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5 6	ANTIMICROBIAL COMPOUNDS (TRICLOSAN AND TRICLOCARBAN) IN SEWAGE
7	SLUDGES, AND THEIR PRESENCE IN RUNOFF FOLLOWING LAND APPLICATION
8	
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25	Abstract
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27	The reuse of treated municipal sewage ('biosolids') on land is an effective method to divert
28	waste away from landfill and to use an alternative, low cost method of fertilisation. While

legislation has mainly focused on the control of nutrient and metal application rates to land, 29 other potentially harmful emerging contaminants (ECs) may be present in biosolids. Up to 30 80% of municipal sewage sludge is reused in agriculture in Ireland, which is currently the 31 highest rate of reuse in Europe. However, unlike other countries, no study has been 32 conducted on the presence of ECs across a range of wastewater treatment plants (WWTPs) in 33 34 this country. This study evaluated the concentrations of two ECs in sewage sludge, the 35 antimicrobials triclosan (TCS) and triclocarban (TCC), and their presence in surface runoff following land application in controlled rainfall simulation studies. In 16 WWTPs, 36 concentrations of TCS and TCC were 0.61 and 0.08 μ g g⁻¹, which is at the lower end of 37 concentrations measured in other countries. The concentrations in runoff post land 38 application were also mainly below the limits of detection (90 ng L^{-1} for TCS, 6 ng L^{-1} for 39 TCC), indicating that runoff is not a significant pathway of entry into the environment. 40

41

42 *Keywords:* Triclosan; triclocarban; biosolids; wastewater treatment plants; surface runoff.

43

44 Introduction

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The reuse of treated municipal sewage sludge ("biosolids") in agriculture provides the necessary nutrients and micronutrients essential for plant and crop growth (Latare et al., 2014; Liu et al., 2015). Biosolids may be used as a soil conditioner, improving its physical (e.g. water holding capacity; Cele and Maboeta, 2016) and chemical properties (e.g. soil test phosphorus; Shu et al., 2016). Their use also addresses European Union (EU) policy on sustainability and recycling of resources (COM, 2014a).

There are several issues associated with the reuse of municipal sewage sludge in agriculture 53 54 (Peyton et al., 2016). While many of these are issues of perception (Robinson et al., 2012), 55 there is considerable concern, which is scientifically based, regarding a number of substances 56 that may be present in biosolids. There are concerns regarding pharmaceutical and personal care products (PPCPs), antimicrobial compounds, and other endocrine-disrupting compounds 57 58 and synthetic compounds in biosolids (Clarke and Cummins, 2014) and the associated risk of contamination of soil, and surface and groundwater (Hanief et al., 2015; Fu et al., 2016). 59 Toxic metals in sludge may accumulate in the soil and crops and enter the food chain 60 61 following continuous applications to land (Stietiya and Wang, 2011; Latare et al., 2014; García-Santiago et al., 2016). Organic and inorganic contaminants may be lost along surface 62 runoff and leaching pathways following land application (Gottschall et al., 2012; Peyton et 63 al., 2016). Furthermore, there is a risk of emission and transport of bioaerosols containing 64 manure pathogens following land application of biosolids (Brooks et al., 2005; Jahne et al., 65 66 2015). These concerns are confounded by the fact that although EU legislation controls the application of biosolids to land by setting limit values for nutrients and metals (EEC, 1986), 67 no safety guidelines currently exist for PPCPs or many emerging contaminants (ECs). 68

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Wastewater treatment plants (WWTPs) cannot fully remove PPCPs or other organic or 70 synthetic compounds from wastewater, the removal of which is affected by treatment 71 technique and operating conditions (Narumiya et al., 2013). Removal pathways include 72 sorption onto sludge (Ternes et al., 2004) and biodegradation/biotransformation (Verlicchi et 73 74 al., 2012). Despite this, several compounds have been measured in digested sewage sludge (Walters et al., 2010; Verlicchi and Zambello, 2015). Therefore, when biosolids are spread on 75 land, there is a risk of indirect exposure to humans through several pathways, including the 76 food chain (consumption of crops, meat, dairy products and drinking water), surface runoff, 77

and leaching to land drainage systems or groundwater used for abstraction by water treatment 78 plants. Clarke et al. (2016) developed a quantitative risk ranking model for human exposure 79 to 16 organic contaminants following biosolids application to land. They found that while 80 nonylphenols had the highest risk, the antimicrobials, triclosan (TCS) and triclocarban 81 (TCC), were considered more of an evolving risk, as these contaminants are emerging and 82 83 have only recently been restricted within the US (US-FDA, 2015) and EU (COM, 2014b). In addition, both compounds are commonly the most abundant contaminants in biosolids 84 (McClellan and Halden, 2010) and both are listed in the top contaminants of concern 85 worldwide (von der Ohe, 2012; Verlicchi and Zambello, 2015). Triclosan, a broadspectrum 86 bacteriostat and fungicide, and TCC, a fungicide and bacteriostat, are known toxins for 87 humans and have been linked to inhibition of muscle function (Cherednichenko et al., 2012), 88 resistance to antibiotics used in human medicine (Yazdankhah et al., 2006), and ecotoxicity 89 in the environment such as the inhibition and killing of algae, crustaceans and fish (Chalew 90 91 and Halden, 2009). On account of this, these compounds are the main focus of the current study. 92

93

94 In the EU there are considerable differences in national policy regarding the reuse of biosolids in agriculture. In some countries, such as Belgium (Brussels and Flanders), 95 Switzerland and Romania, the reuse of biosolids in agriculture is prohibited (Milieu et al., 96 2013a,b,c), whereas in other countries, such as Ireland, up to 80% of municipal wastewater 97 sludge is reused in agriculture (EPA, 2014; Eurostat, 2016). However, despite this, as the 98 country with the greatest reuse of biosolids on land, no study has examined the 99 concentrations of TCS or TCC in biosolids from WWTPs in Ireland. Such national studies of 100 TCS and TCC have been conducted in the USA, Canada, India and South Korea (Table 1), 101 but currently no extensive study across a range of WWTPs exists in the EU. 102

Once applied to land, TCS and TCC in biosolids may either accumulate in plants (Mathews et 104 105 al., 2014); accumulate, biodegrade or biotransform in soil (Wu et al., 2009), or be released in surface runoff during rainfall-runoff or leaching events (Sabourin et al., 2009). The potential 106 107 for loss via surface runoff or leaching depends on their availability in soil, which is a function of their persistence or half-life (Fu et al., 2016). It has been speculated that the persistence of 108 109 TCS or TCC in the soil may be enhanced by the organic content of the soil (Fu et al., 2016), soil temperature (which is positively correlated to half-life), the physicochemical properties 110 111 of the compounds (Wu et al., 2009), and the presence of co-contaminants (Walters et al., 2010), making them potentially more available for loss in surface runoff during rainfall 112 events. Many studies have investigated losses of TCS and TCC in surface runoff from 113 agricultural lands (Table 2), but few, if any, studies have investigated the surface losses from 114 lands which have received sludge applications from the same WWTP having undergone 115 different treatments. Such an experiment may allow the potential for surface water 116 contamination from different sludge treatment methods to be evaluated. 117

118

Therefore, the aim of this study was to (1) characterise, for the first time, the TCS and TCC in biosolids from a range of WWTPs in Ireland, and (2) measure the surface runoff of TCS and TCC under successive rainfall simulations at 1, 2 and 15 days after application of two types of biosolids, originating from the same WWTP.

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124 **2. Methodology**

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126 2.1 WWTP identification and sample collection

In January and February 2015 (Winter in Ireland), biosolids were collected from 16 WWTPs, 128 which had population equivalents (PEs, i.e. the amount of oxygen demanding substances in 129 wastewater equivalent to the demand of the wastewater produced by a single person) ranging 130 from 2.3 million to 6,500. Details of the PE and influent wastewater characteristics of each 131 WWTP are given in Healy et al. (2016a). Most WWTPs received quantities of landfill 132 leachate in low quantities (less than 2% of the influent biochemical oxygen demand (BOD) 133 load), whilst others received industrial, commercial and domestic septic tank sludge 134 comprising up to 30% of the influent BOD load. Anaerobic digestion of sewage sludge was 135 carried out in five WWTPs, thermal drying in eight WWTPs, and lime stabilisation in four 136 WWTPs (one WWTP carried out both anaerobic digestion and thermal drying). Discrete 137 samples (n=8) of biosolids were collected in clean LDPE containers from each WWTP, and 138 were pulverised in an agate ball mill (FritschTM Pulverisette 6 Panetary Mono Mill) with a 139 rotational speed of 500 rpm for 5 min (repeated three times). The metal content of the 140 141 biosolids are reported in Healy et al. (2016a).

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143 2.2 Field study site description and runoff simulations

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Treated municipal sewage sludge from the WWTP in which anaerobic digestion and thermal 145 drying was carried out, was used in this study. Raw, untreated sludge from the same WWTP 146 was modified by the authors with calcium oxide following the method outlined by Fehily 147 Timoney and Company (1999). Therefore, the anaerobically digested (AD), thermally dried 148 (TD) and lime stabilised (LS) biosolids used in this study originated from the same WWTP. 149 The biosolids were applied to replicated (n=3), hydraulically isolated, field-scale micro-plots, 150 each measuring 0.4 m-wide by 0.9 m-long. The slope of each micro-plot ranged from 2.9 to 151 3.7 % and each micro-plot was instrumented with a runoff collection channel, which allowed 152

all surface runoff to be collected over the duration of a rainfall event (Peyton et al., 2016).
The site was planted with ryegrass for over twenty years and the soil pH ranged from 5.9 to 6.
The soil in all micro-plots was classified as loam and the soil organic matter ranged from 8.1
to 9.0%. Full classification of the plots is detailed in Peyton et al. (2016).

157

Anaerobically digested, TD and LS biosolids were applied by hand to the surface of each 158 micro-plot at the maximum legal application rate in Ireland (Statutory Instrument 610 of 159 2010), which is currently 40 kg phosphorus (P) ha⁻¹. As the P content and the dry matter 160 (DM) of the biosolids varied, the application rates were equivalent to 2.6 tonnes DM ha⁻¹ for 161 TD biosolids (97 g per plot), 6.7 tonnes DM ha⁻¹ for AD biosolids (242 g per plot), and 29.5 162 tonnes DM ha⁻¹ for LS biosolids (1063 g per plot). The biosolids used in this experiment were 163 natural and were not spiked with either triclosan or triclocarban. As the experiment was 164 designed to compare the surface runoff from plots amended with three types of biosolids, no 165 study control (grass only) plots were used in this experiment. 166

167

A rainfall simulator was used to apply rainfall to each micro-plot at intensity of 168 approximately 11 mm h⁻¹ (corresponding to a 3 year return period in Ireland; Met Éireann, 169 2016) in time intervals of 1, 2 and 15 d after the time of biosolids application. Each rainfall 170 simulation lasted 30 min from the time of first occurrence of surface runoff. Runoff water 171 samples were collected in solvent washed amber glass Pyrex® bottles with PTFE lined 172 lids and upon returning the laboratory, 4 mol L^{-1} of sulphuric acid (H₂SO₄) was added to 173 174 adjust the water to pH 3 to prevent biodegradation by microorganisms. Samples were then stored and frozen at - 20°C until analysis. 175

176

177 2.3 Analysis of triclosan and triclocarban in biosolids and surface runoff

The method of analysis for TCS and TCC in the biosolids and surface runoff was conducted in accordance with USEPA Method 1694 (USEPA, 2007) for the determination of pharmaceuticals and personal care products in water, soil, sediment, and biosolids by liquid chromatography tandem-mass spectrometry (HPLC-MS/MS) analyses (Mc Clellan and Halden, 2010; Walters et al., 2010; Klosterhaus et al., 2013).

LC-MS-MS analyses was performed using a Waters Xevo[™] TQ MS triple-quadrupole mass spectrometer (Waters Corp., Milford, MA, USA) coupled with a Waters Acquity UPLC[™] system (Waters Corp., Milford, MA, USA) using reverse phase chromatography. Experimentation was based upon the conditions specified in the USEPA Method 1694, and summarised in Table S1 of the Supplementary Material. Details on the methodology employed may also be found in the Supplementary Material.

191 **3. Results and Discussion**

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193 3.1 Triclosan and triclocarban content of biosolids from WWTPs

194

The TCS and TCC concentrations in the biosolids samples are shown in the Figure 1. The concentrations of compounds in sewage sludge are dependent on the influent wastewater characteristics, and up to 50% of the mass of TCS entering a WWTP may remain in the sludge (Heidler and Halden, 2007). Studies have examined the effectiveness of various sludge treatment methods (e.g. Carballa et al., 2007), but there is no indication that one form of treatment is more efficient in TCS or TCC removal.

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One of the WWTPs examined (1 and 2 in Figure 1) had a history of high concentrations of 202 TCS, with concentrations of 25 μ g g⁻¹ previously being reported (EPA, 2009), but the 203 concentrations in this study were below this. Of the previous studies that have carried out 204 testing of TCS and TCC across a number of WWTPs in a given region, the concentrations of 205 both parameters measured in this study ranked the lowest (Table 1). This may be due to the 206 fact that TCS has been restricted in the EU since 2014 (COM, 2014b), whereas its use in 207 'consumer antiseptic wash products' will only be restricted in the USA from September 2017 208 (US-FDA, 2016). The use of TCC has been phased out by pharmaceutical companies in the 209 last number of years (Westervelt, 2014). The biosolids samples used in this study were 210 collected in January and February of 2015, so there may be seasonal variation in the 211 concentrations of TCS and TCC as have been measured in other studies (Martin et al., 2012). 212

213

214 3.2 Triclosan and triclocarban content of surface runoff

215

The surface runoff concentrations of TCS and TCC were below the LOD in all cases, with 216 the exception of TD biosolids at 15 days (0.01 μ g L⁻¹) and LS biosolids (0.02 μ g L⁻¹) one day 217 after application (Table 3). These TCS and TCC concentrations in the surface runoff were 218 lower than values observed in similar studies, and below the concentrations at which biota are 219 considered likely to be potentially impacted (120 ng L^{-1} for TCS, Wilson et al. (2003); 101 ng 220 L⁻¹ for TCC, McClellan and Halden (2010)). The low concentrations in surface runoff may 221 have been a function of the low TCS and TCC concentrations in the biosolids applied to land 222 223 relative to similar studies (Table 2), but more likely were either due to their degradation or transformation to other compounds, or due to the soil characteristics at the study site. Less 224 than 0.5% of the mass of TCS and TCC applied to each plot was lost in each rainfall event 225 (Table 3). Similar results (expressed as a % of mass released versus mass of compound 226

applied) were obtained in a runoff study by Sabourin et al. (2009), who speculated that they 227 remained sequestered in the soil or were leached to groundwater. The relationship between 228 persistence of organic or synthetic compounds and the composition and physico-chemical 229 properties of soil is well established in the literature (Verlicchi and Zambello, 2015). As 230 reported in other studies (Wu et al., 2009), the high soil organic matter content in the micro-231 plots of the current study (8.1 to 9.0%) may have adsorbed some of the TCS and TCC. Unlike 232 other studies which had durations ranging from 46 days (Wu et al., 2009) to 60 days (Fu et 233 al., 2016), it was impossible to determine if the addition of the biosolids altered the soil's 234 physicochemical properties, as soil analysis was only conducted before the experiment 235 commenced and at the end of the experiment (15 days after application of biosolids) (Peyton 236 et al., 2016), which is too short a period to determine if such changes occur. 237

238

Another factor influencing the persistence of organic and synthetic compounds is the manner in which they are applied to the soil. Al-Rajab et al. (2015) reported that dissipation of PPCPs and antimicrobials, including TCS and TCC, increase over a duration of several months if they are placed below the soil surface (i.e. through ground injection systems). Therefore, the surface runoff of TCS and TCC in the current study may have been further reduced if the biosolids were applied in the soil subsurface.

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246 3.3 Impact of the experiment results on reuse of biosolids in agriculture

247

The current study forms part of a larger study in which the metal content of biosolids from a selection of WWTPs were evaluated (Healy et al., 2016a), and the loss of nutrients, metals and pathogens in surface runoff (Peyton et al., 2016) and the uptake of metals by ryegrass following their land application was quantified (Healy et al., 2016b). For the water quality parameters analysed in these studies, the environmental impacts on receiving waters arising
from the landspreading of biosolids are no different to those arising from the landspreading
of, for example, dairy cattle slurry (Brennan et al., 2012). It was also found that the metal
content of ryegrass in biosolids-amended plots and control plots were similar (Healy et al.,
2016b).

257

While the metal, TCS and TCC contents of the biosolids in the WWTPs examined in our 258 studies were below the concentrations measured elsewhere, there may be a possibility that 259 260 this may increase from one season to the next. In addition, until threshold values, based on human or ecological risk, are set, there is no clear basis for determining if the concentrations 261 of TCS and TCC, as measured in the current study, are safe. Furthermore, the current study 262 only examined two types of antimicrobials, which is only a small fraction of the total number 263 of contaminants that may be present in biosolids. Parameter testing of this type is extremely 264 265 expensive, so the costs of routinely testing sludge for all possible contaminants would be prohibitive. While most commentators have stated that the risk to human health following 266 dietary intake of organic contaminants from crops grown on biosolids-amended lands is 267 268 minimal (Verslycke et al., 2016), they acknowledge that a certain amount of uncertainty still exists (Smith, 2009; Oun et al., 2014). On account of these issues, and public perception 269 issues in particular, many countries have prohibited the use of biosolids in agriculture, and 270 even in those countries that permit their use, local restrictions exist (e.g. Bord Bia, 2013). 271 This conservative ethos has a substantial 'buy-in' from major industries, who use products 272 sourced from agricultural land, as there would be reputational damage to a brand if it 273 emerged that biosolids, which could potentially contain ECs, were used in the production of 274 their feedstock. 275

Although legislation has attempted to address public concern by regulating the amount of 277 biosolids applied to land, there are considerable differences in national policy regarding the 278 279 reuse of biosolids in agriculture. In Europe, the application of biosolids to land is governed by EU Directive 86/278/EEC (EEC, 1986), which is based on the nutrient and metal content 280 281 of the biosolids (although more stringent guidelines are enforced in some member states). In 282 comparison, in the majority of states of the USA, biosolids are applied to land based on the nitrogen requirement of the crop being grown and not on a soil-based test (McDonald and 283 Wall, 2011). It would be impossible to fully regulate the application rates of all potential 284 285 contaminants in biosolids, so while legislation has mainly focused on nutrient and metal content of biosolids, the possibility exists that other potentially harmful, unregulated ECs, for 286 which no international standards exist for reuse in agriculture, may accumulate in the soil 287 upon repeated application. 288

289

4. Conclusions

291

There were low concentrations of triclosan and triclocarban in treated sewage sludge across a number of WWTPs of varying PE. In addition, measurements of triclosan and triclocarban in surface runoff following land application were at or below the limits of detection, which indicates that there appears to be no risk of acute biological effects as a result of runoff.

296

There is a need to fully quantify the concentration of pharmaceuticals and other emerging contaminants in treated municipal wastewater, particularly in countries that recycle a high percentage of sludge produced in municipal sewage treatment plants in agriculture. To fully characterise all existing known, as well as emerging contaminants is cost prohibitive.

301	Therefore, any potential economic and practical gains arising from the recycling of sewage
302	sludge in agriculture need to be considered alongside cost and public health issues.
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Table 1. Triclosan and triclocarban concentrations ($\mu g g^{-1}$ dry weight) in national studies of biosolids produced in municipal wastewater treatment plants.

Reference	Country	# WWTPs	Mean co	oncentration	Maximum concentration		
		examined	$(\mu g g^{-1})$		$(\mu g g^{-1})$		
			Triclosan	Triclocarban	Triclosan	Triclocarban	
McClellan and Halden (2010)	USA	94	12.6	36	19.7	48.1	
Subedi et al. (2015)	India	5	1.2	7.0			
Chu and Metcalfe (2007)	Canada	4	4.2	4.3			
Guerra et al. (2014)	Canada	6	6.8	2.9	11.0	8.9	
Subedi et al. (2014)	S. Korea	40		3.1		6.9	
This study	Ireland	16	0.61	0.08	4.9	0.15	

Table 2. Studies examining the surface losses of triclosan and triclocarban from land applied

620 biosolids.

	Reference	Type of	Compound in biosoli		Application		oncentration in surface		
		sludge used	4) (ug g ⁻¹)	rate (expressed as dry matter)	water (ng L ⁻¹)		detection (days post application)	
			Triclosan	Triclocarban	matter)	Triclosan	Triclocarban		
	Edwards et al. (2009)	Anaerobically digested	14	8	~8 Mt ha ⁻¹	240	<loq< td=""><td></td></loq<>		
	Gottschall et al. (2012)	Anaerobically digested	10.9	4.9	$\sim 22 \text{ t ha}^{-1}$	73	40	22	
	Topp et al. (2008)	Not stated			92,500 L ha ⁻ 1	258		1	
	This study ^a	Anaerobically digested	0.27	<2.4	6.7 t ha ⁻¹	<90	<6	15	
		Thermally dried	4.9	0.05	2.6 t ha ⁻¹	<90	<6	15	
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Table 3. Concentrations of triclosan and triclocarban in applied biosolids to field plots ($\mu g g^{-}$

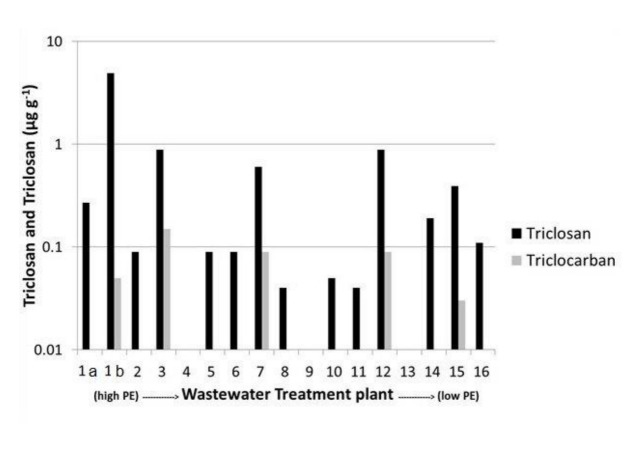
¹; 'Influent') and average concentrations of triclosan and triclocarban in surface runoff (μ g L⁻¹) from field plots. LOD (limit of detection) = 0.09 μ g L⁻¹ (TCS) and 0.006 μ g L⁻¹ (TCC) in this study.

			TT · 1				T · 1	1		
		TCS in	Triclo 1 d	san 2 d	15 d	Т	CC in	Tricloca 1 d	arban 2 d	15 4
		applied	1 u	2 u	15 u		plied	1 u	2 u	15 d
		sludge					udge			
		dose					ose			
		$\mu g g^{-1}$		μg L ⁻¹			μg g ⁻¹		μg L ⁻¹	
	TD	4.9	<lod< td=""><td><lod< td=""><td><lod< td=""><td>0.</td><td>05</td><td><lod< td=""><td><lod< td=""><td>0.01</td></lod<></td></lod<></td></lod<></td></lod<></td></lod<>	<lod< td=""><td><lod< td=""><td>0.</td><td>05</td><td><lod< td=""><td><lod< td=""><td>0.01</td></lod<></td></lod<></td></lod<></td></lod<>	<lod< td=""><td>0.</td><td>05</td><td><lod< td=""><td><lod< td=""><td>0.01</td></lod<></td></lod<></td></lod<>	0.	05	<lod< td=""><td><lod< td=""><td>0.01</td></lod<></td></lod<>	<lod< td=""><td>0.01</td></lod<>	0.01
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Figure 1. Triclosan and triclocarban concentrations (ug g⁻¹) in treated sludge from 16 wastewater treatment plants in Ireland, ranging (numerically in ascending order) from a population equivalent (PE) of 2.3 million to 6,500. Two forms of treatment of sludge are carried out in one WWTP: anaerobic digestion (1a) and thermal drying (1b). WWTPs with no concentrations shown are WWTPs in which triclosan or triclocarban were below the limits of detection (TCS, 0.006 μ g g⁻¹ and TCC, 0.0024 μ g g⁻¹).







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