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The influence of bedrock hydrogeology on catchment-scale nitrate fate and transport in fractured aquifers

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Abstract
Characterising catchment scale biogeochemical processes controlling nitrate fate in groundwater constitutes a fundamental consideration when applying programmes of measures to reduce risks posed by diffuse agricultural pollutants to water quality. Combining hydrochemical analyses with nitrate isotopic data and physical hydrogeological measurements permitted characterisation of biogeochemical processes influencing nitrogen fate and transport in the groundwater in two fractured bedrock aquifers with contrasting hydrogeology but comparable nutrient loads. Hydrochemical and isotopic analyses of groundwater samples collected from moderately fractured, diffusely karstified limestone indicated nitrification controlled dissolved nitrogen fate and delivery to
aquatic receptors. By contrast nitrate concentrations in groundwater were considerably lower in a low transmissivity highly lithified sandstone and pyrite-bearing shale unit with patchy subsoil cover. Geophysical and hydrochemical investigations showed shallower intervals contained hydraulically active fractures where denitrification was reflected through lower nitrogen levels and an isotopic enrichment ratio of 1.7 between $\delta^{15}$N and $\delta^{18}$O. Study findings highlight the influence of bedrock hydrogeological conditions on aqueous nitrogen mobility. Investigation results demonstrate that bedrock conditions need to be considered when implementing catchment management plans to reduce the impact of agricultural practices on the quality of groundwater and baseflow in receiving rivers.

Nitrate isotopic signatures in the groundwater of a freely draining catchment underlain by a karstified aquifer and a poorly draining aquifer with a low transmissivity aquifer.

Key words

Groundwater  Nitrate isotopes  Nitrification  Denitrification
1 Introduction

Excess nitrate ($\text{NO}_3$) is a global environmental problem which is expected to worsen as a result of factors including the increase in human population, political stress and the development of growing economies (Erisman et al. 2011). Surplus nitrate can reflect loss of soil fertility, while its presence in water can cause human and ecological health problems (McLay et al. 2001). To address the impacts of degraded water quality on human and ecological aquatic health, the European Union (EU) introduced the Water Framework Directive (WFD) (Directive 2000/60/EC) requiring member states to achieve good water quality status in inland surface water and groundwater bodies. Effective implementation of the WFD in areas dominated by agricultural activity, while maintaining economic levels of food production, requires understanding of the fate and transport of diffuse agricultural pollutants, such as nitrate, in the wider environment.

Implementing water quality management measures requires an understanding of the factors controlling nitrogen (N) fate and transport. Biogeochemical processes and hydrogeological setting constitute significant factors controlling the fate of nitrogen in groundwater. Previous studies undertaken in various unconsolidated and bedrock aquifers (Burkart & Koplin 1993), in fractured crystalline and Upper Devonian karstified limestone (Nolan 2001) and catchments underlain by sandstone, limestone and Ordovician sediments (Jahangir et al. 2012), show that aquifers having faster travel times and higher permeability were associated with higher nitrate concentrations. In contrast, Puckett & Cowdery (2002) found that longer groundwater residence times can allow time for denitrification to occur thus lowering nitrate concentrations in a glacial outwash aquifer. However, a study undertaken in Cretaceous and Jurassic carbonate rocks and detritic sediments (Stigter et al. 2006) and another study in a catchment underlain by Pleistocene deposits (Debernardi et al. 2008) have shown that the relationship between hydrogeological setting and nitrate fate is less
clear cut. Furthermore, even within aquifers geological heterogeneity can influence contaminant transport (Puls & Paul 1997). Consequently, a better understanding of the biogeochemical processes within different hydrogeological settings is necessary to inform the development of catchment management tools and models. Information relating to processes operating in bedrock is notably scarce. This study investigates the influence of hydrogeological setting on biogeochemical processes, aiming to characterise the dominant processes influencing nitrogen levels in groundwater in two Irish catchments underlain by bedrock aquifers with contrasting (physical and geochemical) hydrogeological properties, but having comparable nutrient loads and thin to no subsoil cover over much of their area. This research considers the spatial heterogeneity of biogeochemical processes within each catchment, both across the catchment and with depth. This has been achieved through monitoring well tracer tests, and analysis of chemical and isotopic signatures of groundwater and surface water. Incorporating the knowledge gained concerning biogeochemical processes into water quality catchment management tools can prove fundamental in reducing the risk posed to groundwater and surface water receptors.

Biogeochemical processes affecting nitrate mobility include nitrification, autotrophic and heterotrophic denitrification, ammonification, anammox and dissimilatory nitrate reduction to ammonium (DNRA) (Rivett et al. 2008), and are summarised by Equations 1 to 8 in Table 1. Ammonification is the process by which ammonium is produced from organic N. This process occurs more rapidly under aerobic conditions, but can also occur within anaerobic conditions (Reddy and Patrick 1984). Nitrification is the microbial oxidation of ammonium to nitrate (Equations 1 and 2), which is considered a critical nutrient in controlling aquatic ecosystem productivity (and eutrophication). By contrast, denitrification has been identified as the most important biogeochemical process in the removal of nitrate from groundwater (Rivett et al. 2008; Korom 1992; Burgin & Hamilton 2007). The emphasis on denitrification over other nitrate removal
processes arises due to the widespread occurrence of denitrifying bacteria, perceived high nitrate removal capacity of denitrification and the relatively limited capacity for alternative processes to remove nitrate (Rivett et al. 2008; Burgin & Hamilton 2007). Denitrification rates are limited by oxygen concentrations and require the presence of N oxides (NO\textsubscript{3}, NO\textsubscript{2}, NO and N\textsubscript{2}O) and electron donors (e.g. dissolved organic carbon or pyrite) as outlined in Equations 3, 4 and 5. Other processes influencing aqueous nitrogen’s mobility in the wider environment include anammox (anaerobic ammonium oxidation), which is the oxidation of ammonium and reduction of nitrite to produce nitrogen gas (N\textsubscript{2}), as outlined in Equations 6 and 7. DNRA has been identified in groundwater using hydrochemical investigative tools (Smith et al. 1991; Jahangir et al. 2012) and is outlined in Equation 8 (Giblin et al. 2013). The presence of anaerobic conditions and a negative correlation between nitrate and ammonium in groundwater has been linked to the occurrence of DNRA (Jahangir et al. 2012).

**Table 1 Chemical equations for biogeochemical processes affecting nitrate fate**

<table>
<thead>
<tr>
<th>Process</th>
<th>Equation</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrification</td>
<td>( \text{NH}_4^+ + 2, \text{O}_2 \rightarrow \text{NO}_2^- + \text{H}_2\text{O} + \text{H}_2\text{O} + \text{H}^+ )</td>
<td>(Willey et al. 2011)</td>
</tr>
<tr>
<td></td>
<td>( \text{NO}_2^- + 0.5, \text{O}_2 \rightarrow \text{NO}_3^- )</td>
<td>(Willey et al. 2011)</td>
</tr>
<tr>
<td>Denitrification</td>
<td>( 4\text{CH}_2\text{O} + 4\text{NO}_3^- \rightarrow 2\text{N}_2 + 4\text{HCO}_3^- + \text{CO}_2 + 2\text{H}_2\text{O} )</td>
<td>(Jørgensena et al. 2004)</td>
</tr>
<tr>
<td></td>
<td>( 5\text{FeS}_2 + 14\text{NO}_3^- + 4\text{H}^+ \rightarrow 5\text{Fe}^{2+} + 7\text{N}_2 + 10\text{SO}_4^{2-} + 2\text{H}_2\text{O} )</td>
<td>(Tesoriero et al. 2000)</td>
</tr>
<tr>
<td></td>
<td>( \text{NO}_3^- + 5\text{Fe}^{2+} + 7\text{H}_2\text{O} \rightarrow 0.5\text{N}_2 + 5\text{FeOOH} + 9\text{H}^+ )</td>
<td>(Kölle et al. 1990; Ottley et al. 1997)</td>
</tr>
<tr>
<td>Anammox</td>
<td>( \text{NH}_4^+ + \text{NO}_2^- \rightarrow \text{N}_2 + 2\text{H}_2\text{O} )</td>
<td>(Mulder et al. 1995)</td>
</tr>
<tr>
<td></td>
<td>( 3\text{NO}_3^- + 5\text{NH}_4^+ \rightarrow 4\text{N}_2 + 9\text{H}_2\text{O} + 2\text{H}^+ )</td>
<td>(Kuenen 2008)</td>
</tr>
<tr>
<td>DNRA</td>
<td>( 2\text{CH}_2\text{O} + \text{NO}_3^- + \text{H}_2\text{O} \rightarrow \text{NH}_4^+ + 2\text{HCO}_3^- )</td>
<td>(Giblin et al. 2013)</td>
</tr>
</tbody>
</table>

The range of processes influencing the fate of aqueous nitrogen, along with the products generated, limit confident identification of the specific processes involved based on the results of
hydrochemical analyses alone. Additional methods, particularly when combined with appropriate hydrogeological conceptual models, can assist in better constraining the range of significant processes. Isotopic analyses of $\delta^{15}$N to $\delta^{18}$O levels in nitrate provide a useful tool for distinguishing nitrate sources and attenuation processes. The identification of nitrate sources using nitrate isotopes proves particularly useful in well oxygenated hydrogeological environments where minimal biogeochemical cycling of N occurs, allowing the source signal to persist (Savard et al. 2010). On the other hand, fractionation caused by other biogeochemical processes can alter the isotopic signatures. Previous groundwater studies have attributed an enrichment of $\delta^{15}$N to $\delta^{18}$O by a factor of between 1.3 and 2.1 to denitrification (Böttcher et al. 1990; Aravena & Robertson 1998; Fukada et al. 2003). Fractionation processes influencing ammonia prior to its ammonification to ammonium and subsequent nitrification to nitrate can have an impact on the nitrate isotopic signature. Volatilisation of ammonia results in isotopic enrichment of the residual NH$_4$ as lighter $\delta^{14}$N is released in the NH$_3$ gas (Mayer 2005). This enrichment of the residual N can be > 20 ‰ (Kendall 1998) and can generate an enriched N signature in the resulting NO$_3$.

Previous studies have used nitrate isotopes to investigate sources and fate of nitrate from field scale to catchment scale (Hiscock et al. 2011; Heffernan et al. 2012; Baily et al. 2011). However, studies have focused on processes operating in soils and subsoils, particularly coarser grained materials (Böttcher et al. 1990; Smith et al. 1991; Baily et al. 2011; Minet et al. 2012). Investigations into the fate and transport of nitrate in competent fractured rock units are far less common, particularly in those units with little to no matrix porosity, with even fewer studies examining the fate of nitrate in bedrock and the variation in attenuation with depth (Fukada et al. 2004). Furthermore many studies focus on an individual study catchment and fail to consider differences between catchments with contrasting bedrock.
Bedrock conditions in the areas considered by this study are representative of conditions encountered over much of the island of Ireland in that they consist of fractured bedrock with negligible primary porosity, overlain by variable thicknesses of subsoil, including large areas where subsoil is less than three metres thick. The bedrock in one catchment is a productive moderately karstified Dinantian (Lr. Carboniferous/Mississippian) limestone lacking a significant conduit network, while that in the other is a highly lithified Silurian sandstone (greywacke) containing beds of pyrite-bearing shale. Previous studies of contaminant transport in catchments underlain by limestone have focused on processes operating in larger conduit networks (Panno et al. 2001; Heffernan et al. 2012), or systems where denitrification is only significant where the aquifer is confined and oxygen depleted (Rivett et al. 2007; Hiscock et al. 2011). Processes operating in less fractured unconfined limestone units remain less studied. Similarly, nitrate fate and transport has been rarely studied in poorly productive (Low Productivity) bedrock units such as Silurian greywacke. Moreover, previous studies of groundwater quality have noted that hydrochemical stratification may occur with depth (Fukada et al. 2004; Böttcher et al. 1990).

2 Materials and Methods

2.1 Site description

2.1.1 Nuenna

The Nuenna Catchment in Co. Kilkenny, Ireland covers an area of 35 km² (Figure 1 and Figure 2). Annual rainfall in the catchment is 985 mm and evapotranspiration is 485 mm (O'Brien et al. 2013). The catchment experiences a cool maritime climate with year-round rainfall and temperatures that rarely drop below freezing. Effective rainfall is greatest in the winter period from late October to early March when evapotranspiration is at a minimum. Climate conditions during the study period were typical for the area investigated.
Teagasc soil and subsoil maps note well drained brown earths and grey brown podzolic soils overlie subsoils consisting of moderately permeable limestone tills, present on the valley slopes, and an alluvial gravel unit (GSI 2012). The alluvial gravel is 8 m thick at the NU2 monitoring wells location (Figure 1 and Figure 2) and occurs in the vicinity of the river from the mid catchment to the catchment outlet.

The groundwater vulnerability maps derived by the Geological Survey of Ireland (GSI) show that approximately 65% of the aquifer is classified as having Extreme vulnerability, with less than 3 m subsoil cover, or X-Extreme vulnerability, with less than 1 m subsoil cover and where rock is at or near surface, or karst is present (Figure 1). The remaining 35% of the aquifer is classified as high vulnerability, with a subsoil thickness of up to 10 m, occurring predominantly in lower lying areas, including groundwater discharge zones. The Nuenna Catchment is underlain by pure bedded Dinantian (Lower Carboniferous or Mississippian) limestone. The groundwater vulnerability maps suggest that over the majority of the Nuenna Catchment diffuse recharge passes through variable thickness of soil and subsoil before entering the bedrock.

Focused recharge can occur in karst features, such as swallow holes (Figure 2). During diffuse infiltration, recharge flows through the transition zone prior to entering the diffuse karst network. This occurs particularly in areas of higher elevation, approaching the catchment divide, where the limestone is overlain by poorly-productive Namurian (Lr. Pennsylvanian) sandstones and pyrite-bearing shales (GSI 2012). Further groundwater in the diffuse karst network drains into the conduits which ultimately discharge to the Nuenna River though a series of springs and, more diffusely, through the alluvial gravel unit in the lower reaches of the catchment. The hydrogeological properties of the rock matrix play a critical role influencing the rate of delivery to the conduit system (Orr 2014).
Transmissivity values, determined from seven pumping tests and nine spring recession events, range from 0.3 m²/day, within the bedrock matrix, to up to 750 m²/day, within the conduit discharge zones, reflecting high levels of hydraulic conductivity heterogeneity (Cawley 1990; Orr 2014). Although no hydrogeological data exists for the Namurian shale unit in the Nuenna Catchment, comparison with other poorly productive Irish aquifers suggests hydrogeological processes in the area are dominated by surface and near-surface runoff (Archbold et al. 2016). This has been corroborated by field observations during hydrologically energetic periods.

Two monitoring well clusters, NU1 and NU2 (Figure 1 and Figure 2), were used for water level and water quality monitoring in the Nuenna catchment. The monitoring wells’ open sections lie within (a) the subsoil, (b) the interval of elevated physical weathering occurring in the vicinity of the bedrock/overburden interface (transition zone), which may be locally absent, (c) the shallower bedrock, and (d) the deeper limestone bedrock (Table 2). The Nuenna River acts as the principal surface water body in the catchment. It has a baseflow index of > 0.87 reflecting the importance of groundwater for stabilising flow (O’Brien et al. 2013). Baseflow at the catchment outlet was determined as being between approximately 0.2 and 0.4 m³/s (Orr 2014).

Landuse across the Nuenna Catchment consists of >70 % pasture, 14 % arable and 8 % woodland, with farmyards, rural settlement and unspecified landuse occupying the remaining land (McAleer 2011). Catchment wide average diffuse N loadings were estimated to be 152 kg N/ha/yr from inorganic fertiliser (52%) and 140 kg N/ha/yr from livestock manure (48%) (Orr 2014). A land use survey of the catchment indicated greater loading of inorganic fertiliser occurred on the valley floor associated with arable land use, while organic N loading occurred in the more elevated southern and northern parts of the catchment as a result of higher livestock density (McAleer 2011).
Figure 1 Catchment locations, groundwater vulnerability (GSI 2012 and Meredith 2010) and monitoring well layout in the Glen Burn and Nuenna catchments

Figure 2 Simplified conceptual hydrogeological models of the Nuenna (left) and Glen Burn (right) catchments. Arrows show flow direction and arrow weight reflects the flow volume. Not to scale.

2.1.2 Glen Burn

The Glen Burn Catchment is located in Co. Down, Northern Ireland and covers an area of 5 km² (Figure 1 and Figure 2). Annual rainfall in the catchment is 843 mm and evapotranspiration is 460 mm (O’Brien et al. 2013). The catchment experiences a cool maritime climate similar to the Nuenna
with temperatures rarely dropping below freezing. Climate conditions during the study period were
typical for the study area investigated.

Where they occur, soils consist of a thin coverage of low permeability rankers, surface water gleys
and groundwater gleys. Localised mounds of glacial till (drumlins) dominate higher elevations, while
very thin glacial till subsoils (less than 1m thick) and rock outcrops occur in areas of lower elevation.
A thin covering of alluvium occurs locally in the vicinity of the river. Figure 1 shows the majority of
the catchment is classified as extreme vulnerability due to the thin subsoil cover, while areas overlain
by drumlins are classified as having moderate to low vulnerability (Meredith 2010).

The Glen Burn Catchment is underlain by greywacke sandstone and shale belonging to the Silurian
Gala Group. Transmissivity values, determined from pumping tests, range from 0.18 m²/day in the
shallow fissured bedrock to 0.07 m²/day in deeper more composite bedrock (Comte et al. 2012).
The aquifer is classified as “Limited Productivity with Fracture Flow” by the Geological Survey of
Northern Ireland (McConvey 2012). Such aquifers typically have low yields, limited regional flow,
shallow local flow paths and few transmissivity values >500 m²/day, some transmissivity values
>50 m²/day and most transmissivity values <50 m²/day (McConvey 2012; GSI 2016). Groundwater
flow occurs predominantly through fractures, with negligible matrix flow. The transmissivity data
suggest that near surface and shallow groundwater pathways act as the dominant hydrological routes
for delivering groundwater to the Glen Burn River. Four monitoring well clusters, located in the
lower Glen Burn catchment, occur between 30 m and 350 m from the river, and have monitoring
points screened against the shallow bedrock and deep bedrock (Figure 1, Figure 1 and Table 2). A
fifth cluster is located in the alluvium. The river has a low baseflow index (BFI) of <0.13 (O’Brien et
al. 2013). Baseflow at the catchment outlet is between approximately 0.002 and 0.008 m³/s (Orr
2014).
Landuse across the catchment consists of 82% pasture, 10% arable and 5% woodland, with farmyards and rural settlement occupying the remaining land (Meredith 2010). Catchment-wide average N loadings were estimated to be 125 kg N/ha/yr from inorganic fertiliser (37%) and 213 kg N/ha/yr from livestock manure (63%).

2.2 Methods

Both physical and chemical methods were carried out in order to characterise the dominant hydrogeological processes influencing nitrogen levels in groundwater within the two catchments and to consider the spatial heterogeneity on biogeochemical processes with depth in both catchments.

2.2.1 Artificial tracer tests

Conventional methods of identifying hydraulically active fractures in bedrock monitoring wells require significant levels of active or induced flow. The threshold levels for these methods often lies below the rates flowing through fractures in poorly productive bedrock units. Artificial tracer testing provides an alternative means of investigating flow regimes in these low-flowing systems. In the current study uranine (sodium fluorescein) tracer tests were carried out to identify hydraulically active fractures in bedrock. A flexible pipe connected to a 50 m long tube was lowered to the base of the monitoring well and the uranine tracer was then poured into the tube to displace monitoring well water as the tube was raised to the surface at an even rate. Following tracer injection, the tubing was lowered and raised to homogenise the tracer within the monitoring well.

Following mixing, a portable GGUN fluorometer FL30 (Aliba, Neuchâtel) was lowered slowly into the monitoring well. This took triplicate measurements of tracer concentration at 0.5 m intervals to a maximum depth of 60 m. The process was repeated three to four times, up to 6 days after the injection, or until the uranine concentrations had fallen below the fluorometer’s detection limit.
Modelling the results of observed tracer concentrations permitted hydraulically active intervals to be identified and reconciled with the results of geophysical monitoring well logging (Nitsche 2014).

### 2.2.2 Water Chemistry Sampling

Groundwater samples in the Nuenna catchment were collected during 20 groundwater sampling rounds between July 2010 and November 2012, while eleven groundwater sampling rounds were carried out in the Glen Burn catchment between February 2011 and November 2012. The sampling strategy for both catchments followed the British Standard BS ISO 5667-11:2009 and BS 6068-6.11:2009 protocols. Whale Mini-Purgers were used to extract groundwater from the monitoring wells with lower yields (all Glen Burn monitoring wells, NU1 Transition and NU2 Shallow), Whale Mega-Purgers were used in higher yielding monitoring wells (NU2 Deep and Subsoil); a Grundfos MP1 pump was used in NU1 Deep as the depth to groundwater exceeded the pumping head of the Whale Purgers. Whole monitoring well sampling regimes can fail to account for vertical flows (McMillan et al. 2014) and changes in biogeochemical processes with depth. Therefore, depth specific samples were taken on one additional sampling round in each monitoring well using a pneumatic double-packer system and Solinst Integra Bladder Low Flow Pump from intervals identified as hydraulically active by artificial tracer testing to investigate potential mixing of waters with contrasting chemistries.

Monitoring well construction details for both catchments are summarised in Table 2. During sampling, the wells were pumped until the groundwater’s field hydrochemical parameters (dissolved oxygen (DO), pH, electrical conductivity (EC), oxidation reduction potential (ORP) and temperature), measured using a TPS 90FL-MV water quality meter and flow through cell, stabilised. Water chemistry samples were collected in HDPE plastic bottles and stored within ice during transport to the laboratory following the protocol set out by the British Standard BS EN ISO 5667-
Table 2 Monitoring well construction details in the Nuenna and Glen Burn catchments

<table>
<thead>
<tr>
<th>Monitoring well ID</th>
<th>Depth of monitoring well (m)</th>
<th>GW body type</th>
<th>Monitoring well type</th>
<th>Depth of open Section/screened interval (m)</th>
<th>Approx. distance from river (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NU1 Deep</td>
<td>74.4</td>
<td>Deep</td>
<td>152 mm open hole</td>
<td>36 to 74.4</td>
<td>620</td>
</tr>
<tr>
<td>NU1 Transition</td>
<td>7.3</td>
<td>Transition</td>
<td>152 mm PVC screen with 2 mm slots</td>
<td>3.3 to 7.16</td>
<td>625</td>
</tr>
<tr>
<td>NU2 Deep</td>
<td>79.5</td>
<td>Deep</td>
<td>152 mm open hole</td>
<td>33.8 to 79.5</td>
<td>20</td>
</tr>
<tr>
<td>NU2 Shallow</td>
<td>27.4</td>
<td>Shallow</td>
<td>152 mm PVC screen with 2 mm slots</td>
<td>11.5 to 26.32</td>
<td>22</td>
</tr>
<tr>
<td>NU2 Subsoil</td>
<td>9.3</td>
<td>Subsoil</td>
<td>152 mm PVC screen with 2 mm slots</td>
<td>5.5 to 8.38</td>
<td>24</td>
</tr>
<tr>
<td>Glen Burn</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2.2</td>
<td>10.2</td>
<td>Shallow</td>
<td>152 mm open hole</td>
<td>4.2 to 10.2</td>
<td>175</td>
</tr>
<tr>
<td>3.1</td>
<td>19.58</td>
<td>Deep</td>
<td>50 mm PVC screen</td>
<td>15 to 21</td>
<td>350</td>
</tr>
<tr>
<td>4.1</td>
<td>34.33</td>
<td>Shallow and deep</td>
<td>152 mm open hole</td>
<td>3.0 to 31.8</td>
<td>50</td>
</tr>
<tr>
<td>4.2</td>
<td>5.7</td>
<td>Shallow</td>
<td>152 mm open hole</td>
<td>1.4 to 5.7</td>
<td>50</td>
</tr>
<tr>
<td>5.1</td>
<td>3.5</td>
<td>Alluvium</td>
<td>152 mm open hole</td>
<td>3.5 to 7.5</td>
<td>30</td>
</tr>
<tr>
<td>5.2</td>
<td>2.5</td>
<td>Alluvium</td>
<td>50 mm PVC screen</td>
<td>1.5 to 2.5</td>
<td>30</td>
</tr>
<tr>
<td>6.1</td>
<td>60</td>
<td>Shallow and deep</td>
<td>152 mm open hole</td>
<td>8.7 to 60</td>
<td>40</td>
</tr>
<tr>
<td>6.2</td>
<td>9</td>
<td>Shallow</td>
<td>152 mm open hole</td>
<td>4.9 to 9</td>
<td>40</td>
</tr>
</tbody>
</table>

2.2.3 Laboratory chemical analyses

Ion chromatography analyses of target anions (NO₂⁻, NO₃⁻, NH₄⁺, SO₄²⁻, Cl) were carried out within 48 hours of collection using a Dionex® DX-500 and ICS-2000. Prior to analysis the samples were filtered through 0.45 μm sterile Millex® MCE membrane filters. Alkalinity was measured using a Hach® digital titrator model 16900. Ferrous iron (Fe²⁺) analysis was carried out using the Hach Lange® DR 2800 Spectrophotometer. Samples collected for dissolved organic carbon analysis were
filtered through 0.45 µm sterile Millex MCE membrane filters and then analysed using a UV
promoted persulfate oxidation method on a TOC analyser (Sievers® 5310C Laboratory TOC
Analyzer).

Groundwater samples used for nitrate isotopic analyses were collected during six of the sampling
rounds in both study catchments including the packer testing. Samples were filtered on site directly
into HDPE containers before freezing. Isotopic analysis for $\delta^{15}$N-NO$_3$ and $\delta^{18}$O-NO$_3$ was carried
out at the Stable Isotope facility in UC Davis, California using the bacterial denitrification assay
method (Casciotti et al. 2002).

3 Results and Discussion

3.1 The Nuenna Catchment

Water level monitoring reveals that shallow bedrock (approximately <25 m BGL) in the Nuenna
catchment is generally unsaturated, although in the areas of lower elevation, close to the Nuenna
River, it may be saturated. Slower flow in the shallow bedrock at NU2 Shallow is reflected by the
lower tracer dilution rates observed over a similar time frame in NU2 Shallow (>80 % over 123
hours), compared to NU2 Deep (below detection limit after 141 hours). The levels of uranine
dilution observed at NU2 Shallow reflect limited levels of groundwater throughflow. However,
slightly lower tracer concentrations observed with depth suggest upward flowing groundwater,
consistent with upwelling discharge close to the river. Therefore, although the transmissivity values
in the diffuse karst of the shallow and deep bedrock units at NU2 lie within the same order of
magnitude, between 0.3 m$^2$/day and 1.5 m$^2$/day (Orr 2014), flow through the shallower bedrock
unit may be slower than in the deep bedrock, reflecting the hydrogeological heterogeneity of the
diffuse karst bedrock.
Results of tracer tests completed at NU1 showed a significant dilution in uranine concentration, suggesting inflow at approximately 35.5 m below ground level (bgl) in NU1 Deep and indicating the presence of a hydraulically active fracture(s). This interval corresponds with a fracture set identified in the geophysical caliper and HiRAT logging (Moran & Crean 2010). Moreover, tracer test results suggest that additional hydraulically active fracture sets may be present at approximately 44-45 m below ground level (bgl), which also corroborate findings of caliper and high resolution acoustic televiewer (HiRAT) geophysical logs. However as dilution is less significant, it is likely that this fracture set was less hydraulically active than the set at 35.5 m at the time of testing.

The responses of uranine tracer test results highlights the heterogeneity of fracture flow within the diffuse fracture network of the karst bedrock and suggests a component of vertical flow within the monitoring well in the form of upwelling close to the river. Such vertical flow may generate sampling bias (McMillan et al. 2014) and therefore point to the need for depth specific packer testing. The hydraulic conductivity of the bedrock matrix at NU2 Deep and Shallow and point to hydraulically active fractures which act as the dominant transport groundwater pathway; processes operating within these fractures therefore are likely to dominate groundwater nitrate concentrations.

3.1.1 Hydrochemistry

Hydrochemical conditions in the two monitoring well clusters in the Nuenna catchment vary both spatially (with depth and laterally) and temporally. Lower median nitrate and DO concentrations are evident in the shallower transition zone groundwater in comparison to the deep groundwater (Table 3 and Figure 3). Occasional relatively low concentrations of nitrate (<9 mg/l) are likely to be as a result of intermittent attenuating reactions, rather than dilution, as they did not coincide with lower chloride concentrations and the NO$_3$/Cl ratio was <0.7. The non-reactive nature of chloride (Davis et al. 1998) provides a basis for evaluating the transformation of reactive species. Variations in
absolute concentrations can confuse evaluation due to differential loading. The use of ratios of reactive species to chloride provides a means of reducing this influence. On the occasion when the lowest recorded concentrations of nitrate (5 mg/l) were detected, anaerobic conditions were noted through the absence of DO (<1 mg/l). This coincided with elevated HCO₃ concentrations (512 mg/l) suggesting heterotrophic denitrification (Equation 3, Jørgensen et al., 2004). Moreover, analysis of field hydrochemical monitoring data demonstrates that redox conditions in the transition zone groundwater fluctuate. This is further evidence that the transition zone can occasionally provide suitable conditions for denitrification to occur. However, as these conditions are sporadic, the potential for nitrate removal in the transition groundwater has very limited capacity to affect nitrate concentrations at a catchment scale.

Overall, higher median nitrate concentrations were observed in the deep groundwater, compared to the samples collected from monitoring wells screened in the transition zone (Table 3 and Figure 3). This suggests that the recharge containing higher concentrations of nitrate bypassed upper layers and entered the deep groundwater body through localised (non-uniform) entry points such as swallow holes or areas where the subsoil is thin or absent (Table 3). Tracer testing carried out in swallow holes and springs within the catchment has demonstrated that focused recharge can enter the limestone aquifer in areas of higher elevation close to the contact with the overlying sandstone and shale units (Hogan 2011); these correspond to areas of higher organic fertiliser application.

In the case of the deep groundwater samples there was a contrast in the median nitrate concentrations between NU1 Deep and NU2 Deep (Table 3 and Figure 3). Slightly higher nitrate concentrations observed in NU1 Deep compared to NU2 Deep are suspected to reflect some heterogeneity in nitrate loading as well as dilution with less impacted water containing lower
concentrations of N closer to the river. The significance of dilution is further highlighted by similar NO$_3$/Cl ratios (Figure 3).

Hydrochemical analyses revealed the lowest median nitrate, DO and ORP values were detected in the shallow groundwater pathway at NU2 Shallow (Table 2). The anaerobic conditions associated with NU2 Shallow create a distinctly different hydrochemical signature compared to the deeper units. DO and ORP conditions observed compare to those suggested in literature as being suitable for denitrification (Jahangir et al. 2012; Thomasson et al. 1991). However, DOC concentrations in the shallow groundwater at NU2 Shallow are relatively low (Table 3) compared to the other monitoring well locations. DOC concentrations at NU2 Shallow are also generally below the suggested threshold for denitrification of 2 mg/l DOC (Thomasson et al. 1991), with the exception of a single spike of 17.3 mg/l DOC. A septic tank connected to a restaurant/pub discharges hydroxide based cleaning agents into the groundwater upgradient of the NU2 monitoring well cluster. The hydroxide results in a higher pH and low HCO$_3$ concentrations at NU2 Shallow monitoring well compared to the other monitoring wells. Therefore, the relationship between nitrate and HCO$_3$ cannot be considered useful regarding heterotrophic denitrification at NU2 Shallow. However, on the basis of the generally low DOC levels it is unlikely that heterotrophic denitrification occurred within the shallow groundwater. Autotrophic denitrification is also unlikely in the shallow groundwater as Fe$^{2+}$ concentrations were below the detection limit (10 µg/l).

Hydrochemical analyses for samples collected at the NU2 cluster suggest that DNRA (see Equation 8) might account for nitrate removal in the shallow groundwater as median ammonium concentrations are relatively high. DNRA is favoured when nitrate availability is the principal limiting factor, while denitrification is favoured when carbon availability is limiting (Korom 1992). However, as both nitrate and DOC concentrations are relatively low in the shallow groundwater it is
unclear which may be limiting. It is also unclear how much the nitrification or denitrification is affected by the high pH in the shallow groundwater. Denitrification can operate at a pH of up to 10.2 (Thomasson et al. 1991) with the optimum pH for nitrification lying between 7 and 8 (Tarre & Green 2004). Additionally, anammox may be occurring as it can take place where nitrate and ammonium coexist and carbon is limited (Burgin & Hamilton 2007). However, it is most likely that elevated NH₄ concentrations is entering the shallow groundwater from the septic tank up-gradient of the NU2 cluster, and due to low DO conditions nitrification is inhibited, allowing the NH₄ to persist.

The similar median nitrate concentrations observed in the alluvial gravel subsoil groundwater and the river water samples are consistent with a strong hydraulic connection between the river and the alluvium. This is supported by in-stream artificial tracer testing which suggested considerable exchange of water between the alluvial gravel aquifer and the river (Hogan 2011). The alluvial gravel aquifer extends from the middle of the catchment to the catchment outlet along the valley floor adjacent to the river. The similar nitrate concentrations in the river and the alluvial deposits suggest that the alluvium may act as a temporary reservoir for nitrate and has limited capacity for nitrate removal.

The results of water quality analyses suggest that very localised processes have a strong influence on the water quality in the NU2 Shallow monitoring well. However, it is unlikely that these conditions are significant throughout the catchment as the nitrate and major ion concentrations of the water within the river at baseflow conditions resemble those observed in the deeper monitoring wells, such as NU1 Deep and NU2 Deep. This is further corroborated by the results of isotopic analyses (below) and highlights the dominance of groundwater delivered via deeper pathways which controls river baseflow and overprints localised processes.
Table 3 Median, maximum and minimum values for hydrochemical parameters in the Glen Burn and Nuenna groundwater pathways and respective surface water bodies.

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\(^1\) Selected samples were analysed for DOC, Fe\(^{2+}\) and nitrate isotopes
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3.1.2 Nitrate Isotopes and nitrate stratification

The artificial tracer single well dilution tests permitted identification of target zones at depth for packer testing to further investigate changes in nitrate fate and transport with depth in the aquifers at a higher resolution. In the Nuenna aquifer the nitrate concentrations are highest in the subsoil and lowest in the shallow groundwater before increasing again in the deep groundwater. Higher resolution depth sampling (Figure 4) showed that nitrate in the deep groundwater does not vary significantly at depth, indicating that the unit is well mixed in the deeper parts of the aquifer and that vertical flow within the monitoring well has negligible influence on the nitrate concentrations. Despite the identification of a hydraulically active fracture at 35.5 mbgl in NU1 Deep and evidence of greater potential for dilution, there was no significant change in nitrate concentrations with depth in the intervals sampled in NU1 Deep. Moreover, relatively little change in NO₃/Cl ratios or nitrate isotopic signature suggest good mixing with depth in the deeper part of the aquifer and that biogeochemical reactions are not a significant factor influencing nitrate fate once the nitrate enters deeper bedrock.

The nitrate isotopic signature of the groundwater and surface water in the Nuenna catchment clusters closely within the ranges for nitrate derived from the nitrification of ammonium fertilisers, soil nitrogen or septic tank waste and manure according to the ranges suggested by Kendall (1998), as shown in Figure 5. The collective groundwater isotopic signatures display an insignificant depletion in δ¹⁵N values with declining nitrate concentration, indicating that at catchment scale denitrification is not a significant process. Furthermore, an enrichment ratio of -2.13 between δ¹⁸O and δ¹⁵N is inconsistent with denitrification.

Nitrate in the transition zone groundwater samples displayed a larger range of variation in δ¹⁵N than in the deeper groundwater samples although the range of δ¹⁸O values is considerably smaller (Table
This finding suggests that, prior to nitrification, volatilisation may have caused the enrichment of the $\delta^{15}$N as ammonia. This corroborates the interpretation based on the hydrochemical data which suggest that denitrification has negligible influence on nitrate concentrations in the groundwater at the catchment scale.

Although nitrate concentrations are lowest in the shallow groundwater, the nitrate isotopic signature is similar to the nitrate isotope signature in the subsoil groundwater, with one exception, which occurred when the lowest measureable concentration of nitrate was detected (0.67 mg/l) in the subsoil groundwater; this displayed depleted isotopic values ($\delta^{15}$N 3.29 ‰ and $\delta^{18}$O -0.67 ‰).

Depleted nitrate isotopic values are indicative of partial nitrification (Kendall & Aravena 2000; Lehmann et al. 2007) and are consistent with the elevated ammonium concentrations (0.4 mg/l) and nitrite concentrations associated with NU2 Shallow. Although nitrite concentrations were below the limit of detection on this occasion (0.1 mg/l), when the lowest measurable nitrate was detected, on other occasions nitrite concentrations in the shallow groundwater reached 2.66 mg/l. This concentration in groundwater far exceeds the Environmental Objectives (Groundwater) Regulations 2010 (European Communities 2010) threshold of 0.375 mg/l and may reflect partial nitrification.

Hydrochemical and isotopic findings suggest that it is unlikely that widespread denitrification occurs across the Nuenna but localised partial nitrification may be intermittently occurring in the shallow groundwater with limited impact on catchment surface water quality.

The groundwater nitrate isotopic signature in the subsoil falls within a similar range to that in the deep groundwater and resembles the river water’s nitrate isotopic signature. Consistent with findings drawn from the hydrochemical analyses, this suggests that nitrification is the dominant process impacting the nitrate in the subsoil and surface water and good hydraulic connectivity exists between the alluvial gravel and the surface water.
Figure 3 Box and whisker plots of NO$_3$, NH$_4$, NO$_3$/Cl, DO, HCO$_3$, DOC, SO$_4$ and Fe$^{2+}$ concentrations in groundwater pathways and rivers in the Nuenna and Glen Burn catchments.
3.2 The Glen Burn Catchment

Uranine (single well) tracer tests carried out in the deep bedrock monitoring well BH 6.1 in the Glen Burn Catchment identified a hydraulically active fracture zone at 14 mbgl and further active fractures at 16, 26, 31, 38 and 52 mbgl (Nitsche 2014), as shown in Figure 6. Flow velocities were too low and/or uniform in the shallow and deep bedrock at BH 4.1, and in the shallow bedrock monitoring well BH 6.2, to identify the dominant hydraulically active fracture sets. Overall however, pumping tests indicate that higher hydraulic conductivity occurs in the shallow bedrock compared to the deep (Comte et al. 2012). This, in turn, suggests that the shallower intervals of the bedrock are more...
important in terms of groundwater transport, despite the identification of hydraulically active fractures at depth; these deeper bedrock fractures appear to have limited connectivity.

3.2.1 Hydrochemistry

Although N loading intensity in the Glen Burn, is similar to that in the Nuenna catchment, nitrate concentrations in the Glen Burn bedrock groundwater are considerably lower (Figure 3 and Table 3), despite comparably thin soil covering (Figure 1). Within the rock more nitrate-rich groundwater was encountered flowing along the shallow groundwater pathway than in the deeper groundwater, although this was found to vary spatially between monitoring well locations across the catchment. Cluster BH6 is located on the south eastern side of a drumlin which is cultivated for tillage, and receives synthetic fertiliser, while BH4 is located on the eastern side of the same drumlin on the boundary between a tillage field and pasture land and is likely to receive both synthetic and organic nitrogen. The area surrounding the drumlin contains a thin (<1m) to absent covering of soil directly over rock. The difference in nitrate concentration in samples collected from shallow monitoring wells at both locations may be due to the difference in landuse practices and to greater synthetic fertiliser application associated with tillage. Thicker (locally derived) till at BH 6.2 (shallow bedrock) (4.90 m) compared to BH 4.2 (deep bedrock) (0.4 m) can contribute to longer travel times (Fenton et al. 2009). However vertical travel time has been shown to have no correlation with groundwater nitrate concentrations in some Irish catchments (Baily et al. 2011). This is corroborated by higher nitrate concentrations at BH 6.2 compared to BH 4.2 which indicates that preferential flow paths allow relatively rapid transport of contaminants to the shallow groundwater with lower removal capacity or that the higher concentrations occurred as a result of lateral groundwater flow from recharge entering upgradient.
Anaerobic conditions and the presence of electron donors, such as organic carbon, suggest that denitrification occurs in groundwater collected from shallow groundwater monitoring points. Bedrock geochemistry has been shown to have an influence on nutrient mobility in Irish catchments where the presence of pyrite may inhibit nutrient mobility in poorly productive units (Donohue et al. 2015). The higher median $\text{Fe}^{2+}$ concentrations in BH 4.2 (shallow bedrock), compared to BH 6.2 (shallow bedrock), indicate a greater capacity for autotrophic denitrification in BH 4.2 (Equation 4). $\text{Fe}^{2+}$ concentrations of up to 1200 ug/l and 1080 ug/l have been detected in BH 4.2 and BH 6.2 respectively, supporting evidence for occasional suitable autotrophic conditions. Pyrite may acts as a potential source of $\text{Fe}^{2+}$ in the denitrification process, which also results in the increase in $\text{SO}_4^{2-}$ (Equation 4). However this is not consistent with reaction stoichiometry and suggests that other factors are influencing $\text{Fe}^{2+}$ and $\text{SO}_4^{2-}$ concentrations in the groundwater. The greywacke bedrock has a high clay content (Comte et al. 2012) which may be a source of $\text{Fe}^{2+}$ as outlined in Equation 5. Other non-pyrite derived Fe sources include detrital silicates and magnetite ($\text{Fe}_3\text{O}_4$) (Davidson et al. 2003.) Alternative sources of $\text{SO}_4$, including atmospheric deposition, fertilisers and the decomposition of organic matter, may contribute to the $\text{SO}_4$ concentrations in the groundwater. $\text{SO}_4$ can be influenced by evapotranspiration, while $\text{Fe}^{2+}$ cannot, however this is unlikely to account for the difference in concentrations as the difference in evapotranspiration is negligible (5%).

Median DOC concentrations of 5.44 mg/l in BH 6.2 compared to 1.87 mg/l in BH 4.2 suggest that heterotrophic denitrification is favoured in BH 6.2 over autotrophic denitrification. Consequently, lower nitrate concentrations in BH 4.2 may suggest autotrophic denitrification is more efficient at reducing nitrate. However, as both locations show potential for denitrification, the difference in nitrate is more likely to be a result of variable loadings patterns related to the pressures or the presence of preferential pathways through the overlying till rather than to the type of denitrification.
Nitrate detected in the deep groundwater samples collected from the Glen Burn catchment were consistently less than 2.6 mg/l; this is considerably lower than in the shallow groundwater. Lower transmissivity values have been calculated in the deeper bedrock than in the shallow bedrock (Comte et al. 2012). This may indicate that lower nitrate concentrations in the deep groundwater occur due to reduced flow rates of younger water containing more elevated levels of nitrate flowing to depth, i.e. having a longer travel time. Nonetheless, the presence of detectable levels of nitrate in the deep groundwater points to a recent recharge flow component with the potential to impact groundwater quality. This is consistent with groundwater age dating carried out in the Co. Down greywacke bedrock sampled nearby, which shows that the groundwater contains a measured modern (post 1953) component (Cronin 2000). Lower Cl concentrations in the deeper part of the bedrock, compared to the shallow groundwater pathways, suggest dilution with groundwater containing lower Cl and nitrate concentrations. However, the contrast in Cl concentrations at deeper and shallower depths is small (ranging between 3 and 38 % with an average reduction of 14 %), compared to the contrast in nitrate concentration (ranging between 3 and 100 % with an average reduction of 86 %), which is frequently below detection limit in the deep groundwater. This highlights that nitrate within the deep groundwater experiences significant additional attenuation through biogeochemical processes; this is consistent with time dependant reactions leading to greater nitrate loss in groundwater with longer residence times (Korom et al. 2012).

Lower ORP and DO conditions in the deep bedrock reflect more reducing conditions than in the shallow bedrock (Figure 3 and Table 3). This further implies that conditions in the deep bedrock are more suited to denitrification. DOC concentrations fluctuate spatially in the deep groundwater. Higher median DOC concentrations are evident in BH 3.1 (deep bedrock) compared to BH 6.1 (deep bedrock). This also coincides with slightly higher HCO₃ concentrations in BH 3.1 compared to BH 6.1. These findings suggest that conditions in the aquifer up-gradient of BH 3.1 have a greater
capacity for heterotrophic denitrification than at BH 6.1 (nitrate was only detected in BH 3.1 on a single occasion at a concentration of 1.04 mg/l.) Overall hydrochemical evidence therefore points to the significant capacity for denitrification in the deep groundwater.

Higher median Fe$^{2+}$ concentrations in BH 6.1 (deep bedrock) compared to BH 3.1 (deep bedrock) suggest greater autotrophic denitrification capacity in BH 6.1 compared to BH 3.1. Furthermore, although median Fe$^{2+}$ concentrations are relatively low, concentrations were higher in the deep groundwater, such as at BH 6.1, compared to the shallow groundwater at BH 6.2; which was below the detection limit. Therefore, heterotrophic denitrification is suspected to be more significant in BH 3.1, while autotrophic denitrification is likely to be more important in BH 6.1. Despite the evidence for autotrophic denitrification at BH 6.1, median DOC concentrations at BH 6.1 are also sufficient to promote heterotrophic denitrification. The considerable fluctuation in Fe$^{2+}$ and DOC concentrations suggest that heterotrophic and autotrophic denitrification may both occur in the vicinity of BH 6.1.

The highest Fe$^{2+}$ concentrations encountered at the Glen Burn site are present in BH 4.1, which intersects both the shallow and deep bedrock. Fe$^{2+}$ concentrations are lower in the shallower monitoring well, BH 4.2, and increase with depth. Similar to BH 6.1, low nitrate concentrations and elevated Fe$^{2+}$ suggest that autotrophic denitrification is occurring. Equation 4 indicates that pyrite mediated denitrification should produce a SO$_4^{2-}$/Fe$^{2+}$ molar ratio of 2:1. SO$_4^{2-}$ is relatively low in both BH 4 monitoring wells and the SO$_4^{2-}$/Fe$^{2+}$ molar ratio is less than 2, indicating that pyrite oxidation is not the dominant source of electron donors for denitrification or there are other processes affecting the Fe$^{2+}$ and SO$_4^{2-}$ in the groundwater as previously discussed. The presence of elevated DOC concentrations suggests that heterotrophic denitrification may also be playing an important role in nitrate reduction.
Hydrochemical data show that the fate of nitrate in the groundwater sampled from the alluvium adjacent to the Glen Burn River differs from the fate of the nitrate in the competent bedrock groundwater. The alluvium groundwater contains higher median nitrate concentrations than the deep and shallow groundwater. Aerobic conditions in the alluvium are believed to inhibit denitrification, while concentrations of ammonium lie below the limit of detection, suggesting that ammonium undergoes nitrification as a result of the aerobic conditions.

Overall, the Glen Burn findings show that the dominant groundwater pathway is the shallow groundwater pathway for delivering nitrate to aquatic receptors. Denitrification is likely to be occurring in the groundwater resulting in lower nitrate concentrations with depth. Water quality data suggest that both autotrophic and heterotrophic denitrification occurs, yet varies spatially across the site according to available electron donors.

### 3.2.2 Nitrate Isotopes and nitrate stratification

Nitrate isotope signatures observed in surface water samples collected from the Glen Burn River, fall within the ranges for ammonium fertiliser and for manure and septic tank waste (Figure 5); these sources of nitrate are widely encountered upstream of the sampling points (Meredith 2010) and suggests that the nitrate in surface water has undergone less signature-altering attenuation than the nitrate in the groundwater. Consequently, in a catchment such as the Glen Burn, which displays evidence of considerable subsurface biogeochemical alteration, the surface water nitrate signature may better reflect the original sources of nitrate than the groundwater. Moreover, comparison of surface water isotopic ratios with those from groundwater indicates that groundwater contributes relatively little to the river's discharge; this is consistent with the relatively low BFI calculated for the river.
Overall nitrate concentration decreases with depth in the Glen Burn aquifer, with a considerable reduction between 7 mbgl and 14 mbgl which also corresponds to a reduction in the NO$_3$/Cl ratio (Figure 6). The decrease in NO$_3$/Cl ratio is suspected to indicate that there is dilution occurring with older water containing lower chloride and nitrate concentrations. However considering the reduction in redox this would suggest that there may be denitrifying conditions. This is supported by the significant enrichment of both the $\delta^{15}$N and $\delta^{18}$O which indicates denitrification in the bedrock.
groundwater. More enriched nitrate isotopic values in the deep groundwater compared to the shallow groundwater suggest that the rate of nitrate removal through denitrification continues at depth as it infiltrates downwards. This is supported by a general trend across the groundwater samples showing lower groundwater nitrate concentrations yet containing more enriched δ¹⁵N. Significantly, these values display an enrichment ratio of 1.7 between δ¹⁵N and δ¹⁸O (Figure 5) which is within the enrichment ratio range of between 1.3 and 2.1 attributed to denitrification (Böttcher et al. 1990; Aravena & Robertson 1998; Fukada et al. 2003).

Figure 6 Variation of NO₃, NH₄, and ORP concentrations and NO₃ isotope (δ¹⁵N and δ¹⁸O) values with depth in BH6 cluster, sampled using a packer system and low flow pump.
Critically, the findings of this study suggest that denitrification processes in the Glen Burn catchment are not restricted to the subsoil as may be the case in the Nuenna catchment, but denitrification is also occurring in the fractured bedrock aquifer. The decrease in nitrate concentration with depth corroborates the findings of the uranine single well tracer tests, which show that the most hydraulically active fractures are in the upper part of the aquifer thus limiting the vertical movement of nitrate into the deeper aquifer. Furthermore the isotopic analysis shows that nitrate which does infiltrate into the bedrock undergoes further denitrification thus preventing more elevated concentrations of nitrate from entering the deeper groundwater body.

3.3 The influence of hydrogeology and biogeochemistry on nitrate fate and transport

Groundwater quality in both the Nuenna and Glen Burn catchments is impacted by agricultural contamination. However, the contrasting hydrogeological settings have a significant influence on the dominant biogeochemical processes influencing nitrate fate and transport. The bedrock in both catchments transports groundwater predominantly through fracture flow. However the variation in groundwater discharge via hydraulically active fracture sets with depth and the transmissivity ranges differ considerably in the two catchments investigated. This has considerable influence on the fate and transport of nitrate in the groundwater bodies. Spahr et al. (2010) reported similar findings in a study of 148 sites across various climatic and geological settings in the United States. The study found that lower nitrate concentrations were associated with impermeable soils and impermeable bedrock, whereas catchments with a combination of permeable soils or permeable bedrock were associated with higher nitrate concentrations.

The bedrock geochemistry also plays a role in nutrient mobility through the presence of electron donors which are capable of supporting iron-mediated denitrification, such as pyrite, reducing
Nitrate mobility. The contrast in water quality between the two catchments is clearly greater than between the monitoring wells within the individual catchments, despite the localised impacts of point sources. This further highlights the significant influence of catchment hydrogeological setting.

Nitrification is the dominant biogeochemical process influencing dissolved N mobility in the Nuenna catchment, where bedrock transmissivity values are higher. Hydrochemical and isotopic data suggest localised anaerobic conditions may occur in a hydrogeological system otherwise dominated by aerobic water that prevents denitrification. Denitrification occurs locally and intermittently in the near surface transition zone, although its influence on a catchment scale is insignificant. The absence of a consistent gradient in hydrochemistry between samples collected from wells in the subsoil and from the deeper bedrock, where lower nitrate was observed in shallow groundwater samples, is consistent with preferential (by-passing) flow mechanisms. Furthermore there is a need for reliable and representative monitoring wells away from point sources in order to determine the target groundwater pathway. In contrast, denitrification is the dominant biogeochemical process affecting aqueous nitrate in the lower transmissivity bedrock of the Glen Burn catchment; the deeper groundwater pathways are more important in higher permeability and karstified bedrock aquifers, such as in the Nuenna catchment.

The nitrate isotopic signature in the karstified bedrock aquifer in the Nuenna catchment reflects the original nitrate source more accurately than the nitrate isotopic signature in the lower permeability Glen Burn aquifer, where there is evidence of fractionation due to biogeochemical processes. Consequently, in low permeability bedrock catchments such as the Glen Burn, the groundwater nitrate isotopic signature is likely to be altered from the original source signature and the surface water samples are more likely to represent original nitrate sources in the catchment.
The importance of hydrogeological setting has implications for the groundwater quality monitoring and implementation of contamination mitigation measures in catchments with differing hydrogeological settings. In karstified aquifers nitrate management strategies should focus on the role played by deep groundwater pathways and diffuse nitrogen sources, whereas in catchments underlain by lower permeability aquifers, the deep groundwater will be a less significant pathway for nitrate to enter surface water bodies and the focus of such management plans should be on pathways nearer the ground surface. Incorporating these considerations, when developing catchment management plans, can assist in addressing the impact of agricultural practices on the water quality of groundwater bodies and contribute toward achieving the aims of the Water Framework Directive.

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