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Perfluoroalkyl substances and brominated flame retardants in landfill-related air, soil, and groundwater from Ireland

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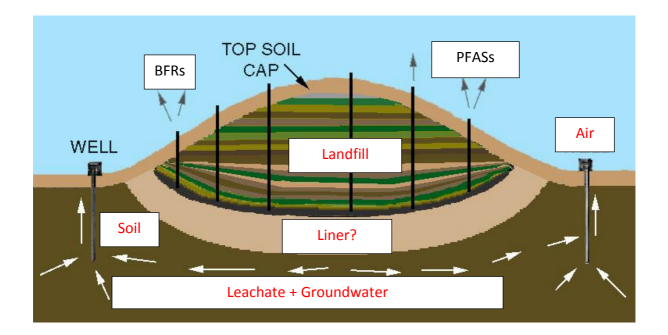
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*Highlights (for review : 3 to 5 bullet points (maximum 85 characters including spaces per bullet point)

HIGHLIGHTS

- BFRs and PFASs measured in air, soil and groundwater near 10 Irish landfills
- First report of DBDPE in groundwater at levels exceeding PBDEs, HBCDD and PFOA
- BFRs and PFASs in air and soil not impacted by landfill
- PFOA in groundwater positively correlated with concentrations in landfill leachate
- PFOA higher in groundwater near unlined than near lined landfills

1	Permuoroalkyi Substances and Drommated Flame Retardants in
2	Landfill-Related Air, Soil, and Groundwater from Ireland
3	
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11	

12 ABSTRACT

13 Between November 2018 and January 2019, samples of air and soil were collected 14 from locations downwind and upwind of 10 landfills across the Republic of Ireland. 15 Samples of groundwater (n=10) were also collected from locations with links 16 traceable to the studied landfills. Concentrations of perfluoroalkyl substances 17 (PFASs), polybrominated diphenyl ethers (PBDEs), hexabromocyclodecane 18 (HBCDD), and decabromodiphenyl ethane (DBDPE) determined in air and soil 19 samples were not significantly different (p>0.05) between downwind and upwind 20 locations. The arithmetic mean concentration of PFOA in groundwater sourced from 21 landfills (n=4) that were not fully lined (69 ng/L) exceeded that in groundwater 22 samples sourced from lined landfills (n=6; 4.1 ng/L), with the difference, however 23 being not statistically significant (p>0.1). A positive correlation (p=0.014) was 24 observed between concentrations of PFOA in groundwater in our study and those 25 reported previously in leachate from the same landfills. However, this correlation was 26 driven substantially by one landfill which displayed the highest concentrations in both 27 groundwater and leachate and no significant correlation (p>0.1) was observed 28 between log-transformed concentrations of PFOA in groundwater and leachate. 29 DBDPE was detected in groundwater for the first time anywhere, in all samples at 30 concentrations (median = 9.4 ng/L; arithmetic mean = 78 ng/L) that exceeded those of 31 any other BFRs or PFASs targeted in this study. This likely reflects its recent use as a 32 "drop-in" replacement for the recently restricted Decabromodiphenyl ether product. 33 Overall, our data suggest that the 10 landfills studied do not exert a discernible 34 influence on local air and soil concentrations of BFRs and PFASs. In contrast, while 35 not of immediate concern, our data suggest that further more detailed study of the 36 impact of landfill emissions on concentrations of DBDPE and PFOA is advisable.

37 **KEYWORDS**

38 PFASs; BFRs; DBDPE; Landfill; Air: Soil; Groundwater

39

40 HIGHLIGHTS

- BFRs and PFASs measured in air, soil and groundwater near 10 Irish landfills
- First report of DBDPE in groundwater at levels exceeding PBDEs, HBCDD and
- 43 PFOA
- BFRs and PFASs in air and soil not impacted by landfill
- PFOA in groundwater positively correlated with concentrations in landfill
- 46 leachate
- PFOA higher in groundwater near unlined than near lined landfills

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INTRODUCTION

- Perfluoroalkyl substances (PFASs) have found widespread application in domestic,
- 51 institutional, and office environments to for example confer stain/oil repellency to
- 52 fabrics and food packaging (BiPRO, 2011, Miralles-Marco and Harrad, 2015).
- 53 Similarly, brominated flame retardants (BFRs) like polybrominated diphenyl ethers
- 54 (PBDEs) and hexabromocyclododecane (HBCDD) have been used extensively in a
- broad range of applications such as soft furnishings and fabrics, as well as electrical
- and electronic equipment (BiPRO, 2011). Concerns about their adverse impacts on
- 57 human and wildlife health have led to restrictions and bans on the manufacture and
- use of PBDEs, HBCDD, and PFASs like perfluorooctane sulfonate (PFOS) and
- 59 perfluorooctanoic acid (PFOA), exemplified by their listing or (in the case of
- 60 perfluorohexane sulfonate (PfHxS)) active current consideration for listing under the

Stockholm Convention on persistent organic pollutants. Despite such concerns, there
remain a substantial quantity of articles treated with such chemicals (Drage et al,
2018). One consequence of such restrictions has been increased use of alternative
flame retardants and PFASs, as evidenced by our recent reports of elevated
concentrations of decabromodiphenyl ethane (DBDPE) and perfluorobutane sulfonate
(PFBS) in indoor air and dust from Ireland (Harrad et al, 2019b; Wemken et al, 2019).
As articles containing BFRs and PFASs become redundant, they enter the waste
stream. One widely-favoured waste disposal method has historically been landfill.
While disposal of waste electrical and electronic equipment (WEEE) to landfill has
been restricted since 2003 within the European Union (EU) as a result of the WEEE
directive (EC, 2003); similar restrictions on landfilling of waste soft furnishings and
fabrics do not exist and moreover, landfills located within the EU that were open
before 2003 may contain WEEE received before this date. Given such material may
contain elevated concentrations of PBDEs, HBCDD, and PFASs; its presence in
landfill is of potential concern.
We recently reported on the ubiquitous presence of PBDEs, HBCDD, and a number
of PFASs in leachate from 40 landfills in the Irish Republic (Harrad et al, 2019a). We
also reported that concentrations of some BFRs and PFASs were significantly higher
in leachate from landfills fitted with impervious liners than in leachate from landfills
that were partially or completely unlined. This was attributed to the higher organic
matter content of leachate from lined landfills (Harrad et al, 2019a), and raised
concern that BFRs and PFASs in leachate from unlined landfills could potentially
contaminate groundwater. Additional concerns have also been voiced that
volatilisation of PBDEs and PFASs from landfilled waste may lead to elevated

85 concentrations in air in the vicinity of landfills (St. Amand et al, 2008; Weinberg et al, 86 2012). This study tests the hypothesis that landfills have caused discernible contamination of 87 88 their surrounding environment. To do so, we report concentrations of PBDEs, 89 HBCDD, DBDPE, and PFASs in samples of air and soil taken both down-and up-90 wind of 10 landfills in the Republic of Ireland. We also report concentrations of the 91 same contaminants in samples of groundwater linked to the same landfills. In addition 92 to facilitating assessment of the influence of landfills on the surrounding environment, 93 the data reported here represent the most detailed database yet on these contaminants 94 in Irish groundwater, outdoor air, and soil.

MATERIALS AND METHODS

Sampling

95

96

Air and soil - Air and soil samples were collected between 6th November 2018 and 97 31st January 2019 from 10 municipal solid waste (MSW) landfill sites across the 98 99 Republic of Ireland (Figure 1). A summary of the methods employed is provided here, 100 with further information about the sampling locations and factors like the passive air 101 sampling rates used to calculate concentrations in air samples (taken from Drage et al, 102 2016 and Goosey and Harrad, 2012) provided as supporting information. 103 Each passive air sampler deployed consisted of a polyurethane foam (PUF) disk (140 104 x 140 x 12 mm) mounted between two stainless steel bowls lined with hexane-rinsed 105 aluminium foil. Each PUF disk was coated with pre-cleaned Amberlite XAD powder 106 by submerging each disk in a solution of the powder crushed to a particle size of 0.75 107 um suspended in hexane. Once dried, the disks were loaded into the passive air 108 samplers, two of which were placed at locations upwind and downwind, respectively,

109	of each landfill site. The locations of sampling sites were based on wind direction data
110	taken from the Irish Meteorological Service (Met Eireann), with slight modification
111	where necessary based on local information from the site operators and ease of
112	access. Sampling points were located between 150 and 500 m of the centre of the
113	landfill.
114	To maximise the effective air volume collected from each location for analysis, two
115	samplers were placed at each sampling point, hung either side of a fence post at a
116	height of 1.2 m above the ground, and left in place for approximately 2 months. At the
117	end of this period, the PUF disks were collected, sealed in a piece of hexane-rinsed
118	aluminium foil and placed in plastic ziplock bags for transportation to the University
119	of Birmingham for analysis.
120	Soil samples were also collected both upwind and downwind of the main landfill
121	body, in the same area as the air samples were taken. These samples were taken
122	within the boundaries of the landfill operational facility, but with care taken to avoid
123	sampling soil used as capping on landfill cells to ensure (i) that soil samples were not
124	collected from soil placed after landfill operations ceased, and (ii) that farming
125	activities would not influence concentrations of target compounds found.
126	Nine sub-samples of soil (100 g each) were taken from each downwind and upwind
127	sampling location, in a "W" formation, with each sub-sample 10 m apart, using a
128	small pre-cleaned stainless steel shovel. For each 100 g sub-sample, the overlaying
129	foliage was removed from a circular area with a diameter of 10-15 cm. A cylinder of
130	soil, 10-15 cm wide and \sim 5 cm deep was loosened and homogenized, and a \sim 100 g
131	sub-sample of soil removed. This was repeated for each of the nine sub-sampling
132	points before each sub-sample was homogenised into a single sample in an amber
133	glass jar, sealed and then transported to the University of Birmingham. On receipt in

134 Birmingham, samples were air-dried in a fume hood at ambient temperature and 135 homogenised further before an aliquot was removed for analysis. 136 137 Groundwater - Groundwater samples were collected from each of the ten sites from 138 two sampling points down-gradient from the main body of the landfills (per landfill 139 zoning maps information). Each sampling point consisted of an approximately 5 cm 140 wide borehole leading down to water reservoirs at a minimum depth of 5 m below 141 ground level. A number of such sampling points are present at every Irish landfill to 142 facilitate quarterly monitoring of water quality by the Irish Environmental Protection 143 Agency. Samples were manually pumped and collected using clean polyethylene 144 tubes and a foot-valve (allowing for the one way flow of water through the tube). 145 Approximately 25 L of water were purged from each of the two sampling points per 146 landfill, before a 500 mL sample was collected in a pre-cleaned polystyrene bottle. 147 The two samples taken per landfill were then combined into a single 1 L pre-cleaned 148 polystyrene bottle, with this combined sample used for analysis. Following transfer to 149 the laboratory, each combined sample was split, with one half analysed for BFRs 150 (passed through a glass fibre filter (GFF) and PUF before extraction) and the other for 151 PFASs (subjected to solid phase extraction (SPE)). 152 153 Target analytes – The following BFRs and PFASs were targeted in this study: PBDEs 28, 47, 99, 100, 153, 154, 183, and 209; α -, β - and γ -HBCDD, DBDPE and PFOS, 154

perfluorobutane sulfonate (PFBS), PFHxS, PFOA, perfluorononanoic acid (PFNA),

perfluoro-1-octanesulfonamide (FOSA), N-methylperfluoro-1-octanesulfonamide

(MeFOSA), N-ethylperfluoro-1-octanesulfonamide (EtFOSA), 2-(N-methylperfluoro-

155

156

- 158 1-octanesulfonamido)-ethanol (MeFOSE), and 2-(N-ethylperfluoro-1-159 octanesulfonamido)-ethanol (EtFOSE). 160 161 Sample Extraction 162 Air Samples - For air samples, extraction and extract purification was carried out in 163 line with previously published procedures used to determine concentrations of the 164 same target BFRs and PFASs in Irish indoor air (Wemken et al, 2019; Harrad et al, 165 2019). Briefly, PUFs and GFFs were packed into 66 mL extraction cells and spiked with known masses of internal standards (BDE-77, BDE-128, ¹³C₁₂-BDE-209, ¹³C₁₂-166 α -HBCDD, 13 C₁₂- β -HBCDD, 13 C₁₂- γ -HBCDD, M8PFOS, M8PFOA, M8FOSA, 167 168 MPFHxS, MPFNA, d-N-MeFOSA, and d-N-EtFOSA (Wellington Laboratories, Canada)). Where possible, native PFASs and BFRs were quantified relative to the 169 170 corresponding isotopically-labeled internal standard. Where this was not possible, 171 BDEs- 28, 47, 99, and 100 were quantified against BDE-77, BDEs- 153, 154, and 183 where quantified relative to BDE-128, while DBDPE was quantified using ¹³C₁₂-172 BDE-209. For PFASs, PFBS, EtFOSA, and EtFOSE were quantified relative to 173 174 MPFHxS, d-N-MeFOSA, and d-N-MeFOSE respectively. Each cell underwent
- Dionex ASE 350. The BFR fraction was obtained using hexane:DCM (3:1, v/v ratio) as the extraction solvent, while the PFAS fraction was obtained with acetone. Both

sequential extraction to produce a BFR fraction followed by a PFAS fraction using a

were extracted at 90 °C with 5 minutes static time, 3 static cycles and 40% flush

volume. Crude extracts were concentrated to ~0.5 mL in hexane using a Turbovap II

180 (Biotage).

175

179

182

181 <u>Groundwater</u> – To determine concentrations of BFRs in groundwater, samples (0.5 L)

were first subjected to filtration via gravity sequentially through a GFF, followed by a

183 PUF cylinder (Harrad et al, 2009). Following filtration and air-drying in a fume hood, 184 the GFF and PUF from each groundwater sample were treated with BFR internal 185 standards and subjected to pressurised liquid extraction as per air samples. In contrast, 186 PFASs were extracted from groundwater samples using solid phase extraction. 187 Briefly, 0.5 L groundwater was spiked with known quantities of PFAS internal 188 standards and passed through an Oasis-WAX cartridge (150 mg/ 6 mL). Cartridges 189 were then dried under vacuum for approximately 30 minutes before target PFASs 190 were eluted with 6 mL methanol (0.1% NH₄OH). 191 <u>Soil</u> – BFRs were extracted from soils following spiking with internal standards, by 192 loading into an ASE cell and extracting as described for air samples. PFASs were 193 extracted from soil samples using ultrasonication. To summarise, 1 g of sample was 194 accurately weighed into a test tube, treated with internal standards and 7 mL of 195 methanol (0.1% NH₄OH) added. Samples were vortexed for 2 minutes followed by 30 196 minutes of ultrasonication at 20 °C. The tubes were centrifuged for 5 minutes at 3500 197 RPM and the supernatant collected into a new tube. The procedure was repeated twice 198 further and the three extracts were combined and concentrated to ~0.5 mL under a 199 gentle stream of nitrogen at 35 °C. 200 201 Clean-up – for BFRs in all matrices, extracts were diluted to ~2 mL in hexane, added 202 to 2 mL >95% concentrated sulfuric acid and vortexed for 2 minutes. The two layers 203 were separated by centrifugation at 3500 RPM for 5 minutes and the supernatant was 204 collected and concentrated to ~0.5 mL. Extracts were further purified by transferring 205 onto a preconditioned (6 mL DCM, 6 mL hexane) ENVI-Florisil SPE cartridge (500 206 mg/3 mL, Sigma Aldrich). Target BFRs were eluted with 6 mL hexane followed by 6

207	mL DCM. Clean extracts were concentrated to incipient dryness and reconstituted in
208	$200\;\mu\text{L}$ in toluene and transferred to autosampler vials ready for analysis.
209	
210	For PFASs, extracts were purified by loading onto an ENVI-Carb (3 mL/250 mg,
211	Sigma Aldrich) SPE cartridge. PFASs were eluted with 2 mL of methanol (0.1%
212	NH ₄ OH) and concentrated to ${\sim}0.5$ mL. Extracts were passed through a 0.22 μm
213	polyether sulfone (PES) syringe filter, concentrated to 200 μL and transferred to
214	autosampler vials ready for analysis.
215	
216	Instrumental Analysis
217	DBDPE and PBDEs -28, -47, -99, -100, -153, -154, -183, and -209 were analysed on
218	a Thermo TRACE 1310 GC coupled to a Thermo ISQ MS as described in Abdallah et
219	al. 2017). HBCDDs (α -, β - and γ -) were analysed on an LC-MS/MS system composed
220	of a Shimadzu LC-20AB Prominence liquid chromatograph coupled to an ABS Sciex
221	API 2000 triple quadrupole mass spectrometer operated in negative ion mode. Full
222	details are provided elsewhere (Abdallah et al. 2008).
223	
224	PFASs (PFOA, PFOS, PFNA, PFHxS, PFBS, FOSA, EtFOSA, MeFOSA, EtFOSE,
225	MeFOSE) were analysed in accordance with the procedures described in Harrad et al
226	(2019a; 2019b) on a Sciex Exion HPLC coupled to a Sciex 5600+ triple TOF MS.
227	Ten microliters of extract were injected onto a Raptor C18 column (1.8 μm particle
228	size, 50 mm length, 2.1 mm internal diameter, Restek). Full details of the method
229	including acquisition parameters and HPLC conditions are provided in the supporting
230	information.
231	

232 Determination of total organic carbon (TOC) in soil samples

TOC was determined in the soil gravimetrically by measuring the loss on ignition after heating the dried samples at 500 °C for four hours. A clean crucible was accurately weighed and the weight recorded (W1). One gram of the soil sample was added to the crucible before being placed in an oven at 105 °C for 24 hours to remove all moisture and the weight recorded again (W2). The crucible was then placed in an oven at 500 °C for 4 hours and left to cool before the weight of the crucible was recorded a final time (W3). TOC was then determined using the following equation:

$$TOC (\%) = \frac{(W2 - W3)}{(W2 - W1)} \times 100$$

241 Quality Control

All samples were processed using procedures that have been previously validated (Harrad et al., 2009, Gallen et al. 2016, Drage et al. 2016). For ongoing accuracy of analysis of BFRs an aliquot of SRM-2585 (NIST) was analysed with every batch of samples (n=4). For PFAS, SRM-2585 was also used for soil and air samples (n=4) while a MilliQ sample spiked with target compounds was also analysed with each batch of groundwater samples (n=2). All target analytes were found to be within 80-120 % of their certified or spiked values, with less than 15% relative standard deviation. Further information can be found in the Supporting Information.

Two field blanks comprising distilled deionised water sampled from the same polystyrene groundwater collection bottles were analysed in identical fashion to groundwater samples. For air samples, clean PUFs were used as field blanks (n = 2), while for soil, clean anhydrous sodium sulfate (n = 2) was the field blank. Concentrations of all target compounds (except for BDE-209) were below detection

256	limits in all blanks. Therefore with the exception of BDE-209, the method limits of
257	detection were based on the compound mass generating a signal to noise ratio of 3:1.
258	While BDE-209 was detected in all blank samples, this was only at an average of 7
259	pg/sample (± 1 pg/sample), which gave a signal to noise (S/N) ratio of 3:1. Therefore,
260	method limits of quantification for all target compounds were estimated based on a
261	S/N ratio of 10:1.
262	
263	Statistical analysis
264	Descriptive statistics, t-tests and Pearson's correlation analyses were calculated using
265	Excel for Mac v.16.28, while Wilcoxon Signed Rank tests were conducted using
266	SPSS for Mac v.25.
267	
268	RESULTS AND DISCUSSION
268 269	RESULTS AND DISCUSSION Concentrations of PBDEs, HBCDDs, and PFASs in Landfill-Related Air and Soil
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269 270 271	Concentrations of PBDEs, HBCDDs, and PFASs in Landfill-Related Air and Soil Samples Background information about the landfills in this study is provided in Table SI-1. In
269 270 271 272	Concentrations of PBDEs, HBCDDs, and PFASs in Landfill-Related Air and Soil Samples Background information about the landfills in this study is provided in Table SI-1. In summary, all of the landfills in this study primarily accepted municipal solid waste,
269270271272273	Concentrations of PBDEs, HBCDDs, and PFASs in Landfill-Related Air and Soil Samples Background information about the landfills in this study is provided in Table SI-1. In summary, all of the landfills in this study primarily accepted municipal solid waste, along with some biomedical, non hazardous industrial, and construction and
269270271272273274	Concentrations of PBDEs, HBCDDs, and PFASs in Landfill-Related Air and Soil Samples Background information about the landfills in this study is provided in Table SI-1. In summary, all of the landfills in this study primarily accepted municipal solid waste, along with some biomedical, non hazardous industrial, and construction and demolition waste. None of the landfills are located near any potential industrial
269 270 271 272 273 274 275	Concentrations of PBDEs, HBCDDs, and PFASs in Landfill-Related Air and Soil Samples Background information about the landfills in this study is provided in Table SI-1. In summary, all of the landfills in this study primarily accepted municipal solid waste, along with some biomedical, non hazardous industrial, and construction and demolition waste. None of the landfills are located near any potential industrial sources of our target compounds. A full list of concentrations of all target PBDEs,

temperatures are lower. It is plausible that sampling in warmer periods of the year

280	would lead to higher concentrations in air due to enhanced volatilisation of our target
281	compounds. Further monitoring in spring and summer months would thus be
282	advisable to verify whether this is the case.
283	$\it Air$ - Table 1 summarises concentrations of BDE-47, BDE-209, Σ HBCDD, and
284	selected PFASs detected in samples of air from locations both downwind and upwind
285	of the landfills in this study, alongside those from other relevant studies. DBDPE was
286	only detected above its LOQ of 0.9 pg m ⁻³ in one air sample taken downwind of a
287	landfill (2 pg m ⁻³). To our knowledge, these are the first reported data on
288	concentrations of PFASs in Irish outdoor air, and represent a very substantial addition
289	to the measurements of PBDEs and HBCDD made at the background Malin Head site
290	as part of the GAPS study (Lee et al, 2016; Rauert et al, 2018). Comparison of our
291	data with those for Malin Head suggest concentrations of BDE-209 and HBCDD
292	around the landfills studied here exceed slightly those at Malin Head. Concentrations
293	of PBDEs, DBDPE, and HBCDD in the vicinity of Irish landfills are well below those
294	reported recently for indoor air in Ireland (Wemken et al, 2019). For BDE-47,
295	concentrations in the current study are lower than those reported previously in the
296	UK, Sweden, Germany, and the USA (Drage et al, 2016; Hoh and Hites, 2005;
297	Newton et al, 2015; Weinberg et al, 2011). With respect to BDE-209, our
298	concentrations exceed those reported for Sweden (Newton et al, 2015), but are
299	markedly lower than those detected in the UK and the USA (Drage et al, 2016; Hoh
300	and Hites, 2005). Meanwhile, concentrations of HBCDD in this study are lower than
301	in the UK (Drage et al, 2016), Sweden (Newton et al, 2015), and the USA (Hoh et al,
302	2005).
303	Concentrations of all our target PFASs in this study are substantially lower than those
304	reported recently for indoor air in Ireland (Harrad et al, 2019). PFOA was detected at

higher concentrations in this study than in the UK (Goosey and Harrad, 2012), at both landfill and reference locations in Germany (Weinberg et al, 2011), and in Canada (Shoeib et al, 2011). This is consistent with our recent report that concentrations of PFOA in Irish indoor air exceed those reported elsewhere (Harrad et al, 2019b) and combined, these observations suggest extensive use of PFOA in Ireland. In contrast, concentrations of PFOS and PFHxS in this study are lower than those reported in outdoor air at a number of UK locations (Goosey and Harrad, 2012). Moreover, concentrations of PFOS, PFNA, PFBS, and PFHxS in this study are all within a similar range to those reported at landfill and reference (i.e. non-landfill-impacted) locations in Germany (Weinberg et al, 2011).

Soil – Concentrations of HBCDD, selected PBDEs and PFASs in soil from locations both down- and up-wind of Irish landfills are summarised in Table 2. Similar to air, DBDPE was detected above its LOQ of 250 pg g⁻¹ (dry weight) in only one sample taken downwind of a landfill (540 pg g⁻¹ dry weight). To the best of our knowledge, our study is the first to report concentrations of our target contaminants in Irish soil and may be the first report of concentrations in soil anywhere for PFBS. To aid comparison with relevant previous studies for PBDEs and HBCDD, data are provided both on a soil dry weight basis as well as normalised to soil organic matter content. Compared to landfill-impacted locations in northern Canada (Danon-Schaffer, 2010), dry weight concentrations in the current study of BDEs- 47, 99, 153, 183, and 209 are approximately one to two orders of magnitude lower. Instead, concentrations in this study resemble more closely those reported by Danon-Schaffer (2010) for soil from reference (non-landfill-impacted) locations in northern Canada. Other pertinent comparisons for PBDEs and HBCDD are with organic matter-normalised

330	concentrations reported for various locations along rural-urban transects in both the
331	UK (Drage et al, 2016) and Sweden (Newton et al, 2015). Overall, concentrations in
332	our study are broadly consistent with those in these earlier surveys.
333	With respect to PFASs, consistent with our observations for air, dry weight
334	concentrations of PFOA in our study are at the high end of the range previously
335	reported for various locations in Europe and North America (Goosey, 2010; Groffen
336	et al, 2019; Rankin et al, 2016). In contrast, concentrations of PFOS, PFNA, and
337	PFHxS in our soils are at the low end of the range of concentrations previously
338	reported. While our data appear the first report of PFBS in soil from non-industrial
339	locations, our concentrations are around 3 orders of magnitude below those reported
340	in the vicinity of a fluorochemical industrial park in China (Bao et al, 2019).
341	Groundwater - Table 3 summarises the concentrations of HBCDD, selected PBDEs,
342	and PFASs detected in samples of groundwater in this study, with data from
343	comparable studies elsewhere provided for reference. As far as we can ascertain, our
344	data are the first anywhere to report concentrations of both HBCDD and DBDPE in
345	groundwater. We detected DBDPE in all 10 groundwater samples analysed at median
346	and arithmetic mean concentrations that exceeded those of any of the other
347	contaminants measured in this study. While we did not measure DBDPE in a previous
348	study of leachate from Irish landfills (including some of those monitored here), our
349	observations in groundwater are consistent with our recent report of elevated
350	concentrations of DBDPE in indoor air and dust in Ireland (Wemken et al, 2019).
351	This suggests strongly that use of DBDPE is more substantial than thought hitherto,
352	likely as a "drop-in" replacement for the Deca-BDE formulation.
353	As expected, PBDE concentrations in our groundwater samples were lower than those
354	reported previously to be present in leachate from Irish landfills (Harrad et al, 2019a).

Moreover, our concentrations of PBDEs are lower than those reported previously for
groundwater from Canada and Taiwan (Levison et al, 2012; Trinh et al, 2019)
Turning to PFASs, our concentrations in groundwater are well below those reported
in Irish landfill leachate (Harrad et al, 2019a), but fall within a similar range to those
reported for Irish tapwater (Harrad et al, 2019b). Consistent with the predominance of
PFOA in air and soil observed in this study, PFOA is the dominant PFAS of those
targeted here, being the only PFAS detected in all samples and at concentrations
exceeding those of other PFASs. Compared to other studies of PFASs in groundwater
the concentrations of PFOS, PFBS, and PFHxS reported here for Ireland are all at the
low end of previous reports for Japan and various European locations (Atkinson et al
2008; Loos et al, 2010; Murakami et al, 2009; Weiss et al, 2012). In contrast, while
concentrations in Irish groundwater of PFOA are within the range reported for Japan,
Germany, and the UK (Atkinson et al, 2008; Murakami et al, 2009; Weiss et al,
2012); they appear higher than those measured in well water from the Netherlands
(Eschauzier et al, 2013) and a survey of various EU locations (Loos et al, 2010)
While we are unaware of previous reports of FOSA in European groundwater, the
concentrations we report here are consistent with those measured previously in Japan
(Murakami et al, 2009).
Downwind and upwind concentrations of HBCDD, PBDEs, and PFASs in air
and soil in the vicinity of Irish landfills
Air - Table 1 shows that with the exception of HBCDD, arithmetic mean and median
concentrations of all target compounds were higher at downwind locations than
upwind. To evaluate whether there was a statistically significant increment at sites
downwind of the landfills in this study, we compared downwind and upwind
concentrations of HBCDD, PBDEs and PFASs at our 10 landfills using a non-

380	parametric Wilcoxon signed-rank test. This revealed that downwind and upwind
381	concentrations were statistically indistinguishable (p>0.1).
382	Soil – To evaluate whether there was any statistically significant difference between
383	concentrations (expressed both as dry- and organic carbon-normalised weight) of our
384	target contaminants in soil from the 6 landfills for which we had samples from both
385	downwind versus upwind locations, we employed a non-parametric Wilcoxon signed
386	rank test. This showed no significant difference between downwind and upwind
387	locations (p>0.1).
388	Overall, our findings suggest no discernible impact of the landfills studied here on
389	concentrations of HBCDD, PBDEs, DBDPE, and PFASs in air and soil surrounding
390	these facilities. This is consistent with the data on concentrations of a similar range of
391	contaminants measured in air in the vicinity of landfills in Germany (Weinberg et al,
392	2011) but appears in contrast with the data of Danon-Schaffer (2010) who detected
393	markedly higher concentrations of PBDEs in soil around landfills in northern Canada
394	than in reference locations.
395	Influence of landfill lining status on concentrations of HBCDD, PBDEs, and
396	PFASs levels in groundwater
397	To minimise contamination by landfill leachate of the surrounding environment
398	including groundwater; modern landfills are fitted with an impervious polymeric
399	liner. In contrast, some older landfills have been operated unlined. In addition, some
400	landfills (categorised here as "mixed") that were in operation before the use of such
401	liners, remained open but were retrospectively equipped with new waste cells that
402	were fitted with liners. In our previous study of Irish landfill leachate, we reported
403	that concentrations of some BFRs and PFASs in leachate from lined landfills
404	exceeded those in leachate from unlined landfills – as the liners are designed to retain

such contaminants within the landfill, and prevent their leacning into the environment
(Harrad et al, 2019a). This is supported by the fact that only the unlined (n=1) and
mixed (n=3) landfills in this study, release leachate to groundwater. We therefore
hypothesised that groundwater impacted by unlined landfills may contain relatively
higher concentrations of BFRs and PFASs. To test this hypothesis, we used a t-test to
compare concentrations of our target contaminants in groundwater sampled near the 6
lined landfills with those in groundwater obtained close to the 1 unlined and 3 mixed
landfills in this study. This revealed no significant difference (p>0.1) in
concentrations between groundwater from lined and mixed/unlined landfills.
However, we note our small sample size, that for PFOA p=0.12, and that arithmetic
mean concentrations of PFOA in groundwater from lined and mixed/unlined landfills
were 4.1 and 69 ng/L respectively.
Relationship between concentrations of HBCDD, PBDEs, and PFASs in
groundwater and landfill leachate
We examined our data on BFR and PFAS concentrations in groundwater in this study
for any correlation with those we recorded in leachate from the same 10 landfills in
our earlier study (Harrad et al, 2019a). Note that DBDPE was not measured in
leachate. We observed only one significant correlation i.e. between concentrations of
PFOA in groundwater and leachate (R=0.74; p=0.014). However, this positive
correlation was driven substantially by samples obtained from/near one landfill which

Conclusions

displayed the highest concentrations of PFOA in both leachate (11400 ng/L) and

groundwater (140 ng/L). Thus, when the correlation was examined for log-

transformed concentrations, it was no longer significant (R=0.54; p>0.1).

This study measured concentrations of BFRs and PFASs in air, soil, and groundwater sampled near 10 landfills in the Republic of Ireland. These were used to test the hypothesis that landfills have caused discernible contamination of their surrounding environment. Our data suggest that concentrations of BFRs and PFASs in air and soil are not discernibly impacted by emissions from the landfills studied. However, while our relatively small sample numbers prevent firm conclusions, our data suggest that leachate from landfills in this study may have some influence on concentrations of PFOA in groundwater, especially for unlined landfills. Moreover, concentrations of DBDPE in groundwater are reported for the first time anywhere and exceed those of all other BFRs and PFASs monitored in this study. Overall, our findings suggest that more detailed study of the possible impact of landfill leachate on groundwater concentrations of DBDPE, PFOA and related contaminants would appear prudent.

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Table 1: Summary^{a,b} of Concentrations (pg m⁻³) of Selected BFRs and PFASs in Air from Locations Downwind and Upwind of Irish Landfills and Comparison with Concentrations Reported in Selected Other Studies

Parameter (Source)	BDE-47	BDE-209	ΣHBCDD	PFOA	PFOS	PFNA	PFBS	PFHxS
Range (Downwind, This Study)	<0.1-0.32	<0.5-9.2	<0.05-3.6	4.4-97	0.23-4.3	0.08-0.52	<0.15-1.4	< 0.15-0.79
Median (Downwind, This Study)	0.22	6.0	< 0.05	23	1.6	0.17	0.35	0.23
Arithmetic Mean (Downwind, This Study)	0.20	6.6	0.62	33	1.7	0.23	0.50	0.34
Range (Upwind, This Study)	<0.1-0.4	<0.5-8.3	<0.05-6.1	2.7-59	<0.2-2.7	< 0.08-0.31	<0.15-1.2	<0.15-0.81
Median (Upwind, This Study)	0.20	5.5	0.07	16	0.60	0.13	0.26	0.08
Arithmetic Mean (Upwind, This Study)	0.20	5.9	1.2	21	0.91	0.15	0.34	0.23
Range (Indoor air, homes, Ireland, Wemken et al, 2019; Harrad et al, 2019b)	<0.43-28	<7.5-5500	0.9-2500	<0.3-390	<0.4-210	<0.3-13	<0.4-270	<0.4-4.6
Range (n=4, Malin Head, Ireland, 2005-06, Lee et al, 2016)	-	-	<0.1-4.6	-	-	-	-	-
Range (n=4, Malin Head, Ireland, 2014, Rauert et al, 2018)	<2	<1	< 0.09	-	-	-	-	-
Range of arithmetic means (UK West Midlands, Drage et al, 2016)	1.6-7.9	92-370	64-130	-	-	-	-	-
Range (Birmingham, UK, Goosey and Harrad, 2012)	-	-	-	<1.9-20	<1.0-6.1			< 1.1–30
Range (Stockholm, Sweden, Newton et al, 2015)	0.19-2.4	0.03-0.55	<0.03-0.58	-	-	-	-	-
Range (Germany, landfill locations, Weinberg et al, 2011)	<loq< th=""><th>-</th><th>-</th><th>0.2-1.2</th><th>0.23-1.3</th><th><loq-0.7< th=""><th><loq-0.46< th=""><th><loq-0.4< th=""></loq-0.4<></th></loq-0.46<></th></loq-0.7<></th></loq<>	-	-	0.2-1.2	0.23-1.3	<loq-0.7< th=""><th><loq-0.46< th=""><th><loq-0.4< th=""></loq-0.4<></th></loq-0.46<></th></loq-0.7<>	<loq-0.46< th=""><th><loq-0.4< th=""></loq-0.4<></th></loq-0.46<>	<loq-0.4< th=""></loq-0.4<>
Range (Germany, reference locations, Weinberg et al, 2011)	<loq-20< th=""><th>-</th><th>-</th><th>0.2-1.8</th><th><loq-1.1< th=""><th><loq-0.3< th=""><th><loq-0.5< th=""><th><loq-0.6< th=""></loq-0.6<></th></loq-0.5<></th></loq-0.3<></th></loq-1.1<></th></loq-20<>	-	-	0.2-1.8	<loq-1.1< th=""><th><loq-0.3< th=""><th><loq-0.5< th=""><th><loq-0.6< th=""></loq-0.6<></th></loq-0.5<></th></loq-0.3<></th></loq-1.1<>	<loq-0.3< th=""><th><loq-0.5< th=""><th><loq-0.6< th=""></loq-0.6<></th></loq-0.5<></th></loq-0.3<>	<loq-0.5< th=""><th><loq-0.6< th=""></loq-0.6<></th></loq-0.5<>	<loq-0.6< th=""></loq-0.6<>
Range (Canada, Shoeib et al, 2011)	-	-	-	<0.47-9.2	<0.02	-	-	-
Arithmetic mean (Germany, Dreyer et al, 2009)	-	-	-	0.3	1.3	-	-	-
Range (USA, Hoh and Hites, 2005; Hoh et al, 2005)	2.7-41	0.2-65	0.2-9.6	-	-	-	-	-

^aonly those BFRs/PFASs with detection frequency ≥20 % shown

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^bfor the purposes of calculating descriptive statistics, "not detects" replaced by 0.5 x LOD, where LOD = the detection limit

Table 2: Summary^a of Concentrations (pg g⁻¹ dry weight) of Selected BFRs (pg g⁻¹ organic matter in parentheses) and PFASs in Soil from Locations Downwind (n=9^b) and Upwind (n=7^b) of Irish Landfills and Comparison with Concentrations Reported in Selected Other Studies

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Parameter (Source) **BDE-47 BDE-99 BDE-153 BDE-183 BDE-209** ΣHBCDD **PFOA PFOS PFNA PFBS PFHxS** <13-480 <13-750 430-63000 6.33-320 7.4-440 80-6200 (<26-(<26-(1100-<1.0-7.7 <1.0-2.9 Range (Downwind, This Study) 150-5800 2.4-140 < 0.6-44 (38-4100)(780-44000)(16-5700)1700) 5300) 640000) 5000 Median (Downwind, This Study) 6.6 77 (260) 61 (400) 13 (180) 48 (330) 560 (4500) 470 4.3 0.60 <1.0 (18000)**Arithmetic Mean** 180 12000 1700 100 (840) 81 (380) 150 (950) 4.5 5.9 1100 26 0.77 (Downwind, This Study) (1300)(100000)(13000)<13-7300 <13-940 <130-5700 3.8-52 10-26 (32-<15-1500 Range (Upwind, This Study) (<60-(<78-(<340-130-7800 3.7-2000 2.9-33 <0.6-2.9 <1.0-3.7 (20-420)(<90-10000) 260) 13000) 38000) 53000) Median (Upwind, This Study) 9.3 6.0 1.2 2.3 14 (140) 14 (96) <13 (<60) <13 (<78) 240 (1000) 180 (1200) 610 Arithmetic Mean 140 1100 1000 20 (170) 15 (140) 520 (4400) 1600 300 14 1.1 1.8 (Upwind, This Study) (1900)(5400)(10000)Range (UK West Midlands, 580-2100 550-1600 <43-440 < 56-1600 940-45000 Drage et al, 2016)^c Range (Stockholm, Sweden, 240-1700 200-1100 <33-190 310-31000 380-12000 Newton et al, 2015)^c Arithmetic mean (Canada, 5000 16000 62000 landfill impacted sites, Danon-8300 12000 Schaffer, 2010)

Arithmetic mean (Canada, reference sites, Danon-Schaffer, 2010)	100	54	5	15	1600	-	-	-	-	-	-
Arithmetic mean (UK, Goosey and Harrad, 2012)	-	-	-	-	-	-	1100	560	-	-	14
Range of arithmetic means (Belgium, Groffen et al, 2019)	-	-	-	-	-	-	<130-1500	<670-8900	1	1	-
Range (North America, Rankin et al, 2016)	-	-	-	-	-	-	22-1800	30-2000	15-1100	1	2.0-37
Range (Europe, Rankin et al, 2016)	-	-	-	-	-	-	15-2700	7.1-3100	9.1-220	1	3.3-100
Range (Near Fluorochemical Industrial Park, China, Bao et al, 2019)	-	-	-	-	-	-	1200-6300	<200-1400	<200- 1100	<200- 42,000	<200- 950

^afor the purposes of calculating descriptive statistics, "not detects" replaced by 0.5 x LOD, where LOD = the detection limit

^bsoil samples from 3 upwind and 1 downwind locations destroyed in transit from field to laboratory

^cconcentrations expressed as pg g⁻¹ normalised to soil organic matter content

Table 3: Summary^a Concentrations (ng L⁻¹) of Selected BFRs and PFAS in Groundwater Samples Linked to Irish Landfills and Comparison with Concentrations Reported in Selected Other Studies

Parameter/Source	BDE-47	BDE-99	BDE-209	DBDPE	ΣHBCDD	PFOA	PFOS	PFBS	PFHxS	FOSA
Range (This Study)	0.17-0.57	<0.1-1.4	5.8-26	1.3-630	<0.1-2.4	1.6-96	<0.1-1.3	<0.1-0.22	<0.1-0.28	<0.1-1.0
Median (This Study)	0.28	<0.1	9.0	9.4	0.81	3.7	<0.1	<0.1	<0.1	0.11
Arithmetic Mean	0.31	0.31	12	78	0.91	30	0.21	<0.1	<0.1	0.22
(This Study)										
Detection frequency (%)	100	50	100	100	80	100	20	20	20	70
Range (Landfill leachate, Ireland, Harrad	<0.1-140	<0.3-58	<1.0-220	-	<0.2-43	9.0-11000	<0.1-7400	<0.1-1700	<0.1-2600	<0.2-65
et al, 2019)										
Range (Tapwater, Ireland, Harrad et al, 2019)	-	-	-	-	-	0.04-18	<0.15-0.76	<0.2-15	-	-
Range (Tokyo, Murakami et al, 2009)	-	-	-	-	-	0.47-60	0.3-130	-	-	<0.1-0.49
Range (UK, Atkinson et al, 2008)	-	-	-	-	-	<24-260	<11-160	-	-	-
Range (Germany, Weiss et al, 2012)	-	-	-	-	-	<loq- 160</loq- 	0.02-8400	<loq-100< td=""><td><loq-2400< td=""><td>-</td></loq-2400<></td></loq-100<>	<loq-2400< td=""><td>-</td></loq-2400<>	-
Arithmetic Mean (EU, Loos et al, 2010)	-	-	-	-	-	3.0	4.0	<0.3	1.0	-
Range (Pumping well water, the						0.5.1.2	<0.01	<0.01	<0.22	-
Netherlands, Eschauzier et al, 2013)	-	-	-	-	-	0.5-1.3				
Range (Taiwan, Trinh et al, 2019)	1.1-390	0.2-77	6.9-3100	-	-	-	-	-	-	-
Range (Canada, Levison et al, 2012)	<2-10	<1-7.2	<2-92	-	-	-	-	-	-	_

^aonly those BFRs/PFASs with detection frequency ≥20 % shown

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bfor the purposes of calculating descriptive statistics, "not detects" replaced by 0.5 x LOD, where LOD = the detection limit

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