Figure 1) was detected in the three microplastics where the 251 element appeared to have been added (or recycled) as a constituent of a brominated 252 flame retardant. Amongst the extractants, mobilisation was greatest in the DA-PBET 253 for polyethylene (samples 5 and 7) but greatest in NaCl solution (i.e. without 254 acidification or digestive additives) for the mixed polycarbonate-acrylonitrile 255 butadiene styrene (sample 4). Cadmium mobilisation (Figure 2) was detected in two 256 microplastics (samples 2 and 8) where the metal had been added as the pigment 257 cadmium sulphide yellow, CdS, or cadmium sulphoselenide red, Cd₂SeS (confirmed 258 from the sample X-ray fluoresence spectra), and in two microplastics (samples 1 and 259 4) where it was present as a contaminant (likely through recycling); significantly, Cd 260 release was not detected in polyethylene (sample 9) where the metal had been added

| 261 | with Hg (presumably as the pigment cadmium mercury red, CdHgS ₂), although |
|-----|--|
| 262 | measureable mobilisation of Hg itself by the DA-PBET was evident (Figure 4). |
| 263 | Cadmium mobilisation was always greater in the PBET and DA-PBET than in NaCl |
| 264 | solution, with maximum release effected by the PBET and the DA-PBET in two cases |
| 265 | each. Chromium mobilisation (Figure 3) was detected from polypropylene (sample 1) |
| 266 | and polyurethane (sample 6) where the metal was likely present as a contaminant or |
| 267 | residual catalyst but not in polyethylene (sample 7) where it was present at the highest |
| 268 | total concentration as a pigment (and most likely chromium oxide green; Cr ₂ O ₃). In |
| 269 | both cases of detectable Cr mobilisation, the DA-PBET released considerably greater |
| 270 | quantities of the metal than the PBET and NaCl solution. |
| 271 | |
| 272 | Lead mobilisation was detected in four cases (Figure 5); in polypropylene (sample 1) |
| 273 | and polycarbonate-acrylonitrile butadiene styrene (sample 4), where the metal was |
| 274 | likely present as a contaminant, release was greatest by the PBET, while for polyvinyl |
| 275 | chloride (sample 3) and polyurethane (sample 6), where the metal was likely present |
| 276 | as part of a stabilising compound or catalytic residue, respectively, release was |
| 277 | greatest for the DA-PBET. Mobilisation of Sb (Figure 6) was detected in the |
| 278 | microplastics where it was co-associated with Br (samples 4, 5 and 7) and in the |
| 279 | microplastic where it was likely present as a contaminant through recycling (sample |
| 280 | 1). Mobilisation was greatest for the DA-PBET in all cases with the exception of |
| 281 | sample 4; here, the PBET mobilised the greatest quantity of the metalloid and the DA- |
| 282 | PBET mobilised the lowest concentration. |
| 283 | |
| | |

3.2. Kinetic modelling of timed data

| 285 | Rate constants and values of C_0 derived from the timed mobilisation data shown in |
|-----|--|
| 286 | Figures 1-6 are given in Table 2. The criterion for assigning a value of either k_1 or k_2 |
| 287 | was based on the shape of the timed profile (approach to equilibrium or a more |
| 288 | continuous increase in concentration with time, respectively) and the model regression |
| 289 | fit that yielded the greater significance. Where k_1 was assigned, the value of C_e , |
| 290 | defined as the concentration measured at the termination of the experiment, is also |
| 291 | shown. Note that any concentrations exceeding this value through the time course |
| 292 | were neglected in the derivation of the rate constant (i.e. $n < 7$ in the regression). |
| 293 | Where k_2 was assigned (and denoted with an asterisk in Table 2), the final |
| 294 | concentration measured in the time course is given but here is defined as the |
| 295 | maximum concentration, C_{max} (and $n = 8$ in the regression). Where neither model |
| 296 | returned a significant fit but mobilisation was detected, C_{max} is shown. |
| | |

Table 2: Constants defining the timed data show in Figures 1 to 6 for elemental
mobilisation from 9 samples of microplastics in NaCl solution, the PBET and the DAPBET. Note that ns = not significant, and values highlighted in yellow or orange
denote, respectively, non-compliance or potential non-compliance with respect to the
European Toy Safety Directive (European Parliament and Council of the EU, 2009).

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| 315 | | | |
| 316 | | | |
| 317 | | | |
| 318 | DA-PBET 0.0283 9.90 0.126 0.978 (7) | 8.50* 112* -0.465 0.998 (8) 35.2 12.0 0.989 (7) 7.83 7.83 0.691 (7) | |
| 319 | 5b PBET 4.10 0.578 0.578 0.938 (7) | 34.1* 35.0* 34.0* 112* 440* 112* 21.5 -0.465 0.987 (8) 0.998 (8) 1.08* 0.0312 14.7* 35.2 0.675 12.0 0.976 (8) 0.989 (7) 0.976 (8) 0.989 (7) 7.20 15.2 0.147 7.83 0.986 (7) 0.691 (7) | |
| 320 | Sb Sb Nacl PBET DA-PBET 0.0160 0.0138 0.0283 1.70 4.10 9.90 0.961 (7) 0.938 (7) | 16:9* 223* 22:9 960 (8) 13:0* 13:0* 996 (8) 996 (8) 0.189 0.189 0.189 | |
| 320 | | | |
| | Pb Pb Nacl PBET DA-PBET 0.0201 0.0141 0.0116 230 445 295 98.2 204 37.5 0.911(7) 0.890 (7) 0.982 (7) ns 0.452 0.0251 ns 0.452 0.0251 99.1 1150 1900 711 660 0.992 (4) 0.900 (7) | ns 0.0199 0.0252 0.60 <u>3.2.5</u> 18.0 10.9 5.97 0.969 (7) 0.886 (7) 0.233 0.228 0.0224 2000 2700 5130 1340 1760 4350 0.793 (7) 0.985 (5) 0.691 (7) | |
| 322 | Pb PBET 0.0141 445 204 0.890 (7) 0.890 (7) 0.892 1150 711 0.992 (4) | 0.0199 10.9 0.969 (7) 0.969 (7) 1.760 0.985 (5) | |
| 323 | NaCl 0.0201 98.2 98.2 0.911 (7) ns 19.1 | ns 0.60 0.0233 2000 1340 0.793 (7) | |
| 324 | | • • • • • • • • • • • • • • • • • • • | 0* 54 (8) |
| 325 | DA-PBET DA-PBET | | 0.140* 2.30* 0.364 0.949 (8) |
| 326 | DA-PBET 0.0487 5.9 5.92 0.968 (7) | 0.472 6.20 5.72 0.844 (3) | |
| 327 | Cr PBET 1 0.0094 7.32 7.71 (7) | n 1.14 | |
| | Cr Cr Nacl PBET DA-PBET 0.0140 0.0094 0.0487 11.4 13.9 25.9 5.58 7.32 5.92 0.746 (7) 0.771 (7) 0.968 (7) | 0.0207 0.500 0.409 0.675 (7) | |
| | | | |
| | DA-PBET 0.857* 10.8* -0.237 0.970 (8) 5.42* 1.88 1.88 0.963 (8) | 0289 0.0193 18.4 18.4 18.4 18.4 18.4 0.983 (7) 0.983 (7) | 0.383* 5.34* 0.0308 0.978 (8) |
| | Cd <u>BET</u> 400* .26* .57 (8) .500 .57 (8) .57 (8) .57 (8) .7.0* .459 .91 (8) | 0289 22.2 64 (7) | 804* 2.1* 0.224 73 (8) |

Rate constants arising from model 1 range from 0.0094 h⁻¹ for Cr in sample 1 exposed

to the PBET to 0.472 h⁻¹ for Cr in sample 6 exposed to the DA-PBET. Values of C_0

344 were usually positive and in many cases exceeded 50% of $C_{\rm e}$ (e.g. Pb in sample 6),

345 suggesting significant instantaneous mobilisation into the extractants. Rate constants

arising from model 2 range from 0.119 μ g [g(h)^{1/2}]⁻¹ for Cd in sample 1 exposed to

347 NaCl solution to 34.1 μ g [g(h)^{1/2}]⁻¹ for Sb in sample 4 exposed to the PBET, and here

348 values of C_0 were always relatively close to the origin.

3.3. Elemental bioaccessibilities

351 Table 3 provides operational measures of elemental avian bioaccessibility, BA (%), or

the percentage of total element that is available in the pre-digestive environment of

353 the oesophagus and crop (NaCl), the digestive environment of the proventriculus and 354 gizzard in the absence of food (PBET) and the digestive environment in the presence 355 of food (DA-PBET) over a timeframe of 168 h. (Note that non-food material may be 356 trapped in the proventriculus-gizzard of many seabirds for weeks to months; Avery-357 Gomm et al., 2012.) Values of BA are calculated from the equilibrium or maximum 358 (i.e., final) concentrations of elements mobilised over the time courses and reported in 359 Table 2 relative to corresponding total elemental concentrations determined by X-ray 360 fluorescence spectrometry and given in Table 1. Where an element was detected by 361 X-ray fluorescence but not detected by an extractant an upper limit is given based on 362 the ICP-MS detection limit in the relevant medium.

363

Table 3: Percentage bioaccessibilities of hazardous elements in the different

365 extractants tested and calculated from $C_{\rm e}$ or $C_{\rm max}$ relative to total concentration.

366

| | | Br | | | Cd | | | Cr | | Hg | | Pb | | | Sb | |
|--------|--------|--------|---------|--------|--------|---------|--------|--------|---------|---------|--------|--------|---------|-------|-------|---------|
| sample | NaCl | PBET | DA-PBET | NaCl | PBET | DA-PBET | NaCl | PBET | DA-PBET | DA-PBET | NaCl | PBET | DA-PBET | NaCl | PBET | DA-PBET |
| 1 | | | | 0.226 | 0.687 | 1.41 | 0.473 | 1.18 | 2.19 | | 2.51 | 4.86 | 3.22 | 1.56 | 3.76 | 9.08 |
| 2 | | | | 0.346 | 0.683 | 0.953 | | | | | | | | | | |
| 3 | <0.600 | <0.039 | <0.004 | | | | | | | | 0.430 | 5.25 | 8.67 | | | |
| 4 | 4.90 | 0.038 | 0.094 | 16.9 | 64.7 | 53.6 | | | | | 0.488 | 26.4 | 14.6 | 2.49 | 4.91 | 1.25 |
| 5 | 0.274 | 1.33 | 1.61 | | | | | | | | | | | 0.660 | 0.746 | 1.8 |
| 6 | <0.880 | <0.058 | <0.006 | | | | 0.431 | 0.980 | 5.34 | | 41 | 56 | 106 | | | |
| 7 | 0.095 | 0.349 | 2.14 | <0.096 | <0.014 | <0.009 | <0.001 | <0.001 | <0.001 | | <0.004 | <0.010 | <0.002 | 0.964 | 1.07 | 2.26 |
| 8 | | | | 0.183 | 0.680 | 0.300 | | | | | | | | | | |
| 9 | | | | <0.001 | <0.001 | <0.001 | <0.004 | <0.004 | <0.004 | 0.048 | | | | | | |

For a given element, values of BA vary considerably amongst the microplastics and extractants. For Br, BA ranges from < 1% for samples 3 and 6 in all extractants to about 5% for sample 4 exposed to NaCl. For Cd, BA is < 1 % in most cases with the exception of sample 4 where values exceed 50% in the PBET and DA-PBET; in all samples where the metal was detected in the extractants, BA was greater in the PBET

373 or DA-PBET than in the near-neutral NaCl solution. For Cr, BA ranges from <

0.005% in samples 7 and 9 to > 2% for samples 1 and 6 exposed to the DA-PBET.

375 Values of BA for Pb range from < 0.01% in sample 7 to about 100% in sample 6

376 exposed to the DA-PBET, and in all cases BA was greater in the PBET or DA-PBET

than in NaCl solution. Regarding Sb, BA ranges from 0.66% in sample 5 exposed to

378 NaCl to over 9% for sample 1 exposed to the DA-PBET, and in all microplastics the

379 maximum value of BA results from exposure to the PBET or DA-PBET.

380

381 *3.4. Mechanisms of element mobilisation*

382 Hazardous elements may be incorporated into the plastic as ions, complexes or

383 compounds, or bound irreversibly to the polymeric backbone (Town et al., 2018). In

addition, at least for the beached plastics, there may be a small amount of element

adsorbed to the surface from the marine environment (Holmes et al., 2012).

386 Neglecting desorption of environmentally acquired elements, the fundamental

387 mechanism of element mobilisation from the microplastics in the present study is

388 diffusion from the plastic matrix into a saline (NaCl) aqueous medium. Free ions and

389 small complexes may diffuse through the particle matrix whereas larger complexes or

those bound irreversibly are immobile, with the permeability (or crystallinity) of the

391 polymer determining the size limit of diffusible complexes. Presumably, therefore, the

392 rapid, instantaneous mobilisation that we observe arises from the release of elements

393 that are located at (but incorporated into) the particle surface and not required to

diffuse through the plastic matrix.

395

396 The rate of mobilisation of elements from the microplastics may be facilitated by

397 altering the composition of the aqueous medium, or, more specifically, making

398 conditions more favourable for the formation of free ions or small complexes and 399 molecules. To this end, the use of an acidic digestive medium is predicted to increase 400 the concentration of metal ions relative to a near-neutral solution. Mobilisation may 401 also be facilitated if a medium is introduced that interacts with the plastic matrix by, 402 for example, exposing a greater surface area to the aqueous phase through polymer 403 chain loosening (Sun et al., 2019). Accordingly, it is possible that the hydrophobic 404 fish oil is able to partially modify the integrity of plastic structure or even act as a 405 solvent for the extraction of relatively hydrophobic organic compounds of bromine 406 (Tanaka et al., 2015).

407

408 The observations in the present study are partly consistent with the assertions above in 409 that mobilisation and bioaccessibility of the metals (Cd, Cr and Pb) are enhanced 410 under the acidic conditions of the PBET and DA-PBET relative to unacidified NaCl 411 solution, with the presence of fish oil usually enhancing but sometimes inhibiting 412 metal release. Acidified conditions also promote the mobilisation of Br and Sb in 413 most cases, but NaCl releases greater quantities of both elements than the PBET 414 and/or the DA-PBET from sample 4. The polymeric composition of this sample 415 (acrylonitrile butadiene styrene-polycarbonate) suggests a rather amorphous structure 416 of relatively low permeability and high thermal stability, at least compared with 417 expanded polyurethane, polyvinyl chloride and the polyolefins (Keller, 2017). 418 Anomalous mobilisation results for sample 4 in the presence of fish oil (for Br and Sb 419 as well as Cd and Pb) may, therefore, reflect the poor penetrability of the relatively 420 large triglyceride molecules into the plastic and, possibly, a propensity to block the 421 migration of other solutes into and out of the matrix.

422

423 3.5. Implications for exposure to seabirds and setting safety guidelines

424 Despite kinetic modelling of the mobilisation of potentially hazardous elements from 425 microplastics under simulated digestive conditions being relatively straightforward, 426 the extent of mobilisation exhibits a complex dependence on the type of plastic, the 427 nature of the additive or reaction residues and the composition of the extractant 428 solution. Mobilisation is, however, considerable in many cases, with a reduction in pH 429 facilitating elemental release for metals and the presence of food oil having a more 430 variable but usually positive effect on the dissolution of both inorganic and 431 brominated compounds. For more meaningful upper estimates of chemical 432 bioaccessibility in microplastics and their risks to fish-consuming seabirds, it is 433 recommended that both a standard avian PBET and a DA-PBET be employed and that 434 the higher result be adopted.

435

436 Currently, there exist no chemical standards for waste environmental plastics that are 437 based on health grounds. The RoHS Directive limits have recently been employed as 438 a screening criterion for compliance of primary and secondary beached plastics (Shaw 439 and Turner, 2019) but the total content of a potentially hazardous chemical does not 440 address its potential for migration or its bioaccessibility to an animal. Accordingly, we 441 propose that the Toy Safety Directive limits on the migration of metals and metalloids 442 (but not brominated compounds) are of most relevance when evaluating the potential 443 health impacts of marine plastic ingestion to mammals. Here, limit values are based 444 on the potential impacts arising from the ingestion of 8 mg of material per day and 445 migration into 0.07 M HCl for 2 h at 37°C (BSI, 1994; European Commission, 2016). 446 Current or proposed migration limits for material that can be "scraped off" toys, including plastics, by biting and sucking are 17 μ g g⁻¹ for Cd, 0.2 μ g g⁻¹ for Cr(VI), 447

94 μ g g⁻¹ for Hg, 23 μ g g⁻¹ for Pb and 560 μ g g⁻¹ for Sb (European Parliament and 448 449 Council of the EU, 2009). Based on these limits, and for an equivalent weekly (168 h) intake of 56 mg of microplastic by a seabird, five of the current samples that include 450 451 beached plastics and both historical and contemporary consumer products would be 452 non-compliant with respect to at least one element and at least one extractant; non-453 compliance occurs for Pb in four cases, Cd in two cases, and Cr in one case (sample 1) if it is assumed that a Pb-Cr association is indicative of lead chromate and the 454 455 higher oxidation state of the metal.

456

457 Clearly, the overall risk to a seabird is more complex as plastic ingestion varies 458 greatly between and among species (Lavers and Bond, 2016; Roman et al., 2019) and 459 the quantities of plastic ingested that contain hazardous elements will depend on 460 availability in the water column, foraging ecology, and any selectivity based on, for 461 example, colour (Kain et al., 2016; Tavares et al., 2017). Nevertheless, microplastics 462 should not be overlooked as a source of harmful additives and residues to animals that 463 digest material for extended periods of time under acidic and oily conditions.

464

465 **4. Conclusions**

Significant quantities of hazardous elements (Br, Cd, Cr, Hg, Pb, Sb) associated with residues and historical additives in plastics can be mobilised under simulated gastric conditions representative of seabirds, and in particular where the diet is considered as part of the digestive chemistry. Although the kinetics of mobilisation can be modelled by simple diffusion equations, the magnitude of the constants associated with these models appear to exhibit a complex dependence on the nature of the additive and the type and condition of the plastic. It is suggested that limits of chemical migration (or

| 473 | mobilisation) stipulated by the European Toy Safety Directive afford a means of |
|-----|---|
| 474 | evaluating the potential risks of microplastics to mammals in the marine environment. |
| 475 | On this basis, five out of nine plastics tested in the present study returned non- |
| 476 | compliant concentrations for at least one element and one digestive fluid simulant. |
| 477 | |
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| 482 | |
| 483 | References |
| 484 | Avery-Gomm, S., O'Hara, P.D., Kleine, L., Bowes, V., Wilson, L.K., Barry, K.L., |
| 485 | 2012. Northern fulmars as biological monitors of trends of plastic pollution in the |
| 486 | eastern North Pacific. Marine Pollution Bulletin 64, 1776-1781. |
| 487 | |
| 488 | BSI, 1994. Safety of Toys - part 3: Migration of Certain Elements. British |
| 489 | Standard BS EN 71-3, London. |
| 490 | |
| 491 | Buchheister, A., Miller, T.J., and Houde, E. 2017. Evaluating Ecosystem-based |
| 492 | reference points for Atlantic menhaden. Marine and Coastal Fisheries Dynamics |
| 493 | Management and Ecosystem Science 9, 457-478. |
| 494 | |
| 495 | Claessens, M., De Meester, S., Van Landuyt, L., De Clerk, K., Janssen, C.R., 2011. |
| 496 | Occurrence and distribution of microplastics in marine sediments along the Belgian |
| 497 | coast. Marine Pollution Bulletin 62, 2199-2204. |

| 498 |
|-----|
|-----|

| 499 | European Commission, 2016. Toy Safety Directive 2009/48/EC. An explanatory |
|-----|--|
| 500 | guidance document (Revision 1.9 CORR). Europese Commissie, Brussel. |
| 501 | |
| 502 | European Parliament and Council, 2003. Directive 2002/95/EC on the restriction of |
| 503 | the use of certain hazardous substances in electrical and electronic equipment. Official |
| 504 | Journal of the European Union L37/19. |
| 505 | |
| 506 | European Parliament and Council, 2011. Directive 2011/65/EU on the restriction of |
| 507 | the use of certain hazardous substances in electrical and electronic equipment (recast). |
| 508 | Official Journal of the European Union L174/88. |
| 509 | |
| 510 | European Parliament and Council of the EU, 2009. Directive 2009/48/EC of the |
| 511 | European Parliament and of the Council of 18 June 2009 on the safety of toys. |
| 512 | Official Journal of the European Union L170/1. |
| 513 | |
| 514 | European Parliament and the Council of the European Union, 2009. Directive |
| 515 | 2009/48/EC on the safety of toys. Official Journal of the European Union L170, 1. |
| 516 | |
| 517 | Hall, N.M., Berry, K.L.E., Rintoul, L., Hoogenboom, M.O., 2015. Microplastic |
| 518 | ingestion by scleractinian corals. Marine Biology 162, 725-732. |
| 519 | |
| 520 | Hansen, E., Nilsson, N.H., Lithner, D., Lassen, C., 2013. Hazardous Substances in |
| 521 | Plastic Materials. Oslo: COWI and the Danish Technological Institute on behalf of |
| 522 | The Norwegian Climate and Pollution Agency, 150 pp. |

| 524 | Holmes, L., Turner, A., Thompson, R.C., 2012. Adsorption of trace metals to plastic |
|-----|--|
| 525 | resin pellets in the marine environment. Environmental Pollution 160, 42-48. |
| 526 | |
| 527 | James, E., Turner, A., 2020. Mobilisation of antimony from microplastics added to |
| 528 | coastal sediment. Chemosphere (submitted). |
| 529 | |
| 530 | Kain, E.C., Lavers, J.L., Berg, C.J., Raine, A.F., Bond, A.L., 2016. Plastic ingestion |
| 531 | by Newell's (Puffinus newelli) and wedge-tailed shearwaters (Ardenna pacifica) in |
| 532 | Hawaii. Environmental Science and Pollution Research 23, 23951-23958. |
| 533 | |
| 534 | Kwon, J.H., Chang, S., Hong, S.H., Shim, W.J., 2017. Microplastics as a vector of |
| 535 | hydrophobic contaminants: Importance of hydrophobic additives. Integrated |
| 536 | Environmental Assessment and Management 13, 494-499. |
| 537 | |
| 538 | Lavers, J.L., Bond, A.L., 2016. Ingested plastic as a route for trace metals in Laysan |
| 539 | Albatross (Phoebastria immutabilis) and Bonin Petrel (Pterodroma hypoleuca) from |
| 540 | Midway Atoll. Marine Pollution Bulletin 110, 493-500. |
| 541 | |
| 542 | Lin, V.S., 2016. Research highlights: impacts of microplastics on plankton. |
| 543 | Environmental Science – Processes and Impacts 18, 160-163. |
| 544 | |
| 545 | Luo, H.W., Xiang, Y.H., He, D.Q., Li, Y., Zhao, Y.Y., Wang, S., Pan, X.L., 2019. |
| 546 | Leaching behavior of fluorescent additives from microplastics and the toxicity of |
| 547 | leachate to Chlorella vulgaris. Science of the Total Environment 678, 1-9. |
| | |

| 549 | Martin, K., Turner, A., 2019. Mobilization and bioaccessibility of cadmium in coastal |
|-----|---|
| 550 | sediment contaminated by microplastics. Marine Pollution Bulletin 146, 940-944. |
| 551 | |
| 552 | Massos, A., Turner, A., 2017. Cadmium, lead and bromine in beached microplastics. |
| 553 | Environmental Pollution 227, 139-145. |
| 554 | |
| 555 | Nakashima, E., Isobe, A., Kako, S., Itai, T., Takahashi, S., Guo, X., 2016. The |
| 556 | potential of oceanic transport and onshore leaching of additive-derived lead by marine |
| 557 | macro-plastic debris. Marine Pollution Bulletin 107, 333-339. |
| 558 | |
| 559 | Ng, K.L., Obbard, J.P., 2006. Prevalence of microplastics in Singapore's coastal |
| 560 | marine environment. Marne Pollution Bulletin 52, 761-767. |
| 561 | |
| 562 | OSPAR, 2008. Background document for the EcoQO on plastic particles in stomachs |
| 563 | of seabirds. London, 355/2007. |
| 564 | |
| 565 | Pritchard, G., 1997. Plastic Additives: An A-Z Reference. Chapman and Hall, |
| 566 | London, 633pp. |
| 567 | |
| 568 | Roman, L., Bell, E., Wilcox, C., Hardesty, B.D., Hindell, M., 2019. Ecological drivers |
| 569 | of marine debris ingestion in procellariiform seabirds. Scientific Reports 9, article no. |
| 570 | 916. DOI: 10.1038/s41598-018-37324-w |
| 571 | |

| 572 | Ruby, M.V. | Davis, A. | Kempton, J.H | I., Drexler, J | .W., Berg | gstrom, P.D., | 1992. Lead |
|-----|------------|---------------|-------------------------------|---------------------------------------|---|---------------|------------|
| 512 | 1.40 , 1.1 | , Duvis, 11., | i so mptom, s . | 1. , D 10/101, 5 | \cdots , $\mathbf{D}\mathbf{C}\mathbf{I}$ | 50tronn, r, | 1772. Lou |

573 bioavailability: dissolution kinetics under simulated gastric conditions. Environmental

574 Science and Technology 26, 1242-1248.

575

Shaw, E.J., Turner, A., 2019. Recycled electronic plastic and marine litter. Science of
the Total Environment 694, 133644.

578

- 579 Sun, B., Hu, Y., Cheng, H., Tao, S., 2019. Releases of brominated flame retardants
- 580 (BFRs) from microplastics in aqueous medium: Kinetics and molecular-size
- 581 dependence of diffusion. Water Research 151, 215-225.

582

- 583 Tanaka, K., Takada, H., Yamashita, R., Mizukawa, K., Fukuwaka, M.A., Watanuki,
- 584 Y., 2015. Facilitated leaching of additive-derived PBDEs from plastic by
- 585 seabirds' stomach oil and accumulation in tissues. Environmental Science and
- 586 Technology 49, 11799-11807.
- 587
- 588 Tavares, D.C., de Moura, J.F., Merico, A., Siciliano, S., 2017. Incidence of marine

debris in seabirds feeding at different water depths. Marine Pollution Bulletin 119, 68-

- 590 73.
- 591
- 592 Town, R.M., van Leeuwen, H.P., Blust, R., 2018. Biochemodynamic features of metal
- 593 ions bound by micro- and nano-plastics in aquatic media. Frontiers in Chemistry
- 594 https://doi.org/10.3389/fchem.2018.00627

595

| 596 | Turner, A., 2018a. Black plastics: linear and circular economies, hazardous additives |
|-----|---|
| 597 | and marine pollution. Environment International 117, 308-318. |

| 599 | Turner, A | ., 2018b. | Mobilisation | kinetics | of hazardous | elements in | n marine | plastics |
|-----|-----------|-----------|--------------|----------|--------------|-------------|----------|----------|
|-----|-----------|-----------|--------------|----------|--------------|-------------|----------|----------|

- 600 subject to an avian physiologically-based extraction test. Environmental Pollution
- 601 236, 1020-1026.

602

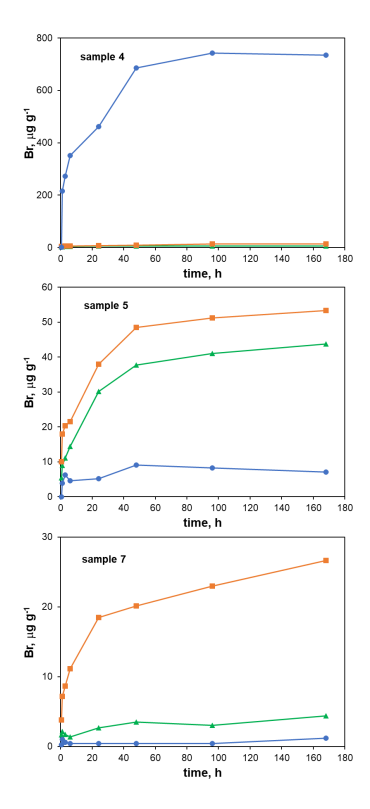
- 603 Turner, 2018c. Concentrations and migratabilities of hazardous elements in second-
- hand children's plastic toys. Environmental Science and Technology 52, 3110-3116.

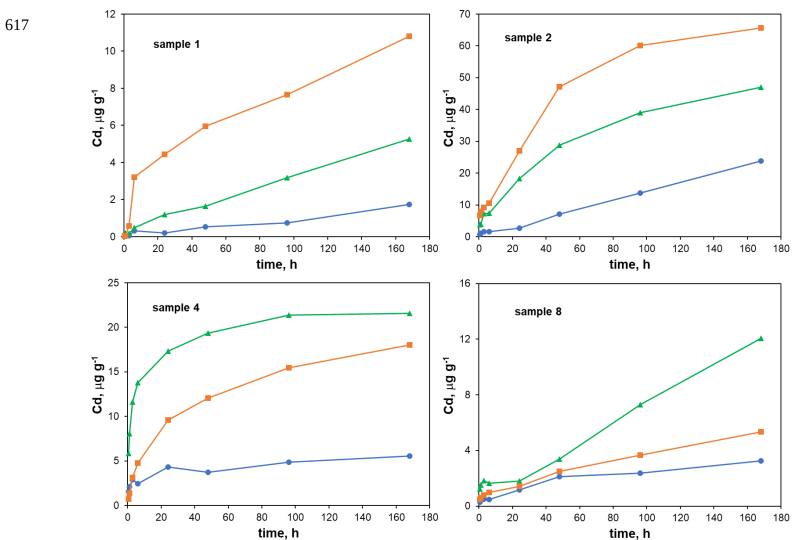
605

- Turner, A., Solman, K.R., 2016. Analysis of the elemental composition of marine
- 607 litter by field-portable-XRF. Talanta 159, 262-271.

608

- 610 Figure 1: Mobilisation of Br from micronised plastic samples as a function of time in
- 611 NaCl solution (blue circles), the PBET (green triangles) and DA-PBET (orange
- 612 squares).
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and DA-PBET (orange squares). 616

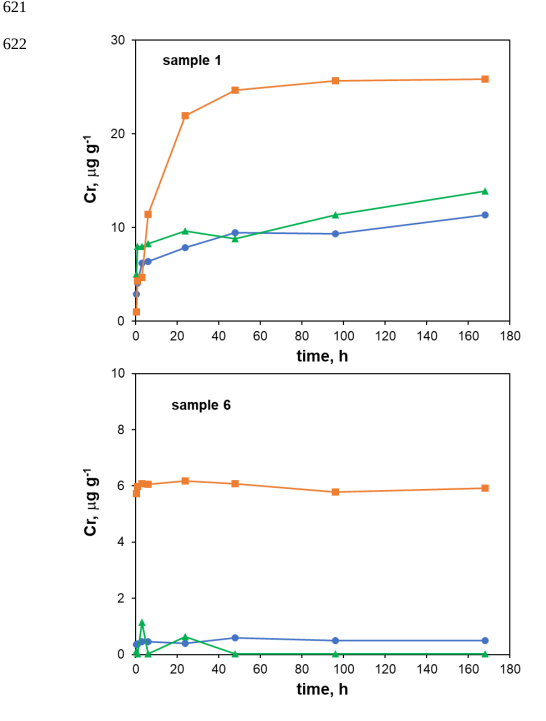
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Figure 2: Mobilisation of Cd from micronised plastic samples as a function of time in NaCl solution (blue circles), the PBET (green triangles)

Figure 3: Mobilisation of Cr from micronised plastic samples as a function of time in

NaCl solution (blue circles), the PBET (green triangles) and DA-PBET (orange

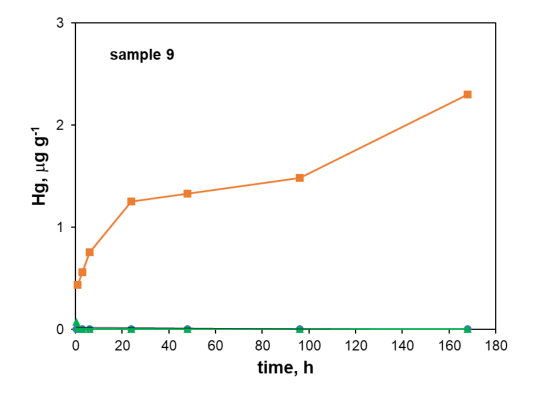
- squares).



623 Figure 4: Mobilisation of Hg from a micronised plastic sample as a function of time in

624 NaCl solution (blue circles), the PBET (green triangles) and DA-PBET (orange

625 squares).



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Figure 5: Mobilisation of Pb from micronised plastic samples as a function of time in NaCl solution (blue circles), the PBET (green triangles)
and DA-PBET (orange squares).

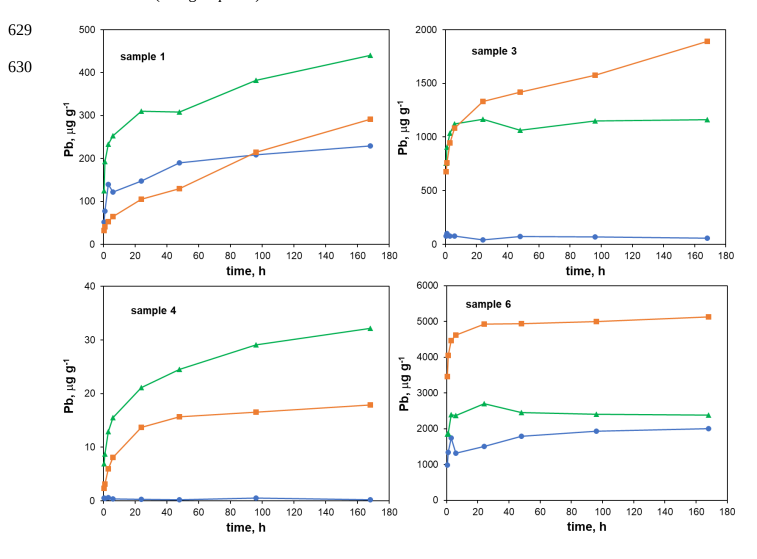
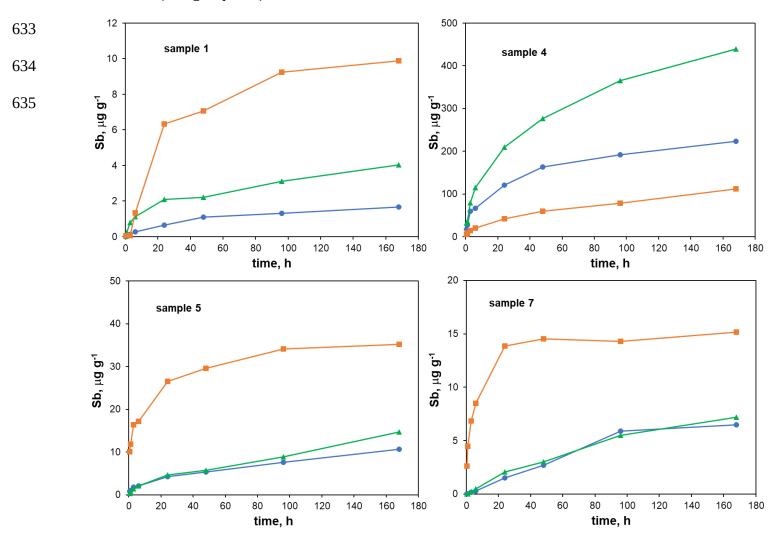


Figure 6: Mobilisation of Sb from micronised plastic samples as a function of time in NaCl solution (blue circles), the PBET (green triangles)



632 and DA-PBET (orange squares).