1	The interplay between transport and reaction rates as controls on nitrate attenuation in
2	permeable, streambed sediments.
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21	Key points:
22	1. Deep sediments (>10cm) are nitrate sinks in groundwater-fed rivers
23	2. Denitrification can be sustained without substantial buried organic matter
24	3. Denitrification in a sand-dominated reach can be transport-controlled.
25	

#### 26 Abstract:

Anthropogenic nitrogen fixation and subsequent use of this nitrogen as fertilizer has greatly 27 disturbed the global nitrogen cycle. Rivers are recognized hotspots of nitrogen removal in the 28 29 landscape as interaction between surface water and sediments creates heterogeneous redox 30 environments conducive for nitrogen transformations. Our understanding of riverbed nitrogen dynamics to date comes mainly from shallow sediments or hyporheic exchange flow pathways with 31 comparatively little attention paid to groundwater-fed, gaining reaches. We have used <sup>15</sup>N 32 33 techniques to quantify in situ rates of nitrate removal to 1m depth within a groundwater-fed riverbed where subsurface hydrology ranged from strong upwelling to predominantly horizontal 34 35 water fluxes. We combine these rates with detailed hydrologic measurements to investigate the interplay between biogeochemical activity and water transport in controlling nitrogen attenuation 36 along upwelling flow pathways. Nitrate attenuation occurred via denitrification rather than 37 dissimilatory nitrate reduction to ammonium or anammox (range = 12 to >17000 nmol<sup>15</sup>N L<sup>-1</sup> h<sup>-1</sup>). 38 39 Overall, nitrate removal within the upwelling groundwater was controlled by water flux rather than reaction rate (i.e. Damköhler numbers < 1) with the exception of two hotspots of biogeochemical 40 41 activity. Deep sediments were as important a nitrate sink as shallow sediments with fast rates of 42 denitrification and short water residence time close to the riverbed surface balanced by slower rates of denitrification and water flux at depth. Within this permeable riverbed >80% of nitrate removal 43 44 occurs within sediments not exposed to hyporheic exchange flows under baseflow conditions, illustrating the importance of deep sediments as nitrate sinks in upwelling systems. 45

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Key words: (up to 6 keywords) hyporheic, nitrate consumption, hot spots, denitrification, residence
time, Damköhler

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- 50

### 51 **1 Introduction**

The global challenge of nitrate saturation of freshwater environments arises from increased nitrogen 52 loading to rivers due to anthropogenic activities such as land use change, domestic and industrial 53 54 wastewater treatment and intensification of agricultural practice [Bernot and Dodds, 2005; Caraco 55 and Cole, 1999]. In the United Kingdom, nitrate concentrations in many rivers and groundwaters have increased since the 1970s [Burt et al., 2011] leading to coastal eutrophication [Maier et al., 56 57 2009], and increasing the costs of drinking water supply in order to meet standards designed to 58 protect the environment [Knapp, 2005; National Audit Office, 2010]. Monitoring data for regulatory purposes indicates that whilst nitrate concentrations in many UK rivers have now 59 60 plateaued, that concentrations in groundwater-fed rivers continue to rise [Burt et al., 2011; Howden 61 and Burt, 2008]. This nitrate legacy has renewed interest in the role that naturally occurring microbially-mediated processes might play in transforming (e.g. dissimilatory nitrate reduction to 62 63 ammonium) and removing (in the case of denitrification and anaerobic ammonium oxidation, 64 anammox) nitrate in riverbeds [Rivett et al., 2008; Stelzer and Bartsch, 2012].

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Considerable attention has been placed on the potential role of the hyporheic zone for nitrate 66 67 removal from surface waters via denitrification [Smith, 2005], and on hyporheic exchange flows 68 (HEFs) as a means of delivering nitrate-rich surface water to the stream bed where microbial 69 activity and denitrification rates are enhanced [Findlay et al., 2003; Fischer et al., 2005]. Problems 70 of nitrate enrichment are particularly pertinent, however, for groundwater-fed rivers in permeable 71 catchments with high N-loading rates where nitrate-rich groundwater will dominate baseflow. The 72 need to understand nitrogen transformations in gaining river settings have led to an alternative 'bottom-up' rather than 'top-down' conceptualization of nitrate removal processes, highlighting the 73 importance of measuring nitrogen transformations in deep stream sediments [Stelzer and Bartsch, 74 2012]. Many experimental studies of nitrogen cycling in stream riverbeds focus on the upper 10cm 75 76 of the riverbed often selecting to conduct experiments ex situ by physically removing sediments,

77 which changes the redox environment and supply of reactants making investigation of the complete nitrogen cycle impossible [Addy et al., 2002; Sheibley et al., 2003]. Likewise, where field studies of 78 nitrogen transformations are attempted, the general approach has been to focus on 'soft' riverbeds 79 80 sediments due to the logistical difficulties associated with working within armored gravel or cobble-81 sized material [Stelzer et al., 2011]. To advance our understanding it is critical that we measure in 82 situ rates of denitrification along with other components of the nitrogen cycle (such as nitrification, 83 anammox and N<sub>2</sub>O production), at depths greater than 10cm in the coarse-grained sediments typical 84 of groundwater-fed systems, so that the relative importance of denitrification in comparison with other nitrate removal processes can be fully evaluated. Application of <sup>15</sup>N-labelled substrates is the 85 86 only method by which multiple pathways of nitrogen cycling can be investigated directly and simultaneously. Injection of <sup>15</sup>NO<sub>3</sub><sup>-</sup> into saturated sediments and recovery of porewaters over time 87 88 [referred to as 'push-pull' sampling; *Istok et al.*, 1997] has been performed at depth within the 89 riverbed and also through groundwater monitoring wells [Addy et al., 2002; Clilverd et al., 2008]. 90 These measurements, however, were focused on quantifying denitrification within large volumes of 91 sediment (10-20L of tracer were injected) and, consequently, had quite wide vertical resolution (e.g. 30-60cm). Finer scale <sup>15</sup>N 'push-pull' investigations have also been performed [Burgin and 92 93 Hamilton, 2008; Lansdown et al., 2014; Sanders and Trimmer, 2006], but to date, not in 94 conjunction with detailed hydrologic measurements.

95

96 The extent to which nitrate is exported from groundwater to surface waters in an upwelling 97 groundwater setting will be controlled by the rate of biogeochemical nitrate removal and the flux of 98 water through the riverbed. The Damköhler number, a dimensionless ratio of reaction rate to 99 transport rate of the solute, can be used to contrast the importance of these two drivers of nitrate 100 removal [*Gu et al.*, 2007; *Ocampo et al.*, 2006]. Damköhler numbers have been widely used in 101 contaminant studies in the hydrogeological literature [*Bahr and Rubin*, 1987] and have also been 102 applied to denitrification in hyporheic zones to distinguish between hydrological and biogeochemical controls on nitrate removal from thalweg and marginal sediments [*Harvey et al.*,
2013]. Recent modelling studies have focused on using residence time analysis to distinguish
between zones of net nitrification and denitrification along hyporheic flow pathways [*Bardini et al.*,
2012; *Marzadri et al.*, 2011; *Zarnetske et al.*, 2012]. Other processes of nitrate reduction, such as
anammox have largely been ignored because their role in nitrate removal is currently thought by
many researchers to be negligible [*Burgin and Hamilton*, 2007].

109

110 Stelzer and Bartsch [2012] have recently developed a conceptual model of nitrate-rich gaining fluvial settings in which nitrate-rich oxic groundwater upwells through deeper riverbed sediments to 111 112 reach a zone enriched with electron donors in the form of particulate organic matter from surface waters. This organically-enriched layer, arising from the deposition and burial of particulate organic 113 matter and varying in thickness (dependent on deposition rate, vertical hydraulic gradient and 114 115 porosity), facilitates the development of hypoxic and anoxic conditions to drive nitrate reduction 116 processes such as denitrification. To date this 'bottom-up' conceptualization of gaining reach settings has focused on the interaction of upwelling groundwater with shallow hyporheic exchange 117 118 flows (HEF). Here, we develop the conceptual model further to evaluate the effect of deeper 119 (> 10cm depth) horizontal subsurface flows on nitrate reduction processes.

120

121 We have previously used measurements of saturated hydraulic conductivity with vertical head gradient from a network of piezometers in a gaining, permeable sandstone reach to show that even 122 123 in a strongly upwelling stream horizontal water fluxes (both lateral and longitudinal; Figure 1a) can 124 influence of on hyporheic zone chemistry [Heppell et al., 2013]. By combining measurement of water flux with an understanding of the spatial variability in redox patterns in the reach we could 125 distinguish nitrate-rich oxic conditions associated with upwelling groundwater from nitrate-poor 126 reducing conditions associated with horizontal flows from hyporheic exchange and/or riparian 127 128 flows [Heppell et al., 2013]. We did not observe nitrate poor, reducing conditions associated with

129 strong groundwater upwelling, probably because the regional aquifer contains little organic carbon 130 [Smith and Lerner, 2008] and, as a result, is oxygenated [Lapworth et al., 2008]. Here, we combine 131 our 3D measurements of spatial variability in vertical and horizontal hydrological fluxes (at a 132 spatial resolution not previously captured in gaining stream settings) with in situ process based 133 measurements of nitrate transformations to investigate the interplay between hydrological and 134 biogeochemical controls on nitrate consumption at the reach scale. We apply the use of Damköhler 135 numbers in order to distinguish between residence time and biogeochemical controls on nitrate 136 reduction in the stream sediments of our gaining reach.

137

138 Specifically, we:

(i) identify the spatial variability in nitrate consumption in a single gaining reach, focusingon the depth distribution of nitrate attenuation.

141 (ii) investigate the factors that controls nitrate consumption in the reach, using Damköhler

numbers to explore the interplay between residence time (hydrological) and biogeochemicalcontrols on nitrate consumption.

(iii) estimate total nitrate consumption within the riverbed using our *in situ* hydrological and
 biogeochemical measurements to quantify the significance of nitrate removal in deep (> 10
 cm) bed sediments of a gaining reach.

147

# 148 **2 Methods**

149 2.1 Site description

150 Our 200m study site, located within the River Leith (Cumbria, UK), receives groundwater from the

151 Aeolian Penrith Sandstone, a major aquifer of the Permo-Triassic Sandstone in the UK [Allen et al.,

152 1997; *Seymour et al.*, 2008]. The gaining reach comprises sandstone bedrock overlain by

153 unconsolidated glacio-fluvial sands and silts (1-2 m) which are topped by sand, gravel and cobbles

154 forming riffle and pool sequences. The catchment of the River Leith is a mixed agricultural

landscape, and the river is a designated Site of Special Scientific Interest (SSSI) and Special Area ofConservation (SAC).

157

158 2.2 Field sampling campaign

159 Riparian and in-stream piezometers (internal diameter = 27mm) were installed in clusters at the site in June 2009 and June 2010 using a percussion drill (see Binley et al. [2013] for a detailed 160 description). Each in-stream cluster comprised three piezometers screened at 100cm, 50cm and 161 162 20cm depth to measure saturated hydraulic conductivity and head gradient. The 100cm in-stream piezometers were fitted with multi-level porewater samplers at target depths of 10, 20, 30, 50 and 163 164 100cm in order to establish porewater chemistry and to enable tracers to be introduced at various depths beneath the riverbed surface. The end of each porewater sampler was wrapped in a fine 165 polyester mesh to prevent ingress of sediment. Collection of porewater samples (total n=72). <sup>15</sup>N 166 167 injections into the multi-level porewater samplers and measurement of vertical hydraulic gradient was performed at 9 points along the study reach (labelled A-I in Figure 1b) from 9-13 August, 2011 168 under low flow conditions ( $<0.5m^3 s^{-1}$ ). At positions A, C and G a transect of three piezometer 169 170 clusters were examined (total number of piezometer clusters = 15) and we were unable to collect porewater from three sampling tubes (G-20cm, H-30cm and I-50cm). 171

172

173 2.2.1 Porewater sampling

Prior to the injection of <sup>15</sup>N-NO<sub>3</sub><sup>-</sup> (*see* below), a 40mL porewater sample was collected from each of
the multi-level samplers via a syringe. A sample of surface water was also collected at each
piezometer cluster. Samples for analysis of chloride and nitrogen species were filtered (0.2μm
polypropylene membrane, VWR International, UK) into plastic vials (polypropylene) in the field
and frozen until later chemical analysis (*see* below). Samples for analysis of dissolved organic
carbon (DOC) were filtered into acid-washed amber glass bottles and acidified to pH<2 with HCl in</li>
the field. For determination of reduced iron (Fe(II)), 1 mL of water was filtered through an oxygen

181	free nitrogen-flushed 0.2µm filter (as above) into 4 mL of phenanthroline-acetate buffer solution
182	and stored in the dark until analysis by UV-spectrophotometry [APHA-AWWA-WPCF, 1976; Grace
183	et al., 2010]. Water samples were also collected to determine the natural abundance <sup>15</sup> N content of
184	nitrogen gas $(N_2)$ and dissolved nitrous oxide $(N_2O)$ and methane concentrations. Gas tight vials
185	(Exetainer, Labco) were overflowed at least two times by gentle discharge of water through a 21-
186	gauge needle to minimize atmospheric exchange and bacterial activity was inhibited by addition of
187	zinc chloride (25 $\mu$ L, 7M). Dissolved oxygen (O <sub>2</sub> ) concentration was measured in the field using a
188	calibrated, fast response electrode (50µm, Unisense, Denmark). Water temperature and pH were
189	measured (pH-100 meter, VWR International, UK) following O <sub>2</sub> determination. For these
190	measurements, water was gently transferred via a three-way stop cock from the collection syringe
191	into an open syringe barrel containing the O <sub>2</sub> electrode or pH probe. We determined the amount of
192	$O_2$ contamination that occurred during sample transfer to be approximately 10 $\mu$ M, and corrected all
193	measured O <sub>2</sub> concentrations accordingly.

194

### 195 2.2.2 In situ measurement of riverbed nitrate reduction

<sup>15</sup>N-labelled NO<sub>3</sub><sup>-</sup> tracer (98 atom % <sup>15</sup>N, Sigma Aldrich) solution was prepared in the laboratory at 196 approximately the same concentration as ambient  ${}^{14}NO_3$  (100, 200, 300, 400 or 500µM  ${}^{15}NO_3$ ) and 197 198 de-oxygenated by bubbling with oxygen-free nitrogen gas (British Oxygen Company). The tracer 199 matrix was artificial river water [Smart and Barko, 1985] tailored to match the major ion chemistry 200 of the River Leith but with added chloride (~4mM KCl) to measure advective flow [Lansdown et 201 al., 2014]. In the field, tracer was drawn into luer-lock syringes under oxygen-free nitrogen or after sparging with air to match ambient  $O_2$  conditions. Sub-samples of the tracer (n=3 per piezometer 202 cluster) were reserved for later chemical analysis and physico-chemical measurements (as above 203 and *see* below). 50mL of <sup>15</sup>N-NO<sub>3</sub><sup>-</sup> tracer was injected into the riverbed via each multi-level 204 205 sampler, with all injections at a piezometer completed within 2.5h. Porewater samples (n=4, 7mL) 206 were collected over time after the dead volume of the sampling tube had been discarded. The first

207 porewater sample was recovered immediately after injection. Recovery of porewaters thereafter 208 occurred according to depth with collection of porewater from10 and 20cm samplers at 5, 10 and 30 209 minutes post injection; 30 and 50cm samplers at 10, 30 and 60 minutes post injection and 100cm 210 samplers at 15, 45 and 120 minutes post injection. Recovered porewater samples were split between 211 gas-tight vials for  $N_2$  analysis and filtered into plastic tubes for anion analysis (using above 212 sampling procedures and analysis methods described below).

213

We worked from downstream to upstream, and from shallow to deep samplers, to ensure that there was no cross-contamination of tracer plumes. Water flux was also sufficiently slow to prevent mixing of tracer injected at different depths within the experimental time frame (*see* Results). Assuming that the injection of the tracer forms a sphere centered at the terminus of the multi-level sampler tube, the magnitude of the <sup>15</sup>NO<sub>3</sub><sup>-</sup> dilution immediately post injection corresponds to a sediment volume of 120 cm<sup>3</sup> (porosity = 0.35). Accordingly, each of our denitrification measurements has a vertical resolution of approximately  $\pm$  3.2 cm.

221

222 2.2.3 Hydrological and sediment analyses

Sediment samples, collected from each core during piezometer installation, were divided into 10cm increments in the field. On return to the laboratory, the sediment samples were air dried and divided for loss on ignition (LOI) and granulometric analysis by sieving and laser diffraction. The < 1mm fraction was digested with 30% hydrogen peroxide to remove organic matter and the samples was dispersed in Calgon before particle size analysis with a Malvern 2000 Mastersizer, Malvern Instruments Ltd., UK). Data from all size distributions were then combined to calculate  $d_{50}$  (mm).

230 Saturated hydraulic conductivity was measured using falling and rising slug tests in the piezometers

at 100, 50 and 20cm depth (see Binley et al., [2013] for detailed description). Head levels in the in-

stream and bank piezometers were measured concurrently with push-pull measurements using an

electronic dip meter. Darcian vertical water flux (m d<sup>-1</sup>) at 100, 50 and 20cm depth was calculated
following the method described in *Binley et al.*, [2013], assuming permeability is isotropic.

235

236 2.3 Laboratory analyses

237 2.3.1 Porewater analysis

238 Nitrate (Limit of detection (LOD) 12 µM, precision 3%) and chloride (LOD 2 µM, precision 1%)

239 were determined using ion exchange chromatography (Dionex ICS2500) whilst ammonium and

240 nitrite were determined by automated colorimetric analysis (Skalar San++) with detection limits and

241 precision of 0.3  $\mu$ M ± 5% and 0.05  $\mu$ M ± 1%, respectively. DOC was analyzed by the non-

242 purgeable organic carbon method (Thermo TOC analyzer; LOD 23µM, precision 5%). N<sub>2</sub>O and

243 methane were determined using gas chromatography (Agilent Technologies) with electron capture

and flame ionization detection, respectively, following addition of a helium headspace (*see* below).

245

# 246 2.3.2 Calculating *in situ* rates of nitrate reduction

A 500 $\mu$ L helium headspace was introduced to each 3mL gas-tight vial and equilibrated with the porewater overnight at 22°C. The <sup>15</sup>N-N<sub>2</sub> content was quantified using mass-to-charge ratios of 28, 29 and 30 measured with a mass spectrometer (Finnigan MAT DeltaPlus) calibrated and corrected for drift following the procedure described in *Trimmer et al.* [2006]. Precision as a coefficient of variation was better than 1%. Production of <sup>29</sup>N<sub>2</sub> or <sup>30</sup>N<sub>2</sub> was quantified as excess above natural abundance, adapted from *Thamdrup and Dalsgaard* [2000]:

253 
$$\Delta N_{2 t=i} (nM N_2) = \left[ \left( \frac{x N_2}{\Sigma N_2} \right)_{t=i} - \left( \frac{x N_2}{\Sigma N_2} \right)_{background} \right] x \Sigma N_2 \text{ sample x } \alpha^{-1} x V_s^{-1}$$
(1)

where  $\Delta^{x}N_{2}$  is the amount of excess  ${}^{29}N_{2}$  or  ${}^{30}N_{2}$  in the recovered porewater at time=i;  ${}^{x}N_{2}/\Sigma N_{2}$ represents the ratio of the  ${}^{29}N_{2}$  or  ${}^{30}N_{2}$  mass spectrometer signal to the total N<sub>2</sub> signal ( $\Sigma N_{2} = {}^{28}N_{2} + {}^{29}N_{2} + {}^{30}N_{2}$ ) for either time series or background samples;  $\alpha$  is the calibration factor (signal: nmol N<sub>2</sub> vial<sup>-1</sup>); and V<sub>s</sub> is the volume of porewater in the gas-tight vial (L vial<sup>-1</sup>). 'Excess' concentrations of  ${}^{29}N_2$  and  ${}^{30}N_2$  in the tracer solution were also calculated via Eq. 1 (where t=i is the tracer) to allow correction for loss through advective flow as follows:

260 
$$\Delta'^{x} N_{2 t=i} (nM N_{2}) = \Delta^{x} N_{2 t=i} + \left[ \left( \frac{\left[ Cl^{-} \right]_{t=i} - \left[ Cl^{-} \right]_{tracer}}{\left[ Cl^{-} \right]_{background} - \left[ Cl^{-} \right]_{tracer}} \right] x \Delta^{x} N_{2 tracer} \right]$$
(2)

where:  $\Delta' {}^{x}N_{2}$  is the concentration of  ${}^{29}N_{2}$  or  ${}^{30}N_{2}$  at time=i corrected for the loss of  ${}^{15}NO_{3}^{-}$  tracer or  ${}^{15}N$  labelled products via advective flow;  $\Delta {}^{x}N_{2 t=i}$  and  $\Delta {}^{x}N_{2}$  tracer are the excess of concentration of  ${}^{29}N_{2}$  or  ${}^{30}N_{2}$  calculated from Eq. 1 in the time series samples and tracer solution, respectively; and [Cl<sup>-</sup>] is the concentration of chloride in the tracer solution (tracer), ambient porewater (background), and porewater collected over time following the injection of  ${}^{15}NO_{3}^{-}$  (t=i).

267 The rate of 
$${}^{29}N_2$$
 and  ${}^{30}N_2$  production ( $p^{29}N_2$  or  $p^{30}N_2$ ) was calculated by linear regression of

268  $\Delta'^{x}N_{2}$  t=i against time. The rate of denitrification was calculated according to *Nielsen* [1992]:

269 Denitrification (nmol <sup>15</sup>N-N<sub>2</sub> L<sup>-1</sup> h<sup>-1</sup>) =  $p^{29}N_2 + 2 x p^{30}N_2$  (3)

Note, as the <sup>15</sup>N-labelling of the N<sub>2</sub> and N<sub>2</sub>O produced after injection of <sup>15</sup>NO<sub>3</sub><sup>-</sup> was the same (*see* Results ) then the contribution of anammox to the production of N<sub>2</sub> gas could be assumed to be negligible [*Trimmer et al.*, 2006] and, as a consequence, Nielsen's original formulation for the isotope pairing technique remained perfectly valid [*Risgaard-Petersen et al.*, 2003].

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The <sup>15</sup>N-labelling of the N<sub>2</sub>O pool following injection of <sup>15</sup>NO<sub>3</sub><sup>-</sup> was determined on a subset of 275 samples (n=49). To quantify <sup>15</sup>N-N<sub>2</sub>O a 100µL sub-sample of the headspace of the gas-tight vial 276 (from above) was injected into an air-filled 12mL gas-tight vial (Exetainer, Labco). The entire 277 278 content of the gas-tight vial was swept, using a two-way needle and analytical grade helium, to a 279 trace gas preconcentrator (Cryo-Focusing; PreCon, Thermo-Finnigan), where the gases are dried 280 and cryo-focused twice in liquid N<sub>2</sub> and before final separation of N<sub>2</sub>O from CO<sub>2</sub> on a PoraPLOT Q capillary column. The sample then passes to mass spectrometer (as above) and the mass-to-charge 281 ratios of 44, 45 and 46 are measured. The amount of dissolved <sup>15</sup>N-N<sub>2</sub>O was calculated by 282

multiplying the total concentration of N<sub>2</sub>O, as measured by gas chromatography, by the proportion of <sup>15</sup>N-label in the N<sub>2</sub>O pool as determined by mass spectrometry (mass-to-charge = 45 / 2 x massto-charge = 46). Concentrations of <sup>15</sup>N-N<sub>2</sub>O were corrected for losses due to advective flow as per equation 2, substituting N<sub>2</sub>O for N<sub>2</sub> values. Rates of <sup>15</sup>N-N<sub>2</sub>O production were then calculated by linear regression of the corrected concentrations against time.

288

# 289 2.5 Data analyses

290 2.5.1 Assigning piezometer clusters to hydrologic setting using porewater chemistry.

291 Each piezometer cluster was assigned to one of three hydrologic settings using chloride

292 concentrations in a two end-member mixing model as follows:

293 Mixing score = 
$$\frac{\left\lfloor Cl^{-} \right\rfloor_{\text{porewater}} - \left\lfloor Cl^{-} \right\rfloor_{\text{surface water}}}{\left\lfloor Cl^{-} \right\rfloor_{\text{surface water}} - \left\lfloor Cl^{-} \right\rfloor_{100\text{cm}}}$$
(4)

294 where "porewater" refers to samples collected between 10 and 50cm depth in the riverbed and 295 "100cm" was porewater recovered from 100cm. Scores can range from -1 to 0. The lower range 296 indicates dominance of upwelling porewater and the higher range indicates maximum surface water 297 influence. Hydrology at piezometer was classed as strong porewater upwelling when scores were -1 298 and hyporheic exchange flows (HEF) when scores vary between -1 and 0. Horizontal water fluxes, 299 for example longitudinal flow along the river channel or lateral inputs from the riparian zone, were 300 inferred when scores were <-1 (no scores were >0). As such, horizontal water fluxes cannot be 301 detected with this method if chloride concentrations of the horizontal source are the same as surface 302 water and upwelling porewater. We are confident however, that the assigned hydrologic settings 303 reflect actual subsurface hydrology as classifications compare favorably with the zones of upwelling, HEF and horizontal fluxes inferred through in situ measurements by Binley et al. [2013]. 304 305

306 2.5.2 Calculations for integrating flux and nitrate removal in sediment via denitrification.

307 Initially, we examined the relative importance of denitrification activity at different depths in the 308 riverbed by simply contrasting rates of reaction, as per *Stelzer et al.* [2012; 2011]. The proportion of 309 denitrification activity at each depth was determined by dividing the individual rate by the sum of 310 all rates within a piezometer cluster (Table 1). We refer to these data as rate-determined 311 proportions.

312

313 Nitrate removal within a riverbed will depend not only upon the denitrification rate (as above) but 314 also on amount of time a parcel of water is exposed to a given denitrification rate (as per Harvey et al. [2013]). In order to explore the effects of spatial variations in upwelling water flux on the extent 315 316 of nitrate removal via denitrification we calculated the residence time of upwelling water in each 317 sediment section (0-10cm, 10-20cm etc.) where residence time was the inverse of the relevant Darcy vertical water flux. We measured saturated hydraulic conductivity at 20, 50 and 100 cm 318 319 depths only, so we estimated residence time of the 0-10cm and 30-50cm depth bands by assuming 320 vertical flux at 10cm was equal to vertical flux at 20cm and the vertical flux at 30cm was the average of fluxes at 20 and 50cm. We then multiplied the *in situ* rate of denitrification (nmol <sup>15</sup>N-N<sub>2</sub> 321  $L^{-1}h^{-1}$ ) by the residence time (h) to calculate the amount of nitrate removed from each sediment 322 323 section as upwelling water passed through it. In order to express this nitrate removal on a sediment volume basis (mmol N m<sup>-3</sup>) we assumed a sediment porosity of 0.35 (-). Relative magnitudes of 324 denitrification activity within each sediment section were calculated as above, however, we refer to 325 these data as depth-integrated proportions. 326

327

Areal rates of denitrification (µmol N m<sup>-2</sup> h<sup>-1</sup>) were estimated by converting measured
denitrification rates (per volume of porewater) through integration of denitrification activity within
depth profiles (*see Laverman et al.* [2007] for similar calculations). Integration was performed
using the trapezium rule and the *in situ* denitrification rate in the 0-10cm depth band was estimated

by extrapolating the trend from measurements at 30, 20 and 10 cm data from shallow sediments to0cm depth.

334

335 2.5.3 Damköhler number calculation

336 The Damköhler number for denitrification (Da<sub>N</sub>) is the dimensionless ratio between a transport ( $\tau_T$ ) and denitrification reaction ( $\tau_R$ ) timescale.  $\tau_T$  is the residence time (units = d) and  $\tau_R$  is the inverse 337 of the first-order reaction rate constant for denitrification,  $K_1$  ( $\tau_R = 1/K_1$ , units for  $K_1$  and  $\tau_R$  are d<sup>-1</sup> 338 and d, respectively) [Harvey et al., 2013]. Our denitrification rates were zero-order, i.e. production 339 340 of <sup>15</sup>N-labelled N<sub>2</sub> was linear with time. To convert our data to first order rate constants we divided the zero-order rate constants (units = nmol  $L^{-1} h^{-1}$ ) by the mean half saturation constant for 341 342 denitrification ( $K_m = 109 \mu M$ ) in rivers of north east England reported by García-Ruiz *et al.* [1998].  $Da_N(\tau_T / \tau_R)$  values < 1 indicate that transport dominates over reaction, whilst values > 1 indicate 343 that reaction processes are occurring faster than advection [Ocampo et al., 2006]. 344

345

346 2.5.4 Statistical analysis

347 All statistics were performed in R [R Development Core Team, 2012]. Differences in chemical 348 parameters across sediment depth bands and hydrologic settings were investigated by two-way analysis of variance (ANOVA). Prior to each ANOVA we tested whether the data were normally 349 350 distributed with a Shapiro Wilk test and attempted to normalize any data where p>0.05. For most 351 variables however, distributions were unable to be normalized using common methods (e.g. log and power transformations) so we rank transformed data prior to the ANOVA. Any significant effects 352 353 detected in the ANOVA (p>0.05) were further examined by pairwise comparisons using a Post-hoc Tukey's test. A paired t-test was used to test for any difference in the <sup>15</sup>N labelling of the N<sub>2</sub> and 354 N<sub>2</sub>O pools following <sup>15</sup>NO<sub>3</sub><sup>-</sup> addition. Relationships between denitrification rate and chemical 355 356 variables were explored using Spearman's rank correlation.

- 358 **3 Results**
- 359

#### 360 **3.1 Summary of reach characteristics**

361 *General description of stratigraphy* 

Median grain size ( $d_{50}$ ) decreased with depth in the riverbed (Table 2); superficial sediments are a mixture of coarse sand, gravel and small cobbles, whilst deep (> 10 cm) sediments are mainly sand. Sediment particulate organic matter content was low, ranging from 0.3-3.3% (loss on ignition), and decayed with depth (ANOVA, F(4,33)=5.6, p=0.002).

366

# 367 Spatial distribution of porewater chemistry and water fluxes

Vertical head gradients were positive throughout the <sup>15</sup>N injections period (range = 0.5 to 28%) 368 indicating upward movement of water from the sediments towards the river. Chloride profiles 369 370 provided further evidence of porewater upwelling but also revealed localized zones of surface water downwelling and horizontal flows (e.g. plots B, F and D, respectively; Figure 1c). We examined 371 372 these water fluxes further using a two end-member mixing model, not to apportion water source in 373 the riverbed, as depth profiles suggested at least three sources of water were present at some sites, 374 but to quantitatively assign piezometer clusters to hydrologic settings. Subsurface hydrology could 375 be described as either strong porewater upwelling, surface water downwelling to ≤20cm (HEF) or horizontal flows (n = 6, 5 and 4 respectively; Figure 2). Water residence time, based on a calculated 376 Darcy flux, varied from 0.11d to 32d for 10cm and 50cm flow pathways, respectively when water 377 378 flux was predominantly vertical (i.e. strong upwelling and/or HEF). Water residence times were 379 slightly longer when upwelling was strong (mean  $\pm$  s.d. = 5 $\pm$ 2d and 3 $\pm$ 1d for upwelling and HEF, respectively) and in deep sediments (mean  $\pm$  s.d. = 0.7 $\pm$ 0.1d and 2.2 $\pm$ 0.5d per 10cm for sediments 380 381 from 0-20cm and 30-100cm, respectively).

Porewater nitrate concentrations were highly variable, ranging from below detection to  $\geq 600\mu$ M at each of the depths examined. Overall, nitrate concentrations within deep sediments (100cm) were higher than those in the river (mean  $\pm$  s.d. = 311 $\pm$ 182  $\mu$ M and 129 $\pm$ 18  $\mu$ M, respectively) and tended to decrease towards the riverbed surface (Table 2). This trend was not statistically significant. Along-reach variation in porewater nitrate was also evident (Figure 3a), and concentrations were typically high in areas characterized by upwelling porewater (Figure 3b; F(2,56)=11.1, p<0.001).

Porewater DOC concentrations did not differ with depth in the riverbed (Table 2) relative to
variation across the study reach (Figure 3c). Shallow sediments at 10 cm depth exposed to HEF
were sites of elevated DOC (*see* points A, F and H in Figure 3c and DOC concentrations
summarized by depth and hydrology in Supplementary Information), although when integrating
data across all depths (0-100 cm), differences between hydrologic settings were not significant
(Figure 3d).

396

397 Porewaters were generally under-saturated in  $O_2$  (mean  $\pm$  s.d. = 49 $\pm$ 21 %) and, like nitrate, oxygen concentrations decreased towards the riverbed surface (Table 2, Figure 3e); although the trend was 398 399 not statistically significant. When considering depth-distributions, dissolved oxygen concentrations 400 were most heterogeneous at 10 cm depth in shallow sediments exposed to HEF (range =  $308\mu$ M, 17 401 to  $325\mu$ M). The next largest range in O<sub>2</sub> concentrations was  $180\mu$ M (57 to  $237\mu$ M) at 50cm under 402 horizontal water flux. Highest median O2 concentrations were associated with sediments dominated 403 by porewater upwelling (Figure 3f; ANOVA, F(2,57)=6.9, p=0.002), and these sediments were also 404 associated with the lowest concentrations of other reduced chemical species such as Fe(II) and 405 methane. Ammonium concentrations were highly variable both with depth in the riverbed and 406 across the reach, ranging from below detection to  $125\mu$ M (mean  $\pm$  s.d. =  $5\pm19\mu$ M).

407

#### 408 **3.2** Factors controlling *in situ* rates of denitrification and overall nitrate removal

### 409 *Quantification of in situ denitrification rates*

Rates of denitrification, as determined through the production of <sup>15</sup>N-N<sub>2</sub>, ranged from 25 to 17,053 410 nmol <sup>15</sup>N L<sup>-1</sup> h<sup>-1</sup> varying both with piezometer cluster and depth across the study reach (Figure 3g). 411 We attribute this <sup>15</sup>N-N<sub>2</sub> production to denitrification rather than anammox as the <sup>15</sup>N-labelling of 412 413 the N<sub>2</sub> and N<sub>2</sub>O pools were not significantly different (t(49)=0.766, p = 0.448; Figure 4a and see 414 below). The rate of denitrification generally decreased with depth in the riverbed (ANOVA, F(4,57) = 4.0, p = 0.006), except where subsurface water flux was horizontal (Table 3). The relationship 415 416 between depth and denitrification was strongest within sediments with upwelling porewater (Table 417 3) but overall, rates of denitrification here were lowest compared to sediments with HEF or 418 horizontal flows (F(2,57) = 3.6, p = 0.034). In the presence of HEF, there was a clear 'step down' 419 between denitrification rates in shallow (<10cm) and deep sediments (Table 3). Rates of denitrification increased with both the DOC concentration of porewater and organic matter content 420 421 of sediment (as LOI, Table 4) and when porewater chemistry was reduced (i.e. low in O<sub>2</sub>, high Fe(II) and CH<sub>4</sub>). We did however, observe denitrification in seemingly oxygenated sediments (see 422 Supplementary Information). Even when the porewater DO concentration was > 200  $\mu$ mol L<sup>-1</sup> O<sub>2</sub> 423 (~60% of air saturation), denitrification could still be measured at up to 3249 nmol  ${}^{15}N-N_2 L^{-1} h^{-1}$ 424 (median = 329 nmol  ${}^{15}$ N–N<sub>2</sub> L<sup>-1</sup> h<sup>-1</sup>, n = 21) 425

426

In the majority of cases, denitrification was complete, i.e.  ${}^{15}$ N-N<sub>2</sub> comprised >99% of the  ${}^{15}$ Nlabelled gas produced, but there were some samples where a considerable fraction of  ${}^{15}$ NO<sub>3</sub><sup>-</sup> reduction stopped at N<sub>2</sub>O (maximum N<sub>2</sub>O/N<sub>2</sub>O + N<sub>2</sub> = 51%, median = 6%; *n*=9 of the 49 examined). Although the patterns were not that strong, higher values for incomplete denitrification were most strongly correlated with high Fe(II) and CH<sub>4</sub> i.e. where sediments were most reduced.

433 The denitrification rates per unit volume were integrated over the top 100 cm to give an estimate of 434 areal activity. Areal activity ranged from 132 to 4597  $\mu$ mol <sup>15</sup>N m<sup>-2</sup> h<sup>-1</sup>, with a median value of 480 435  $\mu$ mol <sup>15</sup>N m<sup>-2</sup> h<sup>-1</sup> (*n*=15). There was no significant difference between areal rates within different 436 hydrological settings (F(1,13) = 3.13, *p* = 0.01; median areal rate = 479, 562 and 1026  $\mu$ mol <sup>15</sup>N m<sup>-2</sup> 437 h<sup>-1</sup> for strong upwelling, HEF and horizontal fluxes, respectively).

438

### 439 Nitrate export using Damköhler analysis

440 Controls on nitrate export were also investigated through calculation of Damköhler numbers, which

441 varied both with depth and piezometer cluster across the study reach (range = 0.003 to 36, n=72).

442 The majority of values however, were below the biogeochemical control threshold of  $Da_N=1$ 

443 (median = 0.14, Figure 5).  $Da_N > 1$  (*n*=14) were generally associated with deeper sediments (e.g.

 $\geq 20$  cm) in two piezometer clusters, sites A and G, that were characterized by HEF and horizontal

- 445 water fluxes, respectively.
- 446

### 447 Nitrate attenuation in riverbed sediments

448 Deep sediments were important sites of nitrate attenuation, however, with ~80% of denitrification, on average, occurring between 10 and 100cm depth in the riverbed (Table 5, depth-integrated data). 449 450 On a per cm basis however, sediments within the 0-10cm depth band were sites of enhanced nitrate 451 removal under strong upwelling or HEF (Table 5, removal data per depth band divided by height of 452 depth band). Total nitrate removal per piezometer cluster was highest in sites identified as biogeochemical hotspots (i.e.  $Da_N > 1$ , see above), similar when water flux was predominantly 453 454 vertical (i.e. upwelling and HEF) and lowest when water flux was horizontal (Table 5). These 455 differences however, were not statistically significant.

456

# 457 **4 Discussion**

458 Here, we have examined nitrate attenuation along upwelling flow pathways in a gaining reach low

459 in organic carbon, measuring nitrogen transformations *in situ*, and simultaneously characterizing

460 subsurface hydrology. Our integrated rates of denitrification (mean  $\pm$  s.e. = 1078 $\pm$ 363 µmol m<sup>-2</sup> h<sup>-1</sup>)

are comparable to those measured by *in situ* "whole stream"  ${}^{15}NO_3$  additions in rivers within 461 agricultural catchments [Mulholland et al., 2009]. Our work adds value to the existing evidence 462 base of riverine nitrogen cycling because we also characterize denitrification below the zone of 463 464 surface water – groundwater mixing. Denitrification occurred throughout the 100cm depth of riverbed we examined, despite the limited amount of organic carbon and the moderate O<sub>2</sub> content of 465 the upwelling porewater, demonstrating that the attenuation of nitrate is not just confined to shallow 466 467 sediments within this groundwater-fed system. Our findings are consistent with those of Storey et 468 al. [2004], Fischer et al. [2005] and Stelzer et al. [2011] and we show here that deep sediments are 469 important sites of nitrate attenuation.

470

#### 471 **4.1 Pathways of nitrate reduction**

472 Removal of nitrate along upwelling flow pathways could occur via a number of different 473 metabolisms, e.g. denitrification, anammox and assimilatory uptake [Burgin and Hamilton, 2007]. 474 The fate of nitrate in the riverbed is important as the benefits of nitrate attenuation could be offset if 475 the removal of nitrate occurs at the expense of production of more bioavailable and potentially harmful forms of nitrogen, e.g. NH<sub>4</sub><sup>+</sup> or N<sub>2</sub>O [Burgin and Hamilton, 2008; Burgin et al., 2013]. In 476 477 agreement with our previous slurry potential incubations [Lansdown et al., 2012] there was no significant anammox activity in situ (proportion of <sup>15</sup>N in N<sub>2</sub> =  $0.57 \approx$  proportion of <sup>15</sup>N in N<sub>2</sub>O = 478 0.54, Figure 4a) and all of the  ${}^{15}N_2$  gas produced could be ascribed to denitrification. The decrease 479 in the proportion of <sup>15</sup>N in N<sub>2</sub> or N<sub>2</sub>O relative to the injected <sup>15</sup>NO<sub>3</sub><sup>-</sup> (98% <sup>15</sup>N) tracer, reflects 480 mixing of the tracer plume with ambient porewater  $^{14}NO_3^{-}$  pool. 481

482

483 For the majority of cases, denitrification was complete, however, for a subset of samples, a

484 significant accumulation of N<sub>2</sub>O was measured (mean  $\pm$  s.d = 15 $\pm$ 20 % N<sub>2</sub>O, *n* = 9) peaking at 51%.

485 Incomplete denitrification was not restricted to individual piezometer clusters, particular hydrologic

486 settings or sediment depths. The production of N<sub>2</sub>O in soils is well characterized but it is poorly

487 constrained in rivers; though the data available suggest a strong influence of hypoxia [*Rosamond et* 488 *al.*, 2012] and here most of the variance in the accumulation of N<sub>2</sub>O production was correlated with 489 patches of low redox environment (accumulated  $NH_4^+$ , Fe(II) and methane) and enhanced microbial 490 activity (low O<sub>2</sub> saturation and fast rates of denitrification). Given N<sub>2</sub>O has a greenhouse warming 491 potential ~280 times that of CO<sub>2</sub> [*Reay et al.*, 2012], the potential environmental trade-off between 492 nitrate attenuation and potent greenhouse gas production via riverbed denitrification warrants 493 further investigation.

494

### 495 **4.2 Interplay between hydrological and biogeochemical controls**

## 496 The role of HEFs in controlling in-situ denitrification

497 Stelzer and Bartsch [2012] described a conceptual model whereby nitrate removal within gaining reaches receiving oxygenated groundwater will proceed only when the upwelling flow path 498 499 interacts with favorable redox conditions created through degradation of deposited and buried 500 particulate organic matter. A similar explanation was given for patterns in nitrate concentrations in 501 the River Tern, except that clay or peat lenses rather than particulate organic carbon derived from 502 ingression, controlled nitrate removal [Krause et al., 2013]. Here, we have quantified denitrification 503 activity to 100cm depth in a riverbed that comprised <1% organic matter, on average (maximum 504 LOI = 3%, cf. average ~12 %, maximum = 50 % in *Stelzer* and *Bartsch* [2012]. We found no 505 evidence for lenses of buried organic matter up-gradient of the flow pathways in this reach despite 506 extensive drilling within the study site (> 100 piezometers within a 200 m reach).

507

508 It would appear that nitrate removal within this gaining reach does not require meter-scale patches 509 of buried particulate organic matter to generate favorable redox conditions for denitrification to 510 occur [sensu *Krause et al.*, 2013]. Rather, we propose that inputs of DOC and particulate organic 511 carbon to the riverbed (i) from HEF [sensu *Stelzer and Bartsch*, 2012] and (ii) via subsurface routes 512 from the floodplain or riparian zone are the key mechanisms driving heterotrophic denitrification in 513 this instance. Conceptually, HEFs could either stimulate denitrification activity by supply of labile 514 organic carbon to the sediments, or suppress denitrification activity as well oxygenated surface 515 water downwells. Here, as with our previous cm-scale investigation [Lansdown et al., 2014], 516 denitrification was observed in porewaters with  $O_2$  concentrations >200 $\mu$ M, although this activity is probably confined to anoxic microsites within oxygenated sediments [Triska et al., 1993]. The role 517 518 that HEFs play in increasing nitrate attenuation capacity of riverbeds [as per *Fischer et al.*, 2005; 519 Harvey et al., 2013; Zarnetske et al., 2011] can be observed in our shallow sediments (< 10 cm 520 depth, Table 3) around piezometer clusters A, C2 (right margin), F, H and I. In these sediments, dissolved O<sub>2</sub> concentrations were elevated, approaching air-equilibrated values in some cases, 521 indicating ingress of well oxygenated surface water (i.e. HEFs). Supply of organic matter from the 522 523 river above through HEFs is inferred from elevated concentrations of DOC in porewaters at 10cm, as well as accumulation of products of mineralization (ammonium, methane see Supplementary 524 525 information). Here, 44% of total depth-integrated nitrate removal (0-100 cm depth) occurred within 526 10 cm of the sediment surface highlighting the increased denitrification capacity of sediments under 527 HEFs.

528

529 It is, however, more difficult to account for the increased denitrification activity in shallow 530 sediments where strong porewater upwelling suppresses groundwater-surface water exchange (as 531 seen in chloride depth profiles of piezometer clusters B, C and D; Figure 1c). Within this hydrological setting approximately 17% of total depth-integrated nitrate removal (0-100 cm depth) 532 533 occurred in the top 10 cm of the riverbed. Here we aimed to characterize riverbed nitrate removal 534 under base flow conditions. A parallel study at this site has quantified the effect of rising river stage 535 on porewater chemistry, finding that stage increase can cause reversal in the vertical hydraulic 536 gradient, potentially allowing surface water to infiltrate areas of the riverbed where no exchange 537 occurs under low flow conditions, altering porewater chemistry [Byrne et al., 2013]. Our finding of 538 enhanced nitrate attenuation capacity within shallow sediments, with little apparent hydrological

539 connection to surface water, or the floodplain, could be explained by groundwater-surface water 540 exchange or horizontal inputs under high flows (e.g. storm events) prior to the sampling campaign. 541 The potential for such event flows to alter biogeochemical cycling, and the timescale over which 542 processes could be affected, is poorly understood in rivers [Zimmer and Lautz, 2014]. Through a 543 combination of modelling and laboratory simulation, Gu et al. [2008] have shown that nitrate attenuation, via denitrification in upwelling groundwater, can be altered simply through changes in 544 545 residence times caused by hydraulic head variation associated with river stage rise. They did not 546 explicitly consider what biogeochemical effects stage variation could have on the subsurface 547 chemistry, but if a change in river stage can alter nitrate attenuation by altering residence times, 548 even after the "event flow" has passed [Gu et al., 2008], porewater chemistry and associated 549 biogeochemical cycling (i.e. rates of denitrification) could be similarly affected.

550

### 551 Are hot spots of denitrification related to horizontal flow pathways?

Variation in the direction and magnitude of water fluxes alter both residence time and subsurface 552 chemistry within the riverbed, thus affecting not only the depth distribution of biogeochemical 553 554 activity, but also spatial zonation of processes such as denitrification across the riverbed. At sites A 555 and G, the total depth-integrated rate of denitrification was two orders of magnitude higher than the rest of the reach (1259 mmol N m<sup>-3</sup>). Here, porewaters were very reduced (low O<sub>2</sub> and high 556 ammonium and methane), denitrification rates were very high and Damköhler numbers were >1; 557 indicating that nitrate export is controlled by reaction-rate rather than residence time. These sites 558 559 were previously identified as biogeochemical hotspots by Heppell et al [2013], where horizontal 560 water fluxes (defined as lateral inputs or HEFs) dominated over groundwater upwelling, supplying 561 organic matter to the subsurface which was then mineralized. Heppell et al [2013] suggested that 562 these sites would exhibit enhanced nitrate removal, and for these patches of riverbed this is indeed the case. From more spatially extensive porewater chemistry obtained through previous work 563 564 [Heppell et al., 2013; Lansdown et al., 2014] we estimate that such biogeochemical hotspots cover

approximately 2.5 % of the study reach ( $\sim 47m^2$ ) but, within which, 8% of the nitrate removal within 565 566 the top 1m of riverbed occurs (average removal within these biogeochemical hotspots divided by average removal across the rest of the reach). These estimates of nitrate removal along horizontal 567 568 flow pathways were performed assuming a maximum flow path of 1m. Where flow pathways were 569 horizontal, rather than vertical, we assume that vertical and horizontal water fluxes were equal, on average [see *Binley et al.*, 2013] and, therefore, that the flow pathway is approximately  $45^{\circ}$ . As 570 571 such, there will be no net effect on residence time over the flow pathway if the direction of flow is 572 horizontal rather than vertical. However, this research also shows that not every area characterized 573 by horizontal flows is a hot spot of nitrate reduction. In fact, for three out of the four zones 574 associated with horizontal water movement, there was less nitrate removed from the top 100 cm of sediments compared to the vertical flow settings (9 mmol N m<sup>-3</sup>). The reasons for this warrant 575 further research but it is likely that variation in the origin and length of horizontal flow paths across 576 577 the reach will influence the quality of the DOC; for example, some horizontal pathways will originate from the nearby riparian and floodplain areas potentially comprising labile DOC whilst 578 579 others will be from deeper groundwater and potentially be characterized by more recalcitrant DOC 580 compounds.

581

582 Through geophysical measurements *Binley et al.* [2013] also identified a preferential discharge zone in this reach at sites B to D, where upwelling porewater flux was very strong and, as a result, there 583 was little exchange with surface water or horizontal water inputs. The short water residence time, 584 585 combined with the high nitrate load in the oxygenated upwelling porewater, led *Heppell et al.* 586 [2013] to propose that nitrate removal would be minimal within this patch of the riverbed. Our results also support this finding as this preferential discharge zone is indeed a cold spot for 587 denitrification: occupying ~20% of the reach area [Binley et al., 2013] but performing <2% of the 588 589 total denitrification that occurs within the reach (average removal from sites B, C and D divided by 590 average removal across the rest of the reach).

591

### 592 The importance of deep sediments for nitrate attenuation.

593 Across the reach as a whole, denitrification within the top 1m of the riverbed removed between 0.3 594 and 32% of nitrate exported from upwelling porewater (median = 9%) but denitrification activity 595 was not equally distributed with depth or hydrologic setting. Considering rates of denitrification 596 alone (see rate-determined denitrification in Table 5), as per Stelzer et al. [2011], deep sediments 597 (>10cm) accounted for 64% of subsurface nitrate removal. However, simply integrating 598 denitrification rates over a given depth to estimate overall nitrate removal within a volume of 599 sediments ignores the potential influence of water residence time on nitrate flux, which can be an 600 important predictor of the fate of nitrate in sediments [Zarnetske et al., 2011]. Damköhler numbers 601 indicate that riverbed nitrate attenuation within this gaining reach is limited by the rate of 602 denitrification (i.e. Da<sub>N</sub><1; Figure 5) and, as a result, nitrate flux to the river above is more strongly 603 controlled by water residence time. When nitrate attenuation is considered as the interaction of hydrology and biogeochemistry (i.e. the product of denitrification rate and water residence time) we 604 605 calculate that, on average, 81% of subsurface nitrate removal occurred within deep sediments 606 (depth-integrated denitrification in Table 5). Nitrate removal of the magnitude presented in Table 5 607 could only occur if the supply of nitrate in the upwelling porewater exceeded the removal capacity 608 of the sediments. Measured porewater nitrate concentrations compare favorably with those 609 predicted from depth-integrated nitrate loss except in the sites identified as biogeochemical hotspots 610 (Figure S2) suggesting data presented in Table 5 are likely indicative of actual rather than potential 611 nitrate removal in the sediments.

612

Maximum nitrate removal will occur when timescales of denitrification and water residence time are well matched [*Gu et al.*, 2007] but nitrate attenuation can also be enhanced when denitrification rate is fast but water residence time is short, or vice versa. *Harvey et al.* [2013] explained equal contributions of fine marginal and coarse thalweg sediments to hyporheic nitrate removal via this 617 mechanism. Here we show such a relationship can also explain nitrate removal with depth in a 618 gaining reach. Where groundwater flux was predominantly vertical, (i.e. strong upwelling or HEF) denitrification rates decayed with depth as the influence of the river on the sediments below 619 620 diminished (see above). Saturated hydraulic conductivity and vertical hydraulic gradients were 621 lower in deep sediments [50cm, 100cm; Binley et al., 2013] resulting in longer water residence times. Consequently, shallow sediments (10cm, 20cm) were characterized by fast denitrification but 622 623 short residence time, whilst in deep sediment denitrification was slow and residence time was long 624 in deep sediments, with a switch between the two scenarios at ~30cm (Figure 6). These results 625 illustrate that both physical and biogeochemical controls on nitrate attenuation, and the interaction 626 thereof, can vary along upwelling flow pathways in a gaining reach.

627

Prior use of Damköhler numbers to investigate nitrate flux within stream beds and the riparian zone 628 629 have assumed denitrification rate and/or water flux to be constant along flow pathways [Gu et al., 2007; Ocampo et al., 2006]. Both our study and that of Harvey et al. [2013] highlight the variability 630 631 of denitrification and water flux timescales across relatively small spatial scales (e.g. m to dm). Here we have shown in a gaining reach, that denitrification is most variable within shallow 632 sediments, whilst high variation in water residence time is associated with deep sediments and, 633 634 therefore, use of a single denitrification rate and water flux value to categorize nitrate transport is 635 simply not appropriate.

636

### 637 **5. Conclusion**

The results of this study provide quantitative evidence for nitrate attenuation within the bed of a groundwater-fed river is controlled by both biogeochemical and hydrologic processes. We have shown that denitrification occurs within carbon-poor, sandy sediments to a depth of at least 1m below the riverbed surface without substantial deposits of buried organic matter, at rates that are generally low, but a similar order of magnitude to the global evidence base of rivers in agricultural 643 landscapes. The majority of nitrate attenuation in our reach is transport-controlled irrespective of 644 the flow pathway (vertical or horizontal). In the case of the River Leith, hyporheic exchange flows 645 and horizontal water fluxes such as shallow groundwater inputs from the floodplain or riparian zone 646 are important because they supply the precursor substrates needed to sustain denitrification. We 647 identified two hot spots of denitrification (which are reaction-rate controlled) located within areas 648 of hyporheic exchange or horizontal water flux, and we estimate that these zones account for 8% of 649 overall nitrate attenuation in the bed sediments.

650

Both reaction rate and water residence times change with depth in the riverbed under stable, low flow conditions. Along an upwelling flow pathway residence time is the most important control on nitrate removal at depth (> 20 cm) whilst the rate of denitrification increases, and leads to enhanced nitrate removal near the sediment surface. Overall, our results highlight the importance of using measurements of both biogeochemical reaction rates and residence time to estimate the extent of nitrate removal from riverbed sediments because, in our reach, ignoring residence time underestimates the importance of deep sediments for nitrate removal by about 20%.

658

We recommend that future work should not only continue to explore factors controlling variation in transport and rate-limited reactions at the reach scale, but also to attempt to up-scale such analysis to consider the effect of different hydro-geological settings on the balance of biogeochemical and hydrological controls influencing nitrate removal.

663

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- 672
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### 836 List of Tables

Table 1: Summary	v of denitrification calculations
Parameter	Method of calculation
Rate-determined	Denitrification rate (nmol ${}^{15}$ N-N <sub>2</sub> L <sup>-1</sup> h <sup>-1</sup> ) <sub>depth=i</sub>
proportion	$\Sigma$ Denitrification rate depth = 10,20,30,50,100 cm
Depth-integrated	Denitrification rate (nmol <sup>15</sup> N-N <sub>2</sub> L <sup>-1</sup> d <sup>-1</sup> ) <sub>depth=i</sub> x residence time (d) <sub>depth=i→i'</sub>
proportion	$\sum \text{Denitrification rate}_{\text{depth}=10,20,30,50,100\text{cm}} \text{ x residence time}_{\text{depth}=0-10,10-20,20-30,30-50,50-100\text{cm}}$
Areal rate <sup>a,b</sup>	$\sum_{n=1}^{\infty} \left[ \text{Denitrification rate } (\mu \text{mol m}^{-3} \text{ h}^{-1})_{\text{depth}=i} \text{ x } (\text{depth}_{i} - \text{depth}_{i'} (m)) \right]$
	$\frac{2}{2}$ + Denit. rate <sub>depth=i'</sub> x (depth <sub>i</sub> - depth <sub>i'</sub> )
Damköhler	Residence time (h) $\div$ $\frac{1}{1}$
number	Denit. rate (nmol $^{15}$ N-N <sub>2</sub> L <sup>1</sup> h <sup>-1</sup> )
	$K_m$ (nM)
adapth_i dapatas a	$a_{a}$ of the denthe complet (10, 20, 20, 50, 100 cm) and $b_{a}$ is the distance

837 Table 1: Summary of denitrification calculations

<sup>a</sup>depth=i denotes any of the depths sampled (10, 20, 30, 50, 100cm) and <sup>b</sup>  $i \rightarrow i'$  is the distance between depth = i and that sampled above (i').

840

Table 2: Median solute concentrations, sediment characteristics and vertical water flux by depth inthe riverbed

	Water	(µM)			Sediment		Vertical water flux	
Depth	NO <sub>3</sub> <sup>-</sup>	DOC	O <sub>2</sub>	Cl	LOI (%) <sup>a</sup>	$d_{50}(mm)^{b}$	$(m d^{-1})$	n=
Surface	125	187	276	701	0.9	6.5	-	7
10cm	128	170	142	433	1.1	2.2	-	15
20cm	240	166	149	408	0.8	0.62	0.12	14 <sup>c</sup>
30cm	227	139	180	418	0.5	0.42	-	14
50cm	200	143	157	410	0.5	0.44	0.02	14 <sup>c</sup>
100cm	303	165	197	409	0.4	0.37	0.04	15 <sup>c</sup>

<sup>a</sup>LOI denotes loss on ignition, a proxy measure of organic matter within sediment. <sup>b</sup> $d_{50}$  represents median grain size. <sup>c</sup>n for vertical flux measurements were 13, 14 and 14 for 20cm, 50cm and 100cm, respectively.

846

847 <u>Table 3: Rates of denitrification measured *in situ* in the riverbed</u>

		Denitri	fication (nmol	$^{15}$ N-N <sub>2</sub> L <sup>-1</sup> h <sup>-1</sup> )					
Depth	Range	Median <sup>a</sup>	Upwelling <sup>b</sup>	Hyporheic	Horizontal <sup>b</sup>				
	C		1 0	exchange <sup>b</sup>					
10cm	184-6314	1081	1075	1486	1062				
20cm	148-10048	539	726	531	406				
30cm	47-17053	362	257	246	644				
50cm	29-4165	341	221	310	666				
100cm	25-2977	178	132	78	509				
0									

<sup>a</sup>n = 15 for 10 and 100cm depth bands and 14 for 20, 30 and 50cm depth bands. <sup>b</sup>Data are median values, n = 6 for all upwelling depth bands, n = 5 for 10, 20 and 100cm hyporheic exchange depth bands with n = 4 for the remaining depths in this setting, n = 3 for the 20cm horizontal depth band with n = 4 for the remaining depths in this setting.

Table 4: Correlation between *in situ* rates of denitrification (D15) and chemical composition of porewater and sediment

poremater	and scannen							
	D15	$O_2$	NO <sub>3</sub> <sup>-</sup>	Fe(II)	$CH_4$	DOC	LOI <sup>a</sup>	
D15	1.00							
$O_2$	-0.406**	1.00						
NO <sub>3</sub> <sup>-</sup>	-0.362**	0.547**	1.00					
Fe(II)	0.416**	-0.532**	-0.541**	1.00				
$CH_4$	0.394**	-0.269*	-0.442**	0.566**	1.00			
DOC	0.357*	-0.219*	-0.126	0.509**	0.297*	1.00		
LOI	0.331*	-0.343*	-0.104	-0.005	0.082	0.152	1.00	

855 \* p<0.05, \*\*p<0.001. Data are spearman's rank correlation coefficients, n = 72 per test except <sup>a</sup>LOI 856 where n = 48.

857

858	Table 5: Co	omparison	of nitrate	removal	calculated	with and	without	inclusion	of water	residence
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time in the riverbed (depth-integrated and rate-determined, respectively) and across different

860 hydrologic settings

Depth	Proportion of Nitrate removal <sup>a</sup>		Nitrate removal (depth-integrated, mmol N m <sup>-3</sup> ) <sup>b</sup>					
Dana (cm)	Rate-	Depth-	Upwelling	Hyporheic	Horizontal	Hot spots <sup>c</sup>		
(CIII)	determined	integrated		exchange				
0-10	0.36	0.19	2.7	7.0	1.2	24		
10-20	0.28	0.12	1.8	1.1	0.5	45		
20-30	0.16	0.05	1.8	0.9	1.1	24		
30-50	0.12	0.25	3.5	4.2	3.8	486		
50-100	0.08	0.39	6.1	2.5	5.0	687		
Median per cluster removal <sup>d</sup> :			16	18	9	1259		

<sup>a</sup>Data are average values, n = 15 per depth band. <sup>b</sup>Data are median values within each hydrological setting and n = 3, 4 and 6 per depth band for horizontal fluxes, hyporheic exchange and upwelling, respectively. <sup>c</sup>Sites A and G are grouped as hot spots of denitrification (*see* text). <sup>d</sup>Median per

cluster removal is the median value of the sum of all depth bands, per piezometer cluster, i.e.

865 median nitrate removal between 0 and 100cm.

Figure 1: Conceptual model of subsurface flow pathways (a), schematic of the study reach showing piezometers used in  ${}^{15}NO_{3}^{-}$  injections (b) and depth profiles of chloride concentrations along the thalweg of the reach (c). Black circles indicate piezometers used in thalweg profiles, white circles are piezometers used for this research but not included in thalweg profiles. Each circle represents a cluster of 3 piezometers (20, 50 and 100cm).



Figure 2: Characterization of subsurface hydrology using chloride concentrations in a mixing model. Mixing score is the output of a two end-member mixing model, modified such that porewater at 100cm and surface water were equal to -1 and 0, respectively. Boxes consist of median values (straight vertical line), the interquartile range (limits of the box), whiskers are the minimum and maximum values and outliers are plotted as crosses (n = 6, 5 and 4 per depth band, for (a) porewater upwelling, (b) hyporheic exchange flows (HEF) and (c) horizontal flows, respectively).



Figure 3 Spatial variation in porewater concentrations of nitrate (a,b), dissolved organic carbon
(c,d), dissolved oxygen (e,f) and denitrification rate (g,h). Thalweg profiles (a,c,e,g) show
individual data points from selected piezometers along the study reach (*see* Figure 1). Boxplots
(b,d,f,h) contrast porewater chemistry and denitrification rate between hydrological settings with
data from all piezometers. Boxes consist of median values (straight horizontal line), the interquartile
range (limits of box), whiskers are the minimum and maximum values and outliers are plotted as
circles.



Figure 4: Proportion of <sup>15</sup>N labelling in the N<sub>2</sub> and N<sub>2</sub>O pools (a) and proportion of <sup>15</sup>N-labelled

- 890 N<sub>2</sub>O in the total nitrogenous gas pool (N<sub>2</sub> + N<sub>2</sub>O, b) produced from injection of  ${}^{15}NO_{3}{}^{-1}$  into the river
- bed (*n*=49 per plot). Boxplots shown in (a) consist of the median value (horizontal line),
- 892 interquartile range (limits of the box), the minimum and maximum values (whiskers) and outliers
- are plotted as circles.

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Figure 5: Timescales of denitrification versus timescales of water transport in a gaining reach (n=72). The solid line shows where the two timescales are equal, i.e. a Damköhler number of 1. Points that plot above the solid line represent sediments where water residence time is more important than denitrification rate in controlling nitrate flux (Da<sub>N</sub> <1). Points that plot above the solid line represent sediments where denitrification rate is the dominant controlling factor of nitrate flux (Da<sub>N</sub> >1).



Figure 6 Variation of denitrification rate and water residence time with depth in the bed of gaining
river, showing an interchange between the importance of factors controlling nitrate removal
between shallow and deep sediments. Data are average values with error bars of one standard
deviation.

