### Response to anonymous reviewer 3

The authors thank the reviewer for their positive comments and helpful suggestions for improvements. Our response to each individual point can be found below marked in red.

Changes to the manuscript as a result of the responses can be found below marked in blue.

# **General comments**

This manuscript is well written with clearly stated objectives and hypotheses. Some specific and technical comments are below. I recommend acceptance of the manuscript for Biogeosciences after addressing the comments below.

Specific comments P16610 L7-10: What could be the reason for no NO3 production. Please elaborate.

The most likely explanation is time, the experiments were carried out over a period of less than two hours. In this time, the soil went from having low soil moisture (around 8%) to be saturated by the rainfall. Both moisture limitation and potential oxygen limitation (due to waterlogging later in the experiment) can inhibit rates of nitrification. It is likely that after the end of the experiment, as the slope began to drain that rates of nitrification would increase. Text explaining this can be added to the manuscript to clarify this point.

Text has been added in lines 166-169, reading "This is most probably due to the soils having a low moisture content (~8%) at beginning the experiments, which then become rapidly saturated. Both low moisture and potential oxygen limitation (due to waterlogging) later in the experiments could inhibit rates of nitrification."

P16617 L5-8: This statement is unclear. Revise it.

The sentence will be clarified to read: "There were no significant increases in 270:360 nm fluorescence from any of the vertical percolated flow pathways in the experiment run at 5° and 120 mm  $h^{-1}$  rainfall, or from the base of the slope (outlet 4) in the 10°, 60 mm  $h^{-1}$  experiment."

Changed as stated above (lines 328-330).

P16617 L19: Suggest including the data as a table/figure

We would prefer to add the median value of each of the tested datasets into the text rather than add as a figure or table. As a result, the data is then visible to the reader but space in the manuscript is not taken up by data which was not statistically significant, as we already have 9 figures in the manuscript.

Values have been added into the text in lines 340 and 341.

P16618 L28: Suggest including the data as a table/figure

A table can be added to display this data to support the text.

Table 3 has been added to the manuscript and is referred to in line 379.

P16621 L20-24: Can the same result be expected in the natural field situation? If not, please add some insights on these lines.

This result would be also expected in a field scenario, providing that the measured material was transported rapidly from the application area. As more time passes or with increasing distance from the application area (i.e. longer transport times), it is more likely that transformations will occur altering the slurry material. Text can be added here to clarify that this could apply in a natural scenario as well as in our experimental setting.

Text has been added on lines 442-443, "This result can also be expected to be observed in a field scenario where material is rapidly transported from an area of slurry application."

P16624 L9-12: Does this dilution effect also reflect in the C and N content of the slurry reported in Table 1

The dilution effect is not reflected in the bulk C and N data presented in Table 1 as this analysis was carried out on freeze-dried samples of the slurry and so all of the data in table 1 is expressed on a dry-weight basis. Conversely, the fluorescence measurements were carried out on the slurry its original form and so any extra water in the samples could influence the results.

# No changes necessary.

# **Technical corrections**

P16605 L20: Replace 'environmental' with 'environment'

Will be changed as suggested

Changed as stated (line 55).

P16608 L13: Change 'is was' to 'was'

Will be changed as suggested

Changed as stated (line 121).

P16615 L24: It appears that 'is' doesn't fit well in this sentence.

Agreed sentence will be amended to "This plot highlights that the difference between the fluorescence spectra of the slurry and the control soil is the magnitude of the peak at excitation wavelength 270 nm...."

Changed as stated (line 293)

P16617 L2: Change 'raised significantly above' with 'significantly higher than'

Agreed, this will be amended to "significantly greater than"

Changed as stated (line 325)

P16619 L1: Something is missing between 'pattern' and 'the'

Agreed, this sentence will be amended to "Results showed a spatial pattern in the distribution of the ratio..."

Changed as stated (line 373).

P16619 L16: Table 1 or Table 2?

Thank you, this should refer to table 2, this will be corrected.

Corrected.

Fig. 4, 5, 6, 8: Suggest increasing the font size of the axis labels and legends.

Font size will be increased as suggested.

Font size has been increased for all of the stated figures.

- 1 Runoff- and erosion-driven transport of cattle slurry: linking molecular tracers to hydrological
- 2 processes
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#### 16 Abstract

The addition of cattle slurry to agricultural land is a widespread practise, but if not correctly managed 17 18 it can pose a contamination risk to aquatic ecosystems. The transport of inorganic and organic 19 components of cattle slurry to watercourses is a major concern, yet little is known about the physical 20 transport mechanisms and associated fluxes and timings of contamination threats. Therefore, the aim 21 of the study was to ascertain the importance of flow pathway partitioning in the transport (fluxes and 22 timing) of dissolved and particulate slurry-derived compounds with implications for off-site 23 contamination. A series of rainfall-runoff and erosion experiments were carried out using the TRACE 24 (Test Rig for Advancing Connectivity Experiments) experimental hillslope facility. The experiments 25 allowed the quantification of the impact of changing slope gradient and rainfall intensity on nutrient 26 transport from cattle slurry applied to the hillslope, via surface, subsurface and vertical percolated 27 flow pathways, as well as particulate transport from erosion. The dissolved components were traced 28 using a combination of ammonium (NH<sub>4</sub><sup>+</sup>) and fluorescence analysis, while the particulate fraction was 29 traced using organic biomarkers,  $5\beta$ -stanols. Results showed that rainfall events which produced 30 flashy hydrological responses, resulting in large quantities of surface runoff, were likely to move 31 sediment and also flush dissolved components of slurry-derived material from the slope, increasing the contamination risk. Rainfall events which produced slower hydrological responses were 32 33 dominated by vertical percolated flows removing less sediment-associated material, but produced leachate which could contaminate deeper soil layers, and potentially groundwater, over a more 34 prolonged period. Overall, this research provides new insights into the partitioning of slurry-derived 35 36 material when applied to an unvegetated slope and the transport mechanisms by which contamination risks are created. 37

40 Contamination of water bodies due to poor agricultural management is a global problem that affects 41 aquatic ecosystems from individual catchments through to estuaries and oceans, and impacts human 42 water supplies (e.g. Mitsch et al., 2001; Diaz and Rosenberg, 2008; Osterman et al., 2009; HELCOM, 43 2009). The ongoing introduction of legislation aimed at improving aquatic ecosystems (e.g. European 44 Union Water Framework Directive) has focussed attention on efforts to decrease inputs of pollutants 45 into surface and groundwater. Transport of inorganic and organic nutrients from livestock slurries to 46 water courses are a particular concern, typically associated with over-application or poorly timed 47 application of organic fertilisers to agricultural land (Dungait et al., 2012;Chadwick and Chen, 2003). The inorganic component of slurries (mainly nitrogen, N) is generally considered as more soluble and 48 49 directly bioavailable compared with the organic fraction, and specific forms such as nitrate are also 50 more liable to immediate loss by leaching into watercourses (e.g. Chambers et al., 2000;Delconte et 51 al., 2014; Quemada et al., 2013). However, studies have shown that the organic fraction can provide a 52 large proportion of the total N load in some catchments and a smaller yet environmentally significant 53 fraction in N-enriched waters (Durand et al., 2011; Willett et al., 2004; Johnes and Burt, 1991). This 54 excess N leads to eutrophication of surface waters, resulting in algal blooms and reduced oxygen levels. Sutton (2011) reported that excess N in the environmental costs the EU between €70-320 55 56 billion per annum, more than double the value that the N fertilisers provide to EU farms in terms of 57 production. Combating and mitigating against these problems is extremely costly; the UK spends up 58 to £300 million each year cleaning water courses, equating to up to 2% of the gross agricultural output 59 (Pretty et al., 2000). Given these severe environmental and financial consequences it is important to 60 better understand the main transport pathways, fluxes and transit times of pollutants from livestock 61 slurries into watercourses to develop effective mitigation strategies. Slurry transport has been 62 quantified mainly in terms of the inorganic fraction (e.g. nitrate and phosphate) and associated 63 pathogens (e.g. coliforms) (e.g. Edwards et al., 2012;Coelho et al., 2012;Eastman et al., 2010), but 64 there is a relative lack of understanding of physical transport mechanisms and the associated fluxes65 and timings of contamination threats from areas treated with livestock slurries.

66 The transport of slurry components on land is primarily controlled by the hydrological and erosion 67 regime operating at the site of application. The dynamics of flow pathway partitioning during storm 68 events and the consequential output of agricultural contaminants has been explored (Delpla et al., 69 2011; Blanchard and Lerch, 2000; Gao et al., 2004; e.g. Zhang et al., 1997; Malone et al., 2004), but there 70 is a lack of experimental data quantifying contaminant export via individual flow pathways due to 71 methodological challenges. In a field context it is challenging to monitor multiple flow pathways 72 without destructive sampling, although new research platforms are now making this type of research 73 more possible (see Peukert et al. (2014) for an example). There are also challenges when choosing 74 field sites that represent transport regimes across different slope angles or to account for other 75 environmental variables. Therefore, laboratory flume experiments have been widely used to 76 investigate questions relating to hydrology and erosion in conjunction with solute and sediment 77 transport, although most studies to date have only measured surface runoff and/or vertical drainage 78 (e.g. Montenegro et al., 2013;Guo et al., 2010;Aksoy et al., 2012;Asam et al., 2012). Our previous work 79 using a one-dimensional soil column revealed the rapid partitioning of livestock slurries in the soil-80 water system into sediment-associated material remaining on or close to the surface and dissolved 81 components which moved rapidly through the soil by leaching (Lloyd et al., 2012). Given this 82 partitioning into surface and subsurface components and its potential importance for contamination 83 of downstream aquatic and soil environments, we aim to quantify the relative fluxes of different slurry 84 compounds driven by surface (overland flow and erosion) and subsurface (throughflow and leaching) 85 flow pathways during a series of experiments in which rainfall rate and slope angle vary. We vary 86 rainfall rate and slope angle in order to simulate hydrological variations which may result in differential 87 partitioning of flow pathways.

88 In this study we use biogeochemical biomarker analysis in combination with controlled, large-scale 89 rainfall-simulation experiments to quantify the relative fluxes of slurry components through different 90 flow pathways within a slope system. In particular, we monitored flow rates in three pathways -91 surface runoff, subsurface throughflow and vertical percolated – over the course of different rainfall 92 simulation experiments in which we varied the slope gradient and rainfall intensity and duration. We 93 also monitored erosion rates in the surface runoff component of the flow. Within samples of water 94 discharging from each flow pathway we measured concentrations of ammonium  $(NH_4^+)$  and the 95 fluorescence spectra as tracers of the dissolved (soluble) component of the slurry. Samples of eroded 96 sediment and in situ soil cores were analysed for total nitrogen (TN), carbon (TC) and 5β-stanols, which 97 have been shown to be an effective and unequivocal biomarker of particulate slurry material (see 98 Lloyd et al., 2012). The aim of the study was to ascertain the importance of flow pathway partitioning 99 in the transport (fluxes and timing) of dissolved and particulate slurry-derived compounds with 100 implications for off-site contamination.

#### 101 **2. Materials and methods**

# 102 2.1 Experimental set-up

103 A series of rainfall-runoff and erosion experiments were carried out using TRACE (Test Rig for 104 Advancing Connectivity Experiments) at the University of Bristol (described in detail in Michaelides et 105 al., 2010). The experimental facility consists of a dual-axis soil slope measuring 6 m × 2.5 m, with a soil 106 depth of 0.3 m. The angle of the two soil containers was manipulated in order to simulate different 107 slope gradients. Beneath the soil layer a wire mesh and geotextile layer separates the soil from a 108 2.5 m<sup>3</sup> gravel layer. The slope was accompanied by a six nozzle rainfall simulator fitted with full-cone 109 nozzles (Lechler, Germany) and suspended 2.5 m above the soil to simulate different rainfall 110 intensities. Water transported via surface, subsurface and vertical percolated pathways was 111 monitored via four pairs of sampling outlets shown in Figure 1. A series of four slurry-treated 112 experiments were carried out varying combinations of slope angle (5° and 10°) and rainfall intensity

113 (60 mm  $h^{-1}$  and 120 mm  $h^{-1}$ ). The intensity and duration of the rainfall simulation were co-varied such that the total volume of rainfall applied was equal between experiments (60 mm h<sup>-1</sup> for 100 min and 114 115 120 mm h<sup>-1</sup> for 50 min). The slope angles were chosen within a realistic range found in agricultural 116 settings. The rainfall intensities were chosen to test the impact of short duration, high intensity rainfall 117 events on slurry transport because extreme storm events have a disproportional impact on the 118 transport of dissolved and particulate contaminants to water courses (Evans and Johnes, 119 2004; Haygarth et al., 2005; Rozemeijer and Broers, 2007; Haygarth et al., 2012). Rainfall simulations 120 using 60 mm h<sup>-1</sup> and 120 mm h<sup>-1</sup> were used so that a systematic doubling of rainfall intensity could be 121 tested in conjunction with a change in slope gradient. 60 mm h-1 was chosen as this is-was the lowest 122 intensity which could be stably simulated that was crucial for ensuring controlled experiments. 123 Therefore we effectively substituted simulation time for intensity.

Before each experiment the slope was packed with the same silt loam soil (34% sand, 37% silt and 27% clay,  $d_{50}$ = 200 µm) and compacted evenly across the surface of the slope. Even compaction was achieved for each experiment by following the method outlined in Michaelides et al. (2010), where layers of soil were added and compacted using a 2.5 m tamping board which was moved up and down the length of the slope. The average final bulk density achieved using this procedure was ~ 1.5 g cm<sup>-3</sup>.

129 Cattle slurry collected from a commercial dairy farm was applied to the top 1 m of the experimental slope at an application rate of 5 L m<sup>-2</sup> mimicking typical field application rates (CSF Evidence Team, 130 131 2011). Table 1 shows the chemical composition of the cattle slurry and the initial soil used for all of the experiments. Two additional control experiments with no slurry treatment were also carried out, 132 133 one at each of the tested slope angles (5 and 10°). Both of the control experiments were carried out 134 using 60 mm h<sup>-1</sup> rainfall intensity. During each experiment rainfall was applied and discharge from the slope was monitored via the three flow pathways: 1. surface runoff, 2. subsurface throughflow and 135 136 3. vertical percolated flow, in order to investigate the transport of slurry-derived compounds. Eroded 137 sediment was separated from the surface runoff using the protocol outlined below and kept for analysis. At the end of the rainfall simulation soil cores were taken from the slope according to the
sampling strategy outlined in Section 2.2. The soil cores and eroded sediment were used to investigate
the erosion and within-slope deposition of slurry-derived particulates.

141 *2.2 Sampling* 

142 Discharge from each flow pathway was determined at regular intervals during the course of the rainfall 143 simulations. Discharge from the surface runoff pathway was monitored by logging water depth using 144 a 'V'-notch weir and capacitance depth probe, at 1 min intervals. The water depth was converted to 145 discharge by a pre-calibrated stage-discharge relationship for the 'V'-notch weir. Additional manual 146 samples of the surface runoff were taken at ~5 min intervals to determine sediment concentration 147 and to obtain eroded sediment samples for analysis. The water and sediment were separated using a 148 centrifuge (2400 rpm, 30 minutes) and the water samples were filtered through 0.45 µm filter 149 (Whatman, cellulose acetate) then stored at -18°C until analysis was carried out. Flow discharge from 150 the subsurface and percolated pathways was monitored manually at ~5 min intervals by timing the 151 flow of a known volume of water.

At the end of the rainfall simulations, 18 soil cores were taken over a regularly spaced 3 × 6 sample grid leaving a buffer of 0.5 m around the slope perimeter to avoid edge effects (samples 0.75 m apart across slope, 1 m apart downslope), to a depth of 5 cm. Only the top 5 cm were sampled because pilot experiments and previous research has shown that this section is the most important in terms of the transport of slurry-derived material (See Lloyd et al., 2012). The soil cores and eroded sediment were freeze-dried and stored at -18°C until analysis.

158 2.3 Laboratory analyses

159 2.3.1 Runoff water analyses

160 All water samples were analysed for  $NH_4^+$  (µg L<sup>-1</sup>) using a continuous segmented flow autoanalyser 161 (AA3; Seal Analytical).  $NH_4^+$  was determined using the Berthelot reaction (Berthelot, 1859), where a 162 blue-green compound was produced and quantified colorimetrically. NH<sub>4</sub><sup>+</sup> was the only inorganic N 163 fraction measured as the cattle slurry used contained no measureable  $NO_3^-$  or  $NO_2^-$  (see table 1). Also, 164 previous work carried out using the same soil and slurry showed that after slurry application and 165 leaching for 8 h, there was no detectable difference between the concentrations of extractable NO<sub>3</sub><sup>-</sup> 166 or  $NO_2^-$  from slurry-treated or control soils (unpublished data). This is most probably due to the soils 167 having a low moisture content (~8%) at beginning the experiments, which then become rapidly 168 saturated. Both low moisture and potential oxygen limitation (due to waterlogging) later in the 169 experiments could inhibit rates of nitrification. As a result, there is no evidence to suggest that 170 nitrification would occur over the time-scales of the current TRACE experiments and therefore NH<sub>4</sub><sup>+</sup> 171 was the sole inorganic compound used to trace slurry in this paper.

172 Fluorescence spectroscopic analysis of all leachate samples was carried out using a HORIBA Jobin Yvon 173 FluoroLog<sup>®</sup>-3 spectrofluorometer (Stanmore, UK); the excitation source was a Xe lamp. Samples were 174 measured in a 1 cm quartz cuvette at room temperature. The methodology of Peuravuori et al. (2002) 175 was adopted to collect synchronous spectra, using 1 nm increments from 250–600 nm, with an 18 nm 176 offset between the excitation and emission monochromators. The spectra were then corrected for 177 the fluorescence of Milli-Q water and for inner-filter effects, based on the sample absorbance. The 178 sample absorbance at 250–600 nm was measured in a 1 cm cuvette, using a Shimadzu UVmini-1240. 179 Milli-Q subtracted absorbance spectra were then used to correct the fluorescence spectra for primary 180 and secondary inner-filter effects using the equation:

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$$I_c = I_m / (10^{-b(A_{ex} + A_{em})})$$

where I<sub>c</sub> is the true fluorescence intensity, I<sub>m</sub> is the measured fluorescence intensity, b is the sample path length and A<sub>ex</sub> and A<sub>em</sub> are the absorbance values at the excitation and emission wavelengths respectively (Lakowicz, 1983;Ohno, 2002). The spectra were then converted to arbitrary Raman units by normalising the fluorescence intensities by the area of the water Raman peak in order to compare the relative magnitude of the peaks present between the experiments. The ratio between the fluorescence intensities of emissions at ~290 nm and ~380 nm was calculated as previous work has shown that it can be used to monitor the presence of slurry-derived compounds in natural waters (Baker, 2002a, b;Naden et al., 2010;Lloyd et al., 2012).

# 190 2.3.2 Soil and eroded sediment analyses

The soil and eroded sediment samples were analysed in triplicate for total carbon (TC), inorganic carbon (IC) and total nitrogen (TN) (%) using a Carlo Erba EA1108 Elemental Analyser. Total organic carbon (TOC) was then calculated by subtracting the IC from the TC.

194 Lipid analysis was used to extract and quantify the stanol concentration within the soil cores and 195 eroded sediment samples, specifically the  $5\beta$ -stanols, which were used as a tracer of slurry-derived 196 particulates (Bull et al., 2002;Leeming et al., 1996;Evershed et al., 1997;Nash et al., 2005;Lloyd et al., 197 2012). As 5 $\beta$ -stanols stigmastanol and its epimer are the biohydrogenation products of sitsosterol (a 198 major plant sterol) and are only produced during rumination, they can be used as unequivocal tracers 199 of the hydrophobic fraction of ruminant faeces. The  $5\alpha$  form is produced by microbial activity under 200 aerobic conditions, i.e. outside the rumen, so the ratio  $5\beta$ : $5\alpha$  is used to investigate the contribution 201 of either source. To achieve this, a total lipid extract (TLE) was obtained from the soil and eroded 202 sediment samples using Soxhlet extraction ( $CH_2Cl_2:Me_2CO 9:1 v/v, 24 h$ , van Bergen et al. (1997)) with 203 an internal standard (preg-5-en-3 $\beta$ -ol). Aliquots of the TLE samples were saponified by adding 1 mL 204 0.5 M methylated MeOH and heating at 70°C for 60 min before acidification to pH 3 by adding 1 mL 205 of 1 M HCl. Then 1 mL of DCM-extracted DDW was added along with 2 mL of DCM and the sample was 206 vortex mixed for 20 s and allowed to settle to give a two-phase sample. The organic layer was 207 extracted from the bottom then the DCM extraction was repeated two more times to ensure the 208 entire organic sample was collected. The composite organic phase was then filtered through a pipette 209 containing anhydrous sodium sulphate (NaSO<sub>4</sub>) in order to ensure any residual water was removed, 210 then the sample was blown down to dryness under N<sub>2</sub>. An aliquot of the saponified TLE was then 211 derivatised using 50 µL of N,O-bis(trimethylsilyl)trifluroacetamide with 1% trimethylchlorosilane and

212 heating at 70°C for 1 h. The samples were then analysed using a Finnigan TRACE GC/MS by injecting 213 1  $\mu$ L on-column, using a HP-1 (50 m, 0.32 mm, 0.17  $\mu$ m) column. The GC temperature programme was: 50°C (2 min), ramping to 245°C at a rate of 15°C min<sup>-1</sup>, then increasing to 250°C at a rate of 0.5°C min<sup>-1</sup> 214 215 <sup>1</sup> followed by a ramp to 300°C at a rate of 6 °C min<sup>-1</sup>, then finally held for 20 min. This GC temperature 216 programme was chosen to maximise the information gained at the time period where the stanols 217 elute. The mass spectrometer was operated at 70eV with the quadrapole mass analyser scanning the 218 range m/z 50-650 (scan time 0.6 s). The distribution of sterols and stanols present were then 219 quantified against the internal standard.

The ratio of  $5\beta$ -: $5\alpha$ -stanols was determined in order to assess the relative contribution of slurryderived insoluble organic matter (IOM) to native soil IOM in the soil cores. This is a useful technique where the  $5\beta$ -stanol concentrations are low, as the ratio allows differences between the samples to be seen more clearly.

### 224 2.4 Statistical analyses

Statistical differences between the slurry treated experiments and the control experiments were tested for each of the measured parameters using either a T-test or a Mann Whitney test depending on whether the specific data sets were normally or non-normally distributed.

228 **3. Results** 

#### 229 3.1 Flow pathway partitioning and erosion

The results show that as rainfall intensity and slope gradient increased a higher proportion of flow was transported via surface and shallow subsurface flows in the slurry treated experiments (Figure 2). Doubling the rainfall intensity had a larger impact on the flow pathway partitioning compared with doubling the slope gradient. Figure 2 summarises the partitioning of water flow through the slope for the slurry-treated experiments. Discharge data for the control experiments are not included as the results mirrored the slurry experiments run under the same experimental conditions. The two 236 experimental end-member scenarios showed a reversal in behaviour of the flow partitioning, where at 5°, 60 mm  $h^{-1}$  ~99% of the flow was transported as vertical percolated flow and at 10°, 120 mm  $h^{-1}$ , 237 238 ~99% of the flow was routed across the soil surface as overland flow. All experiments received the 239 same volume of rainfall, therefore halving the slope gradient and rainfall intensity caused a 60% 240 increase in the storage of water within the soil. At a rainfall intensity of 60 mm h<sup>-1</sup> doubling the slope 241 gradient induced a reduction in the vertical percolated flow (to 26%) and increase in subsurface 242 throughflow (28%) and surface runoff (46%) due to higher surface water velocities and reduced 243 infiltration rates. Therefore, overland flow generation was a combination of infiltration-excess (fast) 244 and saturation excess (slower) overland flow. When the rainfall intensity increased to 120 mm h<sup>-1</sup> the 245 system was dominated by infiltration-excess overland flow. At a slope gradient of 5°, 67% of the discharge was surface runoff and 29% was subsurface throughflow. However, increasing the slope 246 247 gradient to 10° resulted in 99% of the flow transported as surface runoff.

248 Figure 3 shows the corresponding sedigraphs of eroded sediment transported in the overland flow for 249 each of the slurry-treated experiments. Like discharge the data for the control experiments is not 250 included as their behaviour mirrored that of the slurry-treated experiment run under the same experimental conditions. The experiments run at 60 mm h<sup>-1</sup> rainfall produced sediment 251 concentrations that increased gradually until the middle of the experiment before decreasing later in 252 253 the event. At 120 mm h<sup>-1</sup> the 5° experiment produced a similar trend to those run at the lower rainfall 254 intensity (at both 5 and 10° slopes), however the sediment discharges were generally higher. When 255 the slope gradient increased to 10° the shape of the sedigraph changed and the greatest sediment 256 concentrations were at the beginning of the rainfall simulations. This suggests that rainsplash, 257 combined with higher water velocities and higher flow shear stress expected at steeper gradients, 258 could initially rapidly move a large quantity of sediment, after which the system slowed to a sediment 259 export rate similar to that of the lower gradient slope.

260 *3.2 Dissolved chemical species analyses* 

Figures 4, 5 and 6 show a direct comparison of the flow and sediment and the  $NH_4^+$  concentration 262 263 from each monitored flow pathway during each experiment. The control experiments (Figure 4) 264 produced a total of between 12 and 25 mg of leached NH<sub>4</sub><sup>+</sup> originating from the soil (no added slurry), with concentrations never exceeding 0.1 mg L<sup>-1</sup> from any pathway. The total mass of NH<sub>4</sub><sup>+</sup> exported 265 266 from each experiment depended on the relative flow pathway partitioning because the experiments 267 which generated higher volumes of surface runoff contributed a larger flux of NH<sub>4</sub><sup>+</sup>, between 1.8% and 268 11.3% of the total  $NH_4^+$  measured from the slope outlets. An approximately equal proportion of the 269 exported NH<sub>4</sub><sup>+</sup> was leached from the vertical percolated flow outlets in all control experiments, 270 between 10.9 mg (48%) 7.6 mg (62%), with the remainder transported via sub-surface runoff.

271 The addition of slurry to the slope caused a marked increase in the export of  $NH_4^+$  compared to the 272 controls, in some cases by an order of magnitude, indicating that the main source of  $NH_4^+$  leached 273 from the slope was the added slurry. The NH<sub>4</sub><sup>+</sup> concentrations from the slurry-treated experiments 274 were significantly higher than those from the control experiments (Mann-Whitney, control vs. slurry, 275 p=<0.001). As with the control experiments, total  $NH_4^+$  export was affected by the flow pathway 276 partitioning. The slurry-treated experiments which were dominated by surface runoff exhibited the 277 largest export of NH<sub>4</sub><sup>+</sup>, between 66 and 99% of the total. The experiment with the most surface runoff 278 (10°, 120 mm  $h^{-1}$ ) generated a NH<sub>4</sub><sup>+</sup> yield of 404 mg, an order of magnitude larger than any other 279 experiment. NH4<sup>+</sup> was also observed at similar concentrations in the sub-surface flow pathway when 280 activated. During the experiment where the majority of the water flow was via vertically percolated 281 pathways (5°, 60 mm  $h^{-1}$ ), the concentrations of  $NH_4^+$  were similar to those observed in the dominant 282 pathways for other slurry experiments (0.1-0.5 mg L<sup>-1</sup>). The area associated with the largest percolated 283 flux tended to be the top of the slope directly below the slurry application area, illustrating that the 284 slurry can still be rapidly transported during smaller rainfall events but the dissolved fraction is 285 infiltrated rather than routed across the soil surface.

In general, the transport of  $NH_4^+$  was controlled by the flow partitioning, and the concentrations did not increase with contact time with the soil, providing additional evidence that the  $NH_4^+$  was representing the dissolved slurry-derived material. This can be seen very clearly in Figure 6 (c), where the concentration of  $NH_4^+$  observed in the percolated flow pathway is highest after 20 min, showing that the slurry material had reached the bottom of the slope.

291 *3.2.2 Fluorescence spectroscopy* 

292 Figure 7 shows the average fluorescence spectra for each flow pathway for each experiment, along 293 with the mean spectra of the control samples and the slurry. This plot highlights that the difference 294 between the fluorescence spectra of the slurry and control soil is the magnitude of the peak at 295 excitation wavelength 270 nm (emission 290 nm), which has been attributed to 'proteinaceous 296 material' (Baker, 2002a, b; Peuravuori et al., 2002), therefore providing a useful tracer of slurry-derived 297 dissolved organic matter (DOM) (Lloyd et al., 2012). This 'protein' fluorescence can be a useful tracer 298 for slurry as proteins form a significant part of the organic nitrogen pool, which can be derived from 299 plant, animal or bacterial sources (Jones et al., 2005; Vinolas et al., 2001). It is well documented that 300 fluorescence at an excitation at 270 nm can be linked to farm wastes (Baker, 2002a;Hudson et al., 301 2007; Naden et al., 2010) and the lack of a signal in the control samples supports the hypothesis that 302 the slurry is the source in these experiments. The assumed higher molecular weight compounds that 303 fluoresce at longer wavelengths are most likely derived from soil DOM rather than from the slurry. In 304 general, the fluorescence intensity at higher wavelengths (e.g. 360 nm and 446 nm) is lower in the 305 surface runoff and subsurface samples compared with those collected from the percolated pathway, 306 providing more evidence that the source is soil-derived.

The ratio between the fluorescence intensities at excitation at 270 nm and 360 nm can be used to monitor the presence of slurry-derived DOM in natural waters (Baker, 2002a, b;Naden et al., 2010;Lloyd et al., 2012). Figure 8 shows the change in the '270:360' ratio through time for each flow outlet for each of the slurry-treated experiments. The average ratio of the control samples was 0.044

311 with a standard deviation of 0.019, while the slurry had a ratio of 1.72. The slurry-treated water 312 samples had a large variability in ratios with a range of 0.025 to 0.32. The slurry-treated water samples 313 collected from the surface runoff and the subsurface throughflow consistently displayed ratios 314 significantly higher than those from the control slopes (Surface runoff, Mann Whitney, control vs. 315 slurry, p=<0.001; Subsurface, t-test, control vs. slurry, p=<0.001) suggesting that slurry-derived DOM 316 was being transported by those flow pathways. The data from the vertical percolated flow pathways 317 were recorded for each of the four paired outlets (see Figure 1 for locations) so that the spatial 318 distribution of slurry-derived DOM transport could be examined. At the highest slope gradient and 319 rainfall intensity slurry-derived DOM was only detected in percolated outlet 1 which is located directly 320 underneath the slurry application area, there was a lack of infiltration across the rest of the slope. The 321 experiment run with a slope gradient of 5° and rainfall intensity of 60 mm h<sup>-1</sup> produced the highest 322 volume of discharge from vertical percolated flow, however slurry-derived DOM was only significantly 323 detected (Mann-Whitney, control vs. slurry, p=<0.001) via the fluorescence in outlet 1 throughout the 324 experiment and through outlet 2 for the first 30 min. The 270:360 ratio was also raised 325 significantly significantly greater than above the control values in outlets 1, 2 and 4 (Mann-Whitney, 326 control vs. slurry, p=<0.001) at the end of the experiment suggesting slurry-derived DOM was reaching 327 the bottom of the slope by the end of the simulation. There were no significant increases in 270:360 328 nm fluorescence from any of the vertical percolated flow pathways in the experiment run at 5° and 329 120 mm h<sup>-1</sup> rainfall, or from the base of the slope (outlet 4) and in the 10°, 60 mm h<sup>-1</sup> experiment only 330 at the base of the slope, outlet 4 at the end of the rainfall event. Overall, these analyses allowed the 331 identification of the locations where slurry-derived DOM has been transported throughout the rainfall simulations. 332

333

334 *3.3 Soil and eroded-sediment analyses* 

335 3.3.1 Elemental analyses of C and N

336 Our previous work showed that the components of slurry-derived material which are most likely to 337 bind to the soil remain in the top 5 cm even after prolonged periods of leaching (Lloyd et al., 2012). 338 Therefore, elemental analyses of C and N were restricted to the top 5 cm of the soil cores. There was 339 no significant difference between the TOC and TN values in the soil cores for the slurry-treated 340 experiments (median TOC = 1.7%, TN = 0.2%) compared to the experimental controls (median TOC = 341 1.4%, TN = 0.3%) (TOC, Mann-Whitney, control vs. slurry, p=0.052; TN, Mann Whitney, control vs. 342 slurry, p=0.181<del>; data not shown</del>). TOC and TN in the eroded sediment were also determined in samples 343 taken from the slurry-treated slopes. The majority of the experiments showed increases in the 344 concentrations of TOC in the eroded sediment compared to the control slopes (Mann Whitney, control vs. slurry, p=<0.02 for all experiments), with maximum values of up to 4.2%. The experiments which 345 346 did not generate high erosion rates had lower concentrations of 1.5% TOC. Concentrations of TOC in 347 the eroded sediment samples generally decreased through time in most experiments, except when 348 erosion rates where sustained through the experiments whence values also remained stable. TN 349 concentrations of eroded sediment were greater than those in the control soil in only one experiment, 350 however they were not significantly more (Mann Whitney, control vs. slurry, p=0.179). Overall, these 351 data confirm that the elemental analyses alone do not provide a robust method to trace the transport 352 of slurry-derived particulates.

# 353 3.3.2 Lipid analyses on eroded sediment and soil cores

Figure 9 shows example partial chromatograms for a soil core from the top area of the slope, eroded sediment and the control soil. These data clearly illustrate the dominance of the 5 $\beta$ -stanols in the slurry-treated samples relative to the control soil. The concentration of 5 $\beta$ -stanols, a robust biomarker for slurry-derived IOM, in the slurry was 12,982 ng g<sup>-1</sup> compared with an average of 67.7 ng g<sup>-1</sup> in the control soil. The soil cores taken from the slurry application area were significantly enriched with concentrations averaging 2850 ng g<sup>-1</sup> (Mann Whitney, control vs. slurry, p=0.010). Generally, the 5 $\beta$ stanol concentrations in the soil cores decreased exponentially with distance downslope, with average

concentrations at the bottom of the slope of 84 ng g<sup>-1</sup> which were not significantly different from the 361 control soil (Mann Whitney, control vs. slurry, p=0.343). The average concentration of the eroded 362 sediment was 2040 ng g<sup>-1</sup> which was also significantly enriched compared with the control soil (Mann 363 364 Whitney, control vs. slurry, p=0.010). Figure 10 shows the changes in the concentration of  $5\beta$ -stanols 365 in the soil cores in the downslope direction after each individual experiment. Some of the experiments 366 showed increases in the concentrations of soil core  $5\beta$ -stanols at the bottom of the slope, which 367 suggests that deposition of slurry-derived IOM was occurring close to the slope outlet. The error bars 368 in Figure 10 represent the range of values from the three soil cores taken horizontally at each distance 369 downslope. In some locations the range of concentrations was large, possibly due to preferential 370 surface runoff pathways which distributed sediment laterally across the slope as well as downslope.

371 In addition to absolute concentrations the ratio between 5 $\beta$ - (slurry-derived) and 5 $\alpha$ - (soil-derived) 372 stanols was calculated in order to assess the contribution of slurry-derived IOM versus native soil IOM 373 in the soil cores (data not shown). Results showed a spatial pattern in the distribution of the ratio of 374  $5\beta$ -: $5\alpha$ -stanols in the soil across the slope surface for each experiment, exhibiting an exponential 375 decline in 5 $\beta$ -:5 $\alpha$ -stanols ratio in the downslope direction from where slurry was applied. The main 376 difference between the experiments was in the variability of stanol ratios, which was a function of 377 slope gradient. The standard deviation and variance of the stanol ratios was calculated for each 378 experiment (not including the top of the slope where the slurry was initially applied) in order to assess 379 the variation in transported slurry-derived material between experiments (Table 3). The analysis 380 showed that increasing slope gradient resulted in an increase in the standard deviation of the ratios 381 in soil cores across the slope suggesting that surface runoff at lower gradients was flowing more evenly 382 over the soil surface compared to the steeper slopes which may have exhibited more preferential flow 383 routing.

384 The concentrations of 5 $\beta$ -stanols were also quantified in the sediment eroded from the slope during 385 each experiment. In order to obtain an adequate sample of sediment for 5 $\beta$ -stanol analysis, the 386 eroded sediment samples were combined into three time segments, representing the beginning, 387 middle and end of the rainfall simulation (See Table 21 for the experimental time covered by each 388 composite sample). As the sediment flux from the 5°, 60 mm  $h^{-1}$  experiment was relatively low only 389 one analysis was possible which represented the entire event. The average concentrations of 5β-390 stanols are shown in Figure 9 and the temporal variations in concentrations are shown in Table 2. The 391 lowest concentration of 5β-stanols was recorded in the eroded sediment from the experiment with 392 the lowest volume of surface runoff, suggesting that transport of slurry-derived POM was therefore 393 limited, and therefore reflected in the low sediment yield under these conditions. Experiments with 394 higher volumes of surface runoff resulted in an increase in the  $5\beta$ -stanol concentrations by up to a 395 factor of 10. Increasing rainfall intensity appeared to have little effect on the concentrations of 5β-396 stanols exported at the beginning of the rainfall simulations at a slope gradient of 5°. However, 397 increasing slope gradient resulted in a rapid decrease in concentrations during the middle and end of 398 the simulations. The supply of slurry-derived material was exported from the slope faster at steeper 399 slope gradients due to the increased sediment flux, as shown in Table 2. Slope gradient had a larger 400 impact on the total transport of  $5\beta$ -stanols than the increase in rainfall intensity, due to an exponential 401 increase in sediment yield.

#### 402 **4. Discussion**

This study used a series of six controlled laboratory experiments to assess the effect of slope gradient
and rainfall intensity on the transport of slurry-derived compounds. This work also provided an
extension to the proof of concept developed in Lloyd et al. (2012) for using fluorescence and 5βstanols simultaneously to trace both the vertical and lateral transport of slurry-derived material in
both dissolved and particulate forms.

408 4.1 Role of slope gradient and rainfall intensity on transport slurry-derived compounds

409 It was hypothesised that changes in slope gradient and rainfall intensity would alter the relative 410 partitioning of slope flow pathways and erosion rates which would in turn exert a strong control on 411 the flux and yield of slurry-derived compounds. The impact of these variables on slurry transport can 412 be described by two main hydrologically-driven scenarios.

413 1) Overland flow dominated systems, which typically occur on steep slopes and/or high rainfall 414 intensities. High overland flow rates typically result in high erosion rates and the combination of the 415 two enhances transport of slurry-derived compounds via surface pathways (dissolved in runoff and 416 bound to eroded sediment), which are rapid and interact directly with the source of applied slurry 417 material. The concentration of the dissolved components depends on the volume of runoff produced. 418 Higher intensity rainfall events generally transported the overall largest load of slurry-derived 419 dissolved material but the concentrations were lower. This result is supported by Delpla et al. (2011), 420 who showed in a field study that it was the highest intensity rainfall events which generated overland 421 flow that yielded the highest DOC export prior to cattle slurry application. Dissolved components of 422 slurry which have been diluted by increased volumes of discharge will have a lower immediate impact 423 on stream ecology. However, the larger total load exported during high intensity storms will result 424 greater total losses of nutrients from the soil and could result in pressures on receiving waters, such 425 as increased biological oxygen demand (BOD). It is estimated that input of cattle slurry can produce a 426 BOD of between 10,000 and 30,000 mg L<sup>-1</sup>, leading to reduced oxygen levels and ultimately the death 427 of aquatic life (Khaleel et al., 1980; Baker, 2002a; Foy and Kirk, 1995).

The concentrations of slurry-derived particulates were controlled primarily by the slope gradient, however, the overall load increased substantially (between 1 and 3 orders of magnitude) with increased rainfall intensity due to the increased sediment transport by overland flow. This is supported by Michaelides et al. (2010) who showed the importance of erosion events on landscape nutrient loss. Although small amounts of subsurface and percolated flow occurred at the higher rainfall intensity, they do not play a key role in the transport of slurry-derived particulates. In the overland flow 434 dominated experiment (10°, 120 mm  $h^{-1}$ ) over 4 kg of sediment were exported (over an area of 15 m<sup>2</sup>), 435 containing 2.3 mg of 5 $\beta$ -stanols in under an hour of rainfall, compared with a total of ~9.7 mg of 5 $\beta$ -436 stanols added to the soil as slurry before the experiment. The experiment that generated less than 1% 437 of its discharge by overland flow (5°, 60 mm h<sup>-1</sup>) eroded just 14 g of sediment (3  $\mu$ g 5 $\beta$ -stanols) during 438 the experiment. Due to the rapid movement of slurry-derived material in this overland flow scenario 439 there is little time for the slurry derived components to transform, for example the nitrification of 440  $NH_4^+$  to  $NO_2^-$  and  $NO_3^-$  or the mineralisation of the organic fraction to more labile inorganic forms. As 441 a result, over short time scales after slurry application, the transported fractions of slurry tended to 442 reflect the original slurry composition. This result can also be expected to be observed in a field 443 scenario where material is rapidly transported from an area of slurry application.

444 2) A predominance of vertical, percolated flow, which typically occur due to low slope gradients 445 coupled with lower rainfall intensities. Under this regime, the export of slurry-derived material is 446 mainly in dissolved form and the flux tends to be higher directly beneath, or close to the application 447 area. The timing of the dissolved flux also tends to be slower compared with an overland flow 448 dominated regime due to the time needed for percolation to occur through the soil matrix (Kirkby, 449 1969). If surface runoff develops during a rainfall event due to saturation excess, some dissolved 450 slurry-derived material will be transported towards the slope outlet, but the concentrations will be 451 lower than in an overland flow-dominated system. This is because a large proportion of the dissolved 452 slurry-derived material will have already infiltrated into the subsurface in the time taken for the soil 453 to saturate and initiate overland flow. Buda and DeWalle (2009) showed that larger storm events 454 which caused saturation-excess overland were responsible for flushing stored nutrients via shallow 455 subsurface pathways. In addition, overland flow transports sediment-bound components which have 456 been shown to remain in the top layers of the soil regardless of the volume of infiltration (Lloyd et al., 457 2012). Under systems dominated by infiltration and vertical percolated pathways there is a potential longer-term contamination threat to surface sediment and groundwater. The percolated slurry-458 459 derived material will remain in the deeper soil layers and be available for leaching during subsequent

460 rainfall events and potentially assimilated by the microbial community. The NH<sub>4</sub><sup>+</sup> is subject to 461 nitrification resulting in increased NO<sub>3</sub><sup>-</sup> and the organic components will be mineralised over time to 462 add further NO<sub>3</sub><sup>-</sup> to the inorganic nutrient pool. This provides a large supply of accessible nutrients for 463 plant growth, however, if there is a surplus after plant uptake or it is leached below the root zone then 464 these nutrients can be transported into groundwater or via deeper subsurface flows to water courses 465 (Vitousek et al., 2009;Burow et al., 2010;Melo et al., 2012;Morari et al., 2012), thereby posing a longer-466 term contamination threat.

467 Overall, we conclude that the controls on the transport of slurry-derived material (soluble and 468 insoluble) are complex and that flow pathway partitioning induced by slope gradient, rainfall intensity 469 and duration play important roles. Rainfall is an important transport driver for both dissolved and 470 particulate components. Raindrop action is a well-known mechanism for detaching and mobilising 471 sediment particles, but it can also act to eject soil water and therefore release solute into runoff (Gao 472 et al., 2004). In addition, when the rainfall event acts to saturate the soil profile diffusion will occur, 473 allowing diffusion to liberate chemicals from the soil matrix. Experiments run by Gao et al. (2004) 474 showed that at the beginning of a rain event raindrop impact was the main mechanism for liberating 475 solute, followed by diffusion at the latter stages of the storm. In addition, research has also shown 476 that storms which have variable rainfall intensities can increase the transport of solute in surface 477 runoff and via preferential subsurface routes (Zhang et al., 1997; Malone et al., 2004).

478 4.2 Use of a combined tracer approach to monitor the transport of slurry-derived material.

This study also aimed to test efficacy of the use of a combined biomarker approach to monitor the vertical and lateral transport of dissolved and particulate slurry fractions. In the current study the transport of dissolved slurry-derived material within different flow pathways was monitored using NH<sub>4</sub><sup>+</sup> concentrations and fluorescence. Over short time-scales (hours) NH<sub>4</sub><sup>+</sup> concentrations can provide useful information regarding the timing and spatial patterns of movement of dissolved slurry compounds. Field data have shown that areas which have high N-loading, such as those treated with slurry, could receive N at a higher rate than can be incorporated into the organic fraction, resulting in
inorganic-N, especially NH<sub>4</sub><sup>+</sup> becoming available for rapid transfer to watercourses, often via
preferential flow pathways (Heathwaite and Johnes, 1996). However, within days the signal will
degrade due to nitrification of NH<sub>4</sub><sup>+</sup> into NO<sub>3</sub><sup>-</sup>, which would be difficult to distinguish from soil-derived
N, without the use of isotopic enrichment additions.

490 Fluorescence spectroscopy was shown by Lloyd et al. (2012) to be a robust tracer of DOM, which was 491 slurry-derived over short timescales and in relatively small-scale soil columns. However when this 492 technique was applied to the more complex 3D laboratory system the results were not as clear. While 493 the fluorescence intensity at an excitation at 270 nm was not significantly different between the slurry-494 treated and control experiments, using the ratio between '270' and '360' intensities provided a 495 detectable signal. The ratio was able to identify smaller changes in the fluorescence spectra which 496 were not possible using the '270' fluorescence intensity alone. The fluorescence intensities recorded 497 from the TRACE experiments were several orders of magnitude lower than those from the soil column 498 experiments (Lloyd et al., 2012), due to the larger volume of rainfall added in the TRACE experiments 499 which resulted in a significantly smaller slurry:soil ratio. The slurry was only applied to the top 1 m 500 strip of the slope and therefore the signal was rapidly diluted both by the rainfall and the additional 501 signal from the soil-derived DOM. Furthermore, the '270':'360' ratio of the slurry was calculated at 502 1.7, which is lower than the range cited in the literature (~2-5) for cattle slurries (Baker, 2002a). This 503 could be because when the slurry was collected (during April 2008), it had been a very wet spring and 504 as a result the farm slurry store had received higher than average volumes of rainwater, potentially 505 diluting the slurry before application.

Fluorescence spectroscopy has been shown previously to be a powerful technique for characterising DOM, even in very dilute samples (e.g. Barker et al., 2009;Birdwell and Engel, 2010). However, the signal was more difficult to detect in this case because the soil-derived DOM swamped the slurry signal, making it difficult to resolve the two sources. This problem was exaggerated in the case of the 510 higher intensity rainfall experiments due to the additional water being transported with the slurry. If 511 a larger section of the slope had been treated with slurry then it is more likely that the signal could 512 have been seen. Naden et al. (2010) showed that drainage water from field lysimeters treated with slurry could be distinguished from control experiments, even though only a small proportion of the 513 514 applied slurry was thought to have been leached from the system. However, work by Old et al. (2012) 515 showed that fluorescence differences could not be identified in storm runoff samples after an application of cattle slurry to undrained plots. The authors suggest that this was due to rapid 516 517 absorption or microbially-mediated-immobilisation of the slurry material in the soil matrix. Baker 518 (2002b) showed that river water samples from the UK exhibited '270':'360' ratios of 0.37 ±0.41 (n= 242), suggesting that cattle slurries may be difficult to detect once they are diluted compared to the 519 520 background river fluorescence. However, Baker (2002b) conclude that the techniques would be able 521 to detect cattle slurry inputs from large point sources, such as slurry tank failures.

522 The particulate fraction was investigated using bulk elemental analysis, followed by quantifying the 523  $5\beta$ -stanol concentrations. Soil analyses of bulk C and N were found to be generally ineffective at 524 tracing transport pathways of slurry-derived POM. The eroded sediments from all of the experiments 525 showed increases in TOC and TN values compared with the control but the source of origin of the TOC and TN cannot be identified (i.e. slurry or soil). The use of the specific biomarkers 5β-stanols provides 526 527 more robust information about the transport of slurry-derived particulates as they have been shown 528 to be an unequivocal tracer of ruminant slurry (Bull et al., 2002;Elhmmali et al., 2000;Evershed et al., 529 1997; Nash et al., 2005). The concentration of the 5 $\beta$ -stigmastanol in the slurry used in this study was 530 lower than other published values, with an average concentration of ~12  $\mu$ g g<sup>-1</sup> of freeze dried slurry, compared with ~46  $\mu$ g g<sup>-1</sup> (Leeming et al., 1996). However, there was still a large difference between 531 532 the slurry concentration and the initial soil, so this did not cause any issues for using the biomarker as a tracer. On the other hand, the ratios of the  $5\beta$ : $5\alpha$ -stanols determined in the current study were very 533 similar to other published values, where ratios from soil cores taken from the slurry application area 534 535 ranged from 2.4-2.9, compared with an average ratio of ~2 measured in cow manures (Evershed et 536 al., 1997). Data collected here further illustrate that  $5\beta$ -stanols are a robust tracer of slurry-derived 537 IOM, and therefore could be used to monitor the transport of particulate forms of slurry.

538 Overall, the results show that using a combination of tracers and biomarkers for both soluble and 539 insoluble fractions can be very effective for tracing the movement of slurry via multiple transport 540 pathways. This work provides a proof of concept that the methodology works in larger and more 541 complex controlled laboratory systems. While elements of this type of methodology have been tested 542 in field scenarios (e.g. Naden et al., 2010; Granger et al., 2010; Evershed et al., 1997; Bull et al., 543 1998;Nash et al., 2005), there is still a need to test this combined tracer approach in a larger field 544 study.

#### 5. Conclusions

546 This work has provided important and new insights into flow partitioning across a range of controlled 547 hillslope and rainfall scenarios and has allowed quantification of the impact this has on the transport 548 of slurry-derived compounds. Results indicate that the dissolved components of slurry-derived 549 material (traced using  $NH_4^+$  and the ratio between '270':'360' fluorescence intensities) were 550 transported rapidly through the soil system, while the predominant pathway depended on the flow 551 partitioning. When the conditions favoured surface runoff, i.e. high slope gradients and/or high rainfall 552 intensities, larger quantities of slurry-derived material were moved in the surface and subsurface flow 553 pathways. These shallow flow pathways transport water more rapidly to the slope outlet compared 554 with vertical percolated flow which travelled slowly through the soil matrix. The movement of slurry-555 derived particulates (traced using 5 $\beta$ -stanols) is driven exclusively by the erosion rates on the slope.

556 Rainfall events which produced flashy hydrological responses, resulting in large quantities of surface 557 runoff, were likely to move sediment and also flush dissolved components of slurry-derived material 558 from the slope, increasing the contamination risk. Rainfall events which produced slower hydrological 559 responses were dominated by vertical percolated flows removing less sediment-associated material,

but produced leachate which could contaminate deeper soil layers, and potentially groundwater, overa more prolonged period.

562 This work has also provided one of the first examples of using multiple biomarkers to assess the effects

of slope gradient and rainfall intensity on the movement of slurry-derived OM. The results have shown

- that this approach can be successfully applied to more complex 3-D systems (than simple soil columns)
- and can yield valuable data about the interactions between slurry and the soil-water system. Overall,
- this research provides new insights into the partitioning of slurry-derived material when applied to an
- 567 unvegetated slope and the transport mechanisms by which contamination risks are created.
- 568

# 569 Acknowledgements

570 This work was funded by NERC Studentship (NE/F008856/1) to C.E.M.L.

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# 751 Table 1: Chemical characteristics of control soil and cattle slurry (dry weight basis) used in all of

# 752 the experiments.

Initial soil	Cattle slurry
2.7 % ±0.2	39.6 % ±0.3
2.6 % ±0.2	39.0 % ±0.3
0.4 % ±0.09	3.4 9% ±0.004
0.0016 % ±0.0005	0.39 % ±0.0005
0.0022 % ±0.00006	0 %
0.00004 % ±0.00003	0 %
0.4 % ±0.09	3.45% ±0.004
	Initial soil $2.7 \% \pm 0.2$ $2.6 \% \pm 0.2$ $0.4 \% \pm 0.09$ $0.0016 \% \pm 0.0005$ $0.0022 \% \pm 0.0006$ $0.00004 \% \pm 0.0003$ $0.4 \% \pm 0.09$

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# 755 **Table 2: Concentrations of 5β-stanols in the eroded sediment (dry weight basis) through time and**

	60 mm h <sup>-1</sup>						120 n	וm h⁻¹		
		5°		10°			5	0	10	0
	Time	Conc.	Mass	Conc.	Mass	Time	Conc.	Mass	Conc.	Mass
	(min)	(ng g <sup>-1</sup> )	(µg)	(ng g <sup>-1</sup> )	(µg)	(min)	(ng g <sup>-1</sup> )	(µg)	(ng g <sup>-1</sup> )	(µg)
Beginning	0-30	351*	3.3	3477.2	666.7	0-20	306.1	3611.0	2024.4	6912.5
Middle	30-60	-	-	3005.2	716.5	20-35	545.0	1032.2	1426.0	9274.2
End	60-	-	-	2533.1	734.0	35-50	360.7	417.1	1280.2	6725.3
	100									
Total			3.3		2123			5060		22912

# 756 the total exported mass throughout the experiment.

<sup>757</sup> \*This experiment only had one time interval due to the lack of sediment to analyse.

Table 3: Spatial variability in the ratios of 5β:5α-stanols in soil cores across the slope for each
 experiment (excluding samples taken in the slurry application area).

		<u>Standard</u> Deviation	<u>Variance</u>
60 mm h <sup>-1</sup>	<u>5°</u>	<u>0.14</u>	<u>0.02</u>
	<u>10°</u>	<u>0.79</u>	<u>0.62</u>
120 mm h <sup>-1</sup>	<u>5°</u>	<u>0.39</u>	<u>0.15</u>
120 11111 11	<u>10°</u>	<u>0.42</u>	<u>0.17</u>

# 762 List of Tables

- 763 Table 1: Chemical characteristics of control soil and cattle slurry (dry weight basis) used in all of the764 experiments.
- Table 2: Concentrations of 5β-stanols in the eroded sediment (dry weight basis) through time and the
- total exported mass throughout the experiment.
- 767 <u>Table 3: Spatial variability in the ratios of  $5\beta:5\alpha$ -stanols in soil cores across the slope for each</u>
- 768 <u>experiment (excluding samples taken in the slurry application area).</u>

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Figure 1: a) Schematic showing sampling outlets on the TRACE slope and b) photo showing the emptyslope.

Figure 2: Plots showing cumulative discharge from surface runoff, subsurface throughflow and vertical

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Figure 3: Plot showing sedigraphs for each of the slurry-treated slope experiments.

Figure 4: Plots showing water and sediment discharge (a, b) and  $NH_{4^+}$  concentrations (c, d) from each monitored flow pathway for the control experiments (no slurry) at two slope angles (5 and 10°) at 60 mm h<sup>-1</sup> rainfall intensity. Where percolated 1-4 represent vertical transport from each of the pairs of tanks beneath the soil slope as marked in Figure 1.

Figure 5: Plots showing water and sediment discharge (a, b) and NH<sub>4</sub><sup>+</sup> concentrations (c, d) from each
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 of a particular flow pathway.

Figure 6: Plots showing water and sediment discharge (a, b) and NH<sub>4</sub><sup>+</sup> concentrations (c, d) from each
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 of a particular flow pathway.

Figure 7: Plots showing the mean fluorescence spectra from the surface runoff, subsurface throughflow and vertical percolated pathways for each slurry-treated experiment along with the average spectra from all pathways of the control experiments and the spectra from the applied slurry. The shaded area represents 1 standard deviation around the mean of the control soil spectra. Figure 8: Plots showing the ratio of '270':'360' nm fluorescence intensities for each measured flow pathway. The grey lines represent the average value for the control samples and the grey dashed lines are 1 standard deviation around the mean.

Figure 9: Partial chromatogram (m/z= 215) showing examples of characteristic hydrolysed lipid extracts from the top section of the soil slope, the eroded sediment (taken from the 10°, 60 mm h<sup>-1</sup> experiment) and the control soil. Where: 1 = coprostanol, 2 = epicoprostanol, 3 = 5 $\alpha$ -cholestanol, 4 = 5 $\beta$ -campestanol, 5 = 5 $\beta$ -epicampestanol, 6 = 24-ethyl-campestanol, 7 = 24-ethyl-5 $\beta$ -cholestan-3 $\alpha$ -ol, 8 = 5 $\beta$ -stigmastanol, 9 = 5 $\beta$ -epistigmstanol, 10 = 5 $\alpha$ -stigmastanol.

Figure 10: Plots showing the change in average concentration of 5β-stanols in soil cores downslope
 for each of the slurry-treated experiments. The dashed line shows the mean concentration in the
 control soil, and the triangles represent the average concentrations in the eroded sediment.









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815 Figure 3: Plot showing sedigraphs for each of the slurry-treated slope experiments.



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825 Figure 6: Plots showing water and sediment discharge (a, b) and NH<sub>4</sub><sup>+</sup> concentrations (c, d) from each monitored flow pathway for the slurry treated

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