



PEARL

**Polystyrene foam as a source and sink of chemicals in the marine environment:
An XRF study**

Turner, Andrew

Published in:
Chemosphere

DOI:
[10.1016/j.chemosphere.2020.128087](https://doi.org/10.1016/j.chemosphere.2020.128087)

Publication date:
2021

Link:
[Link to publication in PEARL](#)

Citation for published version (APA):

Turner, A. (2021). Polystyrene foam as a source and sink of chemicals in the marine environment: An XRF study. *Chemosphere*, 263(0), 128087-128087. <https://doi.org/10.1016/j.chemosphere.2020.128087>

All content in PEARL is protected by copyright law. Author manuscripts are made available in accordance with publisher policies. Wherever possible please cite the published version using the details provided on the item record or document. In the absence of an open licence (e.g. Creative Commons), permissions for further reuse of content should be sought from the publisher or author.

14 **Abstract**

15 Polystyrene foam (expanded and extruded polystyrene: EPS and XPS, respectively) is a ubiquitous
16 and pervasive type of marine plastic litter whose physical properties, transport and fate are distinctly
17 different to those of other common (unfoamed) types of thermoplastic litter. In this study, a range
18 of fragments of EPS and XPS retrieved from three beaches in southwest England have been
19 characterised by energy-dispersive X-ray fluorescence spectrometry in order to examine the
20 chemical makeup and potential biological and geochemical impacts and interactions of this type of
21 plastic waste. Analyses performed through sample faces and, in some cases and after dissection,
22 through the material core, revealed variable concentrations of Fe, Ti and Zn among the fragments
23 and, in some cases, within the same sample. This likely reflects the presence of reaction residues and
24 pigments arising from the manufacture of polystyrene, and, for Fe and Ti, significant and
25 heterogeneous ion and mineral acquisition from the environment during transport in suspension or
26 while beached. Acquired oxides of Fe are partly responsible for the chemical fouling observed on the
27 face of most samples and are also able to act as an adsorbent for other metals, like Pb. Detection of
28 Br in many fragments up to concentrations of 11,500 mg kg⁻¹ likely results from the incorporation of
29 the flame retardant, hexabromocyclododecane, in EPS and XPS designed for (but not necessarily
30 limited to) the construction sector. These observations suggest that EPS and XPS can act as both a
31 source and sink for contaminants in the marine environment that merit further investigation.

32 **Keywords:** polystyrene foam; expanded; extruded; XRF; flame retardants; additives

33

34

35 **1. Introduction**

36 Because of its low weight, high strength and excellent thermal-noise insulation and shock-absorbing
37 properties, foamed polystyrene, encompassing expanded polystyrene (EPS) and extruded
38 polystyrene (XPS), is widely used across a range of sectors (e.g., construction, transportation,
39 consumer, maritime) (Wünsch, 2000; Block et al., 2017). However, many of the characteristics that
40 make foamed polystyrene such a popular and useful commodity also ensure that it is a significant
41 and pervasive constituent of the waste stream (Nie et al., 2015; Black et al., 2019).

42 In the marine environment, EPS and XPS are among of the most abundant forms of plastic litter
43 (Hinojosa and Thiel, 2009; Davis and Murphy, 2015; Che et al., 2018; Chitaka and von Blottnitz,
44 2018). Sources of polystyrene foam include municipal waste, littering and loss or deterioration of
45 polystyrene-based structures at sea (such as pontoons, floating docks, buoys and boat stands;
46 Hansen et al., 2015; Jang et al., 2020). Unlike other common (unfoamed) thermoplastics, however, a
47 density an order of magnitude lower than that of seawater, coupled with a closed cell structure,
48 results in EPS and XPS waste readily fragmenting in situ and being transported on the ocean surface.
49 Thus, at sea polystyrene foam is subject to considerable windage (Chubarenko et al., 2016),
50 persistent exposure to chemicals resident in the sea surface microlayer (Wurl and Obbard, 2004) and
51 colonisation by rafting organisms (Carson et al., 2013), and when beached material is widely
52 dispersed, abraded, micronised and buried by the action of the wind (Song et al., 2017).

53 Despite its distinctive physical and mechanical properties, pathways and environmental and biotic
54 interactions, however, foamed polystyrene is generally considered or quantified within the more
55 general pool of marine plastic litter in the scientific literature (Fok et al., 2017; Nabizadeh et al.,
56 2019; Tata et al., 2020). Significant from a geochemical and biological-toxicological perspective and
57 that has yet to be documented is its (non-polymeric) chemical composition, including the presence
58 of any additives and residues arising from manufacture and chemicals acquired from the
59 environment during suspension or beaching. Accordingly, the current study reports on an energy-
60 dispersive X-ray fluorescence (XRF) analysis of EPS and XPS fragments retrieved from various
61 Atlantic- and English Channel-facing beaches of southwest England. XRF spectrometry affords a
62 rapid, non-destructive means of quantifying a range of elements in solids and through a complex
63 series of mathematical algorithms solved iteratively is suited for the analysis of materials of low-
64 density like plastics and including foams (Piorek, 2004; Furl et al., 2012). XRF is also able to provide
65 information on the spatial distribution of elements, including the relative significance of those
66 incorporated into the matrix and held at the surface.

67

68 2. Methods

69 2.1. Sample collection

70 As part of a number of research programmes studies aimed at examining marine litter more
71 generally (Turner and Solman, 2016) or targeting samples for experimental studies (Turner and Lau,
72 2016), samples of EPS and XPS had been retrieved manually during late spring or early autumn of
73 2015 from three beaches in southwest England. Namely, Tregantle (50.353, -4.271), a large,
74 southwest-facing sandy beach on the south coast of Cornwall that is backed by high cliffs,
75 Constantine Bay (50.539, -5.027), a large, southwest-facing sandy beach on the north coast of
76 Cornwall that is backed by a dune system, and Mount Batten (50.356, -4.127), a small, sand-gravel
77 beach in the partly-urbanised embayment of Plymouth Sound. Polystyrene fragments were usually
78 abundant on the latest (or highest) strandline and on the back-beach near to the cliff or dune base.
79 In the laboratory, any loosely adherent material was dislodged or brushed off under tap water
80 before samples were dried under desiccation at room temperature for 96 h. Samples were then
81 weighed and photographed and stored in individual polyethylene specimen bags in the dark until
82 required for analysis.

83 2.2. XRF analysis

84 Samples were analysed by energy-dispersive portable XRF spectrometry using a Niton XL3t 950 He
85 GOLDD+ configured in a laboratory accessory stand. The instrument was operated in a customised,
86 standardless 'plastics' mode and because of the low density of foamed polystyrene, a thickness
87 correction algorithm was always applied after measuring the thickness of material through the
88 measurement surface using Allendale digital callipers (Turner and Solman, 2016). Samples were
89 counted for periods ranging from 60 to 180 s, depending on material thickness, that were
90 distributed in a 2:1 ratio between a low energy range (20 kV and 100 μ A) and main energy range (50
91 kV and 40 μ A). Spectra were quantified by fundamental parameter coefficients to yield
92 concentrations on a dry weight basis (in mg kg⁻¹) and with a counting error of 2 σ (95% confidence).
93 Up to 18 elements may be analysed in the plastics mode but the focus of the present study was on
94 those that are known to be used in various plastics (Ba, Br, Fe, Sb, Ti, Zn), act as proxies for
95 hydrogenous and detrital phases (Fe, Ti) or are contaminants of toxicological concern (Br, Cd, Pb).
96 By default, the measurement face was an 8-mm diameter circular area of the central region of the
97 largest or flattest face. For selected samples larger than about 3 cm in diameter, between four and
98 eight measurements were made at different locations over the face to examine the distribution of
99 elements within the matrix. Here, the small-spot facility of the instrument was employed in order to

100 focus the X-ray beam into a 3-mm diameter circular area. For these samples, where possible, material
101 was sliced through the thickest axis with a stainless steel scalpel and the central core analysed.

102 As a performance check, polyethylene reference discs Niton PN 180-619 (Ba = 688 ± 45 mg kg⁻¹; Cd =
103 150 ± 6 mg kg⁻¹; Pb = 150 ± 12 mg kg⁻¹; Sb = 96 ± 10 mg kg⁻¹) and Niton PN 180-554 (Br = 495 ± 20 mg
104 kg⁻¹; Cd = 292 ± 20 mg kg⁻¹; Pb = 1002 ± 40 mg kg⁻¹) were analysed throughout each measurement
105 session, with the instrument returning concentrations that were consistently within 10% of certified
106 values. Detection limits varied depending on counting time, sample thickness and whether the small-
107 spot was engaged but indicative values based on the lowest counting errors returned throughout
108 the study ranged from about 10 mg kg⁻¹ for Br, Pb and Ti to 100 mg kg⁻¹ for Ba and Sb in EPS and
109 about 20 mg kg⁻¹ for Br, Pb and Ti to 250 mg kg⁻¹ for Ba and Sb in XPS. Analytical precision, defined by
110 repeat measurements ($n = 5$) of the same area of sample, was generally better than 10% but could
111 be as high as 25% if concentrations were close to the detection limit and determined with the small-
112 spot facility.

113

114 2.3. Polystyrene confirmatory analysis

115 For sixteen samples that were particularly weathered or discoloured, confirmation of the polymeric
116 matrix was achieved by attenuated total reflectance Fourier Transform Infrared (FTIR) spectrometry
117 using a Bruker Vertex 70. Here, 1-3 mm offcuts of the sample surface were clamped against the ATR
118 diamond crystal and spectra were recorded with 16 scans in the region 4000 to 600 cm⁻¹ and at a
119 resolution of 4 cm⁻¹. After being smoothed, baseline-corrected and normalised via Bruker OPUS 7
120 software, spectra were matched up with spectra of polystyrene contained in various reference
121 libraries.

122

123 3. Results

124 A total of 86 samples, exemplified in Figure 1, were returned to the laboratory for characterisation.
125 Samples ranged in size from individual foamed beads of about 4 mm in diameter and 30 mg in mass
126 to fragments of about 10 cm in length and weighing up to 10 g. EPS samples ($n = 67$) were identified
127 from presence of the distinctive beaded (cellular) structure while XPS samples ($n = 19$) were usually
128 unstructured, flatter and smoother. Where material was heavily fouled or discoloured, FTIR analysis
129 confirmed the polymeric composition of both EPS and XPS from aromatic ring stretches at 1600 cm⁻¹
130 1490 cm⁻¹ and aromatic CH out-of-plane bends at 694 cm⁻¹ and 537 cm⁻¹ (Noda et al., 2007).

131 Most samples of EPS were well-rounded, unidentified fragments that contained surficial cracks or
132 pits, and the polystyrene itself was usually white or off-white. Dissection of samples revealed a
133 “clean” interior that was generally free of fouling or particle intrusion (Figure 1h). Samples of XPS
134 tended to be thinner and more ragged (Figures 1c and 1e), and displayed a broader range of colours
135 that that included white, beige, blue or green. Some samples of XPS had evidently been derived from
136 single-use food or drink containers that likely had a local (littering) origin. With the exception of the
137 latter type of sample, most foamed fragments were visibly discoloured at the face (and relative to
138 the core) by orange-brown chemical fouling, with some EPS soiled by fine particulates trapped in the
139 pore space between weathered beads or stained by oil.

140 Summary statistics for the concentrations of the elements in the samples as determined by XRF are
141 shown in Table 1, with the full dataset shown in Appendix A. Data in Table 1 are pooled for the three
142 locations and a direct comparison is made between the two different types of foamed polystyrene.
143 Note that measurements were made in the central region of the largest face and, because of the low
144 density of the material, values reported signify concentrations that are representative of the entire
145 depth of sample rather than the measurement face. Note also that elemental detection limits for
146 XPS were about double the corresponding detection limits for EPS because of the lower thickness of
147 the former samples. Among the elements considered, Fe and Ti were most commonly detected and
148 exhibited the highest mean, median and maximum concentrations. Bromine and Zn were detected
149 in about 60 cases each, Pb was detected in 14 samples of EPS and Ba was detected in eight samples
150 of EPS; Sb was detected in just one sample of XPS and Cd was never detected ($< 25 \text{ mg kg}^{-1}$). A series
151 of Kruskal-Wallis tests performed in Minitab v19 revealed that median concentrations of Fe and Ti in
152 EPS were significantly greater than respective median concentrations in XPS ($H > 15, p < 0.01$) but
153 there was no significant difference in median concentrations of either Br or Zn between the material
154 types ($H < 1, p > 0.1$).

155 Table 1: Detection frequency for the different elements and summary statistics for their
156 concentrations (in mg kg^{-1}).

	Ba	Br	Cd	Fe	Pb	Sb	Ti	Zn
EPS (<i>n</i> = 19)								
detection, %	9.0	79.1	0	100	20.9	0	98.5	65.7
mean	536	1610		9620	50		2670	392
median	438	224		8170	20		1620	173
min	350	12		282	11		68	42
max	1060	11500		29100	175		50900	3560
XPS (<i>n</i> = 19)								
detection, %	0	31.6	0	89.5	0	5.3	94.7	89.5
mean		890		2300		373	579	193
median		120		1150			518	135
min		27		85			25	36
max		4710		7120			1740	674

157

158 Figure 2 shows the spatial distribution of Br, Fe, Ti and Zn in five samples of EPS (samples of XPS
159 were generally too thin to perform this characterisation (see Figure 1) and Pb was never detected in
160 the core of EPS). Specifically, the variation in concentrations measured on the largest or flattest face,
161 $[]_{\text{face}}$, are shown against single or replicate measurements of concentration of the core, $[]_{\text{core}}$, once
162 the sample had been dissected. For Br, relative standard deviations (rsds) of concentrations are <
163 30% except for the face of one sample (rsd ~ 65%). Overall, however, mean concentrations lie close
164 to unit slope, indicating that concentrations determined through the face and in the core are similar.
165 The distributions of Fe and Ti are rather scattered on the charts, and especially in the y-axis
166 where rsds of up to 50% are observed, and concentrations of both elements measured on the face
167 are similar to (*n* = 2), greater than (*n* = 2) or lower than (*n* = 1) corresponding concentrations
168 measured in the core. Regarding Zn, with the exception of one sample lying well above unit slope,
169 concentrations measured on the face and core are similar and exhibit relatively little variation
170 among replicate measurements. Note that Ba, not plotted, was detected on the face and core of one
171 sample and returned concentrations that were similar ($676 \pm 47 \text{ mg kg}^{-1}$ and $672 \pm 25 \text{ mg kg}^{-1}$,
172 respectively).

173

174 4. Discussion

175 The results of this study are consistent with the reaction residues and functional additives known to
176 be present in foamed polystyrene or in thermoplastics more generally but are significant in providing
177 an insight into the concentration ranges of these chemicals in EPS and XPS fragments encountered in
178 the marine environment. Thus, the detection of Fe in most samples may be attributed to the use of
179 Fe_2O_3 as a catalyst in the production of styrene (Wünsch, 2000) while the presence of Zn in many
180 cases, and at least in EPS, may be attributed to residual Zn stearate that is often used to ensure

181 uniform cell nucleation in the production of the expanded material (European Union, 2008). The
182 detection of Ti in most cases reflects the common usage of TiO₂, and principally rutile, as a pigment
183 that is blended or moulded into the material to provide a high refractive index and chemical stability
184 (Murphy, 2001), while the lack of Ba is due to the avoidance of fillers of high specific gravity, like
185 BaSO₄, in products specifically designed to be lightweight (Turner and Filella, 2020).

186 Detectable Br in a high proportion of EPS samples and about a third of XPS samples is attributed to
187 the presence of brominated flame retardants (Gallen et al., 2014). XRF is unable to discriminate
188 different brominated compounds but the most commonly used in foamed PS, and in particular that
189 destined for the construction sector, was 1,2,5,6,9,10-hexabromocyclododecane (HBCD). Production
190 of HBCD began in the 1980s until it was added to Annex A of persistent organic pollutants in the
191 Stockholm Convention that require elimination (United Nations, 2013) and effectively banned
192 (Schlummer et al., 2017). HBCD was added to foamed polystyrene at concentrations that are low
193 relative to concentrations of other halogenated compounds used to flame-retard plastics (about 0.7
194 to 2.5% by weight; Alaei et al., 2003). Moreover, and unlike flame-retarded plastics more generally,
195 foamed PS impregnated with HBCD did not require the addition of antimony trioxide (Sb₂O₃) as a
196 synergist to meet various building code specifications (Papazoglou, 2004), thereby accounting for
197 the lack of detection of Sb in the current samples (Table 1).

198 Assuming that Br concentrations result from only HBCD and given that the fractional mass
199 contribution of Br to HBCD is 0.75, the maximum concentration of the flame retardant reported in
200 Table 1 is about 15,300 mg kg⁻¹ (or 1.5% of HBCD by weight). This value is in good agreement with
201 the maximum concentration of HBCD measured directly in EPS and XPS fragments retrieved from the
202 north Pacific Ocean (Jang et al., 2017) and is within the content range required for flame-retardancy
203 of construction material (Alaei et al., 2003). This observation, coupled with Br being undetected in
204 samples of XPS that were evidently related to single-use food-drink containment (such as take-out
205 trays and Styrofoam cups) suggests that the majority of samples collected herein originate from the
206 construction industry. However, this may not necessarily be the case because a single grade of HBCD
207 often appears to have been used for wider production (for use in both construction and packaging,
208 for example) (Lassen et al., 2019) and there is evidence for the uncontrolled use or recycling of the
209 flame retardant (Rani et al., 2014; Abdullah et al., 2018). The latter pathway would account for the
210 high variability of Br concentrations among the samples and the occurrence of concentrations well
211 below those required for flame-retardancy in many cases.

212 From a regulatory perspective, the EU has recently introduced a low concentration limit of 0.1%
213 (1000 mg kg⁻¹) by weight for certain brominated compounds, including HBCD, above which items

214 may not be recycled, and a limit of 0.01% (100 mg kg⁻¹) above which products are not permitted for
215 sale in the EU (European Commission, 2016). Converting the Br measurements summarised in Table
216 1 into HBCD concentrations reveals that 18 samples of EPS and one sample of XPS would be non-
217 compliant for recycling and that 43 samples of EPS and three samples of XPS would be non-
218 compliant for resale.

219 Another factor to consider that has a potentially significant impact on elemental concentrations
220 measured in the EPS and XPS samples is the acquisition of chemicals from or their loss to the
221 environment while at sea or once beached. Since the additives and residues identified above are not
222 covalently bonded to the polymeric matrix, there is scope for migration from foamed polystyrene
223 into seawater. Predicted or measured diffusion rates for ions and compounds in virgin plastics vary
224 considerably but are generally very small (Town et al., 2018), with estimates of diffusion half-lives
225 for various brominated flame retardants on the order of tens of thousands of year or more (Sun et
226 al., 2019). That said, physical and chemical processes acting on plastics in the marine environment,
227 including digestion by animals, may considerably accelerate leaching (Tanaka et al., 2015; Smith and
228 Turner, 2020) and result in a more heterogeneous distribution of chemicals in the weathered matrix
229 (Turner et al., 2020).

230 With a specific surface area of about 2 m² g⁻¹ and a point of zero charge of about 4.7, the inherent
231 surface properties of foamed polystyrene are more favourable for the adsorption of metal ions than
232 many other polymers (Zhang et al., 2018). Moreover, the low density of EPS and XPS ensures that
233 material is exposed to elevated concentrations of metals and other contaminants in the sea surface
234 microlayer (Wurl and Obbard, 2004). On aging, the surfaces of EPS and XPS become discoloured and
235 fouled by organic and inorganic precipitates, a process that is visibly and chemically heterogeneous
236 (see Figure 1 and the y-axis distributions of Fe and Ti in Figure 2) and that likely further enhances the
237 adsorption of contaminant metals like Pb. In theory, the degree of metal enrichment at the face may
238 be related to the age of sample or the time spent in suspension in seawater. However, it is
239 important to bear in mind that metal enrichment at or near the polymer surface may also arise from
240 the entrapment of fine detrital particulates within the surficial interstices of the material that evade
241 removal during sample cleaning. Moreover, because of the low density of foamed polystyrene,
242 measurements undertaken through the face are, strictly, more representative of concentrations
243 through the entire sample (and not the measurement face alone).

244 Amongst the elements analysed, there was a strong association between Fe and Ti once two
245 samples with exceptionally high Ti concentrations (> 10,000 mg kg⁻¹) had been excluded (Figure 3a).
246 Results of regression analyses of all data and individual data sets defining the faces of XPS and EPS

247 samples and the central core of EPS samples are shown in Table 2. Thus, concentrations of Fe and Ti
 248 are significantly related in each case, with relatively small intercepts (*c*) and slopes (*m*) that range
 249 from 3.90 for XPS faces to 6.72 for EPS cores. A value of *m* that is greater in EPS than XPS suggests
 250 that residual, catalytic Fe relative to pigmented Ti is higher in EPS, while the greatest value of *m* that
 251 defines the cores of EPS suggests that the ratio of authigenic Fe to Ti acquired from the environment
 252 is lower than the elemental ratio arising from manufacture.

253 In contrast, a lack of association was observed between the concentrations of Fe and Br (Figure 3b).
 254 Here, concentrations of HBCD and residual Fe appear to be unrelated during the manufacturing
 255 process, perhaps because of the uncontrolled use and recycling of the former (Rani et al., 2014).
 256 Moreover, while Fe is readily acquired from the environment, HBCD and other forms of Br are not
 257 taken up appreciably and may be more subject to heterogeneous leaching from the matrix (Tanaka
 258 et al., 2015).

259

260 Table 2: Statistical parameters defining relationships between concentrations of Fe and Ti shown in
 261 Figure 3a.

	<i>n</i>	<i>m</i>	<i>c</i>	<i>r</i> ²	<i>p</i>
all data*	85	5.85	370	0.674	<0.0001
XPS	16	3.90	298	0.752	<0.0001
EPS	64	5.53	1140	0.576	<0.0001
262 EPS-core	5	6.72	-412	0.989	0.0005

263 *After exclusion of two samples where [Ti] > 10,000 mg kg⁻¹.

264

265 5. Conclusions

266 The findings of this study reveal that waste EPS and XPS can act as both a source and sink of
 267 contaminants in the marine environment. Reaction residues and additives in the matrix detected by
 268 XRF include Fe₂O₃, TiO₂ and brominated compounds (presumably dominated by the flame retardant,
 269 HBCD), with the latter of greatest environmental concern. Accumulations of Fe- and Ti-based ions
 270 and minerals on the surface of polystyrene foam reflect the propensity of the material to acquire
 271 chemicals from the environment, with evidence that these phases can also act as adsorbents of
 272 more harmful metals like Pb. Given the ability of EPS and XPS to undergo long-range transport and
 273 ready fragmentation, it is recommended that future research focuses on the geochemical reactivity
 274 and biological interactions of this type of plastic litter.

275 **Acknowledgements**

276 This study was funded by a HEIF Plymouth Marine Institute Grant.

277

278 **Appendix A: Supplementary material**

279 Supplementary data to this article can be found online at...

280

281 **References**

282 Abdullah, M.A.E., Sharkey, M., Berresheim, H., Harrad, S., 2018. Hexabromocyclododecane in
283 polystyrene packaging: A downside of recycling? *Chemosphere* 199, 612-616.

284 Alae, M., Arias, P., Sjodin, A., Bergman, A., 2003. An overview of commercially used brominated
285 flame retardants, their applications, their use patterns in different countries/regions and possible
286 modes of release. *Environment International* 29, 683–689.

287 Black, J.E., Kopke, K., O'Mahony, C., 2019. Towards a circular economy: using stakeholder
288 subjectivity to identify priorities, consensus, and conflict in the Irish EPS/XPS market. *Sustainability*
289 11, 6834 doi:10.3390/su11236834

290 Block, C., Brands, B., Gude, T., 2017. *Packaging Materials 2. Polystyrene for Food Packaging*
291 Applications. ILSI Europe, Brussels, 36pp.

292 Carson, H.S., Nerheim, M.S. Carroll, K.A., Eriksen, M., 2013. The plastic-associated microorganisms of
293 the North Pacific Gyre. *Marine Pollution Bulletin* 75, 126-132.

294 Che, C.L., Kuo, P.H., Lee, T., Liu, C.H., 2018. Snow lines on shorelines: Solving Styrofoam buoy marine
295 debris from oyster culture in Taiwan. *Ocean and Coastal Management* 165, 346-355.

296 Chitaka, T.Y., von Blottnitz, H., 2019. Accumulation and characteristics of plastic debris along five
297 beaches in Cape Town. *Marine Pollution Bulletin* 138, 451-457.

298 Chubarenko, I., Bagaev, A., Zobkov, M., Esiukova, E., 2016. On some physical and dynamical
299 properties of microplastic particles in marine environment. *Marine Pollution Bulletin* 108, 105-112.

300 Davis, W, Murphy, A.G., 2015. Plastic in surface waters of the Inside Passage and beaches of the
301 Salish Sea in Washington State. *Marine Pollution Bulletin* 97, 169-177.

302 European Commission, 2016. Regulation (EC) No 850/2004 of the European Parliament and of the
303 Council of 29 April 2004 on Persistent Organic Pollutants and Amending Directive 79/117/EEC.
304 <https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX:02004R0850-20160930> accessed 6/20.

305 European Union, 2008. Risk Assessment: Zinc distearate. EUR 21168 EN, Luxembourg, 188pp.

306 Fok, L., Cheung, P.K., Tang, G.D., Li, W.C., 2017. Size distribution of stranded small plastic debris on
307 the coast of Guangdong, South China. *Environmental Pollution* 220, 407-412.

- 308 Furl, C., Mathieu, C., Roberts, T., 2012. Evaluation of XRF as a screening tool for metals and PBDEs in
309 children's products and consumer goods. Environmental Assessment Program Report No. 12-03-009,
310 Washington State Department of Ecology, Olympia, WA, 69pp.
- 311 Gallen, C., Banks, A., Brandsma, S., Baduel, C., Thai, P., Eaglesham, G., Heffernan, A., Leonards, P.,
312 Bainton, P., Mueller, J.F., 2014. Towards development of a rapid and effective non-destructive
313 testing strategy to identify brominated flame retardants in the plastics of consumer products.
314 *Science of the Total Environment* 491/492, 255-265.
- 315 Hansen, A.A., Rodbotten, M., Lea, P., Rotabakk, B.T., Birkeland, S., Pettersen, M.K., 2015. Effect of
316 transport packaging and repackaging into modified atmosphere on shelf life and quality of thawed
317 Atlantic cod loins. *Packaging Technology and Science* 28, 925-938.
- 318 Hinojosa, I.A., Thiel, M., 2009. Floating marine debris in fjords, gulfs and channels of southern Chile.
319 *Marine Pollution Bulletin* 58, 341-350.
- 320 Jang, M., Shim, W.J., Han, G.M., Rani, M., Song, Y.K., Hong, S.H., 2017. Widespread detection of a
321 brominated flame retardant, hexabromocyclododecane, in expanded polystyrene marine debris and
322 microplastics from South Korea and the Asia-Pacific coastal region. *Environmental Pollution* 231,
323 785-794.
- 324 Jang, M., Shim, W.J., Cho, Y., Han, G.M., Song, Y.K., Hong, S.H., 2020. A close relationship between
325 microplastic contamination and coastal area use pattern. *Water Research* 171, 115400 DOI:
326 10.1016/j.watres.2019.115400
- 327 Lassen, C., Warming, M., Kjøholt, J., Jakobsen, L.G., Vrubliauskiene, N., Norichkov, B., Strand, J., Feld,
328 L., Bach, L., 2019. Survey of Polystyrene Foam (EPS and XPS) in the Baltic Sea. Danish Fisheries
329 Agency/Ministry of Environment and Food of Denmark, 166pp.
- 330 Murphy, J., 2001. *Additives for Plastics Handbook*, second edition. Elsevier Science, Oxford, 471pp.
- 331 Nabizadeh, R., Sajadi, M., Rastkari, N., Yaghmaelan, K., 2019. Microplastic pollution on the Persian
332 Gulf shoreline: A case study of Bandar Abbas city, Hormozgan Province, Iran. *Marine Pollution*
333 *Bulletin* 145, 536-546.
- 334 Nie, Z.Q., Yang, Z.L., Fang, Y.F., Tang, Z.W., Wang, X.R., Die, Q.Q., Gao, X.B., Zhang, F.S., Wang, Q.,
335 Huang, Q.F., 2015. Environmental risks of HBCDD from construction and demolition waste: A
336 contemporary and future issue. *Environmental Science and Pollution Research* 22, 17249-17252.
- 337 Noda I., Dowrey A.E., Haynes J.L., Marcott C., 2007. Group Frequency Assignments for Major Infrared
338 Bands Observed in Common Synthetic Polymers. In: Mark J.E. (eds) *Physical Properties of Polymers*
339 *Handbook*. Springer, New York, NY. https://doi.org/10.1007/978-0-387-69002-5_22
- 340 Papazoglou, E.S., 2004. Flame retardants for plastics, in: C.A. Harper (ed.), *Handbook of Building*
341 *Materials for Fire Protection*, McGraw-Hill, New York, pp4.1-4.88.
- 342 Piorek, S., 2004. Feasibility of analysis and screening of plastics for heavy metals with portable x-ray
343 fluorescence analyser with miniature x-ray tube. GPEC 2004 Paper abstract #14.
- 344 Rani, M., Shim, W.J., Han, G.M., Jang, M., Song, Y.K., Song, S.H., 2014. Hexabromocyclododecane in
345 polystyrene-based consumer products: an evidence of unregulated use. *Chemosphere* 110, 111-119.

346 Schlummer, M., Mäurer, A., Wagner, S., Berrang, A., Fell, T., Knappich, F., 2017. Recycling of flame
347 retarded waste polystyrene foams (EPS and XPS) granules free of hexabromocyclododecane (HBCD).
348 *Advances in Recycling and Waste Management* 2.2 DOI: 10.4172/2475-7675.1000131

349 Smith, E.C., Turner, A., 2020. Mobilisation kinetics of Br, Cd, Cr, Hg, Pb and Sb in microplastics
350 exposed to simulated, dietary-adapted digestive conditions of seabirds. *Science of the Total*
351 *Environment* 733, 138802.

352 Song, Y.K., Hong, S.H., Jang, M., Han, G.M., Jung, S.W., Shim, W.J., 2017. Combined effects of UV
353 exposure duration and mechanical abrasion on microplastic fragmentation by polymer type.
354 *Environmental Science and Technology* 51, 4368-4376.

355 Tanaka, K., Takada, H., Yamashita, R., Mizukawa, K., Fukuwaka, M.A., Watanuki, Y., 2015. Facilitated
356 leaching of additive-derived PBDEs from plastic by seabirds' stomach oil and accumulation in tissues.
357 *Environmental Science and Technology* 49, 11799-11807.

358 Tata, T., Belabed, B.E., Bououdina, M., Bellucci, S., 2020. Occurrence and characterization of surface
359 sediment microplastics and litter from North African coasts of Mediterranean Sea: Preliminary
360 research and first evidence. *Science of the Total Environment* 713, 136664.

361 Town, R.M., van Leeuwen, H.P., Blust, R., 2018. Biochemodynamic features of metal ions bound by
362 micro- and nano-plastics in aquatic media. *Frontiers in Chemistry*
363 <https://doi.org/10.3389/fchem.2018.00627>

364 Turner, A., Lau, K.S., 2016. Elemental concentrations and bioaccessibilities in beached plastic foam
365 litter, with particular reference to lead in polyurethane. *Marine Pollution Bulletin* 112, 265-270.

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

368 Turner, A., Filella, M., 2020. The influence of additives on the fate of plastics in the marine
369 environment, exemplified with barium sulphate. *Marine Pollution Bulletin* 158, 111352.

370 Turner, A., Arnold, R., Williams, T., 2020. Weathering and persistence of plastic in the marine
371 environment: Lessons from LEGO. *Environmental Pollution* 262, 114299.

372 United Nations, 2013. Stockholm Convention on Persistent Organic Pollutants.
373 C.N.934.2013.TREATIES-XXVII.15 (Amendment to Annex A).
374 <https://treaties.un.org/doc/Publication/CN/2013/CN.934.2013-Eng.pdf> accessed 6/20.

375 Wurl, O., Obbard, J.P., 2004. A review of pollutants in the sea-surface microlayer (SML): a unique
376 habitat for marine organisms. *Marine Pollution Bulletin* 48, 1016-1030.

377 Wünsch, J.R., 2000. Polystyrene: Synthesis, Production and Applications. RAPRA report, Shawbury,
378 UK, 160 pp.

379 Zhang, H., Wang, J., Zhou, B., Zhou, Y., Dai, Z., Zhou, Q., Christie, P., Luo, Y., 2018. Enhanced
380 adsorption of oxytetracycline to weathered microplastic polystyrene: Kinetics, isotherms and
381 influencing factors. *Environmental Pollution* 243, 1550-1557.

382

Figure 1: Examples of the EPS (a,b,d,f,g,h) and XPS (c,e) samples retrieved from beaches of southwest England. Annotated bars are approximately 1 cm.

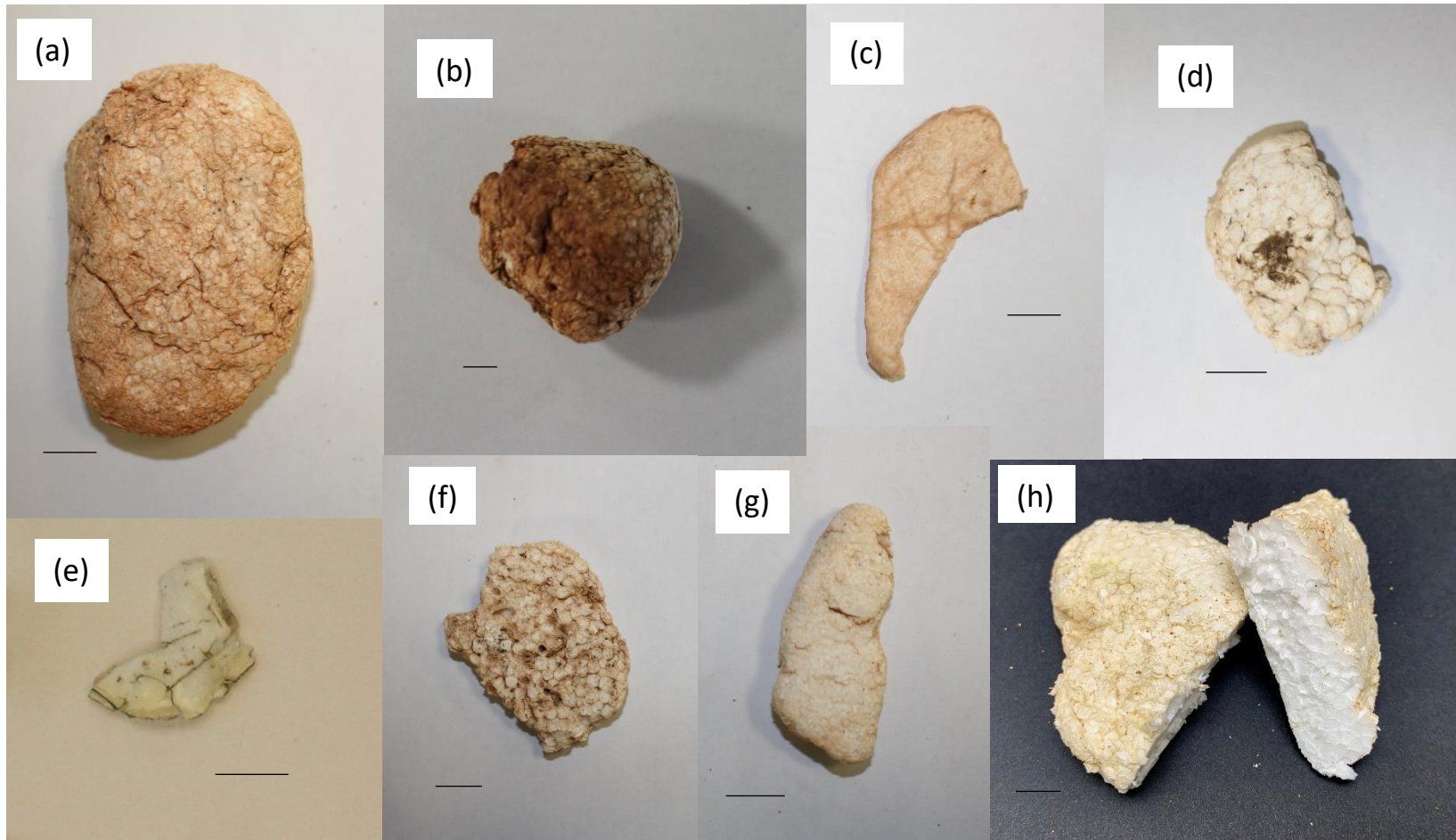


Figure 2: Elemental concentrations measured at various locations on the face versus concentrations measured in the core of five samples of EPS. (a) Br, (b) Fe, (c) Ti, (d) Zn; errors are one standard deviation about the mean ($n = 3$ to 7) and the red line signifies unit slope.

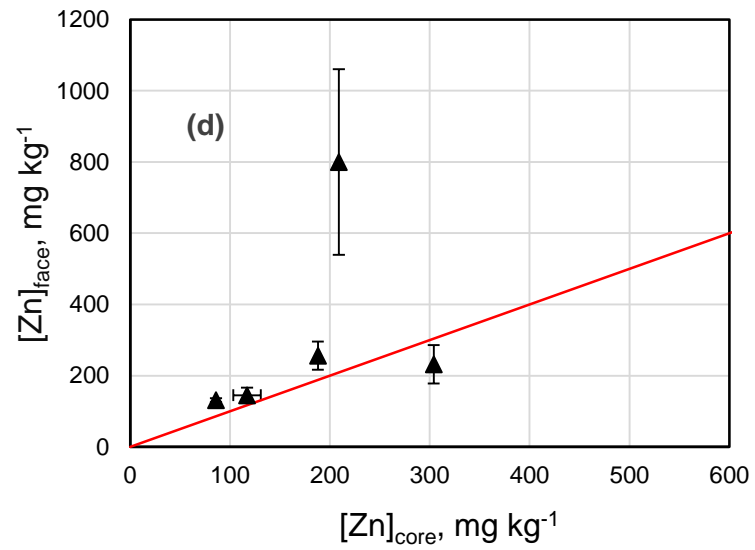
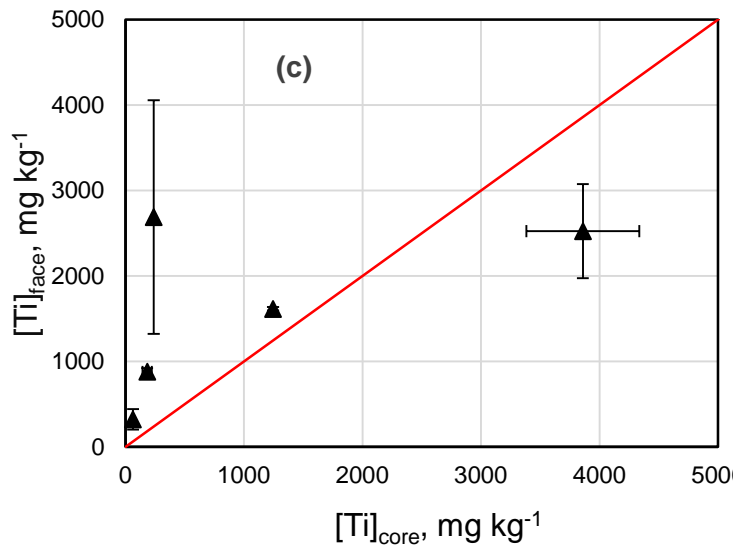
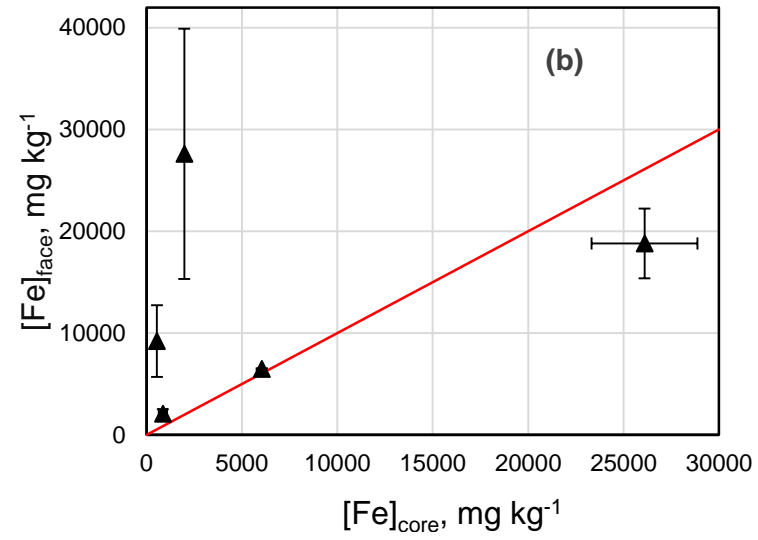
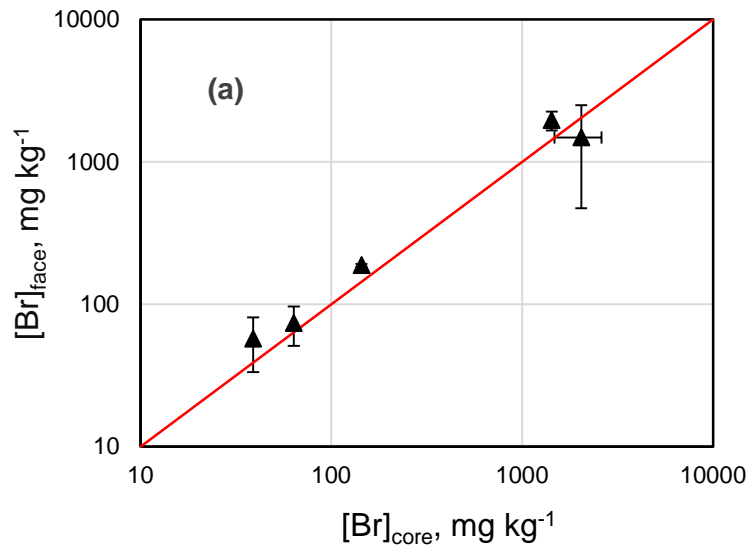


Figure 3: Concentrations of (a) Fe versus Ti and (b) Fe versus Br measured on the central face of XPS and EPS samples and in the core of selected EPS fragments. Parameters defining Fe versus Ti are shown in Table 2.

