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- 1 Tracing of particulate organic C sources across the terrestrial-aquatic continuum, a 2 case study at the catchment scale (Carminowe Creek, South West England)
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21 Abstract

Soils deliver crucial ecosystem services, such as climate regulation through carbon (C) 22 23 storage and food security, both of which are threatened by climate and land use change. 24 While soils are important stores of terrestrial C, anthropogenic impact on the lateral fluxes of C from land to water remains poorly quantified and not well represented in Earth system 25 26 models. In this study, we tested a novel framework for tracing and quantifying lateral C 27 fluxes from the terrestrial to the aquatic environment at a catchment scale. The combined use 28 of conservative plant-derived geochemical biomarkers *n*-alkanes and bulk stable δ^{13} C and $\delta^{15}N$ isotopes of soils and sediments within an inter-disciplinary framework allowed us to 29 distinguish between particulate organic C sources from different land uses (i.e. arable and 30 31 temporary grassland vs. permanent grassland vs. riparian woodland vs. river bed sediments) (p<0.001), showing an enhanced ability to distinguish between land use sources as compared 32 33 to using just biomarkers alone. The terrestrial-aquatic proxy (TAR) ratio derived from n-34 alkane signatures indicated an increased input of terrestrial-derived organic matter (OM) to 35 lake sediments over the past 60 years, with an increasing contribution of woody vegetation 36 over time. This may be related to agricultural intensification, leading to enhanced soil

37 erosion, but also an increase in riparian woodland that may disconnect OM inputs from arable 38 land uses in the upper parts of the study catchment. Spatial variability of geochemical proxies 39 showed a close coupling between OM provenance and riparian land use, supporting the new 40 conceptualization of river corridors (active river channel and riparian zone) as critical zones 41 linking the terrestrial and aquatic C fluxes. Further testing of this novel tracing technique 42 shows promise in terms of quantification of lateral C fluxes as well as targeting of effective 43 land management measures to reduce soil erosion and promote OM conservation in river 44 catchments.

45 **Keywords:** lateral carbon fluxes, sediment fingerprinting, biomarkers, *n*-alkanes, bulk stable 46 13 C and 15 N isotopes

47

48 **1. Introduction**

49 Soils are critical to human wellbeing and deliver crucial ecosystem services, including climate regulation and food security (Adhikari and Hartemink, 2016; Mouchet et al., 2016). 50 51 However, since the onset of agriculture, human activities have greatly altered soil processes 52 at a global scale, with consequences for the essential functions of soils to sequester and store 53 carbon (C), recycle nutrients and resist soil erosion (Amundson et al., 2015). As soils 54 represent the largest terrestrial store of organic C, more than three times as much as either the 55 atmosphere or terrestrial vegetation (Schmidt et al., 2011), these anthropogenic interventions 56 have also impacted the scale of the lateral fluxes of C from land to inland waters (Lauerwald et al., 2015; Tian, 2015; Wohl et al., 2017). 57

58 However, the fluxes of C from land to ocean remain poorly quantified and not fully 59 accounted for in the current generation of Earth system models (Regnier et al., 2013). While over the past decade, the understanding of rivers has been revised from 'inert pipes' simply 60 transporting C from lad to the ocean to 'active agents', which play a crucial role in receiving, 61 62 transporting and processing C equivalent to net terrestrial primary production in their 63 watersheds (Aufdenkampe et al., 2011; Battin et al., 2009; Cole et al., 2007; Wohl et al., 64 2017)., the magnitude, spatiotemporal patterns and controls on C fluxes from land to ocean remain poorly quantified (Regnier et al., 2013; Wohl et al., 2017). While there is a growing 65 66 understanding of the magnitude of global C exports from rivers to the ocean (Li et al., 2017; Ludwig et al., 2011; Tian et al., 2015), estimates of CO₂ evasion from inland waters 67 (Lauerwald et al., 2015; Raymond et al., 2013) and sediment burial in aquatic ecosystems 68 69 (Maavara et al., 2017; Tranvik et al., 2009) are still uncertain. However, the largest 70 uncertainties are associated with the scale of the total lateral C fluxes from land to inland waters, with recent research suggesting that previous estimates may have largely over-71 72 estimated C accumulation in terrestrial ecosystems (the terrestrial C sink) due to under-73 estimation of this lateral C export (Nakayama, 2017). Therefore, there is a need to better 74 understand the scale of the anthropogenic impact on these lateral C fluxes from land to water 75 (Regnier et al., 2013; Wohl et al., 2017), as well as the processes involved in the loss and 76 preservation of C along the terrestrial-aquatic continuum (Marín-Spiotta et al., 2014), to

- 77 properly represent these processes and predict the present and future contribution of aquatic
- 78 C fluxes to the global C budget (Aufdenkampe et al., 2011; Battin et al., 2009; Cole et al.,
- 79 2007; Regnier et al., 2013).

80 Over the past decades, the awareness of the importance of soils in the functioning of many 81 vital ecosystem services, including climate change mitigation, food security, water resource 82 management and flood protection has greatly increased (Schroter et al., 2005). Nevertheless, 83 in many parts of the world where soil erosion rates exceed soil production, the sustainable 84 provision of these ecosystem services is under pressure (Alewell, et al., 2015; Amundson et 85 al., 2015; Panagos et al., 2015; Verheijen et al., 2009). The intensification of agriculture, 86 particularly over the past 60 years, has led to an exponential increase in sediment and organic 87 matter (OM) fluxes within agricultural catchments (Glendell and Brazier, 2014; Graeber et 88 al., 2015), with important consequences for on-site impacts, such as soil productivity, and 89 off-site impacts, in terms of nutrient pollution and sedimentation of water bodies (Tilman et al., 2002). Hence, conservation of soil organic matter (SOM; which contains ~60% SOC) 90 91 remains critical for sustaining soil productivity and food security in a changing world 92 (Amundson et al., 2015) and for mitigating the acknowledged wide-scale impacts of 93 enhanced sedimentation and associated nutrient pollution on the ecological status of water 94 bodies and drinking water quantity and quality (Bilotta and Brazier, 2008; Glendell et al., 95 2014a; Rickson, 2014; Schoumans et al., 2014).

96 Tracing and quantifying the sources of sediment and particulate organic C in the fluvial 97 environment is, therefore, key to supporting sustainable land management decisions and 98 maintaining ecosystem services. To date, most tracing techniques to apportion sediment 99 sources in fluvial environments applied in river management studies have used physical 100 sediment characteristics, geochemical properties, fallout radionuclides or mineral magnetic 101 properties. However, these tracers are not able to distinguish sources between specific land 102 uses, which are essential to inform mitigation measures and catchment management (Guzman 103 et al., 2013; Owens et al., 2016; Smith et al., 2015; Walling, 2013). Conversely, examination of the provenance of sediment-bound OM using plant-specific biomarkers has been 104 105 established in paleo-ecological and marine sciences for some time (e.g. Galy et al., 2011; 106 Meyers and Lallier-Vergès, 1999; Meyers, 2003; Tolosa et al., 2013; Zech et al., 2012). While a number of studies to date have sought to apply one or more biomarkers to understand 107 108 lateral C dynamics at the continental margins (e.g. Feng et al., 2015; Galy et al., 2011; Tao et 109 al., 2016), few studies have applied this approach to inland waters, especially headwaters, 110 which may cumulatively play an important role in lateral C export due to their spatial extent 111 and close terrestrial – aquatic coupling. Therefore the application of biomarkers, especially aliphatic (saturated straight-chained) compounds such as *n*-alkanes (Chen et al., 2016, 2017; 112 113 Cooper et al., 2015; Puttock et al., 2014) and *n*-carboxylic acids (fatty acids) (Alewell et al., 2016; Blake et al., 2012; Reiffarth et al., 2016), is now being examined as a new potential 114 115 tool for attribution of sediment and C provenance in river catchments, with a potential to 116 attribute organic matter sources to specific land uses, such as forest, arable and pasture.

117 *N*-alkanes are naturally occurring hydrocarbons which are relatively recalcitrant and more 118 resistant to microbial decomposition than other functionalized plant-derived lipids, e.g. fatty

- acids or sterols (Ranjan et al., 2015). They are vegetation-specific neutral lipids derived from
- 120 plant waxes with different numbers of C atoms in the aliphatic molecule that are indicative of
- 121 different provenances of OM (Eglinton, 1962). In general, long-chain $(C_{27}-C_{31})$ *n*-alkanes are
- derived from epicuticular plant waxes of terrestrial plants (Galy et al., 2011; Puttock et al.,
- 123 2014), medium chain-length (C_{21} - C_{25}) *n*-alkanes are produces by lower plants and aquatic 124 macrophytes (Fang et al., 2014; Meyers, 2003; Tolosa et al., 2013), while short chain-length
- 125 ($C_{15}-C_{19}$) *n*-alkanes are typically derived from aquatic algae (Meyers, 2003). Both individual
- *n*-alkanes as well as different chain-length ratios have been used in paleo-ecological studies
- 127 to attribute OM sources over decadal to millennial timescales (Ranjan et al., 2015; Zech et
- 128 al., 2013, 2012). As *n*-alkane signatures are altered by land use change, they are ideally suited
- 129 to track changing OM sources from eroded soils and sediments over time (Chen et al., 2016).
- 130 With these naturally occurring biomarkers (and their compound-specific isotopic signatures where vegetation sources with contrasting δ^{13} C values are evident) emerging as the new 131 potential tools for tracing of SOM in catchment studies, a key challenge lies in establishing 132 133 their effectiveness to act as land-management specific tracers of fluvial OM over decadal 134 timescales (Alewell et al., 2016; Cooper et al., 2015). As these new techniques are still in their infancy (Owens et al., 2016) and require further development and testing, in this work 135 136 we aim to investigate the suitability of *n*-alkane biomarkers within an inter-disciplinary 137 context, beyond the traditional confines of soil science alone (Brevik et al., 2015; Owens et al., 2016; Smith et al., 2015). 138
- Therefore, this pilot study aims to evaluate the combined use of *n*-alkanes, bulk stable δ^{13} C 139 and $\delta^{15}N$ isotopes and their ratios (Collins, et al., 2014, 2013; Meyers and Lallier-Vergès, 140 1999; Meyers, 2003; Ranjan et al., 2015) to advance the current understanding of the 141 temporal variability of lateral C fluxes from the terrestrial to 142 the aquatic ecosystem in relation to changing land management practices, over the past century. Our aims were to i) 143 test the ability of conservative *n*-alkane biomarkers and soil and sediment bulk stable δ^{13} C 144 145 and δ^{15} N isotopes to distinguish between terrestrial and aquatic OM sources, ii) understand the impact of land use on the spatial variability of OM provenance in river bed sediments and 146 iii) on OM accumulation in lake bed sediments, iv) test a methodology for quantifying the 147 148 temporal variability and the magnitude of lateral C fluxes from land to water at catchment 149 scales.
- 150

151 **2. Material and Methods**

152 <u>2.1 Study site</u>

The Carminowe Creek study catchment, located in southwest England (50°4' W 5°16'), covers c. 4.8 km² at an altitude range of 0-80 m above sea level (Fig. 1). The catchment outlet drains into a large freshwater lake Loe Pool (50 ha) that is separated from the Atlantic Ocean by a natural shingle barrier, thus creating a relatively closed natural hydrological system. The study catchment comprises two main streams (Northern and Southern subcatchments) with a joint outlet into the south-western branch of Loe Pool. The average total annual rainfall is 159 approximately 1000 mm and mean annual temperature is approximately 11°C 160 (http://www.metoffice.gov.uk/public/weather/climate/). Bedrock geology comprises silt-, sand- and mudstone (http://mapapps.bgs.ac.uk/geologyofbritain/home.html), which is 161 overlaid by freely draining loamy soils (Soil Survey of England and Wales, 1983). Land use 162 on the catchment plateaux is dominated by cropland in rotation of arable crops and temporary 163 164 grassland (also referred to as grass ley), while permanent grassland is found on steeper 165 hillslopes, with riparian vegetation (mostly wet woodland dominated by willow Salix sp., 166 alder Alnus glutinosa and wet grassland) located in the riparian zone within the river corridor.

167

168 <u>2.2 Field sampling</u>

169

78 soil cores were taken 14 hillslope transects across the two sub-catchments (8 cm diameter, 170 171 depth 0-15 cm), covering the topographic sequence from plateaux, convex, steep slope, 172 concave and footslope locations. In total, 31 samples were taken from arable land use, 26 173 from temporary grassland (ley), 14 from permanent grassland and 7 from riparian woodland. 174 Three river-bed sediment samples were collected with a hand trowel and bulked into a single 175 sample on a single occasion at 7 locations along each of the two streams at an (i) upstream (ii) midstream and (iii) downstream location and the joint catchment outlet before the 176 177 confluence with Loe Pool. Two 0.5 m deep lake sediment cores were taken from Loe Pool c. 178 150 m below the outlet of Carminowe Creek using a Mackereth corer (Mackereth, 1969).



Fig. 1 The study site location in south-west England showing land use, the 14 study transects,river bed sediment sampling locations and the lake core.

182

183 <u>2.3 Laboratory analysis</u>

Following sample collection, river bed and lake core sediment samples were stored at 4°C and analysed as soon as possible. Soil samples were oven dried at 40°C and sieved to <2 mm and >2 mm fractions. River bed samples were wet sieved at 250 μ m to separate coarse vegetation debris as in Galy et al. (2011) and the < 250 μ m fraction was oven dried at 40°C and retained for further analysis. The lake sediment core was sliced into 2 cm increments and freeze dried.

All soil, river bed sediment and lake core samples were analysed for total C, N and bulk δ^{13} C 190 and δ^{15} N using a SerCon Integra2 isotope ratio mass spectrometer (SerCon Ltd., Crewe, UK). 191 An in house standard of Alanine (N=16.7%, C=40%, δ^{15} N=-1.68‰_{AIR}, and δ^{13} C=-192 193 19.58‰_{VPDB}) was used in duplicate every 9 samples to provide quality control and to act as an internal reference. This was calibrated within each analytical run by also analysing 194 195 International Atomic Energy Agency (IAEA) reference materials N-1 and N-2 for nitrogen 196 and CH-6 and LSVEC for carbon. Due to systematic blanks data had to be blank subtracted 197 and a linearity correction was applied based on the analysis of the IAEA reference materials. 198 The analytical precision of the internal quality control samples of alanine was determined. 199 The standard deviation of the δ^{15} N measurements was <0.3‰ and of the δ^{13} C measurements 200 was <0.1‰. The values were expressed relative to AIR and Vienna PeeDee Belemnite 201 (VPDB) for nitrogen and carbon respectively. The formula used for presenting δ values is as 202 follows:

203
$$\delta^{15}$$
N X ‰_{AIR} = (R_{sam}/R_{ref})-1)*1000 (1)

204
$$\delta^{13}C X \ \text{\sc vpd} = (R_{\text{sam}}/R_{\text{ref}}) - 1) * 1000$$
 (2)

Where sam is sample and ref is the reference material, R is the ratio of the heavy isotope over the light isotope, X being the isotope ratio expressed in units of per mille (‰).

In order to establish a chronology for the lake core profile, ¹³⁷Cs assay of individual 2 cm core sections was undertaken at 661.67 KeV using an ORTEC GMX co-axial HPGe γ detector, coupled to a multi-channel analyser. Sample count times were generally 24 hours, resulting in analytical precision of c. 5%.

- On the basis of likely hydrological connectivity with the watercourses we selected a sub-set 211 of 50 samples to estimate *n*-alkane concentrations ($\mu g g^{-1} C$) from soil samples Lake core 212 samples were combined into 4 cm increments. . The procedure of total lipid extraction was 213 214 followed by lipid fractionation to isolate the hydrocarbon fraction for analysis using an 215 Agilent 6890 GC instrument coupled to an Agilent 5973 MS instrument and equipped with an Agilent DB-5 ms column (30 m x 250 µm i.d.x 0.25 µm film thickness). The dominant 216 217 fragment ions (base peak) were represented by m/z 57 and the diagnostic ions (m/z) 282 218 (C₂₀), 296 (C₂₁), 324 (C₂₃), 338 (C₂₄), 352 (C₂₅), 366 (C₂₆), 380 (C₂₇), 394 (C₂₈), 408 (C₂₉), 422 (C₃₀), 436 (C₃₁), 450 (C₃₂), 464 (C₃₃) and 478 (C₃₄, internal standard) (Norris, 2013). The 219 220 concentrations of individual n-alkanes were determined relative to the C₃₄ internal standard.
- Interpretation of *n*-alkane results used the percentage of C_{27} , C_{29} and C_{31} calculated as (Torres et al., 2014):

223 %
$$C_i = C_i / (C_{27} + C_{29} + C_{31})$$
 (3)

where C_i stands for the respective *n*-alkane (C_{27} , C_{29} and C_{31}).

Further, we used the ratio between the shorter chain C_{27} (indicative of woody source (Zech et al., 2009) and longer chain C_{31} (indicative of grass source (Eckmeier and Wiesenberg, 2009) to distinguish between respective contributions of OM from woodland and grassland land uses (Puttock et al., 2014).

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230 <u>2.4 Indicators of aquatic versus terrestrial OM sources</u>

To interpret the relative contribution of higher aquatic vs. terrestrial plants to OM in river and lake sediments we used the following formula (Ficken et al., 2000):

233
$$PAQ = \frac{(C23+C25)}{(C23+C25+C29+C31)}$$
 (4)

- where PAQ is the ratio of shorter-chain *n*-alkanes ($C_{23}+C_{25}$) contributed by higher aquatic
- 235 plants (macrophytes) and mosses to the concentration (in $\mu g g^{-1}$) of *n*-alkanes indicative of
- both aquatic and terrestrial vegetation $(C_{23}+C_{25}+C_{29}+C_{31})$.
- The proportion of OM from terrestrial sources in river bed sediments and in the lake core wascalculated using the following formula (Fang et al., 2014; Meyers, 2003)

239
$$TAR = \frac{(C27+C29+C31)}{(C15+C17+C19)}$$
 (5)

where TAR is terrestrial/aquatic ratio of the concentration of *n*-alkanes (in $\mu g g^{-1}$) derived from terrestrial sources (C₂₇+C₂₉+C₃₁) to those indicative of aquatic algae (C₁₅+C₁₇+C₁₉).

Organic matter degradation in the lake core was examined using the odd-over-even
predominance (OEP) *n*-alkane ratio (Zech et al., 2013) as follows:

244 OEP=
$$(nC_{27} + nC_{29} + nC_{31} + nC_{33}) / (nC_{28} + nC_{30} + nC_{32})$$
 (6)

High OEPs point either to an increased OM input and/or to an increased OM preservation
while low OEPs are indicative of accelerated degradation under aerobic conditions (Zech et al., 2013).

- We used δ^{13} C, δ^{15} N and C/N ratio as further geochemical proxies to understand the proportion of OM contributed by algal *vs.* terrestrial plant derived production as used in previous studies (Fang et al., 2014; Hamilton and Lewis, 1992; Meyers, 2003).
- 251

252 <u>2.5</u> Statistical Analysis

Kruskal-Wallis non-parametric test was used to examine the differences between elemental 253 (C/N) and isotopic (δ^{15} N, δ^{13} C) signatures between different sediment sources. All soil, river 254 255 bed sediment and lake core increments were included in this analysis. Principal component 256 analysis (PCA) was used to examine whether elemental and *n*-alkane ratios could be used to 257 distinguish the provenance of sediment sources derived from six potential sources (arable, 258 temporary grassland (ley), permanent grassland, riparian woodland, lake or river bed). All 259 statistical analyses were undertaken in 'R' vs. 3.4.0. Source apportionment was modelled 260 using the Bayesian isotope mixing model of Stable Isotope Analysis in R (SIAR)' (Parnell 261 and Jackson, 2008, R Core Team, 2014).

262

263 **3. Results & Discussion**

264 <u>3.1</u> Distinguishing between terrestrial and aquatic organic matter sources

Statistically significant differences (p<0.001) in the C/N ratio and bulk stable ¹³C and ¹⁵N isotopic composition of terrestrial soils, river bed sediments and lake core sediments (Table 1), were determined. Woodland and river bed sediments exhibited the highest C/N ratios,

268 while the lowest C/N ratios were detected in arable and temporary grassland soils. The high 269 C/N ratio in woodland soils and river sediments is characteristic of more recalcitrant OM sources such as wood, while the low C/N ratio is indicative of more decomposable OM with 270 271 lower lignin content (Brady and Weil, 1999). Percentage C and % N were comparable 272 between the woodland and grassland soils and lake sediments and differed from low % C and 273 % N in cropland soils and river bed sediments, indicating fast OM turnover in cropland 274 rotations and rapid loss of OM and inorganic N from river bed sediments to the downstream 275 lake.

The highest bulk δ^{15} N values were detected in lake core sediments, followed by grassland and 276 arable soils. Enriched $\delta^{15}N$ values in lake sediments may be due to several processes, 277 including significant macrophyte or riparian-aquatic OM inputs (Fang et al., 2014), increased 278 279 denitrification in anoxic lake bottom waters, and increased net primary production (Meyers, 2003). Increased bulk δ^{15} N values on arable land may be indicative of both rapid turnover of 280 OM and long-term application of manure (Glendell et al., 2014b). Bulk δ^{13} C values were 281 relatively uniform between land uses, reflecting the predominance of C3 plants in the study 282 catchment (Puttock et al., 2014). δ^{13} C values were enriched in arable soils and lake core 283 284 sediments, with the former possibly reflecting periodic growing of maize, a C4 plant with a different photosynthetic pathway with natural abundance δ^{13} C values of ~-12‰ (Beniston et 285 286 al., 2015; Puttock et al., 2014) on arable land and the effect of in-lake organic matter 287 production on lake bed sediments (Fang et al., 2014; Hamilton and Lewis, 1992). However, it 288 is important to acknowledge that direct characterisation of the composition of autochthonous 289 OM produced by aquatic plants in the lake ecosystem would allow a more conclusive 290 interpretation of these findings.

Table 1 Elemental and isotopic composition (mean and SD in brackets) of all terrestrial soils, river bed sediment and lake core samples. Values followed by the same letter are not significantly different, while values followed by a different letter are significantly different (p<0.05). N = number of replicates.

Landuse (N)	C/N	δ^{15} N (‰)	δ ¹³ C (‰)	% C	% N
Arable (31)	8.99 (0.7) a	5.5 (0.8) a	-27.6 (0.4) a	2.92 (0.54)a	0.32 (0.05)a
Grass (14)	9.98 (0.8) b	5.6 (1.2) a, c	-28.2 (0.5) b	5.60 (1.21)b	0.56 (0.14)b
Ley (26)	9.23 (0.7) a	5.5 (0.9) a, c	-28.2 (0.40) b	3.60 (0.82)c	0.39 (0.07)c
Woodland (7)	12.84 (2.4) c	4.7 (1.5) a, c	-28.2 (0.3) b	8.06 (1.83)b	0.63 (0.14)bd
River (7)	12.20 (1.0) c	4.5 (1.1) a, d	-28.3 (0.2) b	2.34 (0.84)a	0.19 (0.06)a
Lake (27)	10.76 (0.6) c	6.3 (0.4) b	-27.6 (0.4) a	7.40 (1.08)b	0.69 (0.07)d

295

296 Concentrations of *n*-alkanes of chain lengths C_{15} - C_{33} in the six environments of interest 297 showed a higher concentration of woody- (C_{27} - C_{29}) and grass- (C_{31}) derived OM input in lake 298 sediments, as compared to terrestrial soils. Concurrently, shorter-chain *n*-alkanes indicative 299 of aquatic macrophytes and lower plants such as mosses (C_{21} - C_{25}) (Meyers, 2003) were also 300 apparent in lake sediments and in riparian woodland (Table 2). As expected, C_{31} chain 301 lengths indicative of grasses (forage and cereals) (Eckmeier and Wiesenberg, 2009), were

- 302 more abundant in the soils of arable, temporary ley and permanent grassland land uses, while
- 303 the C₂₇ chain-length *n*-alkanes, indicative of woody vegetation (Zech et al., 2009), were more
- 304 abundant in woodland soils, river bed and lake core sediments.

305	Table 2 n-alkane concentrations (µg g-1 C) in soils and sediments from the six targ	et
306	environments. N – number of replicates.	

N			n-a	lkane cha	in length	concentra	ation µg g	g⁻¹ C						
					Mear	ו (SD)								
Landus	C15	C17	C19	C21	C23	C25	C27	C29	C31	C33				
e														
Arable	0.16	0.22	n/a	0.65	0.22	0.40	0.93	2.68	3.86	1.99				
(11)	(0.11)	(0.12)		(0.73)	(0.11)	(0.12)	(0.34)	(1.27)	(0.91)	(0.48)				
Grass	0.26	0.26 n/a 0.43 0.35 0.76 1.86 3.81 5.52												
(7)	(0.18)	(0.17)		(0.17)	(0.20)	(0.33)	(1.32)	(1.32)	(1.25)	(1.02)				
Ley	0.25	0.32	n/a	0.92	0.46	0.83	1.29	3.10	4.79	2.22				
(9)	(0.25)	(0.29)		(0.98)	(0.70)	(0.79)	(0.74)	(1.09)	(1.46)	(0.91)				
Woodl	0.11	0.22	n/a	0.94	0.92	2.98	12.93	13.89	7.40	4.40				
and (4)	(0.13)	(0.24)		(0.86)	(0.69)	(1.09)	(6.29)	(5.09)	(2.86)	(2.61)				
River	0.24	0.24	0.22	0.46	0.37	1.35	5.70	5.92	2.65	1.10				
(7)	(0.20)	(0.21)	(0.09)	(0.40)	(0.20)	(0.51)	(2.11)	(2.24)	(1.04)	(0.52)				
Lake	0.26	1.28	0.96	0.96	1.90	4.17	16.45	16.62	12.08	4.93				
(12)	(0.18)	(0.85)	(0.42)	(0.43)	(0.70)	(1.42)	(4.27)	(2.91)	(1.55)	(1.23)				

308 Relative contribution of aquatic vs. terrestrial plants to OM was assessed with three 309 indicators, C_{27}/C_{31} ratios, PAQ and TAR. As expected highest C_{27}/C_{31} ratios, indicative of woody sources were found in the woodland soil samples, while low C_{27}/C_{31} ratios in the 310 311 remaining terrestrial soils were indicative of OM origin from grassland vegetation (including 312 cereal crops and forage grass). The river bed sediments were also characterised by higher 313 C_{27}/C_{31} ratios, indicating predominant sediment input from soils under woody vegetation in 314 the well-connected wooded riparian buffer strip (Fig. 2a). The lake sediments appear to be 315 intermediary between these land uses, indicating OM contribution from both woody and grass vegetation, which may be related to OM transport during high-flow events. During 316 317 rainfall events, the less connected arable/grassland sources may make a greater contribution 318 to OM transport, which is likely to be directly routed to the lake, without being deposited in 319 river bed sediments.

Higher median PAQ ratios of > 0.15 in the lake, river and riparian woodland environments were indicative of emergent macrophyte origin, while the lower median ratios <0.1 were indicative of terrestrial plants (Fig. 2b). The higher PAQ ratios in the riparian woodland may reflect a contribution from lower plants including mosses, which were abundant on the ground of this wet woodland. Still there remains a large unexplained variability of observed PAQ, particularly in the grass ley.



Fig. 2 Box plots of *n*-alkane ratios a) C_{27}/C_{31} indicating OM prevalence from woody *vs*. grass dominated sources b) PAQ indicating contribution of OM from aquatic/lower plant *vs*. terrestrial vegetation. The bottom and top of the box refer to the 25th and 75th percentile, the bold line near the middle to the 50th percentile (the median) and whiskers following the default setting of R, respectively.

333 While single tracers and ratios presented above cannot unravel contribution of several sources on their own, a global assessment with a Principal Component Analysis (PCA) was 334 undertaken. PCA revealed that a combined *n*-alkane, elemental and stable ¹³C and ¹⁵N isotopic 335 336 signatures provide a clear separation in sediment fingerprint composition (Fig. 3). The PCA 337 axis 1 can be interpreted as a gradient between grassland and arable land uses with higher % 338 C₃₁ *n*-alkane signature (indicating grass-derived OM) vs. riparian woodland and aquatic environments with a higher % C₂₇, higher C₂₇:C₃₁ and C/N ratios (indicating wood-derived 339 340 OM sources), lower % C₂₉ (indicating lower plants in the woodland ground vegetation) and higher PAQ (Tables 2-4, Fig. 3a). The PCA axis 2 can be interpreted as a gradient between 341 river bed sediments and lake core sediments, with the latter supporting higher δ^{13} C and δ^{15} N 342 isotopic signatures and higher % C and % N content (Tables 2-4, Fig. 3a) and indicating 343 344 different sediment dynamics in the two aquatic environments. This is also reflected in Fig 3b, 345 which shows a clear distinction in sediment composition between lake and river bed sediments and woodland, permanent grassland and cropland sources. However, PCA could 346 347 not distinguish soils from temporary grassland (ley) and arable land, presumably because 348 these two land uses are subject to regular rotations. While other researchers have also found it 349 possible to distinguish between permanent grassland and woodland sediment and OM 350 sources, they were unable to distinguish between arable land use and permanent grassland, based on the use of biomarkers alone (Alewell et al., 2015). In this study, the combined use 351 352 of biomarkers and elements (% C, % N, C/N ratio) allowed us to distinguish between these two land uses as the % N and % C as well as C/N ratio are all higher in grassland soils than in 353 354 arable and temporary grass ley (Tables 3-4), thus acting as further informative tracers in 355 addition to *n*-alkanes.

Land use	C ₂₇ :C ₃₁	PAQ	%	%	%	% N	$\delta^{15}N$	% C	C/N	$\delta^{13}C$
(N)			C ₂₇	C ₂₉	C ₃₁		(‰)			(‰)
	Mean (SI	D)								
Arable	0.25	0.09	0.13	0.35	0.52	0.33	5.3	3.05	9.32	-27.6
(11)	(0.08)	(0.02	(0.04	(0.06	(0.06	(0.05	(1.03	(0.64	(0.97	(0.42)
))))))))	
Grass (7)	0.37	0.10	0.16	0.33	0.51	0.52	5.2	5.40	10.39	-28.0
	(0.34)	(0.04	(0.07	(0.04	(0.11	(0.10	(1.21	(1.08	(0.86	(0.57)
))))))))	
Ley (9)	0.26	0.12	0.13	0.34	0.53	0.40	5.2	3.77	9.21	-28.2
	(0.10)	(0.09	(0.04	(0.01	(0.04	(0.08	(0.63	(1.07	(0.86	(0.33)
))))))))	
Woodland	1.74	0.15	0.37	0.41	0.22	0.62	4.7	7.80	12.67	-28.3
(4)	(0.54)	(0.02	(0.07	(0.05	(0.04	(0.16	(1.97	(2.29	(3.23	(0.31)
))))))))	
River (7)	2.21	0.17	0.40	0.41	0.19	0.19	4.5	2.34	12.20	-28.3
	(0.37)	(0.03	(0.03	(0.02	(0.02	(0.06	(1.06	(0.84	(1.01	(0.24)
))))))))	
Lake (12)	1.37	0.17	0.36	0.37	0.27	0.70	6.4	7.61	10.91	-27.7
	(0.37)	(0.03	(0.05	(0.01	(0.05	(0.07	(0.31	(0.95	(0.46	(0.41)
))))))))	

Table 3 Biogeochemical values (mean and SD) of the 50 source soils, river bed sediments and lake core samples included in the PCA analysis. N = number of replicates.

358

Table 4 Loading scores of ten biochemical tracers, initial eigenvalues and % total variance accounted for by the first three PCA axes with eigenvalues >1. Loading scores >0.3 were used in the interpretation of axes.

	axis 1	axis 2	axis 3
%N	0.33	0.86	-0.34
delta 15N		0.74	0.48
%C	0.49	0.75	-0.45
C/N	0.74		-0.38
delta 13C		0.49	0.34
C27/C31	0.92		
PAQ	0.66		0.35
C27	0.95		
C29	0.63	-0.46	

C31	-0.97		
Arable	-2.17	-0.40	
Grass	-1.22	0.42	-0.81
Lake	1.71	1.80	
Ley	-1.77		
River	2.39	-2.48	0.91
woodland	2.79		-1.28
Eigenvalues	4.49	2.38	1.06
% of			
variance	44.90	23.82	10.65
Cumulative			
% of			
variance	44.90	68.71	79.36





Fig. 3 Two-dimensional plot of a) variable distribution along the first two PCA ordination axes b) sampling site loading scores on the first two PCA axes and 95% confidence ellipses around the categories of land use.

367 <u>3.2 Spatial variability of land use and provenance of OM in river bed sediments</u>

Spatial patterns of OM provenance in relation to land use were examined at each river bed sediment sampling location ("upstream", midstream", "downstream" and "outlet") in the two subcatchments (S and N) (Figs. 1, 4, 5). Organic matter fingerprinting properties reflected some subtle differences in land used between the two subcatchments. While both the S and

372 N subcatchments were characterized by ca. 75 % agricultural land use, 20 % sub-urban land

use and 5 % of riparian vegetation in the riparian corridor, the S subcatchment supported a
higher arable and ley *vs.* permanent grassland ratio (65 % to 10 %, respectively) than the N
subcatchment (48 % to 19 %, respectively) (Fig. 4).





Fig. 4 Proportion of different land uses a) at the outlet sampling location b) the whole studycatchment c) Southern subcatchment and d) Northern subcatchment.



380

Fig. 5 Spatial variability of a) *n*-alkane proxies b) stable isotopes and c) C/N ratio in a downstream direction in the two subcatchments.

This was reflected in river bed OM properties, which had lower C_{27}/C_{31} ratios than in the N 383 subcatchment, with the upstream location supporting the lowest C_{27}/C_{31} *n*-alkane value of 384 1.75 overall (Fig 5a), likely due to a higher proportion of arable and ley land use in this 385 subcatchment (Fig 4c) leading to higher soil erosion rates (Turnage et al., 1997). The 386 observed higher δ^{15} N values at the most upstream location (ca.+7 ‰, Fig 5b) may also be 387 associated with a higher application of farmyard manure and slurry to arable land and ley 388 389 (Bol et al., 2005; Senbayram et al., 2008). Conversely, the higher C_{27}/C_{31} ratio in the N subcatchment and the lack of upstream forested areas indicate a higher contribution of OM 390 391 from wooded vegetation in the riparian zone, as well as a potential buffering of terrestrial OM 392 fluxes from agricultural soils in the vegetated river corridor. Higher PAQ ratios in the N 393 subcatchment, indicating a relatively higher contribution of mosses and macrophytes derived 394 OM, also point towards a greater influence of the riparian zone on lateral C fluxes as 395 compared to the S stream (Fig. 5a).

Higher C_{27}/C_{31} and C/N ratios (Fig. 5a & 5c) at the midstream and downstream locations in both subcatchments indicated an increased contribution of OM from woody vegetation to river bed sediments in these river reaches. The C_{27}/C_{31} *n*-alkane ratio at the joint catchment outlet was lower than in the N subcatchment but similar to the ratios found in the S subcatchment (Fig. 5a), indicating mixing of OM from the two tributaries as well as input from

401 permanent grasslands situated on the steep slopes in the lower reaches of the river corridor.

402

403

3.3 Impact of land use on sediment and C accumulation in lake bed sediments

404

405 Source apportionment modelling of the lake core sediments has shown different results, depending on the composition of the fingerprint signatures. Currently, there is a lack of 406 407 consensus within the sediment fingerprinting community on the most appropriate selection of fingerprint tracers (Sherriff et al., 2015; Zhang and Liu, 2016). Therefore, in this study, we 408 409 firstly included all tracers used in the PCA in the source apportionment analysis (Fig. 6a), 410 followed by just those tracers that encompassed the range of values represented in the 411 mixture (and therefore deemed as conservative) (Fig. 6b). The second approach included 412 C_{27} : C_{31} ratio, % C_{27} , % C_{29} and % C_{31} . As such we had 4 tracers (n) to apportion the contribution of five sources (n+1). In both cases, organic C from riparian woodland was a 413 414 major contributor to the lake sediments. In the second scenario, river bed sediments appeared 415 to make the second largest contribution to lake core sediments over the past 60 years (Fig. 416 6b). However, as river bed sediments are also dominated by woody vegetation, as shown in 417 Fig. 2a above, they can be considered 'equal to' woodland signatures in this apportionment 418 model, due to the restricted number of very selective tracers. However, in both modelling 419 outcomes, the important contribution of organic matter from permanent grassland, which 420 occupies the steep slopes surrounding the lake, is very apparent (Fig. 6a).

421

422 Zhang and Liu (2016) also found that tracer selection greatly impacted the estimated source 423 contributions, due to a number of potential reasons, including i) tracer conflicts ii) tracer 424 measurement error and iii) differences in tracer conservativeness. Therefore, they proposed to 425 use multiple fingerprints to derive 'average' estimated source contribution proportions, 426 instead of just a single fingerprint set. While different sediment contributions can be obtained 427 with different fingerprint selection, recent studies (Palazón et al., 2015; Sherriff et al., 2015) 428 have found that inclusion of more tracers improved the source apportionment results. In this 429 study, modelling results based on the full set of tracers (Fig. 6a) allowed a finer distinction 430 between contributing land uses.

432 Conversely to lake core sediments, it was not possible to model the source apportionment of 433 river bed sediments satisfactorily as all potential tracers in bed sediments appeared to be outside the range of the potential sources. This apparently 'missing source' opens new lines 434 of enquiry for future research. At present we hypothesise that the 'missing source' may either 435 be due to the contribution of petrogenic C originating from the underlying bedrock (Galy et 436 437 al., 2015) or that the *n*-alkane signatures have been altered by autochtonous in-stream 438 production of OM (e.g. from algae) and by in-stream biological processing of river bed sediments (Chen et al., 2016). 439



proportion
 Fig. 6 Probability density function of sediment source apportionment sources from different
 land uses using a) all available tracers b) only tracers that encompass the full range of values
 present in the mixture for the application of the mixing model SIAR.

5

6 <u>3.4</u> Understanding the temporal variability of lateral C fluxes from land to water in 7 relation to land use change

8

9 Caesium-137 (¹³⁷Cs) activity was used to get an approximate dating profile for the lake bed 10 sediment core (Fig. 7a). The depth distribution of ¹³⁷Cs within the core was analysed and the 11 horizon containing peak activity was identified at 26 cm and was assumed to be associated 12 with the peak in bomb derived ¹³⁷Cs fallout attributed to 1963. The offset of bomb testing in 13 1952 was assumed to be associated with lake core depth at 34 cm.

14





Figure 7 Lake core profiles showing a)¹³⁷Cs activity used to date the lake sediment core and
interpret the observed changes in sediment composition over time b-h) sediment
fingerprinting characteristics with depth.

22 The combined use of *n*-alkane ratios and stable isotope signatures shows a variable 23 contribution of terrestrial vs. aquatic sources to sediment accumulation in individual lake core 24 increments over the past 60 years (Fig. 7b-h). The TAR ratio (Fig. 7b) indicates an increasing contribution of terrestrial-derived OM to the lake sediments, while the increasing 25 C_{27}/C_{31} ratio (Fig. 7d) indicates an increase of woody vegetation contribution over the same 26 27 time period. This is corroborated by the fact that the signal from PAQ *n*-alkane proxy lies 28 within the range of 0.01 to 0.23 (Fig. 7c), deemed indicative of terrestrial plants (Silliman and 29 Schelske, 2003) and suggesting mainly allochthonous source of sedimentary OM. The depletion in δ^{13} C values with depth (Fig. 7f) may also indicate an increasing input of 30 isotopically lighter soil-derived dissolved inorganic C (Meyers, 2003) - and thus increasing 31 32 terrestrial input of C from soil erosion. However, it may also be due to the preferential loss of the light isotope $({}^{12}C)$ through microbial respiration over time (Beniston, et al. 2014). In any 33 case correlation between OEP and TAR ratios (p<0.001, $R^2=0.53$) suggests reduced OM 34 decomposition associated with increasing terrestrial contribution of OM over the past 60 35 36 years (Zech et al., 2013), which may be linked to a higher C/N ratio – and therefore lower 37 bioavailability of woodland and grassland derived OM.

While increased $\delta^{15}N$ enrichment (Fig. 7g) may be indicative of increased net primary 38 production associated with the reported eutrophication of Loe Pool since the 1960s (Dinsdale, 39 40 2009), it is not likely to be due to an increase in N fixing cyanobacteria, which directly fix atmospheric N₂ and therefore lead to lower $\delta^{15}N$ signatures in sediments and would be 41 expected to lead to an increased $\delta^{13}C$ signature due to enhanced NPP (Meyers and Lallier-42 Vergès, 1999; Meyers, 2003). In addition, increased δ^{15} N enrichment may be associated with 43 enhanced denitrification (preferential loss of the light ¹⁴N isotope) in anoxic lake bottom 44 waters (Meyers, 2003) or with higher natural abundance ¹⁵N-enriched signatures originating 45 46 from faeces from farmyard manures (Senbayram et al. 2008) and septic tanks (Collins, et al. 47 2013; 2014).

However, it is acknowledged that $\delta^{13}C$ enrichment and C/N ratios are not always indicative of 48 sources as they can be affected by degradation (Ranjan et al., 2015). Laceby et al. (2015) 49 found that while $\delta^{15}N$ bulk isotopic signatures of sediment sources exhibited non-50 conservative behaviour, δ^{13} C signatures appeared to be more stable. Fang et al. (2014) 51 52 observed that significant macrophyte or riparian-aquatic OM inputs may lead to higher $\delta^{15}N$ and δ^{13} C values in lake sediments, thus confounding our ability to distinguish between the 53 terrestrial and aquatic input of OM on the basis of bulk stable ¹³C and ¹⁵N isotopic signatures 54 alone. Therefore, compound-specific stable ¹³C isotope (CSSIA) signatures of plant-derived 55 56 biomarkers are currently explored as more suitable tracers, as the isotopic signatures of 57 individual molecules are likely to be more conservative than bulk stable isotopes alone (Fang 58 et al., 2014; Tao et al., 2016).

In this study, the combined use of *n*-alkanes and bulk stable ${}^{13}C$ and ${}^{15}N$ isotopes detected increased terrestrial input of sediment and increased lake eutrophication over the past 60 years, with terrestrial grass and woody plant-derived *n*-alkanes being more indicative of OM sources, and stable ${}^{13}C$ and ${}^{15}N$ isotopes being more indicative of in-stream and in-lake

processes. The application of compound-specific stable isotope $\delta^{13}C$ and $\delta^{2}H$ signatures of 63 specific *n*-alkane molecules, as opposed to a separate use of *n*-alkane chain length and bulk 64 stable ¹³C and ¹⁵N isotopes, may help to better differential between aquatic and terrestrial 65 plant origin of organic matter in future work (Cooper et al., 2015) and thus quantify the 66 67 autochtonous vs. allochtonous organic matter contribution. Further improvements may be obtained by multi-molecular investigations using simultaneous application of different 68 biomarkers and ¹⁴C isotopes to constrain the transfer of C from land to the ocean (Feng et al., 69 70 2015).

71 The ability to discern OM contribution to aquatic environments from different land uses 72 found in this study, provides an important new tool for the understanding of OM fluxes from 73 land to water at catchment scales. Wohl et al. (2017) proposed a revised paradigm for the 74 understanding of the role of rivers in the transport and processing of terrestrial C, whereby 75 the active river channel and the riparian zone function as one coupled system - a river 76 corridor, in which riparian areas act 'as biogeochemical reactors that facilitate the speciation, 77 transformation, and opportunities for both long-term storage of carbon and mineralization to 78 the atmosphere'. Wohl et al. (2017) posit that while alteration of riparian zone is the most 79 significant and most highly altered aspect of lateral C dynamics, very little is known about 80 the sources and quantities of different kinds of OM stored within river corridors and how C 81 inputs have varied over decadal and millennial timescales as a results of human activities. In this study, we also found very close coupling between the aquatic sediments and the riparian 82 zone and our ability to discern between these sources provides a new opportunity to quantify 83 84 the lateral C fluxes at catchment scales. Coupling the fingerprinting approach explored in this 85 paper with future modelling of soil erosion rates is a promising new tool for quantifying these 86 lateral C fluxes at a range of scales.

87

88 **4.** Conclusions

89 This pilot study tested a new approach to quantify the lateral fluxes of OM from the terrestrial 90 to aquatic environments at a catchment scale. Here we evaluated the combined use of the 91 abundance and ratios of conservative plant-derived biomarkers *n*-alkanes and bulk stable 92 isotopes to distinguish between OM and sediment provenance from different environments. 93 While it was possible to distinguish between arable and temporary grassland, permanent 94 grassland, woodland, river and lake environments, it was not possible to distinguish between 95 arable land and temporary grassland, as these two land uses are part of regular rotations. 96 However, the combined use of biomarkers and stable isotopes allowed us to distinguish 97 between sediment sources from arable and permanent grassland land uses, which has not 98 been previously possible with the use of biomarkers alone. Furthermore, the combined use of 99 biomarkers and stable isotopes enabled us to detect the observed change in the lake trophic 100 status over the past 60 years and attribute this to changing land use, resulting in enhanced 101 sedimentation and nutrient flux from the terrestrial to the aquatic environments. These 102 enhanced lateral OM fluxes can be linked to agricultural intensification, resulting in higher soil erosion rates, over the same period. Moreover, we detected an increased contribution of 103

104 woody vegetation to the OM provenance over time, most likely indicating an increase in the 105 woody vegetation covering the near-stream riparian corridor. The new fingerprinting 106 approach successfully discriminated between terrestrial vs. aquatic C sources and when 107 coupled with quantitative estimates of soil erosion rates, it shows to be a promising new tool 108 for the understanding of lateral C fluxes from land to water at a range of scales. The close 109 coupling between OM provenance and riparian land use observed in this study underlines the 110 importance of the riparian zone for lateral C transfers and thus support the new holistic 111 conceptualization of 'river corridors' as critical zones linking the terrestrial and aquatic C 112 (Wohl et al., 2017).

113

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- 120

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Appendix 1 n-alkane data

			n-alk	ane co	oncent	ration	μgg	¹ soil													
Landus			C1	C1	C1	C1	C1	C2	C2	C2	C2	C2	C2	C2		C2		C3		C3	C3
e	Source	Code*	5	6	7	8	9	0	1	2	3	4	5	6	C27	8	C29	0	C31	2	3
			0.3	0.5	0.3	0.6		0.2	0.3	0.4	0.3	0.2	0.6	0.1		0.3		0.5		0.3	3.8
grass	soil	1.1	6	0	2	1	n/a	8	8	6	2	6	6	6	1.49	3	3.79	9	6.30	2	2
			0.3	0.8	0.5	0.7		0.4	0.7	0.3	0.3	0.1	0.8	0.3		0.5		0.5		0.3	3.8
grass	soil	1.2	6	7	5	7	n/a	3	1	8	8	8	2	6	1.61	0	3.93	7	6.63	5	5
			0.5	0.5	0.3	0.5		0.6	0.5	0.1	0.0	0.0	0.6	0.0		0.2		0.5		0.1	4.1
grass	soil	1.3	7	0	7	9	n/a	9	8	9	6	5	2	8	1.17	4	4.31	1	7.32	8	9
			0.1	0.2	0.2	0.3		0.4	0.3	0.1	0.2	0.2	0.3	0.0		0.4		0.3		0.2	2.1
grass	soil	1.4	8	7	3	7	n/a	0	5	9	7	8	8	3	0.73	8	2.20	4	4.13	5	5
			0.0	0.1	0.1	0.5		0.3	0.3	0.2	0.5	0.1	0.7	0.3		0.5		0.6		0.4	3.3
grass	soil	1.5	7	9	6	0	n/a	4	4	3	3	7	5	7	1.53	5	2.98	1	5.29	5	7
			0.2	0.3	0.2	0.5		0.8	0.4	0.1	0.0	0.3	0.3	0.0		0.2		0.3		0.2	1.9
arable	soil	2.1	8	3	9	6	n/a	2	0	5	9	9	7	9	1.28	7	2.37	5	3.81	4	8
-			0.0	0.0	0.0	0.2		0.1	0.2	0.1	0.2	0.1	0.3	0.1		0.2		0.3		0.2	1.9
arable	soil	2.2	9	8	9	7	n/a	9	1	7	8	1	9	4	0.84	4	2.13	4	3.65	3	9
			0.1	0.2	0.2	0.4		0.4	0.2	0.1	0.3	0.1	0.3	0.1		0.2		0.3		0.2	2.0
arable	soil	2.3	1	8	4	3	n/a	2	6	2	0	8	8	9	0.90	5	2.00	8	3.71	3	0

			0.0	0.1	0.1	0.2		0.3	0.1	0.1	0.3	0.1	0.4	0.1		0.2		0.2		0.2	1.8
arable	soil	2.4	8	0	2	6	n/a	1	7	0	0	8	3	8	1.08	8	1.93	7	3.34	4	1
			0.0	0.0	0.1	0.2		0.2	0.2	0.1	0.2	0.2	0.6	0.1		0.3		0.4		0.2	2.3
arable	soil	3.3	8	9	1	4	n/a	1	6	0	7	0	1	9	1.57	7	3.02	0	4.33	6	0
woodla			0.3	0.5	0.5	0.9		2.4	2.1	0.1	1.7	1.2	4.2	0.6	20.6	1.8	20.4	1.3	11.4	0.6	7.8
nd	soil	3.4	1	8	8	9	n/a	2	7	7	6	0	8	5	8	9	2	8	8	9	7
			0.4	0.4	0.5	0.9		2.3	1.9	0.4	0.3	1.0	2.2	0.2		3.2		0.7		0.4	1.4
ley	soil	5.2	0	9	9	6	n/a	4	1	7	8	4	0	0	0.49	2	3.81	0	6.21	4	9
			0.8	0.9	0.9	1.5		3.2	3.0	0.2	2.2	0.4	1.7	1.2		1.3		0.4		0.3	2.6
ley	soil	5.3	5	8	9	2	n/a	8	8	9	9	0	9	7	1.82	9	3.37	6	5.50	5	0
			0.2	0.2	0.2	0.4		2.0	0.9	0.2	0.1	0.3	0.1	0.3		3.2		0.8		0.4	2.9
ley	soil	5.4	5	3	9	8	n/a	3	9	8	4	7	7	6	1.98	7	4.01	0	5.68	6	9
			0.0	0.1	0.1	0.2		0.2	0.4	0.1	0.5	0.3	1.4	0.7		1.3		1.1		0.5	3.7
ley	soil	5.5	5	2	4	5	n/a	1	6	2	2	0	8	8	2.59	3	5.04	3	6.98	8	5
			0.0	0.1	0.1	0.2		0.2	0.2	0.0	0.2	0.2	0.5	0.2		0.4		0.5		0.4	2.8
ley	soil	6.4	8	2	5	9	n/a	0	4	9	1	5	7	6	1.48	5	3.08	8	5.14	0	4
			0.0	0.2	0.2	0.3		0.2	0.2	0.0	0.1	0.0	0.4	0.2		0.4		0.5		0.2	2.3
ley	soil	6.5	9	1	0	7	n/a	0	2	5	9	8	9	6	1.20	7	2.67	6	4.13	9	2
woodla			0.0	0.1	0.1	0.3		0.2	0.4	0.6	1.1	0.9	3.4	1.4	14.5	2.0	15.3	1.0		0.6	3.4
nd	soil	6.6	6	1	4	5	n/a	8	4	4	9	6	8	8	5	1	1	9	7.12	5	8
grass	soil	8.5	0.1	0.1	0.1	0.2	n/a	0.1	0.1	0.1	0.1	0.2	0.6	0.3	1.71	0.8	3.14	0.7	4.78	0.3	2.1

			5	4	3	1		6	9	1	9	8	4	7		5		7		6	4
arable	soil	9.1	0.0 7	0.1 7	0.1 3	0.2 9	n/a	0.2 8	0.1 2	0.0 6	0.1 8	0.0 9	0.5 2	0.1 3	1.05	0.3 7	3.15	0.4 5	5.66	0.3 3	3.0 3
woodla nd	soil	9.7	0.0 4	0.0 4	0.0 5	0.0 9	n/a	0.1 3	0.2 8	0.1 6	0.4 8	0.3 0	2.0 7	0.6 0	10.7 0	1.3 4	9.02	0.6 0	4.97	0.3 3	4.5 7
arable	soil	10.5	0.0 7	0.1 2	0.1 4	0.3 5	n/a	0.2 4	0.2 5	0.1 7	0.1 1	0.1 0	0.3 2	0.0 7	0.77	0.1 8	1.71	0.2 6	2.83	0.1 8	1.4 1
ley	soil	11.5	0.2 7	0.2 6	0.1 8	0.3 4	n/a	0.4 9	0.4 1	0.1 3	0.1 4	0.1 3	0.0 8	0.1 4	0.44	0.1 3	1.54	0.1 8	2.72	0.1 4	1.0 8
ley	soil	11.6	0.0 7	0.0 8	0.1 1	0.2 5	n/a	0.2 2	0.2 4	0.1 9	0.1 1	0.0 5	0.5 7	0.0 6	0.69	0.2 1	2.10	0.2 5	3.57	0.1 8	1.6 7
ley	soil	11.7	0.2 1	0.1 2	0.1 9	0.3 1	n/a	0.8 4	0.7 2	0.0 5	0.1 9	0.3 3	0.1 4	0.1 8	0.88	0.2 3	2.23	0.5 1	3.21	0.2 2	1.2 2
grass	soil	11.8	0.1 1	0.0 3	0.0 5	0.1 8	n/a	0.1 9	0.4 5	0.1 8	0.6 6	0.5 6	1.4 3	0.6 6	4.75	0.9 0	6.35	0.6 8	4.21	0.3 0	1.6 5
woodla nd	soil	12.5	0.0 4	0.0 8	0.1 1	0.1 8	n/a	2.1 9	0.8 6	0.0 6	0.2 5	0.3 1	2.0 9	0.9 4	5.79	0.1 3	10.8 1	1.0 8	6.01	0.4 7	1.6 7
arable	soil	13.3	0.1 8	0.2 3	0.3 0	0.5 6	n/a	1.2 7	1.3 7	0.1 5	0.2 9	0.1 0	0.2 5	0.4 6	0.32	0.6 8	3.02	0.4 8	4.75	0.2 8	2.5 0
arable	soil	13.4	0.3 9	0.5 6	0.4 5	0.9 2	n/a	2.2 3	2.4 6	0.5 4	0.4 3	0.9 5	0.5 0	0.0 9	1.01	1.4 2	6.18	0.1 6	4.46	0.2 4	1.7 6

			0.2	0.3	0.3	0.6		1.2	1.2	0.1	0.1	0.1	0.4	0.0		0.6		0.3		0.2	1.6
arable	soil	13.5	4	1	8	7	n/a	0	1	5	4	2	6	8	0.84	1	2.25	3	3.51	1	8
			0.1	0.1	0.1	0.3	0.1	0.2	0.4	0.2	0.0	0.1	0.1	0.1		0.0		0.1		0.1	1.4
arable	soil	13.6	5	0	6	7	4	6	5	3	7	5	7	1	0.54	9	1.74	6	2.39	8	0
			0.1	0.0	0.0	0.1	0.0	0.1	0.0	0.1	0.1	0.1	0.5	0.1		0.2		0.1		0.0	0.2
rivor	unstroom	4	0.1	0.0	0.0	0.1	0.0	0.1	0.0	0.1	0.1	0.1	0.5	0.1	2.25	0.2	2 20	0.1	0.00	0.0	0.5
IIVel	upsiteani	4	0	4	5	5	5	0	0	0	/	4	9	/	2.23	1	2.20	9	0.90	0	1
			0.1	0.0	0.0	0.3	0.2	0.3	1.1	0.3	0.2	0.2	1.2	0.3		0.4		0.1		0.1	0.8
river	midstream	6	2	9	9	2	6	3	4	2	7	7	5	5	5.61	7	5.50	2	2.13	7	2
	downstraa		0.0	0.0	0.1	0.2	0.1	0.2	0.2	0.1	0.5	0.2	1.0	0.2		0.7		0.5		0.1	1 1
rivor	m	7	0.0	0.0	0.1	0.2	0.1	0.2	0.5	0.1	0.5	0.2	1.9	0.5	8.00	0.7	7 18	0.5	3 10	0.1	1.1
IIVel	111	/	9	/	0	5	9	5	1	4	5	0	9	4	0.09	4	7.40	5	5.10	4	/
			0.2	0.1	0.5	0.6	0.2	0.2	0.8	0.2	0.2	0.3	1.2	0.3		0.4		0.5		0.2	1.0
river	upstream	9	5	2	2	5	6	6	9	0	9	0	2	1	5.15	9	6.13	6	2.95	4	1
			0.2	0.0	0.1	0.2	0.2	0.2	0.2	0.1	0.7	0.2	2.0	0.7		1.0		0.6		0.1	1.7
	midatacom	11	0.2	0.0	0.1	0.2	0.2	0.2	0.5	0.1	0.7	0.2	2.0	0.7	0 57	1.0	0.42	0.0	1 22	0.1	1./
liver	musueam	11	0	/	1	4	5	0	0	2	5	0	1	0	0.37	/	9.45	5	4.52	2	0
	downstrea		0.6	0.2	0.5	0.7	0.1	0.1	0.1	0.1	0.2	0.3	1.0	0.0		0.2		0.4		0.2	0.8
river	m	13	9	4	2	1	9	8	5	7	1	7	5	6	5.18	0	5.95	4	2.53	6	9
			0.1	0.0	0.2	0.5	0.2	0.2	0.2	0.2	0.2	0.7	1.2	0.2		0.1		0.2		0.2	1.7
	outlat	OUT	0.1	0.0	0.2	0.5	0.3	0.3	0.2	0.3	0.3	0.7	1.3	0.2	5.07	0.1	1 75	0.3	262	0.2	1./
river	outlet	001	5	9	/	0	3	3	9	5	0	1	3	0	5.07	0	4.75	5	2.05	1	5
			0.1	0.4	0.7	0.7	0.5	0.2	0.6	0.3	1.5	0.6	3.8	0.8	18.9	1.1	17.3	1.1		0.4	6.7
lake	1	0-4 cm	0	9	2	0	9	6	4	3	4	1	8	5	4	8	4	5	9.14	5	5
lake	2	4-8 cm																			<u> </u>
			0.5	0.4	1.1	0.9	0.3	0.5	0.5	0.6	0.9	1.1	5.2	1.3	20.4	1.0	20.3	2.6	12.3	0.3	4.7

			4	5	7	3	4	6	4	5	7	8	4	9	8	6	0	6	9	1	5
lake	3	8-12 cm	0.1 6	0.5 0	1.0 4	0.8 7	0.9 8	0.3 3	1.0 5	0.4 2	1.4 3	0.5 7	4.9 6	0.7 5	19.0 2	0.3 1	18.5 9	1.2 9	11.5 2	0.2 7	6.1 8
lake	4	12-16 cm	0.3 1	0.3 3	0.4 3	0.7 0	0.5 4	0.0 8	0.6 2	1.0 3	1.2 0	2.8 6	5.1 5	0.8 9	21.9 7	0.1 3	20.7 4	0.5 0	14.3 9	0.1 6	5.7 0
lake	5	16-20 cm	0.1 3	0.4 4	0.6 6	0.6 0	1.0 9	1.5 1	1.1 4	0.3 6	2.2 3	0.4 6	4.7 6	0.6 4	17.2 7	2.1 5	17.8 2	1.5 9	12.4 0	1.0 3	4.1 2
lake	6	20-24 cm	0.4 2	0.4 8	0.7 0	0.6 2	0.7 2	3.1 6	1.9 7	0.3 9	2.7 3	2.9 1	0.5 7	0.8 0	15.7 2	0.8 0	16.1 1	0.4 4	12.8 5	0.0 5	6.5 8
lake	7	24-28 cm	0.1 1	0.3 1	0.7 0	1.0 6	1.6 2	3.3 5	1.5 0	0.2 2	3.5 2	0.4 0	4.6 0	1.1 6	16.6 8	3.9 7	15.7 8	0.4 5	12.3 2	1.3 8	4.4 7
lake	8	28-32 cm	0.4 1	0.5 9	1.2 7	1.0 5	1.1 6	2.0 7	0.8 1	1.0 0	2.0 3	3.9 4	4.5 0	0.1 7	17.2 1	2.2 6	15.3 6	0.4 5	12.0 9	0.9 9	3.5 5
lake	9	32-36 cm	0.5 6	0.8 2	1.9 6	1.5 8	1.2 1	1.8 5	0.8 9	1.2 6	1.9 8	3.8 7	5.6 2	0.6 3	18.9 9	0.5 9	17.7 7	0.8 9	12.8 2	0.7 9	4.0 3
lake	10	36-40 cm	0.1 9	0.3 8	3.5 4	0.7 5	1.6 8	0.1 6	1.0 4	0.4 0	1.5 8	1.2 1	4.6 9	1.0 5	13.6 0	3.2 5	15.7 3	1.3 6	13.1 0	0.7 7	5.8 0
lake	11	40-44 cm	0.0 3	0.1 2	1.4 9	0.3 1	0.6 2	0.5 0	0.5 3	0.2 0	1.4 7	1.3 5	2.4 5	0.5 6	7.07	1.5 2	9.95	0.9 3	9.05	0.4 3	2.9 7
lake	12	44-48 cm	0.2 0	0.2 7	1.6 7	1.1 0	1.0 0	1.6 6	0.7 8	0.5 6	2.0 8	1.3 8	3.6 2	0.7 8	10.4 2	3.2 3	13.8 8	1.0 6	12.8 8	0.5 4	4.3 3

transect and core number for soil

* samples

location for stream bed sediment samples

depth increment for lake core

Appendix 2 – stable isotope data

				δ^{15} N		$\delta^{13}C$
Landuse	Source	Code	% N	%	% C	‰
grass	soil	1.1	0.42	5.20	4.12	-27.81
grass	soil	1.2	0.49	5.78	5.53	-27.27
grass	soil	1.3	0.57	3.26	6.50	-28.33
grass	soil	1.4	0.44	6.37	4.13	-27.34
grass	soil	1.5	0.61	6.66	6.02	-28.02
arable	soil	2.1	0.40	4.81	4.14	-27.01
arable	soil	2.2	0.30	3.72	2.99	-28.00
arable	soil	2.3	0.38	7.59	3.99	-27.00
arable	soil	2.4	0.37	5.05	3.55	-26.98
arable	soil	3.1	0.39	4.97	3.56	-27.85
arable	soil	3.2	0.37	5.93	3.66	-27.05
arable	soil	3.3	0.25	4.80	2.57	-27.69
woodland	soil	3.4	0.84	5.56	10.36	-28.11
woodland	soil	3.4b	0.69	5.63	8.76	-27.71
woodland	soil	3.4c	0.49	4.70	6.92	-28.35
ley	soil	4.1	0.36	5.07	3.35	-28.27
ley	soil	4.2	0.41	5.35	4.08	-28.13
ley	soil	4.3	0.47	5.25	4.78	-28.30
ley	soil	4.4	0.43	5.09	4.10	-28.43
ley	soil	5.1	0.41	5.10	4.17	-27.95
ley	soil	5.2	0.40	5.28	3.69	-28.37
ley	soil	5.3	0.41	5.69	4.09	-28.01
ley	soil	5.4	0.43	5.20	4.19	-28.52

ley	soil	5.5	0.57	4.90	5.87	-28.58
ley	soil	6.1	0.35	4.87	3.15	-27.99
ley	soil	6.2	0.32	3.82	3.00	-28.72
ley	soil	6.3	0.33	4.63	3.02	-28.14
ley	soil	6.4	0.40	4.85	3.75	-28.41
ley	soil	6.5	0.44	3.96	4.18	-28.52
woodland	soil	6.6	0.64	5.58	7.16	-28.65
arable	soil	7.1	0.35	5.53	3.25	-27.26
arable	soil	7.2	0.21	5.39	1.83	-27.29
ley	soil	7.3	0.23	5.69	2.20	-26.96
ley	soil	7.4	0.31	6.07	2.70	-27.47
ley	soil	7.5	0.38	5.71	3.52	-27.88
grass	soil	7.6	0.59	6.27	5.56	-28.28
grass	soil	7.7	0.48	7.07	5.10	-28.14
ley	soil	8.1	0.47	5.86	4.60	-28.28
ley	soil	8.2	0.38	6.79	3.42	-28.17
arable	soil	8.3	0.32	6.11	2.78	-27.29
arable	soil	8.4	0.32	5.53	2.92	-27.42
grass	soil	8.5	0.68	5.20	6.73	-28.38
grass	soil	8.6	0.71	4.16	6.88	-28.33
arable	soil	9.1	0.35	5.07	3.40	-27.78
arable	soil	9.2	0.32	5.90	2.90	-28.19
grass	soil	9.3	0.92	7.29	8.28	-29.15
grass	soil	9.4	0.60	4.99	5.77	-28.42
grass	soil	9.5	0.48	5.33	4.48	-28.38
grass	soil	9.6	0.47	6.24	4.45	-27.62

woodland	soil	9.7	0.50	5.94	4.98	-28.51
ley	soil	10.1	0.38	6.89	3.22	-28.42
arable	soil	10.2	0.31	6.38	2.64	-27.26
arable	soil	10.3	0.35	5.43	2.90	-27.56
arable	soil	10.4	0.27	5.39	2.34	-27.86
arable	soil	10.5	0.34	4.95	2.98	-27.82
woodland	soil	10.6	0.77	3.80	9.59	-28.25
arable	soil	11.1	0.22	3.62	2.08	-27.85
arable	soil	11.3	0.31	5.20	2.72	-28.00
arable	soil	11.4	0.35	5.17	2.89	-27.84
ley	soil	11.5	0.28	5.95	2.12	-27.69
ley	soil	11.6	0.37	6.01	3.45	-28.09
ley	soil	11.7	0.32	5.33	2.56	-27.79
grass	soil	11.8	0.43	4.10	4.80	-28.84
ley	soil	12.1	0.43	7.64	3.67	-28.82
ley	soil	12.3	0.41	6.66	3.56	-28.46
ley	soil	12.4	0.38	6.03	3.06	-28.23
woodland	soil	12.5	0.50	1.77	8.68	-28.02
arable	soil	13.1	0.31	6.09	2.64	-27.74
arable	soil	13.2	0.38	6.52	3.36	-27.87
arable	soil	13.3	0.30	6.24	2.56	-28.13
arable	soil	13.4	0.29	5.38	2.46	-27.65
arable	soil	13.5	0.29	4.39	2.25	-27.95
arable	soil	13.6	0.32	5.91	2.63	-27.69
arable	soil	14.1	0.29	5.57	2.24	-27.84
arable	soil	14.2	0.31	5.59	2.73	-27.74

arable	soil	14.3	0.34	5.71	2.95	-27.04
arable	soil	14.4	0.37	6.12	3.28	-27.18
arable	soil	14.5	0.38	6.40	3.47	-28.00
river	Upstream	4	0.11	3.41	1.16	-28.15
river	Midstream	6	0.16	3.34	2.08	-28.23
river	Downstream	7	0.21	4.98	2.74	-28.71
river	Upstream	9	0.24	6.48	2.82	-28.34
river	Midstream	11	0.28	4.63	3.68	-28.28
river	Downstream	13	0.19	4.45	2.32	-28.33
river	outlet	OUT	0.13	4.52	1.55	-27.91
lake	0-2 cm	1_1	0.76	6.58	8.32	-28.16
lake	0-2 cm	1_1	0.75	6.31	8.44	-28.20
lake	2-4 cm	1_1	0.76	6.68	8.46	-28.18
lake	4-6 cm	1_2	0.72	6.28	7.84	-28.27
lake	6-8 cm	1_3	0.66	5.76	7.71	-28.24
lake	8-10 cm	1_4	0.69	6.26	7.85	-27.99
lake	10-12 cm	1_5	0.67	6.36	7.53	-28.04
lake	12-14 cm	1_6	0.69	6.17	7.41	-27.99
lake	14-16 cm	1_7	0.68	6.90	7.37	-27.97
lake	16-18 cm	1_8	0.69	6.72	7.65	-27.96
lake	18-20 cm	1_9	0.78	6.94	9.40	-27.73
lake	22-24 cm	1_10	0.77	6.69	9.15	-27.21
lake	24-26 cm	1_11	0.80	6.78	8.50	-27.23
lake	26-28 cm	1_12	0.72	6.33	7.66	-27.33
lake	28-30 cm	1_13	0.78	6.25	8.41	-27.09
lake	32-34 cm	1_14	0.72	6.26	7.58	-27.16

lake	34-36 cm	1_15	0.74	6.63	7.75	-27.15
lake	36-38 cm	1_16	0.73	6.87	7.74	-27.19
lake	38-40 cm	1_17	0.65	6.05	7.06	-27.44
lake	40-42 cm	1_18	0.63	6.58	6.58	-27.68
lake	42-44 cm	1_19	0.57	5.94	5.95	-28.01
lake	44-46 cm	1_20	0.54	6.10	5.72	-27.83
lake	46-48 cm	1_21	0.61	5.88	6.39	-27.58
lake	48-50 cm	1_22	0.58	6.03	6.04	-27.35
lake	50-52 cm	1_23	0.56	6.06	5.51	-27.40
lake	52-54 cm	1_24	0.64	5.76	5.89	-26.86
lake	54-56 cm	1_25	0.62	5.73	5.80	-26.84