

Quantifying the dominant sources of sediment in a drained lowland agricultural catchment: The application of a thorium-based particle size correction in sediment fingerprinting

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1	Quantifying the dominant sources of sediment in a drained lowland agricultural catchment: the
2	application of a thorium-based particle size correction in sediment fingerprinting
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34 Abstract

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36 Soil erosion is one of the main factors influencing land degradation and water quality at the global 37 scale. Identifying the main sediment sources is therefore essential for the implementation of 38 appropriate soil erosion mitigation measures. Accordingly, caesium-137 (¹³⁷Cs) concentrations were used to determine the relative contribution of surface and subsurface erosion sources in a lowland 39 40 drained catchment in France. As ¹³⁷Cs concentrations are often dependent on particle size, specific 41 surface area (SSA) and novel Thorium (Th) based particle size corrections were applied. Surface and 42 subsurface samples were collected to characterize the radionuclide properties of potential sources. 43 Sediment samples were collected during one hydrological year and a sediment core was sampled to 44 represent sediment accumulated over a longer temporal period. Additionally, sediment from tile drains 45 was sampled to determine the radionuclide properties of sediment exported from the drainage network. 46 A distribution modeling approach was used to quantify the relative sediment contributions from 47 surface and subsurface sources. The results highlight a substantial enrichment in fine particles and associated ¹³⁷Cs concentrations between the sources and the sediment. The application of both 48 49 correction factors reduced this difference, with the Th correction providing a more accurate 50 comparison of source and sediment samples than the SSA correction. Modelling results clearly indicate 51 the dominance of surface sources during the flood events and in the sediment core. Sediment exported 52 from the drainage network was modelled to originate predominantly from surface sources. This study demonstrates the potential of Th to correct for ¹³⁷Cs particle size enrichment. More importantly, this 53 54 research indicates that drainage networks may significantly increase the connectivity of surface 55 sources to stream networks. Managing sediment transferred through drainage networks may reduce the 56 deleterious effects of suspended sediment loads on riverine systems in similar lowland drained 57 agricultural catchments.

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Key Words: Sediment tracing, Agricultural lowland areas, Cesium-137, Particle size correction, Tile
 drainage networks

62 **Highlights**:

- Source and sediment caesium-137 activities were used to trace sediment sources
- Two particle size corrections were compared to address enrichment in fine particles
- A thorium-based correction out performed a specific surface area correction
- Sediment in this lowland agricultural catchment is supplied by surface sources
- Drainage networks may increase the connectivity of surface sources to rivers
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72 1. Introduction

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74 Soil erosion by water affects a significant proportion (16%) of agricultural land in Europe 75 (Cerdan et al., 2010; Jones et al., 2012). Although natural, this degradation process is accelerated by 76 land use change and anthropogenic pressures in agricultural landscapes (Bakker et al., 2008; Chartin et 77 al., 2011; Sharma et al., 2011). Around the world, the negative effects of soil erosion are characterized 78 by the decline of crop yields, the reduction of soil water storage capacity and decreases in soil organic 79 matter (Berger et al., 2006; Boardman and Poesen, 2006). The main problems associated with 80 accelerated soil erosion are often not only the actual soil loss itself, rather the adverse effects of 81 elevated suspended sediment yields and adsorbed contaminants downstream (Walling et al., 2003).

The excess of fine sediment particles (e.g., <63 μm) associated with soil erosion are detrimental to water quality and stream environments (Kronvang et al., 2003; Owens et al., 2005; Horowitz, 2008). Elevated fine sediment loads can result in numerous problems such as elevated turbidity and reservoir siltation (Wood and Armitage, 1997; Nakamura et al., 2004). Importantly, fine sediments often transport contaminants such as heavy metals, Polychlorinated Biphenyls (PCB), phosphorus, pesticides, pathogens, and fallout radionuclides (Desmet et al., 2012; Ayrault et al., 2014; Evrard et al., 2014).

89 Currently, there is a limited understanding regarding the dominant sediment source in France 90 in particular, and in drained lowland cultivated areas in general (Russell et al., 2001; Walling et al., 91 2002). The implementation of highly productive cropland agriculture practices in these regions has 92 included the installation of drainage networks that induce a high level of connectivity between eroding 93 soils and river systems. Although these land management changes were clearly associated with 94 increased sediment yields (e.g Foster and Walling, 1994; Dearing and Jones, 2003; Ahn et al., 2008), 95 the identification of the dominant sediment sources, their dynamics, and their pathways in drained 96 landscapes with similar intensive agricultural practices remains poorly understood. Indeed, research 97 has highlighted the limited information available regarding sediment exported through drainage 98 networks in these environments (Kronvang et al., 1997; Walling and Collins, 2005; King et al., 2014).

99 To implement appropriate measures to reduce erosion and sediment production (Evrard et al., 100 2008), it is essential to identify the dominant sediment sources and understand their dynamics. 101 Sediment fingerprinting techniques provide a direct method to identify and quantify sediment 102 contributions from different sources (Collins and Walling, 2002). The technique is based on the direct 103 comparison of sediment and source properties (Walling et al., 1993; Collins et al., 2010). A variety of 104 parameters have been used in sediment fingerprinting studies to discriminate between potential 105 sources, including geochemical composition, sediment color, plant pollen, and fallout radionuclides (Brown, 1985; Walling et al., 2002; Martinez-Carreras et al., 2010). In particular, caesium-137 (¹³⁷Cs) 106

discriminates between surface and subsurface soils, regardless of other soil properties including
catchment geology (Wallbrink et al., 1996; Walling, 2005; Caitcheon et al., 2012).

109 ¹³⁷Cs ($t_{1/2}$ = 30 years) is an artificial radionuclide originating from two main sources in Western Europe: thermonuclear weapons testing (1950-1970s) and the Chernobyl accident. Fallout 110 111 from the Fukushima accident in 2011 was shown to be negligible in this region (Evrard et al., 2012). ¹³⁷Cs is quickly and predominantly fixed to fine particles (He and Walling, 1996; Wallbrink and 112 Murray, 1996; Motha et al., 2002). In undisturbed soils, ¹³⁷Cs is concentrated near the soil surface (i.e. 113 114 within the top 10 cm), with concentrations decreasing exponentially with depth (Matisoff et al., 2002). 115 In cultivated soils, ¹³⁷Cs concentrations are homogenized by tillage (He and Walling, 1997; Matisoff et al., 2002; Chartin et al., 2011). Accordingly, sediment generated from subsoil erosion processes, such 116 117 as channel bank erosion, will have low ¹³⁷Cs concentrations, whereas sediment generated from surface soils will have elevated ¹³⁷Cs concentrations (e.g Caitcheon et al., 2012; Olley et al., 2013). Through 118 119 comparing sediment and source ¹³⁷Cs concentrations, it is possible to determine whether sediment is 120 derived from surface or subsurface sources.

121 Soil erosion processes result in the preferential mobilization and transfer of fine sediment 122 particles (He and Walling, 1996). To reduce the impacts of grain size selectivity, many fingerprinting 123 studies isolate either the <10 µm (Olley and Caitcheon, 2000; Wallbrink, 2004) or the <63 µm fraction 124 (Navratil et al., 2012; Pulley et al., 2015) to mitigate differences in the particle size distributions of 125 source soil and sediment. To further reduce the grain size effect when tracing sediment, a correcting 126 factor has been applied (Collins et al., 1996). This correction is based on the specific surface area 127 (SSA) of sediment and source soils, and it relies on the assumption of positive linearity between SSA 128 and tracer property concentrations. Linear particle size correction factors may be useful when there is 129 a narrow range of particle size differences between source soils and sediment (Koiter et al., 2013). 130 This particle size correction has been applied to various tracers, such as radionuclides (He and Owens, 131 1995; He and Walling, 1996), phosphorus (Owens and Walling, 2002) and extractable metals 132 (Horowitz and Elrick, 1987).

133 Although this particle size correction has been widely applied (Collins et al., 2001; Carter et 134 al., 2003), there are acknowledged limitations (Koiter et al., 2013). For example, Smith and Blake 135 (2014) demonstrated that the most critical assumption, which could have large effects on source 136 apportionments, is the hypothesis of positive linearity between particle size and tracer concentration. 137 Contrary to previous studies, Smith and Blake (2014) demonstrated that this relationship does not 138 apply to all tracer properties and that the assumption of linearity must be routinely tested. Russell et al. 139 (2001) also demonstrated that the linear particle size correction relationship was inappropriate when 140 there were large SSA differences between sources.

Given the potential uncertainty associated with the SSA correction, some authors, such as
Martinez-Carreras et al. (2010), do not apply it. As an alternative to the SSA-derived correction,
Sakaguchi et al. (2006) demonstrated the potential of a Thorium (*Th*) normalization to correct for the

grain size effect in lake sediment cores. Owing to its measurement in gamma spectrometry along with 137 Cs, there is a novel utility in the potential for *Th* to correct for grain size differences that could remove the need for simultaneous *SSA* measurements in sediment fingerprinting studies using fallout radionuclides for source discrimination.

148 The objective of this research is to identify the dominant sediment sources and their temporal 149 variation in the Louroux catchment, France; a small agricultural catchment, representative of the 150 lowland drained landscapes of Western Europe. To determine the main source of sediment in the 151 Louroux catchment, first we examine particle size effects and compare the impacts of Th and SSA corrections on ¹³⁷Cs soil and channel bank concentrations. Second, we characterize the ¹³⁷Cs 152 153 concentrations of surface and subsurface sources. Third, we model the relative contribution of these 154 sources to sediment sampled throughout a year, a sediment core collected in a pond at the outlet of the 155 catchment, and sediment sampled within the tile drainage network.

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157 2. Materials and methods

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159 2.1. Study area

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161 The Louroux pond catchment (24 km²) is a headwater agricultural basin located in the central 162 part of the Loire River basin, France (Fig. 1). The catchment is characterized by a flat topography 163 (average slope 0.44%; altitude ranging from 94 to 129 m). The catchment surface is primarily 164 occupied by arable land (78%), followed by pasture (18%) and forest (4%) (European Environment 165 Agency, 2002). The climate is temperate oceanic with a mean annual rainfall of 684 mm (between 166 1971–2000). The bedrock consists of carbonates, detrital and loess deposits (Rasplus et al., 1982). 167 Soils are classified as Epistagnic Luvic Cambisols, which are predominantly hydromorphic and prone 168 to surface crusting (Froger et al., 1994).

This basin, like the majority of the great western agricultural plain in Europe, has undergone a global modernization of agricultural practices and land use changes. Two land consolidation schemes were implemented in the catchment in 1954 and 1992. Stream networks were created or redesigned, and tile drainage networks were installed to drain the hydromorphic soils.

The Louroux pond, located at the catchment outlet (52 ha; Fig. 1), was created during the Middle-Ages (1000 AD). Recent research indicated that a significant increase in soil erosion in the catchment during the last 70 years resulted in the accelerated sedimentation and eutrophication of the pond (Foucher et al., 2015). Over the last decade, the Louroux pond has received an annual average input of 2500 tons of terrigenous material (Foucher et al., 2015).

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179 2.2 Sampling

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181 2.2.1. Soil and stream bank sampling

182 Soil samples from ground surfaces and subsurface material exposed on actively eroding river 183 banks were collected from January 2013 to February 2014. Sampling was restricted to cropland areas, 184 as soil erosion was shown to be negligible under grassland in similar environments of Northwestern 185 Europe (Cerdan et al., 2002; Evrard et al., 2010). In addition, grassland areas in this catchment are 186 ploughed approximately every 10 years, mixing ¹³⁷Cs in the soil profile similarly to croplands.

Surface sources (n = 34) were sampled by scraping the top 2–3 cm layer of soil with a plastic spatula in potentially eroding areas directly connected to the stream network. Each of these surface source composite samples was comprised of five sub-samples collected within an area of 5 m². Subsurface sources (n = 15) were sampled by scraping a 2–3 cm layer of the actively eroding bank face. In addition, sediment was collected in hillslopes during the rain by placing plastic bottles on the hillslope surface to sample sediment during runoff events (n = 3). These plastic bottles were collected after events along with the deposited sediment.

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195 2.2.2. Suspended sediment sampling

196 Hydro-sedimentary parameters were continuously recorded at five automated monitoring sites 197 (S1-S5, Fig. 1). Water discharge was measured using a v-notch weir. Suspended sediment was continuously measured using a Ponsel[®] calibrated turbidity sensor. Twenty-four liters of river water 198 199 were automatically collected according to the water level at each station during five flood events (n =200 21) and once during the low-water period (n = 4) to examine variability throughout the hydrological 201 year in 2013–2014. All individual samples for each station and for each flood were mixed to prepare 202 composite samples. In addition to the river monitoring sites, three stations were installed at tile drain 203 outlets to characterize the properties of the material transiting through the drainage networks and to determine if sediment exported from this network had ¹³⁷Cs concentrations similar to surface or 204 205 subsurface sources (Fig. 1). To characterize the origin of sediment accumulated over a longer temporal 206 period (2003-2013) at the catchment outlet (Fig. 1), the top 10 cm of a 110 cm long core collected in 207 March 2013 was subsampled in 3 cm increments. This sediment core was sampled in the central pond 208 depression, an area that is representative of sediment deposition in the Louroux Pond. More details on 209 the core sampling and fallout radionuclide dating are provided in Foucher et al. (2015).

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211 2.3. Sample treatment and analysis

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Suspended sediment concentrations for the composite samples were determined by weighing after filtration (40 microns acetate filters). Particle size analyses were performed on all suspended sediment samples as well as on randomly selected source samples (n = 18) after removing carbonates 216 and organic material with hydrogen peroxide. Particle size was analysed with laser granulometry 217 (Malvern Mastersizer[®]) characterizing the textural parameters ranging between 0.01 and 3500 µm.

218 Randomly selected subsamples of subsurface (n = 5) and surface sources (n = 5) were sieved 219 to determine radionuclide activities in different particle size fractions. Samples were mechanically 220 sieved to 63 and 50 μ m. The <20 μ m fraction was then isolated by using the settling velocity 221 relationship predicted by Stokes' Law. These analyses were used to calculate the difference between 222 the bulk samples and the sieved samples, and to examine the utility of the two different corrections.

223 Suspended sediments collected during the flood events were flocculated using calcium 224 hydroxide (CaOH₂), to recover and concentrate radionuclides. For the measurement of radionuclides, 225 ~80 g of material was analysed for source samples and ~10 g for sediment samples. Activities were 226 determined by gamma spectrometry using low background coaxial N and P type GeHP detectors 227 (Canberra/Ortec) at the Laboratoire des Sciences du Climat et de l'Environnement (France) (Evrard et 228 al., 2011). Radionuclides activities were decay-corrected to the sampling date. In addition to the radionuclide measurement, Th concentrations expressed in ppm were calculated from ²²⁸Th activity 229 230 concentrations. Only ¹³⁷Cs is modelled in this study as only one tracer is required to discriminate 231 between two sources.

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 - 2.4. Particle size corrections
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235 To limit the bias potentially introduced by particle size difference between sources and 236 sediment, a correcting factor used by Collins et al. (1996) was applied to radionuclide activities. This 237 approach is based on the ratio of SSA of each individual sediment sample to the mean SSA of each 238 source type, multiplied by the mean activity for each source:

240 SSA correcting factor_i =
$$\frac{(SSA_i)}{(SSA_y)} \times T_y$$

241

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242 where SSA_i = specific surface area for each individual sample (*i*), SSA_v = specific surface area for each 243 source type (y), and T_y = mean tracer concentration for each source type (y).

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245 The Th content correction was calculated based on the ratio of the radionuclide concentration 246 normalized with $\ln(Th)$ of each individual sample, to the mean radionuclide concentration normalized 247 with $\ln(Th)$ of each source type, multiplied by the mean radionuclide concentration of each source:

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(1)

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$$\ln(Th) correcting \ factor_i = \frac{T_i / \ln(Th)_i}{T_y / \ln(Th)_y} \times T_y$$
 (2)

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where T_i = the tracer concentration for each individual sample (*i*), $\ln(Th)_i$ = the logarithm of the *Th* concentration (in ppm) for each individual sample (*i*), and $\ln(Th)_y$ = the logarithm of the *Th* concentration (in ppm) for each source type (*y*).

254 The SSA and Th corrections were applied for randomly selected soil (n = 10) and channel bank samples (n = 7) to test the relationship between the corrected and measured ¹³⁷Cs values. These results 255 were then used to calculate the corrected values of 137 Cs with the SSA and Th correction factors of Eqs. 256 257 (1) and (2). These corrected and measured values are first plotted to deduce equations from these 258 relationships. These relations are then used to correct the entire dataset for each source type and for 259 both correction techniques. To check the applicability of these corrections, sediment samples were 260 collected on hillslopes during different hydrological periods and the ¹³⁷Cs activities measured for these 261 samples were compared to these corrected values.

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- 264 2.5. Distribution modelling
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Distribution modelling approaches have been recently applied to sediment tracing research analyzing fallout radionuclides and elemental geochemistry (Caitcheon et al., 2012; Olley et al., 2013; Laceby and Olley, 2015). This modelling approach incorporates distributions throughout the entire modelling framework, including the relative contribution terms (i.e. not only source and in-stream components).

To determine the relative source contribution to in-stream sediment, it is assumed that the sediment samples are derived from a discrete mixture of sources with surface sources contributing xand subsurface sources contributing 1 - x. With the distribution modelling approach, x is modelled as a truncated normal distribution ($0 \le x \le 1$), with a mixture mean (μ_m) and standard deviation (σ_m). Distributions of ¹³⁷Cs activities in surface (A) and subsurface (B) sources are then modelled when determining the relative source contributions (x) to in-stream sediment based on the following equation:

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$$Ax + B(1-x) = C$$

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where *C* is the in-stream sediment distribution. Normal distributions were modelled for source and sediment distributions as Laceby and Olley (2015) demonstrated that they resulted in more accurate modelling results compared to Student's *t* distributions. Sediment collected from the core sample from the Louroux Pond and sediment obtained from the tile drainage network were first individually

(3)

modelled. Second, distributions were modelled for sediment sample groups for different flood eventsand also the different monitoring stations.

The model was optimized with the Optquest algorithm in Oracle's Crystal Ball software. For more details on the modelling approach, see Laceby and Olley (2015). In general, the optimal value of *x* was determined by simultaneously solving Eq. (3) 2500 times with the Optquest algorithm. During this simulation, 2500 Latin hypercube samples were drawn from the source (*A* and *B*) and in-stream sediment (*C*) distributions while solving Eq. (3) by varying *x*, μ_m and σ_m . This simulation was then repeated an additional 2500 times with the median of *x* for these additional simulations being reported as the surface source proportional contribution to sampled sediment.

The model uncertainty for each sources' proportional contribution is calculated by summing the modelled σ_m , with the median absolute deviation (*MAD*) of σ_m , and also the *MAD* of the modelled source proportional contributions. The latter two components of model error are calculated from the additional 2500 simulations. This model uncertainty combines actual σ_m for each source contribution with the *MAD* of this standard deviation and the *MAD* of the actual source contribution (*x*) for the additional 2500 simulations.

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- 302 3. Results
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304 *3.1. Correction of grain size effects*

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306 Particle size analyses highlight differences between the D_{50} (median particle size) of the 307 surface $(32.2 \pm 7 \,\mu\text{m})$ and subsurface sources $(38.5 \pm 8 \,\mu\text{m})$ in comparison to the in-stream sediment 308 $(6.2 \pm 2 \,\mu\text{m})$. A *t*-test between the two sources did not show a significant difference between their D_{50} 309 (*p*-value = 0.14). Significant differences were found between D_{50} of surface sources and in-stream 310 sediment (p = 0.00) and also between subsurface sources and in-stream sediment samples (p = 0.00). 311 In addition, soil samples collected before and during runoff events clearly highlight the particle size 312 enrichment during sediment generation processes, with an average activity measured before the runoff of 3.7 ± 3 Bq kg⁻¹ compared to 17 ± 3 Bq kg⁻¹ during runoff. These samples demonstrate a difference 313 314 of 78% between surface sources and generated sediment, with an enrichment factor of 4.6.

The significant particle size effect noted above may prevent the direct comparison between bulk sources and sediment samples. To test the impact of this effect on radionuclide activities, several source samples were fractionated. The results from five soil samples highlight an enrichment factor of between the bulk and the <20 μ m particle size fraction for the soil samples and the absence of detectable ¹³⁷Cs in the 20-50 μ m fraction for these samples (Fig. 2). There is an 80% difference between the bulk and sediment particle size which can impact modelling results. For the 63 μ m

- 321 fraction, this difference is reduced to 77% compared to 64% for the 20 μ m fraction. Isolating the <20 322 μ m fraction does not allow for a direct comparison between the sources and the suspended sediment. 323 The particle size enrichment of ¹³⁷Cs concentrations is plotted in Fig. 3 demonstrating clearly that 324 sediment samples collected during the flood events are distinct from the source soils. To address this 325 particle size effect and allow for comparison between sources and sediment samples, source activities 326 were corrected with two distinct particle size proxies.
- SSA, Th and ¹³⁷Cs activities measured for soil (n = 10) and channel bank samples (n = 7) were 327 328 plotted to deduce the relationships between the corrected and measured values (Fig. 4). These plots 329 demonstrate that for the SSA correction a strong linear relationship exists between the corrected values and measured ¹³⁷Cs activities in the soil ($r^2 = 0.89$) and channel bank samples ($r^2 = 0.94$) (Fig. 4). The 330 331 Th particle size correction, based on $\ln(Th)$ normalization, also had a strong exponential relationship 332 for the channel bank samples ($r^2 = 0.98$) and surface soil samples ($r^2 = 0.96$). Both corrections have a 333 similar relationship for the subsurface samples. In contrast, the relationship obtained for the surface samples was different. The ¹³⁷Cs activities after the SSA correction ranged between 3.2 and 20 Bg kg⁻¹ 334 whereas the ¹³⁷Cs activities after the *Th* normalization ranged between 1 and 30 Bq kg⁻¹. For both 335 approaches, the mean values are comparable, 9.3 and 10.2 Bq kg⁻¹ for the SSA and Th corrections 336 337 respectively. The difference was the wider range with the *Th* correction.
- 338 As particle size data were available for only part of the dataset, both correcting factors were 339 applied to the whole data set (subsurface sources n = 15 and surface sources n = 34) by using the 340 equations for the surface and subsurface samples shown in Fig. 4. The corrected data are plotted with ²¹⁰Pb_{ex} in Fig. 5. After the application of both corrections, sediment ¹³⁷Cs activities generally plotted 341 within the ¹³⁷Cs activity range of the potential sources. With the SSA correction, five sediment samples 342 343 remain outside the source range (black squares in Fig. 5), whereas with the Th correction only one 344 sample plots outside the source range. Therefore, the Th correction likely provides a more direct 345 comparison with the sediment samples than the SSA correction.
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- 347 *3.2.*¹³⁷*Cs* concentration in source samples
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- Activities in the samples collected from stream banks range between 0 ± 0.3 and 3.1 ± 0.2 Bq kg⁻¹ with an average of 1.2 ± 0.2 Bq kg⁻¹. Soil surface activities range between 1 ± 0.1 and 7.9 ± 0.2 Bq kg⁻¹ with an average of 3.2 ± 0.2 Bq kg⁻¹. These potential sources are statistically different (p = 352 0.00).
- After the application of the *SSA* correction, the subsurface sample activities range between 0.7 and 5.1 Bq kg⁻¹ (mean: 1.7 Bq kg⁻¹). These sample activities, after the *Th* correction, range between 0 and 6.5 Bq kg⁻¹ (mean: 1.8 Bq kg⁻¹). The surface sample concentrations are more variable, ranging between 3.7 and 23.8 Bq kg⁻¹ (mean: 10 Bq kg⁻¹) after the *SSA* correction and between 0.8 and 35.5 Bq

357 kg⁻¹ (mean: 11.2 Bq kg⁻¹) after the *Th* correction. Summary statistics of 137 Cs concentrations are 358 provided in Table 1.

The fitted normal distributions for surface and subsurface sources with both particle size corrections are plotted in Fig. 6. The source samples are plotted in rank order. The source distribution areas overlap only by 25% with the *SSA* correction compared to 62% with the *Th* correction. These distributions are used to model the proportional contribution of the potential sources to suspended sediment, the core samples, and the drainage network samples.

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365 3.3. ¹³⁷Cs concentration in sediments samples

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367 Activities of ¹³⁷Cs for the 23 sediment samples collected over the hydrological year range between 6.1 ±1 and 36.7 ± 4.7 Bq kg⁻¹ with an average of 18.4 ± 3.4 Bq kg⁻¹. The highest ¹³⁷Cs 368 369 activities have been recorded at the Masnier (station 3) and Grand Bray rivers (station 2) with average ¹³⁷Cs activities of 19 ± 4.9 and 22.4 ± 4.3 Bg kg⁻¹, respectively. For the other stations, ¹³⁷Cs activities 370 371 were similar (17.7 \pm 3.6, 16.7 \pm 2 and 15.5 \pm 2.1 Bq kg⁻¹ for stations 1, 4 and 5). The flood event on January 29, 2014 had the highest activity of all floods, with an average activity of 24.7 ± 2.8 Bq kg⁻¹ 372 373 followed by the flood of April 4, 2014 (19.8 \pm 7.3 Bq kg⁻¹). The two floods sampled on December 29, 374 2013 and February 13, 2014 were similar, with 137 Cs concentrations ranging between 17.1 ± 1.1 and 17 ± 4.3 Bq kg⁻¹ respectively. Samples collected during the low-flow periods are characterized by low 375 137 Cs activities (6.9 ± 0.8 Bg kg⁻¹ on September 10, 2013 and 13.9 ± 12.3 Bg kg⁻¹ on April 30, 2014). 376 The sediment collected at the tile drain outlets (n = 5) were characterized by the presence of high ¹³⁷Cs 377 activities with an average of 23.4 \pm 4 Bq kg⁻¹ (ranging between 8.6 \pm 0.3 and 30.6 \pm 0.5 Bq kg⁻¹) 378 379 indicative of elevated surface source contributions. As shown in Fig. 5, these high concentrations of ¹³⁷Cs in the sediment samples mainly plot within the concentration range of surface sources. Over a 380 10-year period (2003–2013) the ¹³⁷Cs activity of material accumulated in the pond ranged between 381 10.9 ± 0.7 and 11.9 ± 0.7 Bq kg⁻¹ with an average activity at the upper 10 cm of the core being $11.4 \pm$ 382 0.5 Bq kg⁻¹. These values also plot clearly within the surface source range. Summary statistics of ¹³⁷Cs 383 384 concentrations are provided in Table 2 for all sediment samples.

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3.4. Modelling results – source identification

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388 When all sediment samples from the monitoring stations are grouped together (*C* in Eq. (3)), 389 the modelling results for both corrections indicate that sediment transported in the Louroux catchment 390 are almost entirely originated from surface sources (99 \pm 1.2% with the *SSA* correction and 94 \pm 1.5% 391 with the *Th* correction). Corrected ¹³⁷Cs activities in sediment collected at the different monitoring 392 stations clearly plots within the distribution of surface source ¹³⁷Cs activities (Fig. 7). The modelling 393 results averaged for the 5 monitoring stations indicate a surface contribution of 99% (\pm 0.5%) with the 394 *SSA* correction. The results with the *Th* correction are comparable with a mean modelled surface 395 contribution of 98% (\pm 1.9%). The mean difference between surface source contributions with the *SSA* 396 and *Th* corrections modelled for all monitoring stations was 1.2% (\pm 1.6%). Table 3 lists all modelling 397 results.

398 To further examine sediment sources in this catchment, the different events sampled were modelled separately. Sediment samples collected during flood events have elevated ¹³⁷Cs 399 400 concentrations (suspended sediment samples SSE2-6 in Fig. 8). These high activities indicate a major 401 contribution of surface material during the flood events. The modelled results indicate a surface source 402 contribution of 99% (\pm 0.4%) with the SSA correction and 99% (\pm 0.5%) with the Th correction. Samples collected during a low flow event had a ¹³⁷Cs signature clearly lower than during the flood 403 404 conditions representative of a more homogeneous mixture of surface and subsurface sources (SSE1 in 405 Fig. 8). During the low flow period, the sediment distribution plotted closer to the subsurface source 406 distribution (Fig. 8). Surface sources were modelled to contribute 49% (± 3.5%) with the SSA 407 correction compared to 40% (\pm 2.1%) with the *Th* correction. Although there was a 9% difference 408 between the modelled results with the SSA and Th corrections for the low flow event, the mean 409 difference between the modelled surface source contributions for all events was only $2\% (\pm 3\%)$.

The mean ¹³⁷Cs value of sediment at the tile drainage network outlet $(23.4 \pm 4 \text{ Bq kg}^{-1})$ was higher than during the floods $(18.4 \pm 3.4 \text{ Bq kg}^{-1})$, (Figs. 6 and 7). Sediment exported from the drainage network was modelled to originate predominantly from surface sources $(99 \pm 2.5\%)$ with both corrections (Fig. 9). This indicates that these drainage networks potentially facilitate the transfer of surface soils and should be treated as a surface source in this catchment. For the sediment core, results indicate that sediment is mainly derived from surface sources with a modelled contribution of 99% ($\pm 1.5\%$) with the SSA correction compared to 97% ($\pm 6.7\%$) with the *Th* correction.

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418 **4. Discussion**

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420 In this study the SSA and Th corrections reduced particle size enrichment impacts on 137 Cs concentrations. The SSA measured for both sources were similar (subsurface mean: 340 m² kg⁻¹, 421 surface mean: 390 m² kg⁻¹). The significant difference between sediments and source SSA is 422 423 potentially indicative of particle size enrichment during sediment mobilization and transport processes. 424 This enrichment is highlighted by the large observed particle size difference between sources and 425 sediment. The application of both particle size corrections clearly improved the relationship between 426 sediment and their sources. When examining all modelling results, there was only a mean difference of 427 2% (± 3%) between both corrective approaches. There was only one exception, the low flow sediment 428 sample (SSE1) where modelling approaches differed by 9%.

429 The main observed difference between the Th and SSA corrections could be because the 430 application of the *Th* correction allows a broader range of values than the SSA approach. Nevertheless, 431 the high values for the surface sources obtained with the Th correction do not seem to be outliers as the drain samples have higher 137 Cs activities. By comparison at the sample scale, the *Th* correction is 432 more applicable than the SSA correction. The application of the Th correction to the surface source 433 samples (measured ¹³⁷Cs activity: 3.7 ± 3 Bg kg⁻¹) changed the surface source ¹³⁷Cs activity to 14.3 434 Bq kg⁻¹ compared to 11.4 Bq kg⁻¹ with the SSA correction. The average ¹³⁷Cs activities during runoff 435 436 was 17 ± 3 Bq kg⁻¹. The application of both corrections could be improved by collecting additional sediment and source samples covering a wider range of ¹³⁷Cs activities. 437

The modelling results for sediment samples taken over the entire hydrological year (2013/2014) and for the sediment core clearly illustrate that surface sources dominate the supply of sediment in this catchment ($\mu = 97\% \pm 6.7\%$ for the sediments and $99 \pm 1.5\%$ for the sediment core). These results are in agreement with the review on sediment sources in British rivers that indicated surface sources dominate the supply of sediment, accounting for between 60% and 96% of the sediment yields, with 85–95% being typical (Walling, 2005).

The sediment exported by the drainage network is characterized by higher ¹³⁷Cs activities than 444 the soil samples $(3.2 \pm 0.2 \text{ Bg kg}^{-1} \text{ for average soil samples and } 23.4 \pm 4 \text{ Bg kg}^{-1} \text{ for drainage network}$ 445 sediment). These high values are worthy of attention because ¹³⁷Cs activity concentrations are 446 447 expected to decrease below the plough depth in cultivated soil (He and Walling, 1997) and are nearly 448 undetectable in subsurface soil (Matisoff et al., 2002). One possibility could be that there has been 449 ¹³⁷Cs migration down in the soil profile in our catchment. According to previous studies (Sogon et al., 450 1999; Walling et al., 2002; Chapman et al., 2005), macropores are a potential link between the topsoil 451 with high ¹³⁷Cs activities and the drainage networks. These macropores can be induced by the soil 452 properties, agricultural soil works, and vegetal activities (Oygarden et al., 1997), and they can be a preferential pathway of sediment originating from the topsoil and labelled with ¹³⁷Cs (Jagercikova et 453 454 al., 2014). This direct pathway offers the best explanation for the rapid sediment transport observed by Chapman et al. (2005) and potentially the high ¹³⁷Cs activities observed in sediment sampled in our 455 456 studied drainage network.

457 Another explanation is that eroded surface material is being exported through the drainage 458 network. Similarly to Walling et al. (2002), sediment samples from the studied field drains have higher 459 ¹³⁷Cs activities than the soil samples. This radionuclide enrichment at the drainage network outlet has 460 also been described by Sogon (1999) who demonstrated the occurrence of preferential particle size 461 selection in the upper soil and the migration of the finest particles through the tile drainage network 462 during runoff events. A combination of these factors likely explains this particle selectivity between 463 soil and drain material and additional research is required to determine the dominant transfer 464 mechanisms.

465 There are limited analyses of sediment contributions from drainage networks available in 466 France (Penven and Muxart, 1995; Sogon et al., 1999; Penven et al., 2001). Research from other 467 countries demonstrated that drainage networks provide a significant contribution to sediment export. 468 In the UK, drainage networks were clearly an important source to sediment yields accounting for 469 between 27% and 55% of sediment exported (Russell et al., 2001; Walling et al., 2002). Furthermore, 470 Foster et al. (2003) reported that drainage networks contributed more than 50% of sediment. At the 471 global scale, drainage network contributions may be significant. For instance, Macrae et al. (2007) 472 estimated >42% of annual hydrological discharge is originated from the drainage network in a 473 Canadian agricultural catchment. According to King et al. (2014), this hydrological pathway is under-474 studied in agricultural basins. More research is required in the study area to determine whether ¹³⁷Cs 475 has migrated down through the soil profile or whether these drainage networks simply act as conduit 476 for quick transportation of surface soils to the stream network during runoff events.

477 At the global scale, previous sediment fingerprinting studies demonstrated that subsurface 478 contribution to sediment yields varied among catchments depending on several parameters such as 479 morphology and land use. Data compiled by Walling and Collins (2005) indicate that generally bank 480 erosion can contribute between 5% and 15% of sediment exported in British rivers but in some cases it 481 can exceed 40%. In the review of Haddadchi et al. (2013), subsurface erosion from channel bank 482 sources was reported to contribute typically between 15% to 30% of suspended sediment load. In 483 Australian catchments, it is not uncommon for subsurface sources to contribute more than 90% of 484 sediment load (Caitcheon et al., 2012; Olley et al., 2013; Laceby et al., 2015). Our results remain in 485 agreement with European studies, with an average riverbank contribution ranging between 11% to 486 32% for the last decade.

In the Louroux catchment, the majority of the sediments are exported during the flood events. Modelling results indicate the dominance of the surface sources during these events whereas samples collected during the lower flow periods have an increased proportion of sediments derived from subsurface sources. The volume of water and sediment exported during the low flow events is often insignificant compared to the volume of sediment exported during flood events, though more research is required to define the relative proportion of both sources during the entire hydrological year.

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494 5. Conclusions

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This study highlights the potential of Th based particle size corrections. The application of this correcting factor produced globally better results that the *SSA* correction. There remains, however, more experimentation required to test the validity of this approach to particle size correction in various environments, particularly catchments with heterogeneous geologies where Th may be a significant discriminator between different spatial sediment sources. 501 Modelling results indicate that sediment transported during the flood events in the Louroux 502 catchment are almost entirely originated from surface sources, regardless of the correcting factor 503 employed (~ 99%). During these events, two pathways can mobilize this surface source: sheet and rill 504 erosion associated with the runoff events and the sediment exported from the drainage network which 505 was modelled to originate predominantly from surface sources (99% \pm 2.5%). During the low flow 506 period, modelling results correspond to a more homogenous mixture between surface and subsurface 507 sources with contributions of subsurface source ranging between 51% and 60%.

508 Over the last 10 years, surface sources dominated the supply of sediment with both corrections 509 in the pond ($99\% \pm 1.5\%$ to $97\% \pm 1.5\%$). Accordingly, management of deleterious sediments with 510 contaminants accumulated within the Louroux pond should focus on reducing the supply of sediment 511 from surface sources. Future research should examine the efficacy of drainage networks for 512 connecting sediments from surface sources to the stream network in more detail. Managing sediment 513 transferred through drainage networks may reduce suspended sediment loads in similar lowland 514 agricultural catchments.

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- 757 <u>Figure captions:</u>
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- Fig. 1. Catchment and sampling location map with, A) the location of the study site in the Loire River
 basin, and B) the Louroux pond catchment with source sample locations and river monitoring sites
 (S1: Beaulieu River, S2: Grand Bray River, S3: Masnier River, S4: Picarderie River, S5: Conteraye
 River and D1–3: drain stations, H: Hillslope sediment samples collected during runoff events).
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- Fig. 2. ¹³⁷Cs activities in multiple particle size fractions, (bulk soil and fractions ranging 63–50, 50–20, or $<20 \,\mu$ m) and suspended sediment (*SS*).
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Fig. 3. Relationship between ²¹⁰Pb_{ex} and ¹³⁷Cs for surface and subsurface sources along with in-stream
 sediments.

Fig. 4. Relationship between measured and corrected 137 Cs activities for the surface and subsurface samples using the *SSA* (A) and *Th* (B) correcting factors. Error bars represent analytical uncertainties on radionuclide activities (1 sigma).

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Fig. 5. Relationship between 210 Pb_{ex} and 137 Cs for surface and subsurface sources along with in-stream sediments after the *SSA* (A) and *Th* (B) particle size corrections.

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Fig. 6. Probability plots characterizing surface and subsurface sample distributions with the *SSA* (A)
and *Th* (B) particle size corrections. The probabilities were generated for each source with 2500 Latin
hypercube samples from each source distribution.

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Fig. 7. Probability plots for ¹³⁷Cs activity concentrations including sediment samples (black squares) collected from the monitoring sites. In each case, probabilities were generated with 2500 Latin hypercube samples from the source and sediment distributions as well as the mixture distribution. Error bars represent analytical uncertainties of sediment samples equivalent to one standard error of the mean. The solid line is the subsurface source distribution, the dashed line is the surface source distribution and the dotted line is the distribution of sediment from each monitoring station.

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Fig. 8. Probability plots for ¹³⁷Cs activity concentrations including sediment samples (black circles) 789 790 collected during the six sampling survey (SSE1: 09 Sept 2013; SSE2: 30 Dec 2013; SSE3: 29 Jan 791 2014; SSE4: 13 Feb 2014; SSE5: 4 Apr 2014; SSE6: 30 Apr 2014). In each case, probabilities were 792 generated with 2500 Latin hypercube samples from the source and sediment distributions as well as 793 the mixture distribution. Error bars represent analytical uncertainties of sediment samples equivalent 794 to one standard error of the mean. The solid line is the subsurface source distribution, the dashed line 795 is the surface source distribution and the dotted line is the distribution of sediment from each event 796 sampled.

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Fig. 9. Probability plots for ¹³⁷Cs activity concentrations including the core and tile drainage network samples (black circles). In each case, probabilities were generated with 2500 Latin hypercube samples from the source and sediment distributions as well as the mixture distribution. Error bars represent analytical uncertainties of sediment samples equivalent to one standard error of the mean. The solid line is the subsurface source distribution, the dashed line is the surface source distribution and the dotted line is the distribution of sediment from core and tile drainage network.

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805 <u>Table captions:</u>

806

- 807 Table 1. Summary of ¹³⁷Cs activities (Bq kg⁻¹) of the potential sources with the different correction
- 808 techniques (w/o corr: without correction, SSA: surface specific correction, Th: Thorium correction).
- 809
- 810 Table 2. Summary of mean ¹³⁷Cs activities and the standard deviation (σ) (Bq kg⁻¹) for the in-stream
- 811 and drain samples at each monitoring station during flood events.
- 812
- 813 Table 3. Details of modeling results for surface and subsurface contributions.
- 814
- 815

Table 1

Statistic	Subsoil			Surface		
	w/o corr.	SSA	Th	w/o corr.	SSA	Th
Mean	1.3	1.7	1.8	3.2	10.0	11.2
Standard deviation	1.0	1.9	2.2	1.4	4.2	8.3
Mediane	1.0	1.3	0.9	2.9	9.3	9.3
Minimum	0.0	-0.7	0.0	1.0	3.7	0.8
Maximum	3.1	5.1	6.5	7.9	23.8	35.5

Table 2

	Date	¹³⁷ Cs	σ
Station 1	09/09/2013	8.6	0.7
	30/12/2013	21.3	4.2
	29/01/2014	20.3	1.6
	13/02/2014	16.9	0.8
	04/04/2014	10.3	3.2
Station 2	09/09/2013	6.0	0.6
	30/12/2013	24.6	4.4
	29/01/2014	18.3	2.1
	13/02/2014	17.8	1.0
Station 3	30/12/2013	6.3	2.4
	29/01/2014	25.1	3.1
	13/02/2014	14.3	0.9
	04/04/2014	30.2	13.2
	30/04/2014	0.0	17.3
Station 4	09/09/2013	7.1	1.1
	30/12/2013	11.5	5.1
	29/01/2014	23.1	2.4
	13/02/2014	15.4	0.8
	30/04/2014	31.3	8.6
Station 5	09/09/2013	6.1	1.0
	30/12/2013	25.5	5.3
	29/01/2014	36.7	4.7
	13/02/2014	21.3	1.8
	04/04/2014	18.8	5.4
	30/04/2014	24.4	7.7
D1	30/12/2013	13.4	2.3
	29/01/2014	23.5	4.1
D2	16/01/2013	8.6	0.3
D3	30/12/2013	30.6	7.5
	29/01/2014	26.1	2.3

Туре	Surface contribution (%) and standard error (%)			
	SSA		Th	
Average sample	99	±2	94	± 1.5
Sediment Core Drain sediment	99 99	± 1.5 ± 2	97 99	± 6.7 ± 2.5
Station 1	99	± 1	96	±2
Station 2	98 00	± 1.6	95 00	± 1.5
Station 3 Station 4	99 99	± 1 ± 1.2	99 99	±1 ±1
Station 5	99	± 1.1	99	± 1.2
SSE1- 09 sep 13	49	± 3.5	40	±2
SSE2- 30 dec 13 SSE3- 29 jan 14	98 99	±1 ±1	99 99	± 1.4 ± 1
SSE4- 13 feb 14	99	±2	98	± 1.5
SSE5- 04 apr 14 SSE6- 30 apr 14	99 98	± 1 ± 1.5	99 99	± 2 ± 3

Table 3

Fig. 1



Fig. 2

Fig. 3

Fig. 4

Fig. 6

Fig. 2

Fig. 3

Fig. 4

Fig. 6

