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Groundwater nitrate reduction versus dissolved gas production: A tale of two catchments



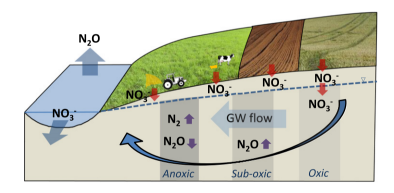
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HIGHLIGHTS

- NO₃ removal capacity was highly variable between and within study catchments.
- Hydrogeological and agronomic factors controlled groundwater hydrogeochemical signatures.
- NO₃ consumption was coupled with excess N₂ and N₂O production.
- Excess N₂ was the dominant denitrification reaction product in near stream groundwater.
- Groundwater N₂O was a net source of greenhouse gas emissions in both catchments.

GRAPHICAL ABSTRACT



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ABSTRACT

At the catchment scale, a complex mosaic of environmental, hydrogeological and physicochemical characteristics combine to regulate the distribution of groundwater and stream nitrate (NO₃⁻). The efficiency of NO₃⁻ removal (via denitrification) versus the ratio of accumulated reaction products, dinitrogen (excess N₂) & nitrous oxide (N2O), remains poorly understood. Groundwater was investigated in two well drained agricultural catchments (10 km²) in Ireland with contrasting subsurface lithologies (sandstone vs. slate) and landuse. Denitrification capacity was assessed by measuring concentration and distribution patterns of nitrogen (N) species, aquifer hydrogeochemistry, stable isotope signatures and aquifer hydraulic properties. A hierarchy of scale whereby physical factors including agronomy, water table elevation and permeability determined the hydrogeochemical signature of the aquifers was observed. This hydrogeochemical signature acted as the dominant control on denitrification reaction progress. High permeability, aerobic conditions and a lack of bacterial energy sources in the slate catchment resulted in low denitrification reaction progress (0–32%), high NO₃ and comparatively low N₂O emission factors $(EF_{5g}1)$. In the sandstone catchment denitrification progress ranged from 4 to 94% and was highly dependent on permeability, water table elevation, dissolved oxygen concentration solid phase bacterial energy sources. Denitrification of NO₃ — to N₂ occurred in anaerobic conditions, while at intermediate dissolved oxygen; N₂O was the dominant reaction product. $EF_{5g}1$ (mean: 0.0018) in the denitrifying sandstone catchment was 32% less than the IPCC default. The denitrification observations across catchments were supported by stable isotope signatures. Stream NO₃ occurrence was 32% lower in the sandstone catchment even though N loading was substantially higher than the slate catchment. © 2016 The Authors. Published by Elsevier B.V. This is an open access article under the CC BY-NC-ND license

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1. Introduction

Anthropogenic application of inorganic and organic nitrogen (N) fertilisers to agricultural landscapes has pervasive consequences including human health implications (e.g. methemoglobinaemia), eutrophication, aquatic acidification, loss of habitat biodiversity and greenhouse gas emissions (Dennis et al., 2012; Gruber and Galloway, 2008; Richards et al., 2015; Weymann et al., 2008). In light of agricultural intensification, the identification of subsurface environments with a natural capacity to attenuate excess N is essential to the development of sustainable management strategies. Globally, denitrification is regarded as the dominant nitrate (NO₃⁻) attenuation mechanism in groundwater (Korom, 1992; Rivett et al., 2008; Seitzinger et al., 2006). Denitrification is a microbially mediated process whereby NO₃ is reduced to dinitrogen (N2) gas. In baseflow dominated catchments, groundwater denitrification has the capacity to mitigate stream water N enrichment by returning N to the long residence time atmospheric pool (Heffernan et al., 2012). Denitrification can represent an environmentally positive nitrate removal process (Schipper and Vojvodić-Vuković, 2001); however such a characterisation is subject to an important caveat. The reaction is sequential and as such there are several intermediary products including nitrite (NO₂), nitric oxide (NO) and nitrous oxide (N₂O). The differentiation between which reaction product is dominant is of key environmental concern: N₂ gas is environmentally benign whereas N₂O is a potent greenhouse gas, while NO contributes to stratospheric ozone depletion, eutrophication and formation and accumulation of surface ozone (Vitousek et al., 1997).

In groundwater, a number of geochemical criteria must be met for denitrification to occur. Studies documenting the relationship between NO₃ concentrations and aquifer physicochemistry are prevalent in the available literature (Brettar et al., 2002; Jahangir et al., 2012a; Rissmann, 2011). A commonality throughout indicates that the presence of denitrifying bacteria, reducing conditions and the availability of bacterial energy sources create zones of enhanced denitrification potential. Traditionally, it has been believed that that groundwater denitrification is predominantly heterotrophic with rates related to the amount of dissolved organic carbon (DOC) coupled with the wide abundance of denitrifiers in the groundwater (Barrett et al., 2013; Rivett et al., 2008). Recent research however suggests that autotrophic denitrification i.e. oxidation of solid phases within an aquifer such as Iron (Fe) and Manganese (Mn) may in fact drive bacterial NO₃ reduction (Green et al., 2008; Heffernan et al., 2012). Great uncertainty surrounds the spatial and temporal distribution of denitrifying zones, owing to a confounding hierarchy of scale. In essence, an aquifer can be visualised as an environmental ecosystem, which is capable of removing between 0 and 100% of reactive N. The geological history of the aquifer (mineralogy, stratigraphy and weathering) at the catchment scale controls the distribution and availability of bacterial energy sources, aquifer flow paths, permeability and connectivity at the sub metre scale (Seitzinger et al., 2006). These physical factors in turn determine the hydrogeochemical signature and N attenuating capacity of the aquifer, while agronomy, soil type, hillslope geometry and meteorology control the temporal N load passing through the ecosystem. In complex geological environments, an entire aquifer or catchment cannot be characterised as having high or low denitrification potential. Denitrification is enhanced in certain spatial zones or hot spots (Jahangir et al., 2012a, 2013) and it is the location and intensity of these hot spots in relation to a receptor e.g. a stream, which is paramount to characterising the potential for natural attenuation of N in an aquifer. Several studies have measured denitrification based upon NO₃ loss (Jahangir et al., 2012a; Thayalakumaran et al., 2008; Tsushima et al., 2002), however NO₃ gradients can result from temporal patterns of source contribution (Seitzinger et al., 2006) and other NO₃ removal pathways such as plant and microbial assimilation, dissimilatory NO₃ reduction to ammonium (NH₄⁺) and anaerobic oxidation of NH₄⁺ (Jahangir et al.,

2016). Studies based solely on NO_3^- dynamics have a capacity to overestimate the contribution of denitrification on NO_3^- removal rates (Green et al., 2008). Directly measured denitrification rates based upon the natural accumulation of denitrification products ($N_2O \otimes N_2$) in groundwater are rare with calculated values spanning orders of magnitude across studies (Green et al., 2008; Heffernan et al., 2012; Jahangir et al., 2013; Weymann et al., 2008). $^{15}N_{NO3}$ isotopic signatures have been used extensively to calculate N sources and processes (Kendall et al., 2007). A dual isotopic approach ($^{15}N_{NO3}$ and $^{18}O_{NO3}$) can be used to infer both the source of NO_3^- to groundwater and also transformational processes such as denitrification (Wassenaar, 1995). Although it is not possible to directly calculate denitrification rates from isotopic signatures, coupled enrichment of $^{15}N_{NO3}$ and $^{18}O_{NO3}$ provides a powerful tool to identify areas of enhanced denitrification (Baily et al., 2011).

Contemporaneous measurements of both N₂O and N₂ in groundwater not only provide evidence of NO₃ removal pathways but also offer an insight into the concept of pollutant swapping of NO₃ for N₂O. Globally it is estimated that agricultural practises are responsible for in excess of 60% of anthropogenic N₂O emissions (Harty et al., 2016). The International Panel on Climate Change (IPCC) subdivides agricultural N₂O into three categories: direct emissions from agricultural land, emissions from animal management strategies and indirect emissions of N₂O that is either volatilised, leached or removed in biomass (IPCC, 1997). Each subcategory is estimated to contribute one third of the total agricultural N₂O source with indirect emission estimations contributing two thirds of the uncertainty (Penman, 2000). There exists a substantial body of research into the contribution N₂O to the global greenhouse gas budget via direct pathways i.e. from soil to the atmosphere (Bouwman, 1990; de Klein et al., 2001; Li et al., 2011, 2013; Soussana et al., 2007). Large uncertainties remain around the contribution of indirect N₂O emission pathways, namely from groundwater and surface drainage, rivers and coastal marine areas (Vilain et al., 2012). Fuelling this uncertainty is a lack of process based understanding regarding the production, consumption and movement of groundwater and stream N2O across a range of hydrogeological settings (Clough et al., 2007; Höll et al., 2005; Jahangir et al., 2013). In 1997, the IPCC published an emission factor of 0.015 for the fraction of agriculturally derived N₂O released from groundwater sources (Mosier et al., 1999). In 2006, the IPCC default value was amended to 0.0025 (de Klein et al., 2006) based upon the combined reviews of Hiscock et al. (2003), Reav et al. (2005) and Sawamoto et al. (2005). While this downward revision indicated that groundwater derived N₂O was less significant than previously proposed, the published range of uncertainty (0.0005-0.025) highlighted the ambiguity surrounding the natural variability of N₂O in groundwater while reinforcing the need for further research to constrain emission factors and reduce uncertainty. Studies combining a complete analysis of N species: organic N, ammonium (NH_4^+-N) , NO_3^--N , nitrite (NO_2^--N) , $N_2O-N \& N_2-N$, with aerobicity (dissolved oxygen and redox potential), electron donors, dual isotopic techniques and aquifer hydraulic properties are rare. It is only through the refinement of scale, from catchment to sub metre, that a process based understanding of groundwater N removal can be developed.

The objectives of this study are (1) to quantify the capacity of hillslope hydrologic systems to naturally attenuate agriculturally derived NO_3^- , 2) to elucidate the extent of denitrification by measuring the accumulation and ratio of reaction products (N_2 and N_2O) and (3) to identify the physical and biogeochemical factors affecting groundwater denitrification rates and indirect N_2O emissions from groundwater.

2. Materials & methods

2.1. Study sites

This research was undertaken along four hillslopes of varying length, geometry and landuse in two agricultural catchments in the Republic of

Ireland (Fig. 1). The catchments, termed herein as sandstone and slate, were chosen to represent the dominant Irish land use categories in landscapes susceptible to potential phosphorous and nitrogen transfer risk (Fealy et al., 2010). The sandstone catchment is characterised by intensive dairy grassland agriculture, while continuous arable crop production dominates the slate catchment. Within both study catchments, two instrumented hillslopes termed Sandstone N/Sandstone S and Slate N/Slate S were targeted to represent general catchment morphology and land use with each exhibiting a range of hydrogeochemical conditions. Each hillslope intersects with a stream channel at its base, representing the receptor or end point for up gradient hydrogeochemical dynamics. A conceptual geological understanding was developed using a suite of geophysical techniques including ground penetrating radar, electromagnetic terrain conductivity, 2D resistivity and seismic refraction (Mellander et al., 2014).

2.2. Monitoring wells

Multilevel monitoring wells were installed in the near stream, midslope and upslope zones of each hillslope (Mellander et al., 2014). In total 12 multilevel wells were installed, each generally comprising three piezometers, yielding a total of 36 sampling intervals. The piezometers (inner diameter: 52 mm) had screened sections ranging in length between one and four metres (typically three metres). The depth to the bottom of each well screened interval ranged in depth from 3.5 to 40 m below ground level (m BGL). The wells screens were targeted to intercept shallow (1–10 m BGL) and deeper bedrock groundwater pathways (10–40 m BGL). Saturated hydraulic conductivity (Ksat) was measured within each screened interval using the slug testing procedure outlined in Butler (1997). Ksat was calculated from the generated timedrawdown data using the Bower and Rice (1976)\ method for unconfined aquifers. Fig. 1 illustrates the experimental design and location

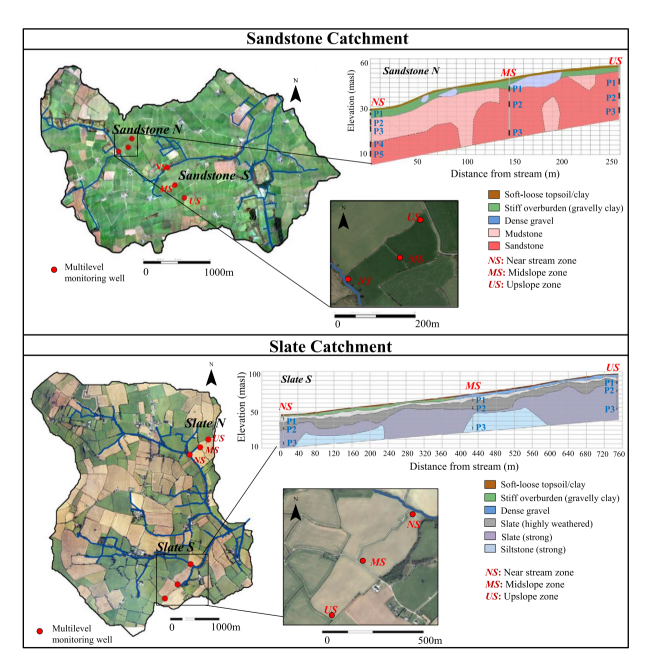


Fig. 1. Location of experimental catchments, stream network, field boundaries and instrumented hillslopes. Also illustrated are geological cross sections of the Sandstone N and Slate S hillslopes showing the location of the multilevel monitoring wells. Note P1, P2 etc. refers to piezometer screened intervals.

of multilevel monitoring wells at the *Sandstone N* and *Slate S* hillslopes. The length and depth of each piezometer screened intervals and hillslope hydraulic properties are presented in Table 1.

2.3. Hillslope characterisation

At the sandstone hillslopes, Quaternary deposits are dominated by free draining loam to clay loam underlain by layers of dense gravel and gravelly clay with weathered mudstone. Bedrock consists of a complex 3 dimensional pattern of mudstone and sandstone with minor siltstone and exhibits varying degrees of weathering (Fig. 1, Supplementary Fig. 1b). The aquifer is characterised as unconfined (GSI, 2016) and exhibits a high baseflow index (BFI) of 73% (mean: 2012–13). BFI describes the proportion of flow in a stream which is supplemented by groundwater discharge and was calculated after Mellander et al. (2012). In situ measurements of Ksat at both hillslopes indicated a layered distribution of permeability (Table 1), becoming less permeable with depth.

The slate hillslopes are characterised by clay to clay loam underlain by dense gravel and firm gravelly clay layers. An extensive zone of highly weathered to moderately weathered slate bedrock is apparent at both hillslope sites. In *Slate N*, the weathered zone ranges in thickness from approximately 4 to 11 m BGL, increasing in magnitude towards the bottom of the hillslope (Supplementary Fig. 2a). In Slate S, the weathered zone is more extensive, ranging in thickness from 5 to 18 m BGL, with greatest magnitudes in midslope and near stream zones (Fig. 1). Underlying the weathered zone is competent slate and siltstone bedrock, showing evidence of localised fracturing. The aquifer in the slate catchment is unconfined (GSI, 2016) with a BFI of 77% (mean: 2013-15), indicating that up-gradient groundwater hydrogeochemistry has a significant effect on stream water quality. The distribution of Ksat revealed that Slate N and Slate S are highly permeable (Table 1), owing to the density and vertical extent of bedrock fracturing. A comprehensive description of hillslope geology and aquifer properties is provided in Supplementary Section 1.

2.4. Water sampling

Groundwater sampling was carried out on a monthly basis from July 2013 until May 2015. Samples were collected from the centre of each well screen using a 200 mL double valve bailer (Solinst, Canada). Stream and drain samples were collected simultaneously with groundwater samples. Samples for nitrogen components (NO₃⁻, NO₂⁻, NH₄⁺),Cl⁻, reduced metals (Fe²⁺ & Mn²⁺) and dissolved organic carbon (DOC) were filtered in-situ through a 0.45 µm hydrophilic membrane filter into polypropylene sample tubes (50 mL). Groundwater temperature, pH, electrical conductivity, redox potential (Eh) and dissolved oxygen (DO) were simultaneously measured using an in-situ multi-parameter (Aquameter) probe. Seasonal groundwater dissolved gas sampling (N2, N2O and DO) was undertaken to complement the monthly hydrogeochemical dataset. Samples for N₂, Argon (Ar) and DO were collected by overflowing groundwater into 12 mL exetainers (Labco Ltd., Wycombe, UK), while groundwater N₂O samples were overflowed into 160 mL glass serum bottles. All gaseous samples were immediately sealed following collection and stored under water (at 4 °C) prior to analysis.

2.5. Hydrochemistry

Total oxidised nitrogen ($NO_3^- \& NO_2^-$), NO_2^- , Cl^- , NH_4^+ were analysed by Aquakem 600 Discrete Analyser (Aquakem 600A, 01621 Vantaa, Finland) following the hydrazine reduction, sulphanilamide diazotisation and dichloroisocyanurate hydrolysis methods respectively. Groundwater NO_3^- was calculated by subtracting NO_2^- from total oxidised nitrogen. DOC and total N were analysed by catalytic combustion using a Shimadzu TOC-L analyser (TOC-V cph/cpn: Shimadzu Corporation, Kyoto, Japan). Reduced metals ($Fe^{2+} \& Mn^{2+}$) were analysed by inductively coupled plasma emission spectrometry (Agilent 5100 + Agilent Technologies, 5301 Stevens Creek Blvd, Santa Clara, CA 95051, United States). Total N_2 , DO and Argon (Ar) were analysed using membrane inlet mass spectrometry (MIMS), according to the average

Table 1Length and depth of piezometer screened intervals, hillslope permeability (Ksat) and hillslope piezometric levels.

Sandstone catch	ment						Slate catchment							
*Hillslope Zone			*Well screen		*Ksat		*Hillslope Zone		metric	*Well screen		*Ksat		
	water level (m BGL)		Subsoil/ Bedrock weathered rock		Subsoil/ weathered rock	Bedrock		water level (m BGL)		Subsoil/ weathered rock	Bedrock	Subsoil/ weathered rock	Bedrock	
Sandstone N (NS)	Min: Max:	0.0 0.7	P1: 2-3.5 P2: 5.5-8.5	P3: 9-12 P4: 16.5-19.5 P5: 22-25	P1: 2.8 P2: 0.8	P3: 0.6 P4: 0.7 P5: 0.3	Slate N (NS)	Min: Max:		P1: 1-4 P2: 9-13	P3: 32-52	P1: 1.2 P2: 0.5	P3: 0.08	
Sandstone N (MS)	Min: Max:	0.1 4.0	P1:4-7	P2: 12-15 P3: 27-30	P1: 2.5	P2: 0.4 P3: 0.3	Slate N (MS)	Min: Max:	0.0 3.9	P1: 1-4	P2: 25-28 P3: 37-40	P1: 2.2	P2: 1.3 P3: 0.15	
Sandstone N (US)	Min: Max:	1.1 7.0		P1: 7-10 P2: 14.5-17.5 P3: 22-25		P1: 1.0 P2: 1.9 P3: 0.09	Slate N (US)	Min: Max	3.0 11.4		P1: 12-15 P2: 27-30		P1: 1.03 P2: 1.4	
Sandstone S (NS)	Min: Max:	0.1 0.5	P1: 4-7	P2: 10-13 P3: 17-20	P1: 0.53	P2: 0.99 P3: 0.35	Slate S (NS)	Min: Max:	0.1 1.0	P1: 3.5-6.5	P2: 14-17 P3: 30.5-33.5	P1: 4.2	P2: 5.2 P3: 0.4	
Sandstone S (MS)	Min: Max:	1.2 12.7		P1: 10.5-13.5 P2: 17-20 P3: 27-30		P1: 2.8 P2: 0.98 P3: 0.54	Slate S (MS)	Min: Max:	3.0 9.8		P1: 10.5-13.5 P2: 17-20 P3: 27-30		P1: 4.2 P2: 4.0 P3: 0.9	
Sandstone S (US)	Min: Max:	5.6 14.0		P1: 13-16 P2: 27-30		P1: 0.40 P2: 0.403	Slate S (US)	Min: Max:	6.3 10.6	P1: 7-11	P2: 16-19 P3: 37-40	P1: n.d	P2: 1.0 P3: 5.3	

^{*}Hillslope zone: NS = near stream, MS = midslope, US = upslope.

^{*}Well screen: metres below ground level of the top and bottom of each screened interval.

^{*}Ksat: Hydraulic conductivity (m/day) calculated after Bower and Rice (1976).

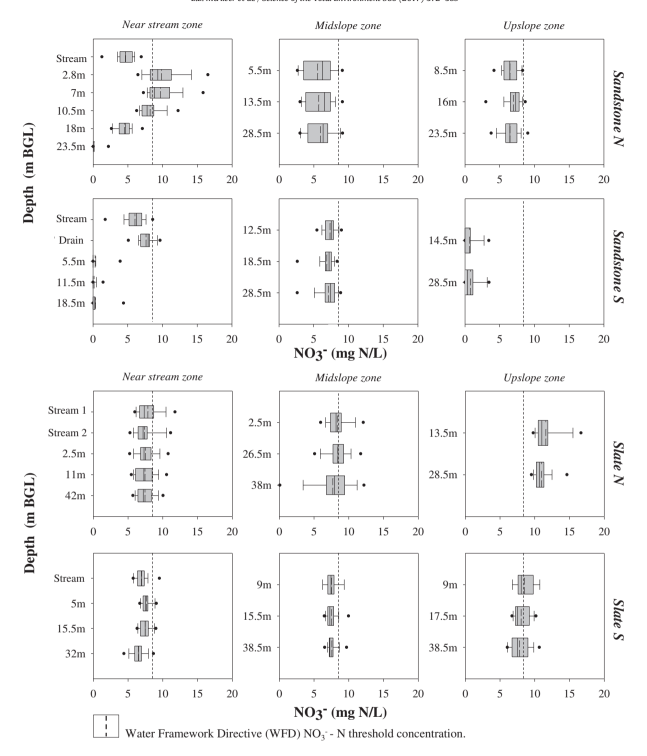


Fig. 2. Boxplots of groundwater and stream NO₃ from 2013 to 2015 in the sandstone and slate catchments, arranged by hillslope, slope position, and piezometer sampling depth.

recharge temperature during sampling (Kana et al., 1994). Denitrified N_2 , termed excess N_2 hereafter, was calculated after Weymann et al. (2008) using the ratio of N_2 /Ar in groundwater and temperature based solubility constants. Preparation of groundwater N_2 O samples was carried out using the headspace extraction technique according to the methodology outlined in Jahangir et al. (2012b). The N_2 O concentration of the extracted headspace gas was analysed using electron capture gas chromatography (CP-3800, Varian, Inc. USA). Frozen samples for natural abundance stable isotopes were packed in dry ice and shipped to the UC Davis Stable Isotope Facility, University of California, USA for

analysis. Samples for $\delta^{15}N_{NO3}$, $\delta^{18}O_{NO3}$ and $\delta^{15}N_{N2O}$ were analysed using the bacterial denitrifier method (Sigman et al., 2001).

2.6. Total initial nitrogen (N_{INI}), denitrification reaction progress (RP), groundwater N_2O emission factors ($EF_{Sg}1$ & $EF_{Sg}2$) and FracLEACH

It is assumed that denitrification along a groundwater flow path results in the reduction of NO_3^- and the production of dissolved N gases. As such, the initial nitrogen concentration ($N_{\rm INI}$) is calculated as the sum of residual N substrates and accumulated N products (Jahangir

et al., 2013) where:

$$(N_{INI}) = NO_3^- - N + NH_4^+ - N + NO_2^- - N$$

+organic N + excess N₂ - N + N₂O-N (1)

Reaction progress (RP) is the ratio between products of a given process and starting substrates (Weymann et al., 2008). RP was used to estimate groundwater denitrification using the following relationship:

$$(RP) = (excess N_2 - N + N_2 O - N)/N_{INI}$$

$$(2)$$

Indirect emissions of N_2O from groundwater were quantified using two methods. The first, termed $EF_{5g}1$ was adopted from Weymann et al. (2008) and takes into account transformations of NO_3^- and N_2O within the aquifer (Eq. (3)). The second $EF_{5g}2$ has been adopted by the IPCC (2006), typically in studies lacking gaseous N data, and assumes no further transformation of NO_3^- and N_2O that has been leached to groundwater (Eq. (4)).

$$EF_{5g}1 = N_2O - N/N_{INI} - N (3)$$

$$EF_{5g}2 = N_2O - N/NO_3^- - N (4)$$

FracLEACH describes the quantity of dissolved N leached from soil to groundwater as a proportion of the total N load applied via agricultural practises. FracLEACH was quantified using the following relationship:

FracLEACH (%)=
$$Q_{GW}(m^3/ha/yr.)*NO_3^--N_s(kg N/m^3)$$
 (5)
/Total N input (kg N/ha/yr.)

Q_{GW} represents groundwater discharge and was calculated as Q_{EF} (m³/Ha) * baseflow index (BFI). Q_{FF} refers to total recharge, measured as effective rainfall (m/yr.) multiplied by field area (ha). Effective rainfall was calculated by subtracting potential evapotranspiration (calculated according to Penman-Monteith) from total rainfall. Total recharge multiplied by the BFI was assumed to equal groundwater discharge. Groundwater discharge was multiplied by NO₃ -N in the shallow groundwater of each hillslope (NO₃-N_s) providing a total load of NO₃ –N per year. Wells exhibiting aerobic conditions were chosen for the FracLEACH calculation to negate the confounding effect of attenuation on leaching rates. Dividing the total load of NO₃ -N_s by the total load of N applied provided a measurement of FracLEACH. The total load of organic and inorganic N applied was calculated from detailed farmer records specific to each hillslope field (2012 – 2013). Hillslope landuse, applied N, values of FracLEACH and the total loads of NO₃ -N_s for each study hillslope are provided in Table 2.

2.7. Statistical analysis

In order to identify statistically significant relationships (p < 0.05) between parameters, a correlation analysis between influential hydrological, hydrogeological and hydrogeochemical variables and NO $_3^-$,

N₂O and excess N₂ was undertaken. The Pearson product moment approach was utilised when data were approximately normally distributed, with appropriate transformation and rescaling of the data where non-normality was apparent. Where it was not possible to transform data to normality, the Spearman rank approach was used. A repeated measures mixed ANOVA analysis (SAS, 2009), including covariance modelling for effects across sampling dates, was used to identify the significant effects of spatial parameters, including sampling depth and hillslope zone on groundwater hydrogeochemistry. The effect of groundwater sampling depth and groundwater residence times on groundwater hydrogeochemistry has been examined in several studies (Jahangir et al., 2012a; Fenton et al., 2009; Weymann et al., 2008). Typically aquifer permeability decreases with increasing depth (Jahangir et al., 2012a), while groundwater residence times increase (Fenton et al., 2011). The use of depth as a grouping factor provides insight into these hypotheses. Each hillslope was split into three zones: near stream, midslope and upslope zone. The depth of the unsaturated zone has been identified as a dominant controlling factor on groundwater nitrate concentrations (Spalding and Exner, 1993). In the near stream zone, the water table is close to the surface. In the midslope and upslope zones, the depth to the water table and depth of the unsaturated zone increases, with greatest thickness in the upslope zone. The statistical model also accounted for between catchment variability (sandstone vs. slate), and within catchment variability e.g. Sandstone N vs. Sandstone S. Given the extensive range of sampling depths across catchments, depth was grouped into six categories namely 0-5, 5-10, $10-15\ 15-20$, 20-30 and >30 m BGL. The interactions between categorical variables were also scrutinised within the model, while a validation of the model assumptions was achieved using a residual analysis.

3. Results

3.1. NO_3^- , NH_4^+ and NO_2^- concentrations

The spatial distribution of groundwater and stream NO_3^- in the sand-stone and slate catchments, organised by hillslope, hillslope zone and sample depth are illustrated in Fig. 2. Mean groundwater NO_3^- concentration was lower (p < 0.001) in the sandstone catchment (4.9 mg N/L) compared to the slate catchment (8.1 mg N/L). This pattern was echoed in stream concentrations with NO_3^- in the slate catchment streams (mean: 7.3 mg N/L) exceeding the sandstone streams (mean: 5.4 mg N/L) by 32% (p < 0.001).

The coefficient of variation (CV) was used as a measure of spatial variability specific to each experimental hillslope, calculated using the overall hillslope mean, and standard deviation from the mean at each spatial sampling interval. A high CV value indicates spatial heterogeneity of a given parameter associated with different sampling depths and hillslope zones i.e. upslope vs. midslope vs near stream. A low CV indicates spatial uniformity. The spatial distribution of groundwater NO_3^- in the sandstone catchment was complex (CV *Sandstone N*: 42% vs. *Sandstone S*: 117%). Significant relationships were identified between hillslopes (p < 0.0001) and sample depths (p < 0.05), with significant

Table 2Hillslope landuse, surface applied N, N leached to groundwater and FracLEACH.

Sandstone cat	chment				Slate catchment							
Hillslope Landuse		*Applied N (kg N/ha/yr.)	*NO ₃ ⁻ -N _s FracLEACH (kg N/ha/yr.) (%)		Hillslope	Landuse	*Applied N	*NO ₃ -N _s (kg N/ha/yr.)	FracLEACH (%)			
Sandstone N	Grassland, dairy production	Inorganic: 295.3 Organic: 0	39.1	14	Slate N	Arable, spring barley production.	Inorganic: 122.0 Organic: 21.0	69.0	41			
Sandstone S	Grassland, dairy production	Inorganic: 301.5 Organic: 78.6	72.5	21	Slate S	Arable, spring barley production.	Inorganic: 154.8 Organic: 0	62.4	44			

^{*}Applied N: Mean applications of inorganic and organic N (2012–2013).

^{*}NO₃_-Ns: Mean load of NO₃_ leached to shallow groundwater (2012–2013).

interactions between hillslope and hillslope zone (p < 0.001), hillslope and depth (p < 0.05) and hillslope zone and depth (p < 0.0001). An analysis of the slope of the interaction terms indicated that depth had a greater effect on NO $_3^-$ concentration in *Sandstone N* vs. *Sandstone S*, while in both hillslopes, a combination of near stream hillslope zone

and depth had the greatest baring on groundwater NO_3^- concentrations. In *Sandstone N*, a negative correlation was observed between depth and NO_3^- concentration. In the near stream zone, there was a clear concentration gradient in NO_3^- moving deeper down the depth profile (Fig. 3a). The gradient in NO_3^- concentration mirrored the distribution

Sandstone N

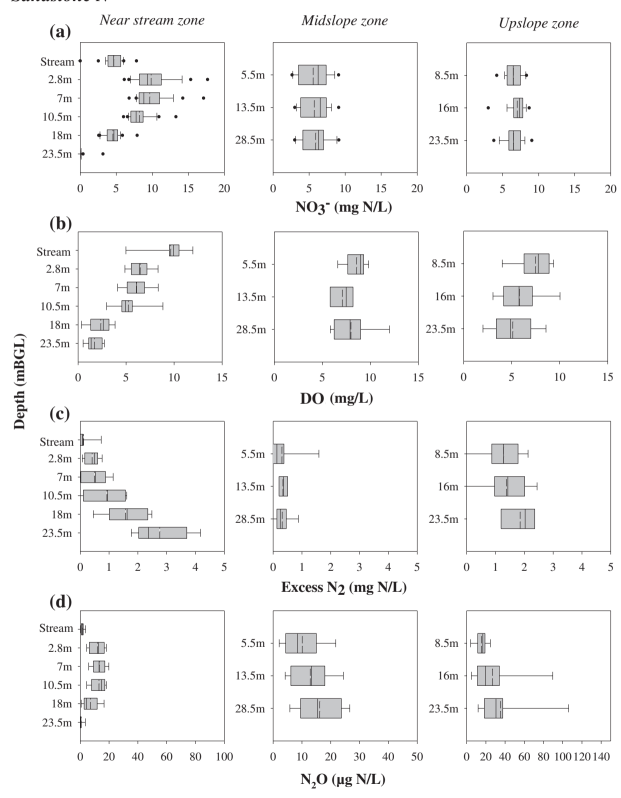


Fig. 3. Boxplots of groundwater and stream NO₃₋, DO, excess N₂ and N₂O from 2013 to 15 in Sandstone N, arranged by slope position and piezometer sampling depth.

of groundwater DO (Fig. 3b), NO₃ concentrations in the near stream zone were significantly lower than the midslope (p < 0.001) and upslope (p < 0.005) zones. Despite elevated NO₃ concentrations in the near stream shallow groundwater of Sandstone N (mean 9.8 mg N/L between 2 and 8.5 m BGL), stream concentrations were significantly lower (mean 4.6 mg N/L), most closely reflecting deeper groundwater. In Sandstone S, groundwater NO₃ in the near stream zone was lower than the midslope zone (p < 0.0001), showing no significant difference with the upslope zone. At all sample depths in the near stream and upslope zones, groundwater NO₃ was either below or marginally exceeded the limit of detection (0.25 mg N/L) (Fig. 2). Fig. 4a and b illustrate the relationship between NO₃ and hydraulic conductivity (Ksat) and NO₃ and the water table ratio. The water level ratio refers the ratio between the depth of the unsaturated zone and the total depth of sampling. The nearer the ratio is to 1, the closer the water table is to the ground surface. The ratio also takes the depth of the saturated water column into account. Therefore the greater the depth of the overlying water column, the closer the ratio is to 1. Water level ratios are highest where a deep well has a shallow water table. In the sandstone catchment, groundwater NO₃ increased significantly with increasing Ksat values (r = 0.54, p < 0.05) and decreased significantly with increasing water table ratios (r = 0.58, p < 0.05). Typically, Ksat decreased with increasing depth BGL corresponding to less weathered bedrock, while the water level ratio decreased moving upslope corresponding to increases in the depth of the unsaturated zone. Stream NO₃ in Sandstone S was significantly higher (p < 0.05) than the Sandstone N. The mean stream concentration of 6.0 mg N/L closely reflected midslope groundwater and water sampled from an upstream tile drain.

In the slate catchment, there was a significant interaction between hillslope and hillslope zone (p < 0.0001). As such, while there was a clear effect of hillslope zone on NO $_3^-$ concentration, the effect was specific to each hillslope. Slate N exhibited higher groundwater NO $_3^-$ than Slate S with mean concentrations of 8.7 and 7.6 mg N/L respectively. Slate S was characterised by spatial uniformity throughout the hillslope (CV: 6.9%) with no significant differences identified as a function of hillslope zone or sampling depth (Fig. 2). In Slate N, there was greater spatial variation than Slate S (CV: 18.9%) with a hillslope zone effect (p < 0.0001). Highest NO $_3^-$ concentrations occurred in the upslope

zone, with lower concentrations in the midslope and near stream zones. In the slate catchment, no significant relationships were identified between NO_3^- and Ksat or NO_3^- and water table ratio (Fig. 4). In both slate hillslopes, stream NO_3^- concentrations were similar to shallow groundwater.

Groundwater NH $_4^+$ concentrations where marginally lower in sand-stone catchment compared to the slate catchment (mean sandstone: 0.03 mg N/L, mean slate: 0.04 mg N/L). In the sandstone catchment there were no significant differences between hillslopes means, with limited spatial variability (CV: 13%). Similarly, in the slate catchment, no significant relationships were identified between hillslopes with limited spatial variation across hillslope zones and sample depths (CV: 25%). Groundwater NO $_2^-$ concentrations were typically either at or below the limits of detection (0.006 mg N/L) with mean concentrations of 0.002 and 0.007 mg N/L in the slate and sandstone catchments.

3.2. Groundwater hydrochemistry

3.2.1. Aquifer aerobicity

Groundwater DO was lower (ANOVA; p < 0.001) in the sandstone catchment compared to the slate catchment (mean sandstone: 5.5 vs. mean slate: 8.9 mg/L). In the sandstone catchment, mean DO concentrations were 5.8 and 4.1 mg/L in Sandstone N and Sandstone S. respectively. Spatial variation was high in both hillslopes (CV Sandstone N: 38% vs. CV Sandstone S: 54%) and there was a significant effect of sampling depth (p < 0.01) and hillslope zone (p < 0.0001) on groundwater DO. In both hillslopes, near stream zones had the lowest DO (p < 0.0001) with the exception of the upslope monitoring well in Sandstone S, which showed comparably low concentrations (Table 3). DO typically decreased with increasing depth, while a distinct geochemical gradient was apparent in the near stream zone of Sandstone N (Table 3). The sandstone catchment was characterised by anaerobic conditions at a number of spatial sampling points. Screened intervals with DO <3 mg/L included the near stream zones of Sandstone N (18.5 and 23.5 m BGL) and Sandstone S (5.5, 11.5 and 18.5 m BGL) and the upslope zone of Sandstone S (14.5 m BGL). Fig. 4c and d illustrate the relationship between DO and hydraulic conductivity (Ksat) and DO and the water table ratio (as described in Table 1). In the sandstone catchment, groundwater DO

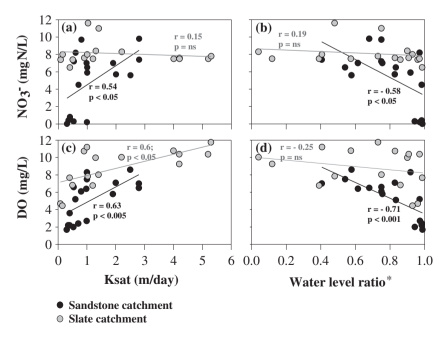


Fig. 4. a & b Linear regression analysis of hydraulic conductivity (Ksat) and water level ratio vs. NO3 — in the sandstone and slate catchments. (Fig. 4c & 4d) Regression analysis of hydraulic conductivity (Ksat) and water level ratio vs. DO in the sandstone and slate catchments. P < 0.05 = significant correlation; ns = not significant. Black regression line = sandstone catchment. Grey regression line = slate. Note: water level ratio* refers to the depth of the water table BGL, relative to the total depth of the well screen BGL.

Table 3
Mean groundwater and stream NO_3^- (\pm standard error SE) vs. aerobicity, bacterial energy sources, denitrification reaction products and N_2O emissions arranged by hillslope, hillslope positions and piezometric sampling depth in the sandstone catchment.

Hillslope zone*/ Depth* (m BGL)	Reactive N			Aerobicity		Bacterial energy sources			Denitrification rate & products			N ₂ O emissions	
	NO ₃ ⁻ (mg N/L)	N _{INI} * (mg N/L)	NO ₃ -/Cl- (ratio)	DO (mg/L)	Eh (mV)	DOC (mg/L)	Fe ²⁺ (μg/L)	Mn ²⁺ (μg/L)	RP*	excess N ₂ (mg N/L)	N ₂ O (μg N/L)	N ₂ O−EF _{5g} 1* Above IPCC 2006: ✓, below: ४	
						Sar	ndstone N						
Stream	4.7 (0.3)	5.5 (0.4)	0.16 (0.01)	9.6 (0.7)	94.9 (11.8)	6.3 (1.4)	18.0 (3.3)	11.6 (2.9)	0.02 (0.02)	0.1 (0.1)	1.5 (0.4)	0.0003 (0.0001) x	
NS (2.8 m)	9.8 (0.5)	12.5 (1.0)	0.35 (0.01)	6.5 (0.4)	65.6 (7.3)	2.0 (0.2)	10.5 (6.8)	6.2 (1.3)	0.04 (0.01)	0.4(0.1)	12.1 (1.8)	0.0010 (0.0001) x	
NS (7 m)	9.7 (0.4)	12.1 (0.9)	0.35 (0.01)	6.1 (0.5)	62.9 (11.3)	2.0 (0.3)	5.7 (2.1)	6.6 (1.7)	0.04 (0.01)	0.5 (0.2)	13.3 (1.6)	0.0011 (0.0002) x	
NS (10.5 m)	8.2 (0.3)	10.1 (0.8)	0.30 (0.01)	5.2 (0.7)	39.5 (12.2)	1.9 (0.4)	4.7 (2.1)	11.6 (2.6)	0.09 (0.02)	0.9 (0.2)	13.0 (1.8)	0.0013 (0.0002) x	
NS (18 m)	4.5 (0.2)	6.5 (0.4	0.17 (0.01)	2.4 (0.8)	-21.2(16.1)	1.9 (0.3)	3.4 (1.1)	54.6 (14.9)	0.25 (0.03)	1.6 (0.2)	7.1 (1.9)	0.0010 (0.0002) x	
NS (23.5 m)	0.0 (0.02)	3.0 (0.3)	0.01 (0.00)	1.7 (0.6)	-73.7(30.0)	2.5 (0.9)	79.3 (28.0)	633.2 (64.2)	0.95 (0.01	2.9 (0.3)	0.7 (0.4)	0.0002 (0.0001) x	
MS (5.5 m)	5.6 (0.4)	7.4 (0.6	0.23 (0.01)	8.6 (0.4)	107.4 (7.8)	2.6 (0.3)	3.2 (1.3)	4.0 (0.9)	0.05 (0.03)	0.3 (0.2)	10.1 (2.2)	0.0014 (0.0003) x	
MS (13.5 m)	5.7 (0.4)	7.6 (0.6)	0.24 (0.01)	7.1 (0.5)	102.0 (8.9)	2.9 (0.3)	4.1 (1.9)	12.5 (9.4)	0.05 (0.01)	0.3 (0.1)	12.9 (2.3)	0.0017 (0.0003) x	
MS (28.5 m)	5.9 (0.4)	7.6 (0.7)	0.24 (0.01)	8.3 (0.9)	120.1 (15.4)	2.7 (0.3)	15.7 (12.2)	6.1 (1.3)	0.04 (0.01)	0.3 (0.1)	16.2 (2.5)	0.0022 (0.0003) x	
US (8.5 m)	6.5 (0.2)	8.3 (0.5)	0.28 (0.01)	7.5 (0.6)	120.7 (9.6)	2.7 (0.6)	6.8 (3.8)	3.1 (0.5)	0.16 (0.03)	1.3 (0.2)	15.5 (2.0)	0.0019 (0.0003) x	
US (16 m)	7.0 (0.3)	8.9 (0.5)	0.27 (0.01)	5.8 (0.9)	117.2 (9.6)	2.6 (0.5)	3.4 (1.5)	14.0 (6.8)	0.18 (0.02)	1.6 (0.2)	26.8 (8.6)	0.0038 (0.0016) 🗸	
US (23.5)	6.5 (0.3)	8.6 (0.4)	0.26 (0.01)	5.1 (2.2)	123.4 (12.1)	2.4 (0.5)	18.9 (14.6)	16.2 (11.4)	0.23 (0.03)	1.9 (0.2)	35.0 (9.4)	0.0047 (0.0015) 🗸	
						Sai	ndstone S						
Stream	6.0 (0.3)	7.1 (0.4)	0.19 (0.01)	8.7 (0.7)	99.5 (11.4)	6.6 (1.4)	22.5 (5.1)	15.1 (2.9)	0.03 (0.02)	0.2 (0.1)	5.4 (1.3)	0.0008 (0.0002) x	
NS (5.5 m)	0.3 (0.2)	2.6 (0.5)	0.02 (0.008)	2.0 (1.4)	18.4 (19.8)	2.1 (0.8)	2910.3 (484.7)	2997.3 (210.1)	0.88 (0.05)	2.4 (0.5)	1.1 (0.8)	0.0004 (0.0002) x	
NS (11.5 m)	0.2 (0.1)	2.4 (0.4)	0.01 (0.002)	2.7 (1.0)	20.1 (11.7)	1.9 (0.6)	2111.6 (422.6)	2407.0 (295.6)	0.89 (0.03)	2.2 (0.4)	1.0 (0.4)	0.0005 (0.0002) x	
NS (18.5 m)	0.4 (0.3)	2.7 (0.5)	0.02 (0.01)	2.2 (1.2)	20.9 (20.1)	1.9 (0.6)	2560.1 (394.6)	1860.9 (135.1)	0.89 (0.02)	2.4 (0.4)	1.4 (0.9)	0.0016 (0.0014) x	
MS (12.5 m)	7.4 (0.2)	9.7 (0.7)	0.27 (0.007)	7.0 (0.5)	58.9 (6.7)	3.5 (1.7)	6.7 (2.1)	11.2 (4.1)	0.16 (0.03)	1.7 (0.5)	21.6 (4.4)	0.0023 (0.0004) x	
MS (18.5 m)	7.0 (0.3)	9.4 (0.9)	0.24 (0.01)	6.4 (0.5)	52.5 (21.9)	2.6 (1.2)	8.4 (3.0)	14.7 (3.8)	0.18 (0.03)	1.8 (0.5)	20.3 (4.2)	0.0021 (0.0005) 🗶	
MS (28.5 m)	7.2 (0.3)	9.2 (1.2)	0.26 (0.01)	6.6 (0.9)	56.3 (29.2)	2.6 (0.9)	61.9 (26.0)	89.5 (31.1)	0.17 (0.04)	1.6 (0.4)	22.8 (5.0)	0.0025 (0.0005) ✓	
US (14.5 m)	0.6 (0.2)	4.4 (1.1)	0.03 (0.008)	2.2 (1.2)	23.7 (11.5)	2.0 (0.5)	2910.2 (765.0)	2061.3 (180.1)	0.82 (0.07)	3.4 (0.8)	4.9 (2.4)	0.0015 (0.0005) 🗴	
US (28.5 m)	0.8 (0.2)	4.8 (0.9)	0.03 (0.007	3.6 (1.9)	29.8 (22.1)	3.8 (2.5)	1425.0 (469.7)	1691.4 (177.9)	0.73 (0.07)	3.3 (0.8)	7.1 (4.6)	0.0013 (0.0005) x	

Hillslope zone *: NS = near stream, MS = midslope, US = upslope. Depth*: metres below ground level to the centre of each well screen.

 $N_{INI}^*: NO_3^- + NH_4^+ + NO_2^- + \text{organic N} + \text{excess N}_2 + N_2O.$

Reaction progress (RP)*: excess $N_2 + N_2O/N_{INI}$.

EF_{5g}1*: IPCC N₂O emission factor: N₂O/N_{INI}.

 Table 4

 Mean groundwater and stream NO_3^- (\pm standard error SE) vs. aerobicity, bacterial energy sources, denitrification reaction products and N_2O emissions arranged by hillslope, hillslope positions and piezometric sampling depth in the slate catchment.

Hillslope zone*/	Reactive N			Aerobicity		Bacterial energy sources			Denitrification rate & products			N ₂ O emissions	
Depth* (m BGL)	NO ₃ (mg N/L)	N _{INI} * (mg N/L)	NO ₃ /Cl – (Ratio)	DO (mg/L)	Eh (mV)	DOC (mg/L)	Fe ²⁺ (μg/L)	Mn ²⁺ (μg/L)	RP*	Excess N ₂ (mg N/L)	N ₂ O (μg N/l)	N ₂ O−EF _{5g} 1* Above IPCC 2006: ✓, below: ४	
						Slate	N						
Stream I	7.8 (0.2)	8.7 (0.8)	0.41 (0.01)	10.56 (0.3)	110.6 (8.0)	2.5 (0.6)	14.6 (4.9)	4.0 (1.4)	0.00 (0.11)	0.02 (0.5)	0.69 (0.5)	0.0001 (0.00008) 🗶	
Stream II	7.4 (0.2)	8.9 (1.0)	0.40 (0.02)	9.93 (0.6)	111.2 (7.2)	2.3 (0.6)	16.9 (5.6)	3.6 (1.2)	0.01 (0.01)	0.10 (0.1)	0.56 (0.1)	0.0001 (0.00001) 🗶	
NS (2.5 m)	7.5 (0.2)	10.8 (0.9)	0.37 (0.02)	6.78 (1.2)	139.8 (10.5)	2.0 (0.8)	27.3 (5.9)	13.1 (2.8)	0.09 (0.02)	0.87 (0.1)	6.57 (1.8)	0.0006 (0.0002) 🗶	
NS (11 m)	7.4 (0.2)	9.4 (0.9)	0.38 (0.02)	6.83 (1.1)	139.6 (9.3)	1.7 (0.6)	21.1 (7.3)	14.7 (4.5)	0.08 (0.02)	0.73 (0.1)	5.89 (1.8)	0.0006 (0.0001) 🗶	
NS (42 m)	7.4 (0.2)	8.9 (0.8)	0.37 (0.02)	4.69 (1.2)	128.0 (10.7)	1.7 (0.6)	12.7 (4.4)	12.7 (9.1)	0.06 (0.02)	0.62 (0.2)	6.30 (1.8)	0.0007 (0.0001) 🗶	
MS (2.5 m)	8.3 (0.2)	9.6 (0.9)	0.39 (0.01)	10.03 (0.7)	119.8 (15.2)	1.9 (0.8)	11.0 (3.1)	21.8 (8.5)	0.06 (0.03)	0.68 (0.3)	1.78 (1.5)	0.0002 (0.0002) 🗶	
MS (26.5 m)	8.4 (0.2)	8.6 (0.8)	0.40 (0.02)	9.98 (0.5)	131.9 (9.5)	1.7 (0.6)	6.9 (2.2)	24.0 (11.7)	0.10 (0.03)	0.76 (0.3)	1.39 (0.3)	0.0001 (0.00003) 🗶	
MS (38 m)	8.0 (0.3)	7.3 (0.5)	0.37 (0.02)	4.45 (1.1)	126.9 (9.9)	2.0 (0.6)	9.9 (2.9)	19.8 (9.3)	0.32 (0.13)	2.50 (0.7)	3.56 (0.9)	0.0004 (0.0001) 🗶	
US (13.5 m)	11.6 (0.2)	13.57 (1.3)	0.46 (0.01)	7.85 (0.8)	150.5 (8.7)	2.0 (0.7)	9.7 (3.2)	6.6 (2.9)	0.05 (0.02)	0.86 (0.2)	3.26 (0.6)	0.0002 (0.00004) x	
US (28.5 m)	11.0 (0.3)	12.46 (0.4)	0.46 (0.02)	8.02 (0.5)	138.8 (9.5)	1.8 (0.7)	13.4 (6.5)	7.2 (3.4)	0.09 (0.02)	1.12 (0.2)	3.43 (0.6)	0.0002 (0.00004) 🗶	
						Slate	e S						
Stream	6.9 (0.1)	7.3 (0.9)	0.37 (0.01)	10.72 (0.2)	131.5 (8.3)	3.9 (2.4)	27.1 (13.2)	6.1 (3.5)	0.00 (0.02)	0.0 (0.2)	0.7 (0.5)	0.0001 (0.00004) x	
NS (5 m)	7.6 (0.1)	8.5 (0.3)	0.43 (0.01)	10.38 (0.4)	157.6 (9.7)	1.3 (1.1)	26.7 (3.9)	14.7 (1.7)	0.09 (0.02)	0.8 (0.2)	2.6 (0.5)	0.0003 (0.00005) 🗶	
NS (15.5 m)	7.5(0.1)	8.2 (0.2)	0.41 (0.01)	10.37 (0.4)	161.4 (9.1)	2.0 (1.1)	8.3 (3.9)	5.0 (1.7)	0.07 (0.02)	0.6 (0.2)	2.5 (0.4)	0.0003 (0.00005) 🗶	
NS (32 m)	6.5 (0.1)	8.3 (0.8)	0.36 (0.01)	7.67 (0.5)	145.8 (9.1)	0.8 (0.1)	31.6 (15.2)	8.0 (2.6)	0.12 (0.02)	1.1 (0.2)	2.3 (0.4)	0.0003 (0.00005) 🗶	
MS (9 m)	7.5 (0.2)	7.3 (1.0)	0.48 (0.02)	9.26 (0.4)	149.4 (11.4)	2.3 (1.2)	24.6 (10.6)	3.6 (0.9)	0.02 (0.03)	0.2 (0.3)	13.7 (5.6)	0.0019 (0.0008) 🗶	
MS (15.5 m)	7.5 (0.1)	7.8 (0.2)	0.47 (0.02)	10.78 (0.2)	163.8 (8.8)	2.4 (1.0)	34.6 (20.4)	7.6 (4.3)	0.03 (0.02)	0.2 (0.1)	2.5 (3.9)	0.0003 (0.0006) 🗶	
MS (38.5 m)	7.6 (0.1)	8.4 (0.4)	0.47 (0.01)	10.74 (0.6)	155.5 (9.6)	2.0 (1.1)	9.3 (2.3)	4.2 (0.9)	0.07 (0.02)	0.7 (0.2)	2.9 (0.4)	0.0003 (0.00006) 🗶	
US (9 m)	8.5. (0.4)	11.2 (1.1)	0.55 (0.03)	11.82 (0.1)	161.9 (17.6)	1.3 (0.3)	13.1 (3.0)	7.9 (2.0)	0.07 (0.03)	0.8 (0.4)	3.6 (0.9)	0.0003 (0.00009) 🗶	
US (17.5 m)	8.0 (0.2)	9.3 (0.8)	0.52 (0.02)	11.20 (0.4)	168.7 (11.3)	1.9 (0.6)	10.6 (3.2)	4.4 (0.8)	0.10 (0.03)	0.9 (0.2)	2.3 (0.6)	0.0002 (0.00007) 🗴	
US (38.5 m)	7.8 (0.2)	8.7 (0.6)	0.53 (0.02)	11.75 (0.6)	164.7 (9.2)	1.7 (0.7)	17.1 (8.3)	5.9 (1.3)	0.11 (0.03)	1.0 (0.3)	3.4 (0.7)	0.0003 (0.00007) 🗶	

Hillslope zone *: NS = near stream, MS = midslope, US = upslope. Depth*: metres below ground level to the centre of each well screen.

 $N_{INI}^*: NO_3^- + NH_4^+ + NO_2^- + organic N + excess N_2 + N_2O$.

Reaction progress (RP)*: excess $N_2 + N_2O/N_{INI}$.

EF_{5g}1*: IPCC N₂O emission factor.

increased significantly with increasing Ksat values (r = 0.63, p < 0.005) and decreased significantly with increasing water table ratios (r = -0.71, p < 0.05).

Within the slate catchment, *Slate N* had lower DO (p < 0.05) than *Slate S* with mean values of 7.3 and 10.5 mg/l (Table 4). DO was uniformly high in *Slate S* across groundwater sampling intervals (CV: 12%), whereas *Slate N* exhibited greater spatial variability (CV: 29%). In both hillslopes there existed weakly negative relationships between DO and sampling depth (p = 0.07). Groundwater in the slate catchment was consistently aerobic, with limited development of anaerobic zones. A significant positive correlation (r = 0.6, p < 0.05) was identified between DO and Ksat (Fig. 4c), while no significant relationship was identified between DO and the water table ratio (Fig. 4).

Eh in the sandstone catchment (mean: 54.9 mV) was significantly lower (p < 0.001) than the slate catchment (mean: 171.4 mV). Spatial variation was high with CVs of 94% and 50% in Sandstone N and Sandstone S respectively. Hillslope zone and depth effects mirrored those of DO owing to a strongly positive correlation between Eh and DO (r = 0.72; p < 0.001) (Table 5). Low Eh (< 10 mV) occurred in the near stream zones of Sandstone N (18.5 and 23.5 m BGL) and Sandstone S (11.5 and 18.5 m BGL) and in the upslope zone of Sandstone S (14.5 m BGL) (Table 3). The slate catchment had low spatial variation (CV: 6%) in both hillslopes and elevated Eh (>100 mV) across all hillslope zones and sampling depths.

3.2.2. Bacterial energy sources

Mean groundwater DOC in the sandstone catchment was significantly higher (p < 0.001) than the slate catchment (2.5 vs. 1.8 mg/L). In both catchments, stream DOC concentrations were greater than groundwater (mean sandstone: 6.5 vs. mean slate: 2.9 mg/L). In the sandstone catchment, groundwater Mn^{2+} concentrations (mean:

626.3 μg/L) were higher (p < 0.001) than the slate site. Within the sandstone catchment, *Sandstone S* had higher Mn²⁺ (p < 0.05) than *Sandstone N* with mean concentrations of 1391.7 and 69.8 μg/L respectively (Table 3). In both hillslopes, spatial variability was high (CV *Sandstone N*: 268% vs. *Sandstone S*: 85%). Screened intervals with highest Mn²⁺ (>600 μg/L) were coincident with lowest DO and Eh and included the near stream zones of *Sandstone N* (23.5 m BGL) and *Sandstone S* (5.5, 11.5 and 18.5 m BGL) and the upslope zone of *Sandstone S* (14.5 and 28.5 m BGL). Fe²⁺ was significantly greater (p < 0.001) in the sandstone catchment (mean: 639.5 μg/L) compared to the slate site. Analogous to the distribution of Mn²⁺ spatial variability was high with CV values of 157% and 87% in *Sandstone N* and *Sandstone S* respectively. Sampled intervals with highest Fe²⁺ concentrations (>1000 μg/L) followed the distribution of DO, Eh and Mn²⁺ (Table 3).

In the slate catchment, groundwater Mn²⁺ and Fe²⁺ concentrations were typically low (Table 4). Mn²⁺ ranged from 2.3 to 13.3 μ g/L, with a mean value of 5.3 μ g/L and a CV of 57%. Fe²⁺ ranged from 0.98 to 16.58 μ g/L, with a mean groundwater concentration of 7.2 μ g/L. Spatial variation was similar between *Slate N* and *Slate S* with CV values of 65% and 62% respectively.

3.3. Excess N₂ concentrations

Excess N_2 in the sandstone catchment was significantly higher than the slate catchment (p < 0.05). Mean groundwater concentrations of 1.62 and 0.74 mg N/L were measured in the sandstone and slate catchments respectively. In the sandstone catchment, significant relationships were shown between excess N_2 and hillslopes i.e. Sandstone N vs. Sandstone S (p < 0.001), hillslope zones (p < 0.05) and sampling depths (p < 0.05). Significant interactions were identified between hillslope and depth (p < 0.05) and hillslope zone and depth (p < 0.05).

Table 5 Correlation coefficient and significance matrix between NO_3^- , the drivers of denitrification, denitrification rate and reaction products, stable isotope signatures and N_2O emissions for the sandstone and slate catchments.

		Aerobicity		Bacterial 6	energy sourc	es	Denitrification rate & products			N ₂ O emissions	Stable isotopes		
	NO_3^-	DO	Eh	DOC	Fe ²⁺	Mn ²⁺	RP	Excess N ₂	N ₂ O	N ₂ O-EF _{5g} 1	$\delta^{15}N_{NO3}$	$\delta\ ^{18}O_{NO3}$	δ $^{15}N_{N2O}$
Sandstone ca	tchment												
NO_3^-													
DO	0.78**												
Eh	0.56^{*}	0.72**											
DOC	0.05 ^{ns}	0.39*	0.19 ^{ns}										
Fe ²⁺	-0.83**	-0.72**	-0.43^{*}	-0.13^{ns}									
Mn^{2+}	-0.90**	-0.87**	-0.56^{*}	-0.15^{ns}	0.93**								
RP*	-0.94**	-0.87**	-0.64**	-0.16^{ns}	0.90**	0.95**							
excess N ₂	-0.77**	-0.79^{**}	-0.54^{*}	0.09 ^{ns}	0.76**	0.84**	0.85**						
N_2O	0.83**	0.77**	0.72**	0.35 ^{ns}	-0.71**	-0.79**	-0.87**	-0.52^{*}					
N ₂ O - EF _{5g} 1	0.54^{*}	0.58**	0.79**	0.30 ^{ns}	-0.40^{ns}	-0.52**	-0.60**	-0.28^{ns}	0.85**				
¹⁵ N _{NO3}	-0.51	-0.47^{*}	-0.22^{ns}	-0.15^{ns}	0.42ms*	0.44 ^{ms*}	0.57^{*}	0.49^{*}	$-0.44^{\text{ms}*}$	-0.21^{ns}			
¹⁸ O _{NO3}	-0.47^{*}	-0.50^{*}	$-0.41^{\text{ms}*}$	-0.07^{ns}	0.38 ^{ns}	0.45 ^{ms*}	0.58**	0.57^{*}	$-0.44^{\text{ms*}}$	-0.30^{ns}	0.95**		
$^{15}N_{N2O}$	-0.78**	-0.68**	-0.59**	0.14 ^{ns}	0.80**	0.86**	0.86**	0.79**	-0.71**	-0.55^{*}	0.37 ^{ns}	0.46*	
Slate catchm	ent												
NO_3^-													
DO	-0.03^{ns}												
Eh	-0.09^{ns}	0.69**											
DOC	0.19 ^{ns}	0.00 ^{ns}	0.01 ^{ns}										
Fe ²⁺	-0.41^{ns}	-0.05^{ns}	0.22 ^{ns}	-0.16^{ns}									
Mn^{2+}	-0.05^{ns}	-0.43^{ns}	-0.75**	-0.23^{ns}	-0.07^{ns}								
RP*	-0.08^{ns}	-0.45^{ns}	-0.33^{ns}	-0.15^{ns}	-0.22^{ns}	0.40 ^{ns}							
excess N ₂	0.14 ^{ns}	-0.48^{ns}	-0.35^{ns}	-0.21^{ns}	-0.27^{ns}	0.36 ^{ns}	0.96^{*}						
N_2O	-0.12^{ns}	-0.41^{ns}	-0.04^{ns}	0.27 ^{ns}	0.39 ^{ns}	-0.27^{ns}	-0.15^{ns}	-0.15^{ns}					
N ₂ O - EF _{5g} 1	-0.42^{ns}	-0.37^{ns}	-0.03^{ns}	0.23 ^{ns}	0.50 ^{ns}	-0.24^{ns}	-0.12^{ns}	-0.18^{ns}	0.93**				
$^{15}N_{NO3}$	-0.20^{ns}	-0.77**	-0.66^{*}	0.00 ^{ns}	0.00 ^{ns}	0.52 ^{ns}	0.22 ^{ns}	0.20	0.80**	0.74**			
¹⁸ O _{NO3}	-0.09^{ns}	-0.67^{*}	$-0.52^{\text{ms*}}$	-0.10^{ns}	0.05 ^{ns}	0.35 ^{ns}	0.05 ^{ns}	0.08 ^{ns}	0.87**	0.72**	0.94**		
¹⁵ N _{N2O}	-0.23^{ns}	0.58 ^{ms*}	0.44 ^{ns}	-0.45^{ns}	0.41 ^{ns}	0.06 ^{ns}	-0.52^{ns}	-0.54^{ns}	-0.29^{ns}	-0.21^{ns}	-0.35^{ns}	-0.19^{ns}	

^{**} Correlation is significant at the 0.01 level.

^{*} Correlation is significant at the 0.05 level.

ms* Correlation is marginally significant at the 0.1 level.

ns Correlation is not significant.

Sampling depth was positively correlated with excess N_2 . Analogous to the zonal relationship described with NO_3^- , a combination of near stream zone and sampling depth had the greatest effect on groundwater excess N_2 (Fig. 3c). Highest excess N_2 corresponded to lowest NO_3 — (Fig. 3a) and DO (Fig. 3b). In *Sandstone N*, near stream excess N_2 concentrations were higher than midslope (p < 0.0001) and upslope (p < 0.05) (Fig. 3c) with lowest values in the midslope zone. In *Sandstone S*, near stream (p < 0.05) and upslope (p < 0.05) excess N_2 concentrations were higher than midslope values. In stream excess N_2 was lower than groundwater with mean concentrations of 0.1 and 0.2 mg N/L in *Sandstone N* and *Sandstone S*.

Within the slate catchment, no statistically significantly significant differences were identified between excess N_2 and hillslopes, hillslope zones or sample depths. Spatial variation was relatively low (CV: 34%) in both hillslopes, with the exception of the midslope zone of *Slate N* (38 m BGL) were a mean excess N_2 concentration of 3.8 mg N/L was identified. In stream excess N_2 concentrations were similar to groundwater with mean values of 0.12 and 0.22 mg N/L in *Slate N* and *Slate S*.

3.4. N₂O and N₂O emission factors

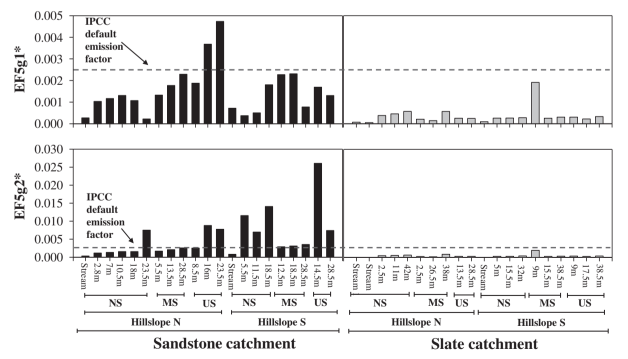
Groundwater N₂O was significantly higher (p < 0.0001) in the sand-stone catchment compared to the slate catchment (mean sandstone: 13.0 vs. mean slate: 4.0 µg N/L). In the sandstone catchment, significant relationships were identified between N₂O and hillslopes (p < 0.01) with greatest concentrations in *Sandstone N* (Table 3). In *Sandstone N*, near stream N₂O was lower than midslope (p < 0.05) and upslope concentrations (p < 0.05) with highest values in the upslope zone. In *Sandstone S*, near stream N₂O was lower than midslope (p < 0.005), which had the highest N₂O concentrations. No significant difference was shown between the near stream and upslope zones (p = 0.61). Depth had contrasting effects on N₂O in both hillslopes which were contingent upon hillslope position. In the near stream zones, depth exhibited a

typically negative correlation, however in the midslope and upslope zones the relationship was typically positive (Fig. 3d; Table 3).

In the slate catchment, no significant relationships were identified between N_2O and hillslope (p=0.64), hillslope zones (p=0.96) or sample depths (p=0.67). Spatial variability in both hillslopes was relatively high (mean CV: 72%). This CV was however influenced by four comparative outliers in the dataset namely the near stream zone of *Slate N* (2.5, 11.5 and 42 m BGL) and the midslope zone of *Slate S* (9 m BGL). In both catchments, in-stream N_2O concentrations were lower than groundwater. Mean stream N_2O in the sandstone catchment exceeded the slate catchment by factor of five.

Fig. 5 illustrates the spatial distribution of groundwater and stream water N₂O emission factors (EF_{5g}1 and EF_{5g}2). EF_{5g}1 in the sandstone catchment was significantly higher than the slate catchment (p < 0.0001) with the sandstone mean (0.0018) exceeding the slate mean (0.00043) by a factor of three. Mean emission factors for both sandstone hillslopes were comparable (Sandstone N: 0.0017 vs. Sandstone S: 0.0019) with similar spatial variation (62% vs. 67%). In the sandstone catchment, a significant relationship was found between $EF_{5g}1$ and hillslope zone (p < 0.001). In both hillslopes, near stream zones had lower emission factors than midslope and upslope zones (p < 0.05). While a statistically significantly relationship with depth was not identified, an analysis of the data, revealed a depth wise relationship analogous to N₂O. In Sandstone N, lowest N₂O emission factors were observed at greatest sampling depths (18 and 25 m BGL) in the near stream zone (0.001 and 0.0002). Conversely, highest emission factors were measured at greatest sampling depth (16 and 23.5 m BGL) in the upslope zone (0.0035 and 0.0045) (Fig. 4). In Sandstone S, lowest EF_{5g}1 corresponded to near stream groundwater, whereas highest values were calculated in the midslope zone.

In the slate catchment, the distribution of $EF_{5g}1$ mirrored the relative abundance of N_2O , with highest emission factors in the near stream zone of *Slate N* (2.5, 11.5 and 42 m BGL) and the midslope zone of *Slate S* (9 m BGL). Spatial variability amongst the remaining groundwater sampling locations was low (CV: 24%). Calculated $EF_{5g}2$ emission factors



 $\mathbf{EF_{5g}1^*}$: IPCC N₂O emission factor: N₂O/N_{INI} (NO₃⁻ + NH₄⁺ + NO₂⁻ + organic N + excess N₂ + N₂O) $\mathbf{EF_{5g}2^*}$: IPCC N₂O emission factor: N₂O/NO₃⁻

Fig. 5. Distribution of N₂O emission factors, mean EF_{5g}1 and EF_{5g}1 (2013–15), in the sandstone and slate catchments organised by hillslope, slope position and sampling depth.

exceeded corresponding $EF_{5g}1$ values by a factor of 1.3, 1.1, 1.8 and 4.4 in *Slate N, Slate S, Sandstone N* and *Sandstone S* respectively. $EF_{5g}2$ values were significantly higher (p < 0.0001) in the sandstone catchment vs. the slate catchment. $EF_{5g}2$ emission factors were significantly greater than corresponding $EF_{5g}1$ values in the near stream zones of both *Sandstone N* and *Sandstone S*, with lowest values in the midslope zones. Conversely the spatial distribution of $EF_{5g}2$ in the slate catchment was similar to the distribution of $EF_{5g}1$.

3.5. Denitrification reaction progress (RP)

Denitrification reaction progress (RP) is shown Tables 3 and 4 for the sandstone and slate catchments respectively. Groundwater RP in the sandstone catchment was significantly higher (p < 0.005) than the slate catchment ranging from 0.04–0.94. In the sandstone catchment significant interactions were identified between hillslope and hillslope zone (p < 0.05) and hillslopes zone and depth (p < 0.05). A combination of near stream zone and increasing depth had the greatest effect on RP values (p < 0.05). In *Sandstone N*, RP values in the near stream zone were significantly higher (p < 0.05) than up gradient values. There was a significant positive correlation between RP and sampling depth. In *Sandstone S*, the near stream and upslope zones had greater RP than the midslope zone (p < 0.001; p < 0.05) with no significant difference between near stream and upslope (p = 0.91). RP values were elevated at all sample depths in the near stream zone (Table 3).

In the slate catchment, RP ranged from 0.02 to 0.34, with a mean value of 0.11 (Table 4). No significant relationships were identified between RP and hillslope, hillslope zone or sampling depth.

3.6. Denitrification vs. dilution

To differentiate between denitrification and dilution, the distribution Cl⁻ (conservative) versus NO₃ (non-conservative) was examined. In the sandstone catchment, the mean Cl⁻ concentration was 31.38 mg/L. Spatial variation across hillslope zones and sampling depths was low, with CV values of 6% and 3% in *Sandstone N* and *Sandstone S*, respectively. Conversely, the ratio of NO₃ to Cl⁻ showed considerable spatial variation (CV *Sandstone N*: 38%; CV *Sandstone S*: 110%).

In the slate catchment, the spatial mean Cl⁻ concentration was 19.27 mg/L, approximately 40% lower than the sandstone catchment (Table 4). Spatial variation across both hillslopes was low (mean CV: 16.6%). In contrast to the sandstone catchment, the ratio of NO₃⁻ to Cl⁻ was relatively uniform in both hillslopes with a range of 0.35 to 0.55 and a CV of 14%.

3.7. Factors affecting denitrification

3.7.1. Dissolved oxygen

The relationships between groundwater DO and RP, NO $_3^-$, excess N $_2$ and N $_2$ O for both study catchments are presented in Fig. 6. A breakdown of catchment specific relationships are presented as a correlation matrix in Table 5. In the sandstone catchment, strongly negative correlations were identified between groundwater DO and RP (r=-0.87, p<0.0001). DO was negatively correlated with excess N $_2$ (r=-0.79, p<0.0001) and positively correlated with NO $_3^-$ (r=0.78, p<0.001) (Fig. 6). The relationship between N $_2$ O and DO in the sandstone catchment, while statistically positive (r=0.77, p<0.05) was complex. Highest N $_2$ O occurred at DO between 4 and 8 mg/L with significantly lower concentrations between 0 and 3 mg/L and 8 to 10 mg/L (Fig. 6). In contrast to the sandstone catchment, groundwater in the slate catchment was consistently aerobic, with no significant correlations identified between DO and RP, NO $_3^-$, excess N $_2$ or N $_2$ O.

3.7.2. Bacterial energy sources

In the sandstone catchment, concentrations of reduced metals (Mn²⁺ and Fe²⁺) were highly dependent on aquifer aerobicity

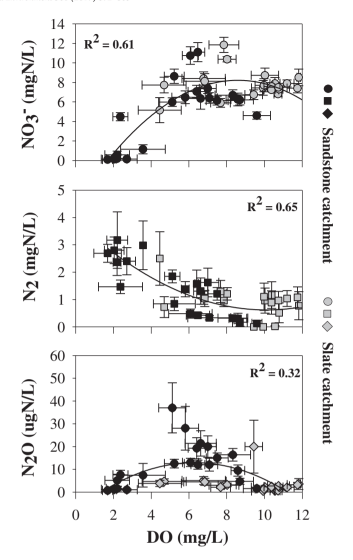


Fig. 6. Groundwater dissolved oxygen (DO) plotted against NO_3^- , excess N_2 and N_2O for the sandstone (black symbols) and slate (grey symbols) catchments.

(Table 5). Mn^{2+} was negatively correlated with DO (r = -0.87; p < 0.0001) and Eh (r = -0.56; p < 0.001). Similarly, Fe²⁺ was negatively correlated with DO (r = -0.72; p < 0.0001) and Eh (r = -0.43; p < 0.05), and positively correlated with Mn²⁺ (r = 0.93;p < 0.0001) (Table 5). Significant positive relationships were discovered between Mn²⁺ and RP (r = 0.95, p < 0.0001) and Fe²⁺ and RP (r = 0.90, p < 0.0001) p < 0.0001). Strongly negative relationships were shown between Fe²⁺ and NO_3^- (r = -0.83, p < 0.005) and between Mn^{2+} and NO_3^- (r = -0.90, p < 0.0001). N₂O was negatively correlated with Fe²⁺ (r = -0.71, p < 0.005) and Mn^{2+} (r = -0.79, p < 0.005). Conversely, excess N_2 was positively correlated with Fe^{2+} (r = -0.76, p < 0.0001) and Mn²⁺ (r = 0.84, p < 0.0001). No significant relationships were observed between DOC and excess N2 or N2O. DOC did however exhibit a weakly negative correlation with DO (r = -0.39, p =0.087). In the slate catchment, no significant relationships were identified between reduced bacterial energy sources (DOC, Mn²⁺ & Fe²⁺⁾ and RP, DO, NO_3^- , N_2O or excess N_2 .

3.7.3. Denitrification reaction progress (RP)

DO concentrations and the abundance of reduced metals (Mn^{2+} and Fe^{2+}) were the dominant factors affecting denitrification RP. The spatial relationships between groundwater RP and NO_3^- , excess N_2 and $\mathrm{N}_2\mathrm{O}$ for both study catchments are presented in Fig. 7 and Table 5. In the sandstone catchment, RP exhibited a strongly negative correlation with NO_3^-

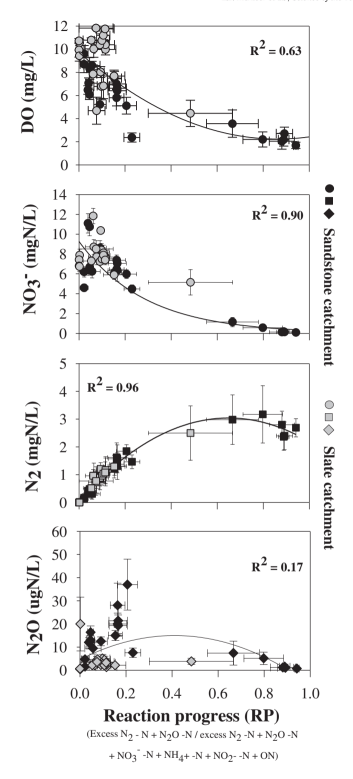


Fig. 7. Groundwater reaction progress (RP) plotted against dissolved oxygen (DO), NO_3^- , excess N_2 and N_2O for the sandstone (black symbols) and slate (grey symbols) catchments.

(r=-0.94, p<0.0001) and N_2O (r=-0.87, p<0.0001). Conversely, there was a significantly positive relationship between RP and excess N_2 (r=0.85, p<0.0001). Lowest NO_3^- concentrations corresponded to RP >0.60, which was consistent with the highest measured excess N_2 . N_2O concentrations were low at RP <0.10 and >0.60, with highest concentrations occurring between 0.10 and 0.3 (Fig. 7). In slate catchment, no significant correlations were identified between RP and NO_3^- or N_2O

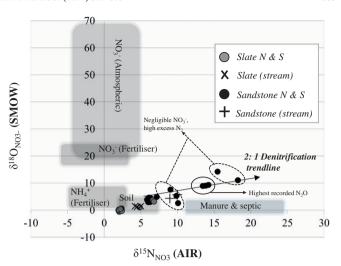


Fig. 8. δ^{18} O versus δ^{15} N values of NO $_3^-$ for groundwater samples collected in September 2014 Also shown are the δ^{18} O and δ^{15} N ranges typical of NO $_3^-$ sources and a 2:1 trendline of δ^{18} O versus δ^{15} N which is typical of denitrification progress (after Kendall et al., 2007).

3.8. Stable isotope values

 $\delta^{15} N_{NO3}$ and $\delta^{18} O_{NO3}$ were significantly higher in the sandstone catchment compared to the slate site (Fig. 8) (p < 0.0001). In the sandstone catchment, $\delta^{15}N_{NO3}$ ranged from +6.0 to $18.2^{\circ}/_{oo}$ with a mean ratio of $9.6^{\circ}/_{oo}$. $\delta^{18}O_{NO3}$ ranged from 3.0 to $11.6^{\circ}/_{oo}$, with a mean value of $6.3^{\circ}/_{oo}$. Highest values of both $\delta^{15}N_{NO3}$ and $\delta^{18}O_{NO3}$ occurred in the near stream zones of Sandstone N and Sandstone S and in the upslope zones of both hillslopes. $\delta^{15} N_{NO3}$ was negatively correlated with NO_3^- (r = -0.51, p < 0.05) (Fig. 9), DO (r = -0.47, p < 0.05) and N₂O (r = -0.44, p < 0.1) and positively related to RP (r = 0.58, p < 0.01)and excess N₂ (r = 0.49, p < 0.05) (Table 5). δ^{15} N_{N2O} was significantly higher in the sandstone catchment versus than the slate catchment (p < 0.05). In the sandstone catchment, $\delta^{15}N_{N20}$ ranged from -13.6 to $9.7^{\circ}/_{\circ \circ}$, with a mean value of $4.9^{\circ}/_{\circ \circ}$ and exhibited both positive and negative enrichment ratios (Fig. 9). $\delta^{15}N_{N2O}$ was negatively correlated with NO₃⁻ (r = -0.78, p < 0.01) (Fig. 8), DO (r = -0.68, p < 0.01)and N₂O (r = -0.71, p < 0.01) (Table 5). Conversely δ^{15} N_{N2O} was positively correlated with RP (r = 0.86, p < 0.05) and excess N₂ (r = 0.79,

In the slate catchment, $\delta^{15} N_{NO3}$ ranged from + 1.9 to $6.8^{\circ}/_{oo}$ with a mean value of $3.3^{\circ}/_{oo}$. $\delta^{18} O_{NO3}$ ranged from - 0.5 to + 3.8°/ $_{oo}$ with a mean enrichment ratio of $0.8^{\circ}/_{oo}$. $\delta^{15} N_{NO3}$ was not correlated with NO_3^- , RP or excess N_2 but did exhibit significant relationships with DO (r=-0.77, p<0.05) and N_2O (r=0.8, p<0.05). $\delta^{15} N_{N2O}$ was consistently negative, ranging from -20.8 to $-4.6^{\circ}/_{oo}$ (mean: $-3.2^{\circ}/_{oo}$) (Fig. 9). No significant correlations were identified between $\delta^{15} N_{N2O}$ and NO_3^- , RP, excess N_2 or N_2O . A marginally significant correlation was shown between $\delta^{15} N_{N2O}$ and DO (r=0.58, p<0.1).

4. Discussion

The study highlighted a complex mosaic of NO₃⁻ removal capacity between study catchments, between catchment hillslopes, and within catchment hillslopes. A range of spatially variable, physical and hydrogeochemical parameters where identified to regulate 1) NO₃⁻ removal capacity, 2) gaseous accumulation and 3) indirect N₂O emissions. Previous research has documented that low permeability aquifers coupled with the presence of denitrifying bacteria, reducing conditions and the availability of bacterial energy sources create zones of enhanced denitrification potential (Brettar et al., 2002; Jahangir et al., 2012a; Rissmann,

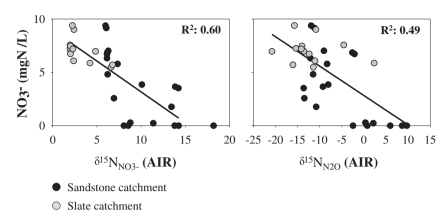


Fig. 9. $\delta^{15}N_{NO3}\delta^{15}N_{N2O}$ values vs. NO_3^- in the sandstone and slate catchments.

2011; Thayalakumaran et al., 2008. While a number of these criteria may be met within a given catchment, the absence of one can arrest the denitrification process and resultant NO_3^- attenuation. The combined analysis of physical and hydrogeochemical factors and stable isotope signatures versus the natural abundance of NO_3^- , excess N_2 and N_2O revealed a tale of two catchments. The slate catchment was characterised by hydrogeochemical uniformity with limited groundwater NO_3^- reduction. The sandstone catchment exhibited spatially heterogeneous zones of incomplete and complete NO_3^- removal. Despite significantly higher applications of organic and inorganic N in the sandstone catchment, mean groundwater and stream NO_3^- concentrations were significantly lower. The positive mitigation effect of NO_3^- removal in the sandstone catchment was however concomitant with substantially greater emissions of N_2O .

4.1. Denitrification vs dilution

The ratio between NO $_3^-$ and Cl $^-$ was used as an indirect indictor to differentiate between denitrification and dilution processes, with Cl $^-$ acting conservatively and NO $_3^-$ acting non-conservatively along a flow path. In the sandstone catchment, low concentrations of NO $_3^-$ with unaffected Cl $^-$ resulted in low NO $_3^-$ /Cl $^-$ ratios in various zones throughout each hillslope (Table 4). Similar to Fenton et al. (2009), this suggests that groundwater denitrification was the dominant factor regulating groundwater NO $_3^-$ occurrence. In the slate catchment high hillslope NO $_3^-$ /Cl $^-$ ratios and a proportional decrease of NO $_3^-$ and Cl $^-$ progressing downslope, suggest dilution as the dominant NO $_3^-$ attenuating mechanism.

Stable isotope signatures were used to identify transformational processes in the study catchments. The bacteria involved in denitrification preferentially metabolise the lighter NO₃ isotopes. As such, an enrichment of the heavier NO₃ isotopes in groundwater provides direct evidence of denitrification. In the sandstone catchment, contemporaneous isotopic enrichment of $\delta^{15} N_{NO3}$ and $\delta^{18} O_{NO3}$, along an approximate 2:1 regression trend line signified denitrification (Fig. 8). Previous authors reported a comparable trend in the relative proportion of $\delta^{15}N_{NO3}$ vs. $\delta^{18}O_{NO3}$ as representative of denitrification with ratios of 1.5:1 identified in Baily et al. (2011) and 2:1 in Aravena and Robertson (1998), Mengis et al. (1999) and Panno et al. (2006). Correlations between $^{15}N_{NO3}$ and NO_3^- and $\delta^{15}N_{N2O}$ and NO_3^- indicated both reduction of NO₃ and production of N₂O in groundwater (Fig. 9). In highly denitrifying zones, N2O was further reduced to excess N2, as indicated by positive $\delta^{15} N_{N2O}$ values e.g. Well et al. (2012). N_2O reduction was coincident with high RP, low NO₃ and high excess N₂. In the slate catchment a lack of correlation between $\delta^{15}N_{NO3}$, $\delta^{15}N_{N2O}$ and NO_3^- (Table 5) indicated that denitrification was unlikely. Uniformity in groundwater NO₃ concentrations was coincident with low RP. Moreover, $\delta^{15}N_{NO3}$ and $\delta^{18}O_{NO3}$ ratios (Fig. 5) were reflective of source rather than transformational signatures supporting the contention that dilution rather than denitrification dominated in the slate catchment.

4.2. Catchment N dynamics & denitrification capacity

In spite of a 79% greater surface application of available N to the sandstone hillslopes (Table 2), mean stream NO₃ concentrations were 32% lower than the slate sites (Fig. 2). This is part due to the presence of long term grassland cover in the sandstone catchment versus tillage in the slate catchment. Tillage catchments undergo periodic ploughing and reseeding with associated bare soils resulting in higher N leaching losses from soil to groundwater and lower DOC loss which limits subsurface denitrification (Jahangir et al., 2014). The greater propensity for N in loss in arable catchments was reflected in a mean FracLEACH value of 42.5% in the slate catchment versus 17.5% in the grassland catchment (Table 2). While FracLEACH was considerably higher at the slate hillslopes, the associated N loads reaching the water table were comparable with 55 and 60 kg N/ha/yr. reaching the water table in the sandstone and slate catchments respectively. These are similar to leaching losses reported by Huebsch et al. (2013) in intensively managed grassland and Premrov et al. (2012) in spring barley. Comparing shallow groundwater NO₃ with stream NO₃ concentrations in both catchments revealed a 43% reduction in NO₃ from groundwater to stream in the sandstone catchment versus a 7% reduction in the slate site. It is likely therefore that a combination of agricultural practises and hydrogeochemical transformations acted to mitigate stream N enrichment in the denitrifying sandstone catchment.

Analogous to the findings of Hinkle et al. (2007), Weymann et al. (2008) and Jahangir et al. (2012a) the sandstone catchment was characterised by highest NO₃ in shallow groundwater, typically decreasing with depth (Fig. 2). Aligned to Fenton et al. (2009), highest NO₃ concentrations were observed in high permeability quaternary deposits. Underlying the Quaternary deposits in both sandstone hillslopes, a layered distribution of bedrock permeability (Ksat) was evident, becoming less permeable with depth (Table 2). In the sandstone catchment, weathered bedrock zones with high Ksat allowed a fast migration of NO₃ contaminated groundwater with limited scope for microbial attenuation. Lower Ksat with depth resulted in a longer residence time of both groundwater NO₃ and DO. With a longer residence time in the sandstone catchment, denitrification was promoted, resulting in lower NO₃ concentrations. This contention was supported by significant negative correlations between Ksat and NO₃ and Ksat and DO (Fig. 4). High permeability weathered zones ranging in thickness from 4 to 11 m and 5 to 18 m in Slate N and Slate S respectively (Table 1), aligned with a lack of correlation between Ksat and NO₃ or DO suggested that rapid groundwater flow restricted denitrification progress.

Groundwater DO concentration was the dominant control on RP, NO₃, excess N₂ and N₂O. Analogous to field studies undertaken in highly denitrifying aguifers (Blicher-Mathiesen et al., 1998; Well et al., 2012), significant correlations were identified between DO, NO₃, N₂O and excess N₂. Previous studies have documented a DO range between <1 and 4 mg/L as supportive of denitrification (Rivett et al., 2008). In the sandstone catchment, negligible groundwater NO₃ concentrations corresponded to DO < 3.5 mg/L and Eh < 30 mV. In low oxygen environments, denitrification results in the sequential reduction of NO₃⁻ to N₂O to excess N2. Near stream zones exhibited a significant capacity for complete denitrification to excess N2, where low DO and high RP corresponded to negligible NO₃, low N₂O and high excess N₂. The correlation between the relative depth of the water table and DO (Fig. 4), indicated that a water table close to the ground surface promoted lower groundwater DO and NO₃ concentrations. In near stream zones where water tables are typically < 1 m BGL, the development of anaerobic conditions, and as a result denitrifying environments are promoted. With the exception of the upslope zone of Sandstone S which was a relative outlier, the range of DO (5.1–8.6 mg/L) in the midslope and upslope zones of the sandstone catchment inhibited complete denitrification to excess N2, instead arresting the denitrification reaction at the N₂O production stage. Maximum water table depths in the midslope and upslope zones of the sandstone hillslopes ranged from 4 to 14 m BGL. Higher DO in the midslope and upslope hillslope zones of the sandstone catchment support the results of Young et al. (2013), where substantially greater diffusive and advective transport of DO into groundwater underling unsaturated zones in excess of 5 m thickness was observed. Similarly, Fenton et al. (2009) and Jahangir et al. (2012a, 2013) described highest NO₃ and N₂O in groundwater underlain by thick unsaturated zones. Results of the statistical model and correlation analysis suggest therefore that near stream zones promote complete reduction of NO₃⁻ to excess N₂ whereas in midslope and upslope zones, N₂O is the dominant denitrification reaction product.

Whereas the sandstone catchment exhibited a complex interplay between groundwater DO, RP, NO₃⁻, N₂O and excess N₂, the slate catchment was characterised by a lack of significant correlation. In the slate catchment, aerobic conditions at all hillslope zones and depths restricted the development of denitrifying hotspots, with correspondingly low RP values throughout (Fig. 7). This resulted in uniformly high NO₃⁻, low excess N₂ and low N₂O. Although permeability was typically high, low Ksat zones in *Slate N* did not promote complete denitrification. Moreover intermediate DO concentrations which were comparable to the sandstone catchment resulted in significantly lower N₂O production.

Reduction of both DO and NO_3^- requires a bacterial energy source. Under low DO concentrations, NO_3^- is the most energetically favourable electron acceptor for bacterial metabolism. Energy sources include surface derived DOC and/or solid phase carbon, Fe^{2+} , Mn^{2+} and S^- (Rivett et al., 2008) dissolved under anaerobic conditions. Mean DOC of 2.5 and 1.8 mg/L in the sandstone and slate catchments respectively was aligned with the low concentrations described in previous studies (<5 mg/L) (Rivett et al., 2008). A lack of bioavailable DOC is a dominant limiting factor on groundwater denitrification rates (Starr and Gillham, 1993; Wassenaar, 1995). In both catchments there was a lack of correlation between DOC and RP, NO_3^- , excess N_2 or N_2 O (Table 5). The required stoichiometry of 1:1.25 between NO_3^- and DOC (Dimkić et al., 2008; Thayalakumaran et al., 2008) indicated that the levels of DOC in the saturated zones of both catchments could not support heterotrophic denitrification.

Typical of grassland agriculture, inputs of organic N to the sandstone hillslopes were high (Table 2). Previous research (Pabich et al., 2001) documented an exponential decrease of DOC with depth below the water table. Conversely, in the sandstone catchment, spatial variability in DOC was uniformly low in both shallow and deeper groundwater pathways. During transport through the unsaturated zone DOC is biodegraded via oxidation to CO_2^- , which in turn reduces DO. Mean DO of 5.0 mg/L in shallow groundwater (<10 m BGL) of the sandstone

catchment, aligned with low spatial variation of DOC, suggested DOC consumption and DO reduction prior to reaching the water table. A significantly negative correlation between DOC and DO (Table 5) within the saturated zone suggested that a proportion of the leached DOC was bioavailable (Chapelle et al., 2012). Organic N input to the sandstone hillslopes was on average 116% greater than the slate sites. Conversely, mean shallow groundwater (<10 m BGL) DOC concentrations were only 15% greater. DOC consumption prior to reaching the water table in the sandstone catchment was therefore a likely factor. Lower lability soil organic carbon is typical of long term arable cropping systems compared to grassland dominated catchments (Premrov et al., 2012). Organic inputs were not sufficient to support DO reduction in the slate catchment, as indicated by low DOC and high DO in both hillslopes.

When labile DOC is not sufficient in supply or has been biodegraded during DO reduction, solid phase electron donors must be present to support denitrification. When anaerobicity prevails, electron donors such as Mn²⁺ and Fe²⁺ can accumulate along a flow path (Tesoriero and Puckett, 2011). Several studies have hypothesised autotrophic denitrification in DOC limited aguifers (Green et al., 2008; Heffernan et al., 2012; Weymann et al., 2008). In the sandstone catchment, strongly positive correlations were identified between Fe²⁺/Mn²⁺, RP and excess N₂ (Table 5). While RP and excess N₂ increased with increasing Fe²⁺ and Mn²⁺; NO₃, N₂O, DO and Eh decreased signifying autotrophic denitrification. If an aquifer lacks solid phase electron donors (Liao et al., 2012), DOC concentrations in recharge are too low to support DO reduction (Thayalakumaran et al., 2015) or residence times are too short to support significant solid phase dissolution (Vidon and Hill, 2005), denitrification is suppressed. It is likely that this was the case in the slate catchment.

4.3. Catchment N₂O concentrations & emissions

The spatial mean groundwater N₂O concentrations (sandstone: 13 μg N/L; slate: 4 μg N/L) observed in the present study were comparable those observed by other authors. Vilain et al. (2012) measured mean concentrations of 37.4, 11.1 and 9.5 µg N/L in the upslope, midslope and near stream zones of a limestone aquifer, while Mühlherr and Hiscock (1998) described mean N_2O concentrations 33, 12.5 and 4.3 µg N/L in three limestone aguifers. A commonality throughout the literature indicates that N₂O accumulation is spatially variable with CVs of 217% (Yanai et al., 2003), 258% (von der Heide et al., 2009) compared with 72–99% in slate and sandstone catchment. At the average sampling temperature (11 °C), the expected air equilibrium concentration of N₂O in groundwater is 0.33 µg N/L (Mühlherr and Hiscock, 1998). Mean groundwater N₂O was 39 and 12 times greater than atmospheric equilibrium in the sandstone and slate catchments respectively. Furthermore, the maximum recorded N₂O concentration in the sandstone catchment exceeded atmospheric equilibrium by a factor of 300. Any N₂O in excess of atmospheric equilibrium will degas to the atmosphere once groundwater rises to the surface, enters the stream or diffuses through the unsaturated zone. The described concentrations in both catchments therefore represent potential greenhouse gas sources to the atmosphere. Directly measured N₂O concentrations however do not represent the actual emissions from the aquifer to the atmosphere (Weymann et al., 2008). N₂O may increase or decrease with greater residence time prior to its eventual release point. Von der Heide et al. (2009) measured negligible fluxes of N₂O vertically from groundwater to the atmosphere in spite of high N2O concentrations in shallow groundwater. Given that highest N₂O accumulation occurred at depth, it is likely that advective rather than diffusive transport of groundwater N₂O dominated in the present study. In stream N₂O concentrations provide a measurement groundwater derived N2O degassing to the atmosphere. In the sandstone catchment, mean near stream groundwater N₂O was 95% lower than up gradient concentrations while mean stream N₂O concentrations were 68% lower than near stream groundwater. Analogous to Höll et al. (2005) it is likely that groundwater N₂O was

consumed during passage from groundwater to the stream. While substantially lower than up gradient groundwater concentrations, stream N₂O in Sandstone N and Sandstone S were 4.5 and 16 times atmospheric equilibrium respectively, indicating a net contribution to atmospheric greenhouse gas emissions. In the slate catchment, mean stream N₂O was 164% less than the sandstone catchment. In the present study, two available methods were used to calculate EF_{5g}, producing contrasting results. EF_{5g}1 takes into account all forms of reactive N including the products of denitrification. The results of the EF_{5g}1 analysis support the IPCC (2006) downward revision from 0.015 to 0.0025 with mean emission factors of 0.0018 and 0.0004 in the sandstone and slate catchments respectively. Emissions above the IPCC default were consistent with highest N₂O concentrations in the midslope and upslope zones of the sandstone hillslopes (Fig. 5); however as demonstrated, actual N2O transport to the streams did not reflect the up gradient maxima. The calculation of EF_{5g}2 considers NO₃⁻ and N₂O only. In the sandstone catchment, where there was significant denitrification, $EF_{5g}2$ overestimated emissions, with the mean value of 0.005 exceeding EF_{5g}1 by 113%. Highest EF_{5g}1 values were consistent with lowest N₂O concentrations (Fig. 5), and reflected low NO₃ (resulting from conversion to excess N₂) rather than high N₂O. Conversely, in the slate catchment, EF_{5g}2 exceeded $EF_{5g}2$ by 15%. As such, $EF_{5g}2$ can only be considered as an accurate measurement in aquifers with limited denitrification.

5. Conclusion

The capacity of hillslope hydrologic systems to naturally mitigate groundwater and stream NO₃⁻ is site specific. There exists a hierarchy of scale whereby physical factors including agronomy, water table elevation and permeability determine the hydrogeochemical properties of the aquifer. The hydrochemical signature (DO, Eh and bacterial energy source availability) can in turn either support or suppress denitrification and subsequent NO₃ reduction. DO concentration was the dominant control, explaining 87%, 78% and 77% of the variance in groundwater RP, NO₃ and N₂O respectively in the denitrifying sandstone catchment. Information on aquifer geochemistry can therefore be used as a predictor of denitrification capacity, NO₃ and N₂O concentrations in different hydrological settings. In catchments with sufficient organic N inputs, DOC and water table elevation regulate the concentration of DO reaching the water table surface. In the saturated zone, permeability distribution controls the dissolution of solid phase electron donors, which in turn drives further DO reduction and the development of denitrifying conditions. In catchments were organic inputs are not sufficient to promote DO reduction, and residence times are too short to facilitate solid phase electron donor dissolution, denitrification progress is arrested, with correspondingly higher stream and groundwater NO₃ abundance. An entire aguifer or catchment cannot be characterised as having high or low denitrification potential. In the sandstone catchment, complete removal of NO₃ was enhanced in near stream denitrifying hot spots. In base flow dominated catchments, all groundwater entering the stream must first pass through these denitrifying zones, which has a positive effect on stream NO₃ mitigation. Near stream zones should therefore be prioritised and protected, with consideration given to the location of land drainage, which can act to bypass near stream NO₃- removal zones. Highest N₂O abundance occurred in up-gradient zones of the sandstone hillslopes, greatly exceeding atmospheric equilibrium concentrations, while also in excess of the IPCC EF_{5g} threshold. The upslope N_2O maxima were not however manifested in the sandstone streams, highlighting that groundwater discharge through near stream zones was not only paramount to NO₃ reduction, but also vital to N₂O mitigation. While stream N₂O was substantially lower than groundwater, mean values were significantly greater than atmospheric equilibrium, particularly in the sandstone catchment. The positive environmental effect of NO₃⁻ reduction was therefore concomitant with a net source of harmful greenhouse gas emissions. A review of the IPCC methodology (EF $_{\!5g}1$ and EF $_{\!5g}2$ revealed that $EF_{5g}2$ can only be used effectively in non-denitrifying catchments, while mean $EF_{5g}1$ calculations supported the downward IPCC revision of N_2O emissions from groundwater.

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Appendix A. Supplementary data

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