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1	Dominance of biologically produced nitrate in upland waters of
2	Great Britain indicated by stable isotopes
3	
4	Chris J. Curtis <sup>1</sup> , Timothy H.E. Heaton <sup>2</sup> , Gavin L. Simpson <sup>1</sup> , Chris D. Evans <sup>3</sup> , James
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1	intercept a large proportion of deposited N. The dominance of microbial sources of NO3 <sup>-</sup> in upland
2	waters suggests that reduced and oxidised N deposition may have similar implications in terms of
3	contributing to $NO_3^-$ leaching.
4	
5	
6	Keywords
7	Nitrate leaching, nitrification, <sup>18</sup> O, <sup>15</sup> N, uplands, lakes, N deposition
8	
9	Running title:
10	Dominance of biological nitrate in upland waters
11	
12	Introduction
13	
14	The environmental problems associated with anthropogenic nitrogen (N) deposition to
15	semi-natural, upland catchments include surface water acidification and
16	eutrophication of both aquatic and terrestrial ecosystems (Curtis et al. 2005a; Maberly
17	et al. 2002; Galloway et al. 2003; Bergström and Jansson 2006; Emmett 2007; Elser et
18	al. 2009). However, despite many biogeochemical studies of N in upland catchments
19	over the last 10-15 years, the mechanistic links between N deposition and nitrate
20	$(NO_3)$ leaching are still not well understood. A key factor is that inorganic forms of N
21	are readily available for biological uptake in N-limited terrestrial upland ecosystems,
22	so that only a small proportion of deposition inputs ever reaches surface waters.
23	Simple empirical observations of NO <sub>3</sub> outputs versus N deposition inputs show that
24	there may be a deposition threshold below which NO <sub>3</sub> -leaching rarely occurs, but
25	above which there may be a wide range of responses from very low to very high $NO_3^-$
26	leaching (Dise et al. 1998). Several factors related to cumulative N deposition effects

1	have been proposed to explain these patterns, including soil C:N ratios (e.g. Dise and
2	Wright 1995; Gundersen et al. 1998; MacDonald et al. 2002; van der Salm et al.
3	2007) and total catchment carbon pools or proportion of organic soils (Evans et al.
4	2006; Helliwell et al. 2007). Spatial variations in catchment sensitivity to N
5	deposition and associated NO <sub>3</sub> <sup>-</sup> leaching have also been linked to vegetation type
6	(Rowe et al. 2006) and certain physical catchment attributes such as altitude, presence
7	of bare rock and steep slopes (Kopácek et al. 2005; Helliwell et al. 2007).
8	

9 A simple, initial hypothesis is that, since ammonium concentrations in upland waters 10 are generally negligible, and elevated NO<sub>3</sub><sup>-</sup> concentrations only occur where there are 11 high deposition loads of oxidized N (NOx), the NO3<sup>-</sup> observed in impacted upland 12 waters must derive directly from NO<sub>3</sub><sup>-</sup> in precipitation (cf. NEGTAP 2001; Durka et 13 al. 1994; Emmett 2007). Ammonium in soilwaters is subject to cation exchange and is less mobile than NO<sub>3</sub>, with a longer residence time in soils and hence greater 14 15 potential for microbial or vegetation uptake, so NO<sub>3</sub><sup>-</sup> is more likely to bypass these 16 biological sinks for N to be leached into surface waters. This hypothesis is 17 superficially supported by the observation that steeper, rockier catchments may leach 18 more NO<sub>3</sub><sup>-</sup>, leading to the usage of the term "hydrological NO<sub>3</sub><sup>-</sup>" (e.g. Moldan et al. 19 1995; Curtis et al. 2005a, b) and the idea of the "Teflon basin" in early studies of 20 alpine lakes in North America, whereby rapidly flushed NO<sub>3</sub><sup>-</sup> is not retained in alpine 21 basins (later questioned by Campbell et al. 2002). Such observations have resulted in 22 a common perception that enhanced  $NO_3^-$  leaching is associated primarily with NOx 23 deposition in upland catchments.

1	Several studies, primarily from North America, have more recently challenged the
2	perception that leached $NO_3^-$ is of direct, atmospheric origin, through the use of stable
3	isotope techniques to identify the source of $NO_3^-$ isolated from surface water samples
4	(e.g. Durka et al. 1994; Williard et al. 2001; Spoelstra et al. 2001; Campbell et al.
5	2002, 2006; Hales et al. 2007; Sebestyen et al. 2008). Many of these studies have
6	demonstrated that even in alpine catchments, a very large proportion of leached NO <sub>3</sub> <sup>-</sup>
7	is of microbial origin, as indicated by large differences in the abundance of the
8	naturally occurring heavy isotope of oxygen, <sup>18</sup> O, between NO <sub>3</sub> <sup>-</sup> collected from
9	precipitation and surface water samples. The direct implication is that leaching of
10	unaltered, atmospheric $NO_3^-$ is a minor source of surface water $NO_3^-$ in the uplands
11	and that other factors linked to the nitrification of ammonium are much more
12	important.
13	
14	The aim of the present study is to determine the relative importance of atmospheric
15	versus microbial sources of NO3 <sup>-</sup> across four semi-natural, non-forest upland
16	catchments in the UK with differing levels of $NO_3^-$ leaching, deposition loads and
17	different land cover / physical attributes. The UK uplands experience very high
18	nitrogen deposition loads compared with most of the semi-natural watersheds in North
19	America where dual isotope studies have been carried out; total wet + dry N
20	deposition in 2006 exceeded 24 kgN ha <sup>-1</sup> yr <sup>-1</sup> across many upland regions (RoTAP in
21	press). Furthermore, many upland waters in these areas experience chronic NO <sub>3</sub> <sup>-</sup>
22	leaching for much or all of the year (Curtis et al. 2005a). The hypothesis that leached
23	NO <sub>3</sub> <sup>-</sup> is primarily of atmospheric origin suggests that policy measures to control NOx
24	emissions should be the main priority to protect upland waters from the adverse
25	effects of enhanced NO <sub>3</sub> <sup>-</sup> leaching. If leached NO <sub>3</sub> <sup>-</sup> is mainly microbially produced,

then all external sources of inorganic N are likely to contribute to observed patterns of
 NO<sub>3</sub><sup>-</sup> leaching.

3

#### 4 *Site description*

Four study sites were selected on the basis of the availability of co-located long-term 5 6 water chemistry data from the UK Acid Waters Monitoring Network (AWMN: Monteith and Shilland 2007) and bulk deposition chemistry from the UK Acid 7 8 Deposition Monitoring Network (ADMN) since 1999 when the sites were 9 instrumented (Lawrence et al. 2008; Table 1). All four sites occupy semi-natural 10 headwater catchments (Fig. 1) with atmospheric deposition as the only major source 11 of anthropogenic pollution. The Afon Gwy and Scoat Tarn catchments comprise 12 mainly acid grassland with sheep grazing. The River Etherow catchment is mainly 13 Calluna vulgaris dominated moorland on blanket peat, which is managed as a grouse 14 moor by burning, with some sheep grazing. The blanket bog is affected by gully 15 erosion and historical loss of Sphagnum moss cover, which has been attributed to acid 16 deposition (Tallis 1987). The catchment of Lochnagar comprises alpine/subalpine 17 heath with a large proportion of bare rock and scree. The Afon Gwy and River 18 Etherow are streams, while Scoat Tarn and Lochnagar are headwater lakes with smaller catchments (Fig.1, Table 1). All four sites are acidified, showing exceedance 19 20 of critical loads for acidity (Curtis et al. 2005a), with moderate NO<sub>3</sub><sup>-</sup> leaching at the 21 Afon Gwy, high NO<sub>3</sub> leaching at Scoat Tarn and Lochnagar and extremely high NO<sub>3</sub> 22 (for a semi-natural upland site) in the River Etherow (Table 1). These patterns reflect 23  $NO_3^-$  concentrations in bulk deposition, with comparable concentrations between 24 surface waters and bulk deposition at all sites except the River Etherow, where mean 25  $NO_3^-$  concentrations are much higher in the stream (Table 1). This observation

- appears to support the hypothesis that surface water NO<sub>3</sub><sup>-</sup> may be largely dictated by
   concentrations in rainfall, with some additional source of NO<sub>3</sub><sup>-</sup> at the Etherow.
- 3

5 Methods

- 6
- 7

A pilot study to assess the feasibility of the dual isotope  $({}^{15}N/{}^{14}N \text{ and } {}^{18}O/{}^{16}O)$  method 8 9 was first carried out at the Afon Gwy from September 2004 to August 2005, on the 10 basis of previous studies at the site and the development of a conceptual model of 11 NO<sub>3</sub><sup>-</sup> leaching zones (Evans et al. 2004). A high volume bulk deposition collector with a small collecting roof (c.  $0.5m^2$ ) was installed adjacent to the streamwater 12 13 sampling point. Three zero-tension tray lysimeters were installed in an exposed 14 hillslope cutting approximately 700m from the bulk deposition collector on the 15 adjacent hillslope (grid ref. SN81967, 86100), at depths of c. 10cm (bottom of rooting 16 zone), 20cm (base of O horizon) and 50cm (B horizon, above an impermeable 17 ironpan). Streamwaters, bulk deposition and tray lysimeters were all sampled monthly 18 for isotopes. Additional streamwater samples were obtained from a number of 19 subcatchments on a quarterly basis for comparison with the main channel, with 20 subcatchments selected on the basis of dominant soils and previous water chemistry 21 data from Evans et al. (2004). Only data from tributary LB7a are presented here.

22

Following successful application of the techniques at the Afon Gwy, monthly isotopic sampling commenced at the other three sites in August 2005, ending in July 2006 (Table 1). At these sites, bulk deposition collectors were located close to surface water

1 sample points and two tray lysimeters were deployed nearby in deeper organic 2 horizons at c. 10cm and 20cm depths. The high-volume bulk deposition collectors at 3 the three new sites comprised a large diameter HDPE funnel (c. 32cm diameter) 4 attached to a pole about 2m above ground level and connected via a length of neoprene tubing to a sealed 5L LDPE carboy buried under c. 10cm of soil. At the 5 6 River Etherow, streamwater samples were obtained from two tributary streams (Rose Clough and Swan Clough) as well as the main river channel at the weir (Table 1; Fig. 7 8 1). At Scoat Tarn, the two major inflow streams were sampled as well as the lake 9 outflow, while at Lochnagar a spring-fed inflow stream was sampled as well as the 10 lake outflow. 11 12 13 Sample collection and analysis 14 15 Surface water and bulk deposition sampling and analysis for isotopes 16 For isotopic analysis, monthly surface water and bulk deposition samples were 17 collected and a subsample analysed in the field using a portable spectrophotometer 18 (Hach DR/2400) for concentrations of the acid anions  $NO_3^-$ , sulphate and chloride to 19 provide approximate estimates of required sample volumes for isotopic analysis 20 (target was at least 100 µeq NO<sub>3</sub><sup>-</sup>). All samples were collected in acid-washed, DIW-21 rinsed LDPE carboys. For surface waters at least one 20L carboy was filled after 22 rinsing three times with sample (up to 40L in two carboys in summer when low  $NO_3^{-1}$ concentrations  $<5 \mu$ eq l<sup>-1</sup> were found). Bulk deposition samples were collected in the 23 24 original 5L LDPE carboys which were then replaced with clean carboys following 25 rinsing of the funnel and tubing with DIW.

2	Streamwater and bulk deposition $NO_3^-$ samples were then filtered to 0.45 µm using
3	high capacity groundwater filtration cartridges or 142mm diameter disc filters
4	(Whatman GF/F 0.7 $\mu$ m pre-filter and Pall Supor-450 0.45 $\mu$ m membrane filter) and
5	pre-treated through cation exchange resins (AG50W-X8) prior to collection on anion
6	exchange resins (Dowex AG1-X8) according to the method of Chang et al. (1999).
7	Where possible, at least 100 $\mu$ eq of NO <sub>3</sub> <sup>-</sup> was passed through the exchange columns
8	but care had to be taken not to exceed the exchange capacity of the anion resins (6000
9	$\mu$ eq) to avoid possible isotope fractionation of the collected NO <sub>3</sub> <sup>-</sup> sample. In practice,
10	it was not always possible to collect sufficient $NO_3^-$ for isotopic analysis due to very
11	low concentrations in streamwaters relative to other anions, especially chloride. The
12	anion resins were refrigerated prior to being transported to the NERC Isotope
13	Geosciences Laboratory, Keyworth, where the NO <sub>3</sub> <sup>-</sup> was converted to silver nitrate
14	(Silva et al., 2000; Heaton et al., 2004). ${}^{15}N/{}^{14}N$ and ${}^{18}O/{}^{16}O$ ratios were analysed by
15	combustion to $N_2$ in a Flash EA Elemental Analyzer, or thermal conversion to CO in a
16	TC-EA, respectively, with gases passed on-line to a Delta Plus XL mass spectrometer
17	(all ThermoFinnigan, Bremen, Germany). Sample purity and freedom from organic
18	contamination was checked by ensuring that N/O ratios were close to those of pure
19	$NO_3^-$ standards (Heaton et al. 2004). Nitrate isotope ratios were calculated as $\delta^{15}N$
20	values versus air (atmospheric $N_2$ ) and $\delta^{18}O$ versus VSMOW by comparison with
21	standards IAEA N-1 and N-2, and USGS 34 and 35. Water $^{18}O/^{16}O$ ratios were
22	determined following equilibration with CO <sub>2</sub> in an Isoprep-18 on-line to a SIRA II
23	mass spectrometer (VG Isotopes, Middlewich, England), and calculated as $\delta^{18}O$
24	values versus VSMOW by comparison to IAEA standards VSMOW and SLAP.

#### 1 <u>Soilwater sampling and analysis</u>

Monthly tray lysimeter soilwater samples were filtered and analysed for acid anions in the field using the methods described above. Where sufficient NO<sub>3</sub><sup>-</sup> was measured, the sample was loaded onto ion-exchange resin columns for dual isotope analysis. Where sample volumes were insufficient for isotopic analysis, samples were kept refrigerated and bulked with subsequent months until sufficient sample volume was obtained.

7

#### 8 <u>Water chemistry sampling programmes</u>

9 Water chemistry data were obtained from separate ongoing sampling programmes at 10 the study sites. Separate surface water and bulk deposition samples were obtained at 11 the same frequency (or greater) as isotope samples (except at Scoat Tarn) and 12 analysed for water chemistry in the laboratory according to the established protocols 13 of the AWMN for surface waters (Monteith and Shilland 2007) and ADMN for bulk 14 deposition (Lawrence et al. 2008), using ion chromatography with detection limits of c. 1  $\mu$ eq l<sup>-1</sup> for NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup> and ion-balance checks as part of routine analytical 15 16 qualitical control. Surface water samples were taken weekly from the Afon Gwy, 2-17 weekly from Lochnagar, monthly at the River Etherow and quarterly at Scoat Tarn. 18 Bulk deposition sampling for water chemistry was carried out 2-weekly at all sites 19 using collectors following the design of Hall (1986). Only these chemistry data are 20 reported here; monthly field results obtained using the portable spectrophotometer are 21 considered to be approximate only.

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24 Data analysis

1 <u>Calculation of the theoretical  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> of bacterial NO<sub>3</sub><sup>-</sup></u>

Determination of the proportions of atmospheric and microbial NO<sub>3</sub><sup>-</sup> using  $\delta^{18}$ O relies on the measurement of atmospheric  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> in bulk deposition, and the theoretical calculation of microbial  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> based on measurement of soilwater  $\delta^{18}$ O-H<sub>2</sub>O. This calculation has traditionally been based on the assumption that autotrophic microbial NO<sub>3</sub><sup>-</sup>derives one part of its oxygen from atmospheric O<sub>2</sub> ( $\delta^{18}$ O = +23‰) and two parts from soilwater H<sub>2</sub>O, which is measured directly (Amberger and Schmidt 1987; Kendall 1998):

9

10 
$$\delta^{18}$$
O-NO<sub>3</sub><sup>-</sup> = (2/3  $\delta^{18}$ O-H<sub>2</sub>O soilwater) + (1/3  $\delta^{18}$ O-O<sub>2</sub> atmosphere) (1)

11

12 This theoretical calculation makes a number of assumptions which may be valid for many, but not all environments (Mayer et al. 2001; Kendall et al. 2007; Spoelstra et 13 al. 2007; Snider et al. 2010). Thus, from experimental results Mayer et al. (2001) 14 15 suggested that there may be some circumstances under which heterotrophic 16 nitrification might yield NO<sub>3</sub><sup>-</sup> oxygen derived from two parts atmospheric oxygen and only one part soilwater oxygen. In contrast, recent incubation experiments with <sup>18</sup>O-17 18 labelled waters have suggested that in some cases over 90% of the NO<sub>3</sub><sup>-</sup> oxygen might be derived from soil water (Snider et al. 2010). Here we calculate the soil microbial 19 20  $NO_3^-$  end-member using the commonly used equation above, assuming the ratio of 21 atmospheric-O to soilwater-O equals 1 to 2; but comment later on the validity of this. 22 Future isotope studies may be able to overcome these uncertainties by using techniques for measuring NO<sub>3</sub><sup>-</sup>  $\delta^{17}$ O in addition to  $\delta^{18}$ O (Michalski et al. 2004; Curtis 23 24 et al. 2011).

1 <u>Statistical analysis</u>

2 A series of linear models fitted by ordinary least squares was used to formally address 3 a set of key hypotheses; i) there are differences between study sites in the isotopic signature of deposited NO<sub>3</sub><sup>-</sup> (i.e. both  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> and  $\delta^{15}$ N-NO<sub>3</sub><sup>-</sup>), ii) there are 4 differences between sites in  $\delta^{18}$ O-H<sub>2</sub>O of soilwaters, used in the derivation of 5 theoretical microbial  $\delta^{18}$ O-NO<sub>3</sub>, iii) there are isotopic differences between deposition 6 NO<sub>3</sub><sup>-</sup> and surface water NO<sub>3</sub><sup>-</sup> within study sites (for both  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> and  $\delta^{15}$ N-NO<sub>3</sub><sup>-</sup>), 7 iv) there are within-site differences in both  $\delta^{15}$ N-NO<sub>3</sub><sup>-</sup> and  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> between inflow 8 9 streams and lake outflows indicating differences in the contribution of atmospheric NO<sub>3</sub><sup>-</sup> leaching, v) there are differences between sites in  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> of streamwaters 10 11 indicative of differing levels of atmospheric NO<sub>3</sub><sup>-</sup> leaching. The assumptions of 12 ordinary least squares were checked via exploratory plotting of data and via Fligner-13 Kileen tests for homogeneity of variances. Where the homogeneity of variances 14 assumption was not met by the data, a sandwich estimator of the standard errors of 15 model parameters was used, providing heteroscedastic-consistent standard errors. 16 Post-hoc pair-wise comparisons were conducted according to the method of Tukey's Honest Significance Differences (Tukey's range test) at a 95% family-wise confidence 17 18 level (Bretz et al. 2010). All analyses were performed using R (version 2.11-patched 19 R Core Development Team, 2010) with the multcomp (version 1.2-2, Hothorn et al. 20 2008) and sandwich (2.2-6, Zeileis 2004) packages.

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23 Results
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25 Deposition and surface water chemistry

1	Concentrations of $NO_3^-$ and $NH_4^+$ in bulk deposition were significantly related
2	(ANCOVA: likelihood ratio 109.9045, 4 d.f., $p < 0.0001$ ; Fig. 2). The highest
3	concentrations in precipitation occurred at the River Etherow, regularly exceeding 50
4	$\mu$ eq l <sup>-1</sup> for both NO <sub>3</sub> <sup>-</sup> and NH <sub>4</sub> <sup>+</sup> , while lower peaks in the range 20-30 $\mu$ eq l <sup>-1</sup> occurred
5	at the Afon Gwy. At Lochnagar, concentrations of NO <sub>3</sub> <sup>-</sup> in precipitation were
6	generally higher than $NH_4^+$ while at other sites $NH_4^+$ concentrations regularly
7	exceeded those of $NO_3^-$ .
8	
9	Mean concentrations and deposition fluxes of NO <sub>3</sub> <sup>-</sup> , ammonium and dissolved
10	inorganic N (DIN) are presented in Table 2. While concentrations of all ions in bulk
11	deposition were greatest at the River Etherow site, the greatest deposition fluxes
12	occurred at Scoat Tarn, where rainfall was more than double that at the Etherow.
13	Furthermore, while mean concentrations of $NO_3^-$ and $NH_4^+$ were lowest at the Afon
14	Gwy, much lower rainfall at Lochnagar meant that deposition fluxes were lowest at
15	the latter site.
16	
17	Concentrations of $NO_3^-$ in bulk deposition were generally higher than or similar to
18	those in surface waters for most of the year except at the River Etherow, where $NO_3^-$
19	concentrations in the stream exceeded those in precipitation on about 50% of
20	sampling occasions (Fig. 2). Surface water concentrations of $NH_4^+$ were negligible at
21	all sites except for occasional measurable levels at the River Etherow and are not
22	considered further.
23	
24	Surface water $NO_3^-$ concentrations over the study period were very similar to the
25	longer-term means in Table 1 and showed the same pattern relative to bulk deposition

1	(Table 2). Mean $NO_3^-$ concentrations were similar in the stream and bulk deposition at
2	the River Etherow while at the other three sites, mean concentrations in surface waters
3	were slightly lower than in bulk deposition. Although flow was not measured at the
4	River Etherow or Scoat Tarn, flow data for the Afon Gwy were used to calculate a
5	flow-weighted mean concentration of 6.8 $\mu$ eq l <sup>-1</sup> which is slightly higher than the
6	unweighted mean of 6.0 $\mu$ eq l <sup>-1</sup> and closer to the volume weighted mean of 10.1 $\mu$ eq l <sup>-</sup>
7	<sup>1</sup> in bulk deposition. At Lochnagar, monthly scaling factors based on historical flow
8	data were used to calculate a flow-weighted mean $NO_3^-$ concentration of 15.2 µeq $l^{-1}$ ,
9	which is slightly lower than the unweighted mean of 15.8 $\mu$ eq l <sup>-1</sup> but close to the
10	volume-weighted mean value in bulk deposition of 17.7 $\mu$ eq l <sup>-1</sup> . In the absence of
11	standard flow data for all four sites, catchment leaching fluxes of NO3 <sup>-</sup> were estimated
12	from modelled runoff data. Runoff estimates were provided by the Centre for Ecology
13	and Hydrology, Wallingford, and were derived from interpolated long-term mean
14	rainfall measurements (1941-70) and modelled evapotranspiration on a 1km grid for
15	the UK, based on meteorological data obtained from the UK Met Office.
16	
17	
18	Dual isotope analysis
19	
20	All isotope data are summarised in the Supplementary Tables S2 to S5.
21	
22	Summary box plots of the monthly dual isotope analysis of bulk deposition $NO_3^-$ at
23	the four study sites are shown in Figs. 3-4. The range of $\delta^{15}$ N-NO <sub>3</sub> <sup>-</sup> varied from -2.3 to
24	+3.0‰ (n=9, mean= -0.3‰, SD= 1.6‰) at the Afon Gwy, -4.5 to +3.1‰ (n=12,
25	mean=+0.5‰, SD= 2.3‰) at the River Etherow, -4.3 to +1.0‰ (n=11, mean= -

1.5‰, SD= 1.5‰) at Scoat Tarn and -5.7 to +1.0‰ (n=11, mean= -1.8‰, SD= 1.8‰)
 2 at Lochnagar.

4	Analysis of variance indicated that there were significant differences in bulk
5	deposition $\delta^{15}$ N-NO <sub>3</sub> <sup>-</sup> between some sites (F <sub>(3,38)</sub> =3.41, p=0.027; Fig.3). Post hoc
6	comparison of pairwise differences of means showed a significant difference in $\delta^{15}N\text{-}$
7	$NO_3^-$ between Lochnagar and the River Etherow (t=-2.89, p=0.030) while the
8	difference between Scoat Tarn and the River Etherow was not significant (t=-2.39,
9	p=0.093). While there was a scatter around 0.0‰ at the Afon Gwy and River
10	Etherow, most samples of bulk deposition NO <sub>3</sub> <sup>-</sup> from Scoat Tarn and Lochnagar were
11	depleted in <sup>15</sup> N and only the River Etherow had a (slightly) positive mean value of
12	$\delta^{15}$ N.
13	
14	At three sites the range of $\delta^{15}$ N-NO <sub>3</sub> <sup>-</sup> values in surface waters largely overlapped with
15	that in atmospheric deposition. The exception was the River Etherow, where
16	streamwater $\delta^{15}$ N-NO <sub>3</sub> <sup>-</sup> values were significantly higher than those in bulk deposition,
17	especially for the two tributary streams sampled (Tukey's HSD: Rose Clough, t=6.64,
18	p<0.001; Swan Clough t=7.67, p<0.001; River Etherow t=3.92, p=0.002). Mean
19	values of $\delta^{15}$ N-NO <sub>3</sub> <sup>-</sup> were +4.6‰ (n=12, SD= 2.0‰) for Rose Clough and +5.2‰
20	(n=12, SD=1.1%) for Swan Clough, in both cases exceeding the maximum values
21	recorded in bulk deposition for the River Etherow catchment (+3.1‰ in December
22	2005). Mean $\delta^{15}$ N values for surface waters in the Etherow catchment were 2.4 to
23	4.7‰ higher than those in bulk deposition. Despite the greater overlaps in $\delta^{15}N$ values
24	at other sites, surface water mean $\delta^{15}N$ values were also higher (by 0.9 to 1.5‰) than
25	those in bulk deposition.

2	Highly positive values of $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> in bulk deposition were found at all four sites,
3	with a minimum value of +48.3‰ and a maximum of +82.1‰, both at Lochnagar
4	(Fig. 4). Much smaller ranges were found at the Afon Gwy and River Etherow sites
5	(+60.4 to +71.6‰). Mean values from all sites were very similar, from +66.5 (River
6	Etherow, $n=12$ , SD=3.6‰) to +69.6‰ at Lochnagar ( $n=11$ , SD= 9.5‰). Analysis of
7	variance showed no significant differences in bulk deposition $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> between sites
8	(F <sub>(3,37)</sub> =1.02, p=0.394). However, there were large and significant differences in $\delta^{18}$ O-
9	$NO_3^-$ between surface waters and bulk deposition at each site (Tukey HSD, p<0.001
10	for all sampled water bodies; see Fig.5). The maximum surface water $\delta^{18}$ O value of
11	+22.4‰, observed at Lochnagar in April 2006, was lower than the minimum value
12	observed in deposition. This allowed the use of a simple two end-member mixing
13	model to estimate the proportion of untransformed atmospheric $NO_3^-$ in surface waters
14	(see below). In the few soilwater samples which yielded sufficient $NO_3^-$ for isotopic
15	analysis, the $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> values were comparable to surface waters at Scoat Tarn (+9.1
16	to +19.8‰), but intermediate between surface waters and bulk deposition at both the
17	River Etherow ( $\delta^{18}O = +17.2$ to +28.7) and Lochnagar ( $\delta^{18}O = +35.5$ to +53.6‰ in
18	bulked samples; see Fig. 5, Table S6).

19

20 While the differences in  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> between bulk deposition and surface waters were 21 much greater than for  $\delta^{15}$ N-NO<sub>3</sub><sup>-</sup>, there was some evidence of inverse relationships 22 between the  $\delta^{18}$ O and  $\delta^{15}$ N values of surface water NO<sub>3</sub><sup>-</sup> (i.e. decreasing  $\delta^{15}$ N with 23 increasing  $\delta^{18}$ O) in the Etherow tributaries Rose Clough and Swan Clough, inflow 2 at 24 Scoat Tarn, and both inflow and lake at Lochnagar (all p<0.05; see Fig. 5).

1 Theoretical  $\delta^{18}O$ -NO<sub>3</sub><sup>-</sup> of microbially produced NO<sub>3</sub><sup>-</sup>

2	Measurement of $\delta^{18}$ O-H <sub>2</sub> O at the four sites showed very similar mean values of -5.9
3	(Afon Gwy), -6.0 (River Etherow), -6.4 (Scoat Tarn) and -7.4‰ (Lochnagar). The
4	only significant difference between sites was between Lochnagar and the River
5	Etherow (t=-2.77, p=0.035). Mean theoretical $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> values for microbially
6	produced NO <sub>3</sub> <sup>-</sup> were +3.7‰ (Afon Gwy, River Etherow), +3.4‰ (Scoat Tarn) and
7	+2.8‰ (Lochnagar). The annual range of theoretical microbial $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> for each
8	site is shown in Fig. 5; monthly data are provided in Supplementary Information.
9	
10	
11	Derived contribution of untransformed atmospheric $NO_3^-$
12	The simple two end-member mixing model employs an interpolation between the
13	theoretical microbial value and the measured bulk deposition $\delta^{18}\text{O-NO}_3^-$ (in each case
14	referring to $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> ):
15	
16	% atmospheric = $(\delta^{18}O_{surface water} - \delta^{18}O_{microbial}) / (\delta^{18}O_{deposition} - \delta^{18}O_{microbial}) \times 100\% (2)$
17	
18	Calculated values for each monthly sample are presented in supplementary Tables S2-
19	S5 but here we present annual means based on % values calculated monthly from
20	measured $\delta^{18}O_{surface water}$ , $\delta^{18}O_{microbial}$ and $\delta^{18}O_{deposition}$ .
21	
22	
23	Afon Gwy
24	At the Afon Gwy, most measurements of $\delta^{18}\text{O-NO}_3^-$ in the stream lay within the
25	theoretical range for microbially produced $NO_3^-$ (Fig. 5), suggesting that almost all the

1 NO<sub>3</sub><sup>-</sup> observed in the stream had been microbially produced, and that direct leaching 2 of untransformed atmospheric  $NO_3^-$  was negligible (Fig. 6). Very low  $NO_3^-$ 3 concentrations in the main channel at the Afon Gwy regularly prevented the collection 4 of sufficient  $NO_3^-$  on the anion resins for isotopic analysis. The highest proportion of 5 atmospheric NO<sub>3</sub><sup>-</sup> was recorded from a tributary of the Afon Gwy (stream LB7a) with 6 slightly higher NO<sub>3</sub><sup>-</sup> concentrations, indicating 7% untransformed NO<sub>3</sub><sup>-</sup> in June 2005; 7 the annual mean value was only 3% (Table S2; SE= 1.4%). Due to bad weather, this 8 site could not be accessed for sampling during February-March 2005. In order to 9 investigate the possibility that peaks in both NO<sub>3</sub><sup>-</sup> concentration and atmospheric 10 contribution had been missed as a result, the site was resurveyed as part of a subsequent study in February-April 2009. In this later study, bulk deposition  $\delta^{18}$ O-11 12  $NO_3^{-1}$  fell within the range reported here while the proportion of atmospheric  $NO_3^{-1}$  was 13 3% in February 2009 and <1% in March 2009. Insufficient NO<sub>3</sub><sup>-</sup> was collected in 14 April 2009 for dual isotope analysis.

15

#### 16 <u>River Etherow</u>

17 At the River Etherow and its two sampled tributaries, all samples analysed had low  $\delta^{18}$ O-NO<sub>3</sub> values, close to or within the range for microbially produced NO<sub>3</sub> (Fig. 5). 18 19 Less than 10% of streamwater  $NO_3^-$  was untransformed atmospheric  $NO_3^-$ , with peak 20 and mean values of only 9% and 6% (SE = 0.7%) respectively in the main channel (Fig. 6). In the Rose Clough and Swan Clough tributaries the  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> signal was 21 22 barely distinguishable from the theoretical range for microbially produced  $NO_3^-$  (Fig. 23 5), with mean values for both streams indicating <3% (SE = 0.5 and 0.7%) 24 respectively) atmospheric NO<sub>3</sub><sup>-</sup> contribution (Table S3). Bulked samples from shallow 25 soilwater lysimeters did show a larger proportion of atmospheric NO<sub>3</sub><sup>-</sup> in the upper 1 peat horizons, up to 40% for the period November 2005 to February 2006 (Fig. 5), but

2 there was no evidence that this atmospheric NO<sub>3</sub><sup>-</sup> reached surface waters.

3

#### 4 <u>Scoat Tarn</u>

Most outflow samples from Scoat Tarn showed  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> values much greater than 5 6 the range for microbially produced  $NO_3^-$  at the site. The range of values was greater in 7 the inflow streams than in the lake outflow, but the annual mean value was greater in 8 the outflow (Fig. 5; Table S4). The proportion of untransformed atmospheric  $NO_3^{-1}$ indicated by  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> values varied from 9-21% (mean 15%, SE= 1.0%) in the lake 9 10 outflow, with a greater range of 5-23% (mean 11%, SE= 1.6%) in Inflow 1 and 1-24% 11 (mean 9%, SE= 1.8%) in Inflow 2 (Figs. 5-6). The tray lysimeters at Scoat Tarn produced the greatest number of samples of sufficient size for isotopic analysis and 12 indicated a similar range in  $\delta^{18}$ O values as surface waters, with inferred proportions of 13 14 atmospheric  $NO_3^-$  from 8-26% (Fig. 5, Table S6).

15

#### 16 <u>Lochnagar</u>

Values of  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> in the loch outflow at Lochnagar were all much higher than the 17 18 range for microbial  $NO_3^-$  (Table S5) although some of the inflow spring values overlapped the microbial range (Fig. 5). The calculated proportions of atmospheric 19 20  $NO_3^-$  showed a greater range and higher mean values than other sites; 14-32% (mean 21 21%, SE= 1.7%) in the loch outflow and 0-30% (mean 13%, SE= 3.3%) in the inflow 22 spring (Fig. 6). Lysimeter samples from this site were all very small and the only data 23 obtained were for samples bulked over several months. However these samples had very high  $\delta^{18}$ O values of +35.5‰ and +53.6‰ in the two lysimeters, indicating 43 24 and 69% atmospheric  $NO_3^-$  in the shallow tray lysimeters over this period. 25

#### 2 Temporal patterns

3 Temporal variations in the direct contribution of atmospheric  $NO_3^-$  to surface water 4 concentrations are illustrated in Fig. 6. While there were too few samples to identify temporal patterns at the Afon Gwy, the other sites showed distinct seasonal patterns in 5 6 the contribution of atmospheric  $NO_3^-$  which were much more pronounced in streams and lake inflows than in lake outflows (Fig. 6). The contribution of atmospheric NO<sub>3</sub><sup>-</sup> 7 8 was lowest in the autumn or early winter and increased to a maximum in the late 9 winter or spring, when seasonal peaks in NO<sub>3</sub><sup>-</sup> concentration are normally observed. 10 At Scoat Tarn both minimum and maximum atmospheric contributions occurred later 11 in the lake outflow relative to inflow streams. Comparisons with rainfall data from the 12 bulk deposition collectors (supplementary Fig. S3) show that while fewer high rainfall 13 events occurred in summer there was no real pattern in rainfall seasonality 14 corresponding with seasonal atmospheric contributions to surface waters.

15

16

#### 17 **Discussion**

18

19 Consideration of deposition input fluxes and leaching fluxes alone suggests that the 20 four study catchments leached varying proportions of bulk deposited N; from 57% at 21 the Afon Gwy to 156% at the River Etherow as a proportion of bulk deposited  $NO_3^-$ , 22 or from 26% at the Afon Gwy to 71% at the River Etherow when expressed as a 23 proportion of total inorganic N (NH<sub>4</sub><sup>+</sup> + NO<sub>3</sub><sup>-</sup>) in bulk deposition.

1 While surface water  $NO_3^-$  concentrations were comparable to those in bulk deposition 2 at three of the four study sites, the dual isotope data indicated that most of the  $NO_3^{-1}$ 3 reaching surface waters had been microbially cycled and was not simply atmospheric 4 NO<sub>3</sub><sup>-</sup> transported hydrologically through catchments into surface waters. These results 5 correspond with previous studies using the same technique in North America 6 (summarized in Curtis et al. 2011). This study also confirms the findings of previous authors that the  $\delta^{18}$ O signature of NO<sub>3</sub><sup>-</sup> is much more useful than  $\delta^{15}$ N for separating 7 atmospheric and microbial sources of surface water NO<sub>3</sub><sup>-</sup> in upland catchments where 8 9 there are no other important inputs of inorganic N (Kendall 1998; Hales et al. 2004; 10 Piatek et al. 2005). A major implication of this study is that where microbially produced  $NO_3^-$  is found, it may originate from both atmospheric  $NO_3^-$  and  $NH_4^+$ 11 deposition which contribute to the overall biological N pool (cf. Durka et al. 1994; 12 13 Campbell et al. 2002), and which were of very similar magnitude and significantly 14 related in the sites studied here (Fig. 2). Nitrification is thus a key process controlling  $NO_3^-$  concentrations in upland lakes and streams. Excess  $NH_4^+$  availability not only 15 promotes nitrification (for which it is the substrate) but may inhibit  $NO_3^{-1}$ 16 immobilisation, either indirectly through preferential utilization of NH<sub>4</sub><sup>+</sup> by plants and 17 18 microbes, or directly (Bradley 2001; Rennenberg and Gessler 1999). Nevertheless, the  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> data showed that even if NO<sub>3</sub><sup>-</sup> immobilisation was reduced by NH<sub>4</sub><sup>+</sup> 19 20 availability, cycling of NO<sub>3</sub><sup>-</sup> must have occurred at one or more locations within the soil-water continuum to drastically alter the  $\delta^{18}$ O values of deposited NO<sub>3</sub><sup>-</sup>. 21

- 22
- 23

The  $\delta^{15}$ N values of surface water NO<sub>3</sub><sup>-</sup> tended to be slightly higher than those of bulk 1 deposition NO<sub>3</sub><sup>-</sup>. While the  $\delta^{15}$ N value for bulk deposition NH<sub>4</sub><sup>+</sup> was not measured in 2 this study,  $\delta^{15}N$  values of  $NH_4^+$  in atmospheric deposition tend to be lower than those 3 of  $NO_3^-$  (Heaton et al. 1997) so a possible  $NH_4^+$  deposition source for N in  $NO_3^-$ 4 produced by nitrification is unlikely to account for the elevated  $\delta^{15}$ N-NO<sub>3</sub><sup>-</sup> in surface 5 waters. Instead, the predominant control on the  $\delta^{15}N$  value of the microbial NO<sub>3</sub><sup>-</sup> is 6 7 more likely to be the large pool of soil organic N. Although the primary inputs of N to 8 soils in uncultivated areas (atmospheric deposition and N fixation) both have  $\delta^{15}N$ values close to 0‰, soil  $\delta^{15}$ N values are commonly higher than this: up to +10‰ or 9 10 more in some environments (Kendall 1998). The reasons for this are the subject of 11 debate, but largely focus on the fact that isotope fractionation associated with processes which remove N from the soil (assimilation by plants, leaching losses of 12 organic and inorganic N, denitrification and volatilization) tends to favour loss of <sup>14</sup>N. 13 leaving residual soil N enriched in <sup>15</sup>N (Handley et al. 1999; Amundson et al. 2003; 14 Kramer et al. 2003; Inglett et al. 2007; Conen et al. 2008). As a result, soils saturated 15 with N, and more 'open' or 'leaky' with respect to N loss, may in time develop high 16 soil total N  $\delta^{15}$ N values (Handley et al. 1999; Amundson et al. 2003; Inglett et al. 17 2007). This may explain why the largest difference between the  $\delta^{15}$ N values of NO<sub>3</sub><sup>-</sup> 18 in surface water and atmospheric deposition are found in the River Etherow 19 20 catchment, where the high concentrations of  $NO_3^{-1}$  in surface water may reflect a 21 greater degree of soil N saturation. The abundance of peat in the Etherow catchment could also be relevant: <sup>15</sup>N preferentially accumulates during humification of soil 22 23 organic matter (Kramer et al. 2003; Conen et al. 2008), so that mineralisation of 24 degraded peat might produce NO<sub>3</sub><sup>-</sup> with elevated  $\delta^{15}$ N.

## 2

## 3 $\delta^{18}O-NO_3^{-}$ and nitrogen biogeochemistry

Mean bulk deposition  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> values from all sites were very similar (+66.5 to +69.6‰) and fell within the ranges published from studies elsewhere (e.g. Kendall et al. 2007; Granger et al. 2008). The two end-member mixing model (Equation 2) showed that at all sites in this study a large proportion of leached NO<sub>3</sub><sup>-</sup>, which had much smaller  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> values than deposition, was therefore microbially produced by nitrification (monthly range 68-100%, annual mean 79-98%).

10

11 At the River Etherow site, this result appears to contradict a previous study on 12 nitrification potentials in catchment soils, which were very small in the highly acidic 13 upper horizons of the degraded peat soils at this site (Curtis et al. 2004).  $NO_3^{-1}$ 14 production at the Etherow must therefore be occurring elsewhere within the 15 catchment, either deeper in the soil profile, in microbial hotspots (e.g. riparian zones) 16 or perhaps in-stream (Curtis et al. 2011). At the two other sites common to the 17 previous study (Afon Gwy, Scoat Tarn) much higher nitrification potentials were 18 found in catchment soils, which is consistent with the isotope results reported here 19 showing the dominance of microbially produced NO<sub>3</sub>.

20

The relatively high proportion of atmospheric  $NO_3^-$  found in soilwater lysimeters (up to 41% at the Etherow and 69% at Lochnagar) supports the idea that progressive transformation may occur deeper in the soil profile or in-stream, particularly at the River Etherow where there is little evidence of atmospheric  $NO_3^-$  in surface waters. Other studies have also found a high proportion of atmospheric  $NO_3^-$  in lysimeters (e.g. Sebestyen et al. 2008) while increasingly "microbial" δ<sup>18</sup>O values with soil depth
 were reported by Ohte et al. (2004).

3

4 The results presented here are consistent with the  $NO_3^-$  flushing hypothesis (e.g. 5 Williard et al. 2001) whereby a large proportion of the  $NO_3^-$  observed in surface 6 waters during rainfall events has been displaced from soil waters where it was 7 produced by microbial nitrification. The relative importance of different hydrological 8 flowpaths is key to determining the proportion of untransformed atmospheric  $NO_3^{-1}$ 9 reaching surface waters, e.g. by overland flow or through preferential flowpaths (cf. Curtis et al. 2005b, 2011). A tracer <sup>15</sup>NO<sub>3</sub><sup>-</sup> addition study at the Afon Gwy (Evans et 10 11 al. 2008) showed however that NO<sub>3</sub><sup>-</sup> immobilisation could occur within hours along 12 preferential flowpaths, implying that there is potential for microbial cycling (and thus 13 isotopic transformation) of atmospheric  $NO_3^-$  even in water passing quite quickly 14 through the catchment. This could be considered analogous to the 'nutrient spiralling' 15 concept described for streams (Curtis et al. 2011).

16

#### 17 Microbial nitrification model

The inverse relationship between  $\delta^{15}N$  and  $\delta^{18}O$  values of surface waters at a number 18 19 of sites provides additional evidence that greater leaching of untransformed atmospheric NO<sub>3</sub><sup>-</sup> is related to both higher  $\delta^{18}$ O and lower  $\delta^{15}$ N values, i.e. is 20 21 consistent with the simple two end-member mixing model. As noted in the Methods section, our mixing model assumed a theoretical  $\delta^{18}$ O value for the microbial NO<sub>3</sub><sup>-</sup> 22 23 end-member, calculated assuming stoichiometry in which NO<sub>3</sub><sup>-</sup> derives one O atom from atmospheric O<sub>2</sub>, and two O atoms (i.e 67%) from water (H<sub>2</sub>O). Experiments on 24 soils incubated with <sup>18</sup>O-labeled H<sub>2</sub>O have sometimes suggested a very different 25

1	relationship, with the proportion of O derived from $H_2O$ ranging from extremes of
2	32% for possible heterotrophic nitrification (Mayer et al. 2001), up to 96% in systems
3	subject to significant O exchange with nitrite (Snider et al. 2010). However, the
4	general applicability of laboratory experiments, and prevalence of high nitrite
5	concentrations in natural systems may be questioned (Snider et al. 2010). For our
6	waters, moreover, calculations assuming 32% O or 96% O derived from $H_2O$ would
7	yield theoretical $\delta^{18}$ O values for the microbial NO <sub>3</sub> <sup>-</sup> of +14 to +16‰ or -7 to -3‰,
8	respectively. If the former range was applicable we would not be able to explain the
9	measured $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> of the great majority of our surface waters, whose values are
10	much lower than this. If the latter range was applicable it would imply that all of our
11	waters had a minimum of 10% atmospheric nitrate (i.e. none were purely microbial
12	nitrate), a feature we consider highly unlikely. In contrast, the assumption that
13	microbial NO <sub>3</sub> <sup>-</sup> derives 67% (two-thirds) of its O from water would yield theoretical
14	$\delta^{18}$ O values of +2 to +5‰, which exactly correspond to the lowest end of the
15	measured range of $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> values in our surface waters. Being mindful of the
16	uncertainties involved, we therefore calculated the $\delta^{18}$ O value of our microbial NO <sub>3</sub> <sup>-</sup>
17	from Equation 1.

19 Between site differences in streamwater  $\delta^{18}O-NO_3^{-1}$ 

20 While comparison of data from lake and stream sites must consider residence times in 21 lakes and direct atmospheric deposition to lake surfaces (see below), differences in the 22 transport of atmospheric NO<sub>3</sub><sup>-</sup> to streams may be assessed using inflow data for the 23 lake sites to compare with the stream catchments, i.e. comparing streams in each 24 catchment (Fig. 7). Values of  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> were significantly lower in Rose Clough and 25 Swan Clough than all other streams including the River Etherow (p<0.05), into which

1	they flow. Scoat Inflow 1 had a significantly greater $\delta^{18}$ O-NO <sub>3</sub> <sup>-</sup> than the River
2	Etherow. No other significant differences were found. The greatest proportions of
3	atmospheric NO <sub>3</sub> <sup>-</sup> were found in streams at Lochnagar (13%), then Scoat Tarn (Inflow
4	1 = 11%, Inflow $2 = 9%$ ) and the main channel of the Etherow (6%). The mean
5	proportion was only 2-3% the Afon Gwy and the Etherow tributaries, Rose Clough
6	and Swan Clough.

7

8	While the differences between sites are not significant, the higher proportions of
9	atmospheric $NO_3^-$ leaching are associated with the highest altitude, steepest
10	catchments of Lochnagar and Scoat Tarn (Table 1). Lochnagar also has a particularly
11	high proportion of bare rock compared with the other catchments. The streamwater
12	data are therefore consistent with previous studies showing a greater amount of
13	hydrological bypass transportation and/or lower microbial immobilisation of
14	atmospheric NO <sub>3</sub> <sup>-</sup> associated with these physical catchment attributes (e.g. Evans et
15	al. 2004, 2006; Helliwell et al. 2007).
16	

17 Importance of streamflow and hydrological flowpaths

18 Very detailed isotopic studies in gauged catchments in North America have

19 demonstrated the importance of stream flow conditions and the timing of sampling

20 relative to baseflow and extreme flow events in affecting the proportion of

21 atmospheric NO<sub>3</sub><sup>-</sup> in surface waters (e.g. Sebestyen et al. 2008, 2009). In particular,

- 22 several studies in snowmelt dominated systems have shown that a relatively large
- 23 proportion of annual NO<sub>3</sub><sup>-</sup> fluxes are transported during very high flow events e.g.
- 24 during snowmelt (Campbell et al. 2002; Schiff et al. 2002; Ohte et al. 2004; Pardo et
- 25 al. 2004; Sebestyen et al. 2008, 2009; Goodale et al. 2009), and at these times the

- proportion of atmospheric NO<sub>3</sub><sup>-</sup> may be elevated. Sampling regimes which do not
   include these high flow events may therefore underestimate both total NO<sub>3</sub><sup>-</sup> fluxes and
   the proportion of untransformed atmospheric NO<sub>3</sub><sup>-</sup> exported from catchments.
- 4

5 In our study, only one of the four catchments (Afon Gwy) was instrumented for flow 6 measurement throughout the study period. At the Afon Gwy, flow was measured 7 every 15 minutes and streamwater sampled weekly. The flows sampled during the 8 monthly isotope sampling programme cover a wide range of the mean flows observed 9 on a daily basis but do not represent the most extreme conditions (see supplementary Fig. S1). However, a sample from 11<sup>th</sup> November 2004 was taken during very high 10 11 flow (only 3% of daily mean flows were higher over the sampling year) and yet 12 yielded insufficient NO<sub>3</sub><sup>-</sup> for dual isotope analysis. Furthermore, high NO<sub>3</sub><sup>-</sup> 13 concentrations were seen under both high and low flow conditions (Fig. S2). Rainfall 14 data for the study catchments show that high rainfall may occur throughout the year 15 and does not appear to account for seasonal variations in the proportion of 16 atmospheric NO<sub>3</sub><sup>-</sup> (Fig. S3). At Scoat Tarn, the highest rainfall recorded for a twoweek period during the study (20.5mm per day for sample dated 15<sup>th</sup> November 2005) 17 18 corresponded with the lowest proportion of atmospheric NO<sub>3</sub><sup>-</sup> recorded for the site 19 (Table S4).

20

Therefore, while our monthly sampling regime could not capture extreme flow events and may therefore underestimate annual mean  $NO_3^-$  fluxes, there is no evidence from the highest resolution data at the Afon Gwy that any underestimate of either  $NO_3^$ fluxes or the contribution of atmospheric  $NO_3^-$  is likely to be large. Indeed, the highresolution event-based study of Sebestyen (2009) found that although atmospheric

contributions to streamwater NO<sub>3</sub><sup>-</sup> could increase greatly during high flow events, the
effect of this increase on total annual contributions was small. Hence while our study
cannot provide detailed information about atmospheric contributions to NO<sub>3</sub><sup>-</sup> leaching
fluxes on an event basis or during extremes of flow, it does provide robust estimates
of the importance of direct atmospheric NO<sub>3</sub><sup>-</sup> leaching on an average annual basis.

Finally, the issue of short-lived, very high flow events is less important for lakes with
long residence times compared to streams. The average residence time of water in
Lochnagar is 242 days (Jenkins et al. 2007) while at Scoat Tarn it is 49 days. Hence
lakes are ideal integrators of both varying bulk deposition inputs and flow conditions
in inflow streams and show a damped temporal pattern in both NO<sub>3</sub><sup>-</sup> concentrations
and atmospheric contributions relative to streams (Fig. 6).

13

14 Lochnagar is the most snow-dominated site in our study at the highest altitude, with 15 approximately 20% of precipitation falling as snow in an average year (Jenkins et al. 16 2007). Periods of significant snowmelt usually occur in late April/early May but short 17 periods of extremely high flow (1-2 days) occur throughout the year (Jenkins et al. 18 2007). This study does indeed show that both NO<sub>3</sub><sup>-</sup> concentrations in the lake and the 19 greatest proportion of atmospheric  $NO_3^-$  occur at this time of year, i.e. the monthly 20 sampling regime has successfully captured this seasonal pattern. It must however be 21 recognized that our data could underestimate the total contribution of directly leached 22 atmospheric NO<sub>3</sub><sup>-</sup> and especially the maximum contributions which may be attained 23 during extreme hydrological events.

24

### 2 Lakes versus streams

3 It might be expected that a higher proportion of untransformed atmospheric  $NO_3^{-1}$ would be found in lakes relative to streams because of direct deposition to lake 4 5 surfaces, bypassing the terrestrial processing of deposition inputs. At the two lake 6 sites in this study, Scoat Tarn and Lochnagar, both lake outflows and major inflow streams were sampled at the same time. Mean values of  $\delta^{15}$ N were slightly lower in 7 lake outflows relative to inflow streams (Fig. 5). Values of  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> were 8 9 significantly higher in Scoat Tarn (mean=+12.6%) than in Inflow 2 (mean=+9.0%, 10 Fig. 8: t=3.31, p=0.009) and while also higher than in Inflow 1 (+10.3‰) on average, this difference was not significant (t=2.35, p=0.095). At Lochnagar  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> was 11 significantly higher in the loch outflow (mean=+16.7%) than in the inflow spring 12 13 (mean = +10.9%, t=3.36, p=0.005).

14

15 These data provide isotopic evidence that there was indeed a greater proportion of 16 untransformed atmospheric NO<sub>3</sub><sup>-</sup> in the two lakes compared with their inflow streams, 17 presumably due to direct deposition to the lake surface. For Scoat Tarn the annual 18 mean proportion of atmospheric  $NO_3^-$  in the lake was 15% compared with 11% in 19 inflow 1 and 9% in inflow 2, i.e. was greater by 4-6% in the lake outflow. The surface 20 area of Scoat Tarn is 4.3 ha and represents 5% of the total catchment area. Likewise, 21 at Lochnagar the mean proportion of atmospheric  $NO_3^-$  in the loch was 21% compared 22 with 13% in the inflow spring i.e. 8% greater in the outflow, while the loch has a 23 surface area of 9.9 ha representing 9% of total catchment area. These figures are 24 remarkably consistent with the idea that direct deposition to lake surfaces contributes 25 to catchment scale leaching of untransformed atmospheric  $NO_3^{-1}$  in direct proportion to

1	lake:catchment area ratios. There appears to be relatively little cycling of directly
2	deposited $NO_3^-$ within the lakes themselves since all of the directly deposited $NO_3^-$ is
3	recovered in the lake outflows. An alternative hypothesis is that the greater proportion
4	of untransformed NO <sub>3</sub> <sup>-</sup> in lake outflows relative to their inflow streams simply reflects
5	elevated, event-based inputs of atmospheric NO3 <sup>-</sup> in streams which are not captured in
6	the monthly streamwater sampling but which increase the overall proportion of
7	atmospheric $NO_3^-$ in the receiving lakes. Further work on a greater number of lakes
8	and associated inflows would be required to test these hypotheses.
9	
10	While catchment hydrology is a key factor determining the delivery of untransformed

10 While catchment hydrology is a key factor determining the delivery of untransformed 11 atmospheric  $NO_3^-$  to upland streams, the surface area of upland lakes relative to their 12 catchments is also an important factor controlling their exposure to direct inputs of 13 atmospheric  $NO_3^-$ . Lake:catchment ratio must therefore also be an important 14 determinant of exposure to other pollutants subject to terrestrial retention and 15 processing, e.g.  $NH_4^+$  deposition.

16

17 Although microbially produced  $NO_3^-$  dominates in the four sites studied here, the 18 persistent contribution of atmospheric NO<sub>3</sub><sup>-</sup> all year round in lakes and especially 19 inflow streams indicates a chronic atmospheric NO<sub>3</sub><sup>-</sup> leaching problem which has seldom been seen in other isotopic studies, where streamwater  $\delta^{18}$ O-NO<sub>3</sub><sup>-</sup> falls within 20 21 the microbial range for much of the year (e.g. Piatek et al. 2005; Campbell et al. 2006; 22 Hales et al. 2007; Barnes et al. 2008; Sebestyen et al. 2008; Burns et al. 2009; 23 Goodale et al. 2009). Notable exceptions showing persistent contributions of 24 atmospheric NO<sub>3</sub><sup>-</sup> (though still predominantly microbial) through most of the year 25 include the studies of Spoelstra et al. (2001) in forested catchments in the Turkey

Lakes Watershed, Canada (10-15% atmospheric) and Campbell et al. (2002) in alpine
watersheds in the Rocky Mountains (up to 50% atmospheric during snowmelt, but
much less at other times - and these catchments had >80% bare rock). Hence the
results of the present study show a greater (and more persistent) atmospheric
contribution to NO<sub>3</sub><sup>-</sup> leaching in terms of annual fluxes than most other isotopic
studies.

7

#### 8 Conclusions

9

10 The major proportion of  $NO_3^-$  observed in surface waters in upland catchments, for 11 which atmospheric deposition is the only source of anthropogenic N inputs, is derived 12 from microbial production. While mindful of the uncertainty in knowing the precise  $\delta^{18}$ O value for microbial NO<sub>3</sub><sup>-</sup>, our best estimates for late winter / early spring peak 13 14 contributions of untransformed atmospheric NO<sub>3</sub><sup>-</sup> at the four study sites ranged from 15 5% to around 30%, but annual means varied from just 2-13% for streams and 15-21% 16 for lakes. Crucially, this means that deposition of reduced N compounds cannot be 17 ignored as a possible source of the N leached as NO<sub>3</sub>, since only a small proportion of 18 surface water NO<sub>3</sub><sup>-</sup> derives directly from NO<sub>3</sub><sup>-</sup> in bulk deposition. For upland streams, 19 other studies have shown that catchment hydrology and linked physical attributes such 20 as slope, bare rock and amount of organic matter in soils, are key determinants of the 21 delivery of atmospheric  $NO_3^-$  to surface waters, while we show here that for lakes an 22 important additional source is direct deposition to lake surfaces. In the two lakes 23 studied here, all the directly deposited  $NO_3^-$  appeared to be recovered from the 24 outflows. Microbial NO<sub>3</sub><sup>-</sup> production is however the major overall determinant of 25 surface water  $NO_3^-$  concentrations in upland catchments, and hence future trends in

NO<sub>3</sub><sup>-</sup> leaching will be intimately linked with global change impacts on microbial
 macronutrient cycling.

3

4 While the greatest proportion of untransformed atmospheric NO<sub>3</sub><sup>-</sup> was found in the 5 headwater lakes, the greatest leaching flux of NO<sub>3</sub><sup>-</sup> in both absolute terms and as a 6 proportion of deposition inputs was found at the River Etherow, which has the lowest 7 altitude, gentlest slopes and greatest proportion of organic soils. Hence, there is a 8 clear distinction to be made between the physical attributes of a site which may allow 9 direct leaching of atmospheric inputs to surface waters, and the nitrogen saturation 10 status of a site. In the case of the River Etherow, the catchment is a net source of  $NO_3^{-1}$ 11 despite very high NO<sub>3</sub><sup>-</sup> concentrations in deposition and has negligible scope for 12 direct leaching of deposition inputs without biological cycling.

13

14

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22

23

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2 Figure 1: Location, catchment outlines and sampled subcatchments streams at a) Scoat Tarn, b)

- 3 Lochnagar, c) Afon Gwy and d) River Etherow. Subcatchments of sampled streams are shaded.
- 4 Contour intervals in metres above sea level.





3 Figure 2: Inorganic nitrogen concentrations in bulk deposition (NH<sub>4</sub><sup>+</sup>, NO<sub>3</sub><sup>-</sup>) and surface waters

- 4 (NO<sub>3</sub><sup>-</sup>)
- 5
- 6



Figure 3: Box and whisker plot of δ<sup>15</sup>N-NO<sub>3</sub><sup>-</sup> in bulk deposition at the four study sites. Boxes
represent median and inter-quartile range, whiskers extend to 1.5 × inter-quartile range and
individual points represent data outside this range





1 Figure 5: Results of dual isotope analysis ( $\delta^{15}$ N vs SMOW and  $\delta^{18}$ O vs Air, ‰) of NO<sub>3</sub><sup>-</sup> in surface



2 water, soilwater and bulk deposition samples



3 Figure 6: Seasonal variation in contribution of atmospheric NO<sub>3</sub><sup>-</sup> to surface waters









- 1 Table 1: Description of study sites with selected long-term mean surface water
- 2 chemistry, April 1998 March 2006 (from Monteith and Shilland 2007) and bulk
- 3 deposition chemistry (1999-2006; source Lawrence et al. 2008)

Site:	Afon Gwy	<b>River Etherow</b>	Scoat Tarn	Lochnagar
Sampling location	Flume	Main channel	Outflow	Outflow
OS grid Ref	SN82400, 85350	SK11557, 99691	NY15819, 10344	NO25317, 86268
Altitude (m)	385	280	595	788
Max. altitude	741	633	841	1155
Catchment area (ha)	389	1295	87	109
Bare ground	<2%	4%	<2%	24%
Sampled inflows /	LB7a:	Rose Clough:	1 (Scoat Fell):	Spring:
tributaries	SN81450, 86500	SK12181, 99532	NY15866, 10423	NO25279, 85770
		Swan Clough:	2 (Red Pike):	
		SK11909, 99453	NY15935, 10503	
Rainfall (mm)	2021	1004	2313	1279
Concentration (µeq l <sup>-1</sup> )				
NO3 <sup>-</sup> (water body)	6.9	42.0	15.5	18.5
NO <sub>3</sub> <sup>-</sup> (bulk depn.)	9.5	28.3	14.7	20.8
NH4 <sup>+</sup> (bulk depn.)	11.8	31.8	18.3	17.0
Deposition flux (kgN ha <sup>-1</sup> yr <sup>-1</sup> )				
NO <sub>3</sub>	2.7	3.9	4.8	3.8
DIN $(NO_3^- + NH_4^+)$	6.0	8.4	10.6	6.7

1 Table 2: Mean concentrations (µeq I<sup>-1</sup>) and fluxes (kgN ha<sup>-1</sup> yr<sup>-1</sup>) of N species in surface waters

2 and bulk deposition. Figures in parenthese indicate flow-weighting (surface waters) or volume

3 weighting (bulk deposition). No flow data were available at the River Etherow or Scoat Tarn.

4 NH<sub>4</sub><sup>+</sup> is negligible in surface waters. See text for further details.

5

	S	Stream /	Rainfall	ЕТ	NO -	Bulk deposition			Deposition flux		
	Sampling				NO <sub>3</sub>	concentration					
Site	Period	lake NO <sub>3</sub> -	(mm)	(%)	flux	$\mathbf{NH_4}^+$	NO <sub>3</sub> -	DIN	NH4 <sup>+</sup>	NO <sub>3</sub> -	DIN
Afon Gwy	07/9/04-	6.0	2050	15	1.4	19.4	15.5	22.8	3.6	2.9	6.5
	06/9/05	(6.8)				(12.7)	(10.1)				
River Etherow	10/8/05-	39.6	984	29	3.9	45.2	37.8	56.1	4.2	3.5	7.7
	08/8/06	(n/a)				(30.7)	(25.4)				
Scoat Tarn	9/8/05-	13.8	2316	19	3.6	28.4	24.1	36.5	6.9	5.0	11.9
	04/8/06	(n/a)				(21.1)	(15.4)				
Lochnagar	10/8/05-	15.8	1099	20	2.0	15.3	20.4	32.1	2.2	2.8	5.0
	10/8/06	(15.2)				(14.4)	(17.7)				

6

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