- 1 Stable isotopic evidence for the excess leaching of unprocessed atmospheric
- 2

nitrate from forested catchments under high nitrogen saturation

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3 Abstract

4	Owing to the elevated loading of nitrogen through atmospheric deposition, some
5	forested ecosystems become nitrogen saturated, from which elevated levels of nitrate
6	are exported. The average concentration of stream nitrate eluted from upstream and
7	downstream of the Kasuya Research FK forested catchments (FK1 and FK2 catchments)
8	in Japan wereas more than 90 μ M, implying that these forested catchments were under
9	nitrogen saturation. To verify that these forested catchments were under the nitrogen
10	saturation, we determined the export flux of unprocessed atmospheric nitrate relative to
11	the entire deposition flux (M_{atm}/D_{atm} ratio) in these catchments, because the M_{atm}/D_{atm}
12	ratio has recently been proposed as a reliable index to evaluate nitrogen saturation in
13	forested catchments. Specifically, we determined the temporal variation in the
14	concentrations and stable isotopic compositions, including $\Delta^{17}O$, of stream nitrate in
15	the FK catchments for more than 2 years. In addition, for comparison, the same
16	parameters were also monitored in the Shiiba Research MY forested catchment (MY
17	catchment) in Japan during the same period, where the average stream nitrate
18	concentration was low, less than 10 μ M. While showing the average nitrate
19	concentrations of 109.5, $90.994.2$, and 7.1 μ M in FK1, FK2, and MY, respectively, the
20	catchments showed average Δ^{17} O values of +2.6, +1.57, and +0.6 ‰ in FK1, FK2, and
21	MY, respectively. Thus, the average concentration of unprocessed atmospheric nitrate
22	([NO _{3⁻atm}]) was estimated to be 10.8, $5.16.1$, and 0.2 µM in FK1, FK2, and MY,
23	respectively, and the M_{atm}/D_{atm} ratio was estimated to be <u>14.143.9</u> , <u>6.67.9</u> , and 1. <u>32</u> %

in FK1, FK2, and MY, respectively. The estimated Matm/Datm ratio in FK1 (14.113.9%) 24 was the highest ever reported from temperate forested catchments monitored for more 25 than 1 year. Thus, we concluded that nitrogen saturation was responsible for the 26 enrichment of stream nitrate in the FK catchments, together with the elevated NO3⁻atm 27 leaching from the catchments. While the stream nitrate concentration ([NO₃⁻]) can be 28 affected by the amount of precipitation, the Matm/Datm ratio is independent of the amount 29 of precipitation; thus, the Matm/Datm ratio can be used as a robust index for evaluating 30 31 nitrogen saturation in forested catchments.

32

33 1 Introduction

Nitrate is important as a nitrogenous nutrient in the biosphere. Traditionally, forested 34 ecosystems have been considered as nitrogen limited (Vitousek and Howarth, 1991). 35 36 However, owing to the elevated loading of nitrogen through atmospheric deposition, some forested ecosystems become nitrogen saturated (Aber et al., 1989), from which 37 elevated levels of nitrate are exported (Mitchell et al., 1997; Peterjohn et al., 1996). 38 39 Such excessive leaching of nitrate from forested catchments degrades water quality and 40 causes eutrophication in downstream areas (Galloway et al., 2003; Paerl and Huisman, 41 2009). Thus, evaluating the stage of nitrogen saturation in each forested catchment including its temporal variation, is critical for sustainable forest management, 42 especially for forested ecosystems under high nitrogen deposition. 43

44 Both concentration and seasonal variation of stream nitrate have been used as indexes

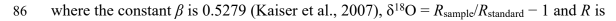
to evaluate the nitrogen saturation of each forested catchment in past studies (Aber, 1992; Rose et al., 2015; Stoddard, 1994). A forested stream eluted from Fernow Experimental Forest USA, for instance, showed an elevated average nitrate concentration of 60 μ M, along with the absence of a seasonal variation in the stream nitrate concentration, so the forest was classified into stage 3, the highest stage of nitrogen saturation (Rose et al., 2015).

However, using both the concentration level (high or low) and seasonal variation 51 (clear or absent) of stream nitrate as indexes to evaluate nitrogen saturation has 52 limitations, including the following (1) seasonal variation of soilstream nitrate can be 53 buffered by groundwater within forests long residence time under humid, temperate 54 climates such as Japan, so the, so that the seasonal variation in stream nitrate 55 concentrations is unclear in stream nitrate concentration in Japan, even in normal forests 56 57 under the nitrogen saturation stage of 0-(Mitchell et al., 1997); and (2) the stream nitrate concentration can be enriched or diluted depending on the volume of rainfall, so the 58 59 concentration level can be high in low precipitation area irrespective of the stage of 60 nitrogen saturation.

Nakagawa et al. (2018) lately proposed that the M_{atm}/D_{atm} ratio, the export flux of unprocessed atmospheric nitrate (M_{atm}) relative to the deposition flux of $NO_3^{-}_{atm}$ (D_{atm}), can be an alternative, more robust index for evaluating nitrogen saturation in each forested catchment, because the M_{atm}/D_{atm} ratio directly reflects the demand for atmospheric nitrate deposited onto each forested catchments as a whole, and thus reflect the nitrogen saturation in each forested catchment. That is, we can expect high M_{atm}/D_{atm} ratios in forested catchments under nitrogen saturation and low M_{atm}/D_{atm} ratios in forested catchments with nitrogen deficiency.

69 To estimate the M_{atm}/D_{atm} ratio accurately and precisely in each forested catchment, the fraction of unprocessed atmospheric nitrate (NO3⁻atm) in the stream needs to be 70 estimated accurately and precisely. In recent, tTriple oxygen isotopic compositions of 71 nitrate (Δ^{17} O) have <u>recently</u> been used as a conservative tracer of NO₃⁻_{atm} deposited 72 onto each forested catchment (Inoue et al., 2021; Michalski et al., 2004; Nakagawa et 73 al., 2018; Tsunogai et al., 2014; Ding et al., 2022), showing distinctively different Δ^{17} O 74 from that of remineralized nitrate (NO_{3^{re}}), derived from organic nitrogen through 75 general chemical reactions, including microbial N mineralization and microbial 76 nitrification. While NO_{3^{re}}, the oxygen atoms of which are derived from either 77 78 terrestrial O₂ or H₂O through microbial processing (i.e., nitrification), always shows the relation close to the "mass-dependent" relative relation between ¹⁷O/¹⁶O ratios and 79 ¹⁸O/¹⁶O ratios; NO_{3⁻atm} displays an anomalous enrichment in ¹⁷O reflecting 80 81 oxygen atom transfers from atmospheric ozone (O_3) during the conversion of NO_X to NO_{3⁻atm} (Alexander et al., 2009; Michalski et al., 2003; Morin et al., 2011; Nelson et 82 al., 2018). As a result, the Δ^{17} O signature defined by the following equation (Kaiser et 83 al., 2007) enables us to distinguish NO_{3⁻atm} (Δ^{17} O > 0) from NO_{3⁻re} (Δ^{17} O = 0): 84 $1 + \delta^{17}$ O

85
$$\Delta^{17}O = \frac{1+0}{(1+\delta^{18}O)^{\beta}} - 1$$
 (1)



87	the ¹⁸ O/ ¹⁶ O ratio (or the ¹⁷ O/ ¹⁶ O ratio in the case of δ^{17} O or the ¹⁵ N/ ¹⁴ N ratio in the case
88	of $\delta^{15}N$) of the sample and each standard reference material. In addition, $\Delta^{17}O$ is almost
89	stable during "mass-dependent" isotope fractionation processes within terrestrial
90	ecosystems. Therefore, while the $\delta^{15}N$ or $\delta^{18}O$ signature of $NO_{3^{-}atm}$ can be overprinted
91	by the biological processes subsequent to deposition, $\Delta^{17}O$ can be used as a robust tracer
92	of unprocessed $NO_3^{-}_{atm}$ to reflect its accurate mole fraction within total NO_3^{-} ,
93	regardless of the progress of the partial metabolism (partial removal of nitrate through
94	denitrification and assimilation) subsequent to deposition (Michalski et al., 2004;
95	Nakagawa et al., 2013, 2018; Tsunogai et al., 2011, 2014, 2018).
96	Past studies reported that the maximum concentration of stream nitrate was 58.4 μ M
97	in the KJ forested catchment in Japan, with the maximum value of the M_{atm}/D_{atm} ratio
98	was 9.4 % (Nakagawa et al., 2018; Sase et al., 2022). Whether the index of the $M_{\text{atm}}/D_{\text{atm}}$
99	ratio can be applied to forested catchments, where the leaching of stream nitrate is much
100	higher than the KJ forested catchment, remained unclarified. Besides, the advantages
101	of the Matm/Datm ratio within the past indexes of nitrogen saturation have not been
102	discussed.
103	In recent, Chiwa (2021) has recently reported the enrichment of nitrate of more than
104	90 μ M on the annual average in forested streams eluted from the FK catchments (FK1
105	and FK2) in Kasuya Research Forest, Kyushu University, Japan (Figs. 1a and 1b). The
106	observed enrichment of stream nitrate implied that these forested catchments were

107 under nitrogen saturation. Thus, in this study, we determined the M_{atm}/D_{atm} ratio in the

108	FK1 and FK2 forested catchments by monitoring both the concentration and Δ^{17} O of
109	stream nitrate for more than 2 years to verify that these forested catchments were under
110	nitrogen saturation. For comparison, the MY forested catchment in Shiiba Research
111	Forest, Kyushu University, Japan (Figs. 1a and 1c), was also monitored during the same
112	period, where the average stream nitrate concentration was low (less than 10 μM).
113	Furthermore, the $M_{\text{atm}}/D_{\text{atm}}$ ratios in these forested catchments were compared with
114	those reported in past studies to verify the reliability of the M_{atm}/D_{atm} ratio as an index
115	of nitrogen saturation.
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117	2 Methods
110	2.1 Study sites
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MY forested catchment, respectively. Details on the studied forested catchments have
been described in the past studies (Chiwa, 2020, 2021).

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132 2.2 Sampling

The stream water eluted from the FK1 (14 ha), FK2 (62 ha), and MY (43 ha) 133 forested catchments were collected about once every month in principle from 2019/11 134 to 2021/12 (Fig. 1). At the FK catchments, stream water was collected at upstream 135 136 (FK1station A) and downstream (FK2station B) locations (Fig. 1b). At the MY catchment, stream water was collected at station C (Fig. 1c). Samples of stream water 137 to determine the concentration and stable isotopic compositions (δ^{15} N, δ^{18} O, and Δ^{17} O) 138 of stream nitrate were collected manually in bottles washed with deionized water before 139 sampling and then rinsed at least twice with the sample before sampling at each 140 141 sampling site.

142

143 2.3 Analysis

All the stream water samples were passed through a membrane filter (pore size 0.45 μ m) within two days after sampling and stored in a refrigerator (4 °C) until analysis. The concentrations of nitrate were measured by ion chromatography (Prominence HIC-SP, Shimadzu, Japan). To determine the stable isotopic compositions of nitrate in the stream water samples, nitrate in each sample was chemically converted to N₂O using a method originally developed to determine the ¹⁵N/¹⁴N and ¹⁸O/¹⁶O ratios

150	of seawater and freshwater nitrate (McIlvin and Altabet, 2005) that was later modified
151	(Konno et al., 2010; Tsunogai et al., 2011; Yamazaki et al., 2011). In brief, 11 mL of
152	each sample solution was pipetted into a vial with a septum cap. Then, 0.5 g of spongy
153	cadmium was added, followed by 150 μL of a 1 M NaHCO3 solution. The sample was
154	then shaken for 18-24 h at a rate of 2 cycles s^{-1} . Then, the sample solution (10 mL) was
155	decanted into a different vial with a septum cap. After purging the solution using high-
156	purity helium, 0.4 mL of an azide-acetic acid buffer, which had also been purged using
157	high-purity helium, was added. After 45 min, the solution was alkalinized by adding
158	0.2 mL of 6 M NaOH. Then, the stable isotopic compositions ($\delta^{15}N$, $\delta^{18}O$, and $\Delta^{17}O$) of
159	the N ₂ O in each vial were determined using the continuous-flow isotope ratio mass
160	spectrometry (CF-IRMS) system at Nagoya University. The analytical procedures
161	performed using the CF-IRMS system were the same as those detailed in previous
162	studies (Hirota et al., 2010; Komatsu et al., 2008a). The obtained values of $\delta^{15}N$, $\delta^{18}O$,
163	and Δ^{17} O for the N ₂ O derived from the nitrate in each sample were compared with those
164	derived from our local laboratory nitrate standards to calibrate the values of the sample
165	nitrate to an international scale and to correct for both isotope fractionation during the
166	chemical conversion to N_2O and the progress of oxygen isotope exchange between the
167	nitrate derived reaction intermediate and water (ca. 20 %). The local laboratory nitrate
168	standards used for the calibration had been calibrated using the internationally
169	distributed isotope reference materials (USGS-34 and USGS-35). In this study, we
170	adopted the internal standard method to calibrate the stable isotopic compositions of

171	sample nitrate (Ding et al., 2022; Nakagawa et al., 2013, 2018; Tsunogai et al., 2014)
172	Specifically, three kinds of the local laboratory nitrate standards were used in this study,
173	which were named to be GG01 ($\delta^{15}N = -3.07 \%$, $\delta^{18}O = +1.10 \%$, and $\Delta^{17}O = 0 \%$),
174	<u>HDLW02 ($\delta^{15}N = +8.94 \%$, $\delta^{18}O = +24.07 \%$), and NF ($\Delta^{17}O = +19.16 \%$), which the</u>
175	GG01 and the HDLW02 were used to determine the $\delta^{15}N$ and $\delta^{18}O$ of stream nitrate,
176	and the GG01 and the NF was used to determine the Δ^{17} O of stream nitrate. The GG01,
177	HDLW02, and NF had been calibrated using the internationally distributed isotope
178	reference materials (USGS 34 and USGS 35). The oxygen exchange rate between
179	nitrate and water during the chemical conversion was calculated through Eq. (2):
180	$\underline{\text{Oxygen exchange rate (\%)}} = \Delta^{17} O(N_2 O)_{\text{NF}} / \Delta^{17} O(NO_3)_{\text{NF}} $ (2)
181	where the $\Delta^{17}O(N_2O)_{NF}$ denote the $\Delta^{17}O$ value of N_2O that convert from the NF
182	nitrate, the $\Delta^{17}O(NO_3)_{NF}$ denote the $\Delta^{17}O$ value of NF nitrate ($\Delta^{17}O = +19.16$ ‰)
183	(Tsunogai et al., 2016; Nakagawa et al., 2013, 2018; Ding et al., 2022).
184	The $\delta^2 H$ and $\delta^{18} O$ values of $H_2 O$ of the stream water samples were analyzed using
185	the cavity ring-down spectroscopy method by employing an L2120-i instrument
186	(Picarro Inc., Santa Clara, CA, USA) equipped with an A0211 vaporizer and
187	autosampler. The errors (standard errors of the mean) in this method were $\pm 0.5\%$ for
188	$\delta^2 H$ and $\pm 0.1\%$ for $\delta^{18} O.$ Both the VSMOW and standard light Antarctic precipitation
189	(SLAP) were used to calibrate the values to the international scale. The $\delta^{18}O$ values of
190	$\rm H_2O$ were used to calibrate the differences in $\delta^{18}O$ of $\rm H_2O$ between the samples and
191	those our local laboratory nitrate standard samples (Tsunogai et al., 2010, 2011, 2014).

To determine whether the conversion rate from nitrate to N₂O was sufficient, the 192 concentration of nitrate in the samples was determined each time we analyzed the 193 isotopic composition using CF-IRMS based on the N_2O^+ or O_2^+ outputs. We adopted 194 the δ^{15} N, δ^{18} O, and Δ^{17} O values only when the concentration measured via CF-IRMS 195 correlated with the concentration measured via ion chromatography prior to isotope 196 analysis within a difference of 10 %. We repeated the analysis of δ^{15} N, δ^{18} O, and Δ^{17} O 197 values for each sample at least three times to attain high precision. All samples had a 198 nitrate concentration of greater than 3.5 µM, which corresponded to a nitrate quantity 199 greater than 35 nmol in a 10 mL sample. Thus, all isotope values presented in this study 200 have an error (standard error of the mean) better than ± 0.2 % for δ^{15} N, ± 0.3 % for δ^{18} O, 201 and ± 0.1 ‰ for Δ^{17} O. 202 Nitrite (NO_2^{-}) in the samples interferes with the final N₂O produced from nitrate 203 204 because the chemical method also converts NO₂⁻ to N₂O (McIlvin and Altabet, 2005). Therefore, it is sometimes necessary to remove NO_2^- prior to converting nitrate to N_2O . 205 In this study, however, we skipped the processes for removing NO₂⁻ because all the 206 207 stream samples analyzed for stable isotopic composition had NO₂⁻ concentrations lower than the detection limit (0.05 μ M). 208

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210 2.4 Deposition rate of atmospheric nitrate

The annual deposition rate of atmospheric nitrate (D_{atm}; total dry and wet deposition rate of atmospheric nitrate) in each catchment was estimated using the annual "bulk"

213	deposition rate of atmospheric nitrate (D _{bulk}) calculated in Chiwa (2020) at each
214	catchment by multiplying the volume-weighted mean concentration of nitrate in the
215	bulk deposition samples collected every 2 weeks at each catchment for 10 years (from
216	2009/1 to 2018/12) by the annual amount of precipitation. The bulk deposition samples
217	were samples those accumulated in a plastic bucket installed in an open site of each
218	catchment 55 cm above the ground. The concentrations of nitrate in these samples were
219	measured by ion chromatography. The distances between the monitoring sites of bulk
220	deposition in the FK1, FK2, and MY forested catchments and the stations of stream
221	water sampling (stations A, B, and C) were 3.9, 2.9, and 4.5 km, respectively. The
222	concentrations of nitrate in the bulk deposition samples were measured by ion
223	chromatography.

The D_{bulk} determined through this method, however, is less than D_{atm} (Aikawa et al., 2003) because the dry deposition velocities of gases and particles on the water surface of the plastic bucket are smaller than those on the forest (Matsuda, 2008). Thus, we corrected the differences by using Eq. (32) to estimate D_{atm} from D_{bulk} :

228
$$D_{atm} = D_{bulk} - D_{dry}(W) + D_{dry}(F)$$
 (32)

where D_{dry}(W) and D_{dry}(F) denote the annual dry deposition rates onto water and forest,
respectively.

The $D_{dry}(W)$ and $D_{dry}(F)$ at each catchment were determined using an inferential method (Endo et al., 2011) through Eqs. (<u>43</u>) and (<u>54</u>), respectively:

233
$$D_{dry}(W) = [NO_3^- a_{tm}]_{gas} \times V_{gas}(W) + [NO_3^- a_{tm}]_p \times V_p(W)$$
 (43)

234
$$D_{dry}(F) = [NO_3^- atm]_{gas} \times V_{gas}(F) + [NO_3^- atm]_p \times V_p(F)$$
 (54)

where $[NO_3^{-}atm]_{gas}$ denotes the concentration of gaseous nitrate in air; $[NO_3^{-}atm]_p$ 235 denotes the concentration of particle nitrate in air; Vgas(W) and Vgas(F) denote the 236 237 deposition velocities of gaseous nitrate on the water surface and forest, respectively; and $V_p(W)$ and $V_p(F)$ denote the deposition velocities of particulate nitrate on the water 238 surface and forest, respectively. Those determined by Chiwa (2010) using the annular 239 denuder method from 2006/5 to 2007/4 were used for the $[NO_3^-]_{gas}$ and $[NO_3^-]_p$ in the 240 FK catchments. Those determined by the National Institute for Environmental Studies 241 (Environmental Laboratories Association of Japan, 2017) using the filter-pack method 242 at Miyazaki (31°83'N, 131°42'E) from 2011 to 2017 were used for the [NO3-]gas and 243 $[NO_3^-]_p$ in the MY catchment. The $V_{gas}(F)$, $V_{gas}(W)$, $V_p(F)$, and $V_p(W)$ of each 244 catchment were determined by applying the estimation file for dry deposition (Matsuda, 245 2008; 246

http://www.hro.or.jp/list/environmental/research/ies/katsudo/acid_rain/kanseichinchak u/kanseichinchaku.html), where V_{gas} and V_p were calculated using the meteorological data of wind speed, temperature, humidity, radiation, and cloud amount and land use. The meteorological data monitored by Japan Meteorological Agency at the nearest Fukuoka station (33°34'N, 130°22'E) and Miyazaki station (31°56'N, 131°24'E) from 2009 to 20<u>21</u>48 were used for the FK and MY catchments, respectively. The forested land use of 100 % was chosen for each area.

255 2.5 Flux of stream water

The flux of stream water (F_{stream}) in each catchment was not measured <u>directly fully</u> in this study. Instead, the water balance in each catchment was used to estimate F_{stream} , assuming that the outflux of water from the study catchments to deep groundwater was negligible:

$$260 F_{stream} = P - E (65)$$

where P denotes the annual average precipitation and E denotes the annual evapotranspiration flux of water in each catchment. In this paper, the equation obtained by Komatsu et al. (2008) was used to estimate the E of the FK and MY catchments. Details on this equation are shown below.

Komatsu et al. (2008) compiled the annual flux of evapotranspiration determined in 43 forested catchments in Japan and found that E shows a positive correlation with the average temperature (T_{avg}) of each catchment. Thus, they proposed the modeled relation of E (mm) = 31.4 T_{avg} (°C) + 376 to estimate E in each forested catchment in Japan, where the standard error of 162.3 mm was included in the estimated evapotranspiration flux (E).

271

272 2.6 Concentration of unprocessed NO_{3⁻atm} in each water sample

273 The Δ^{17} O data of nitrate in each sample was used to estimate the concentration of 274 NO_{3⁻atm} ([NO_{3⁻atm}]) in each water sample by applying Eq. (<u>76</u>):

275
$$[NO_3^-] = \Delta^{17}O/\Delta^{17}O_{atm}$$

-(<u>76</u>)

276	where $[NO_3^-]$ and $[NO_3^-]$ denote the concentrations of $NO_3^-]$ and nitrate (total) in
277	each water sample, respectively, and $\Delta^{17}O_{atm}$ and $\Delta^{17}O$ denote the $\Delta^{17}O$ values of
278	$NO_{3\ atm}$ and nitrate (total) in the stream water sample, respectively. In this study, we
279	used the annual average Δ^{17} O value of NO ₃ ⁻ _{atm} determined at the Sado-Seki monitoring
280	station in Japan (Sado Island; Fig. 1a) from April 2009 to March 2012 ($\Delta^{17}O_{atm}$ =
281	+26.3 ‰; Tsunogai et al., 2016) for $\Delta^{17}O_{atm}$ in Eq. (72) to estimate [NO _{3⁻atm}] in the
282	stream. We allow for an error range of 3 ‰ in $\Delta^{17}O_{atm}$, where the factor changes in
283	$\Delta^{17}O_{atm}$ from +26.3 ‰ caused by both areal and seasonal variations in the $\Delta^{17}O$ values
284	of NO3 ⁻ _{atm} have been considered (Nakagawa et al., 2018; Tsunogai et al., 2016; Ding et
285	al., 2022).
286	The annual export flux of unprocessed $NO_3^{-}_{atm}$ per unit area of the catchment (M _{atm})
287	was determined by applying Eq. ($\underline{87}$):
288	$M_{atm} = [NO_3^{-}_{atm}]_{avg} \times F_{stream} $ (87)
289	where $[NO_3^-a_{tm}]_{avg}$ denotes the annual average $[NO_3^-a_{tm}]$ in each stream. The index of
290	nitrogen saturation (M_{atm}/D_{atm} ratio) was calculated by dividing M_{atm} with D_{atm} in each
291	catchment.
292	
293	2.7 Concentration and isotopic compositions of stream nitrate eluted only from the FK2
294	catchment
295	The concentration and isotopic compositions ($\delta^{15}N$, $\delta^{18}O$, and $\Delta^{17}O$) of stream nitrate
296	determined at the station B were the mixture of those eluted from FK1 and FK2

297	catchments (Fig. 1b). Assuming that the stream nitrate eluted from FK1 catchment was
298	stable during the flow path from station A to station B. The concentration of stream
299	nitrate eluted from the FK2 catchment was determined by applying Eq. (9):
300	$[NO_{3}^{-}]_{FK2} = ([NO_{3}^{-}]_{FK1+FK2} * F_{FK1+FK2} - [NO_{3}^{-}]_{FK1} * F_{FK1}) / F_{FK2} $ (9)
801	where F_{FK1} , F_{FK2} , and $F_{FK1+FK2}$ denote the flux of stream water eluted from the FK1,
302	FK2 (only), and FK1+FK2 catchment, respectively. [NO ₃ ⁻] _{FK1} , [NO ₃ ⁻] _{FK2} , and
303	[NO ₃ ⁻] _{FK1+FK2} denote the concentration of stream nitrate eluted from the FK1, FK2
304	(only), and FK1+FK2 catchment, respectively. In this study, the flow rates measured at
305	stations A and B on 2021/01/15 by using the salt dilution method (Sappa et al., 2015)
306	was used for F_{FK1} (0.85 L/s) and $F_{FK1+FK2}$ (4.75 L/s), respectively, and the measured
307	[NO ₃ ⁻] at stations A and B was used for [NO ₃ ⁻] _{FK1} and [NO ₃ ⁻] _{FK1+FK2} , respectively.
308	Because the relation between the measured flow rates was comparable with the relation
309	between the catchment area of FK1 (14 ha) and that of FK1+FK2 (76 ha), we concluded
310	that the measured flow rates of 0.85 L/s and 4.75 L/s were reasonable as for those
311	representing the F_{FK1} and $F_{FK1+FK2}$, respectively. According to the mass balance of water,
312	we can estimate the F_{FK2} eluted from the FK2 catchment only to be 3.90 L/s.
313	Assuming that the stream nitrate eluted from FK1 catchment was stable during the
314	flow path from station A to station B, the δ^{15} N, δ^{18} O, and Δ^{17} O values of stream nitrate
315	eluted from the FK2 catchment only were determined by applying Eq. (10):
516	$\underline{\delta_{FK2}} = (\underline{\delta_{FK1+FK2}} * [NO_3^-]_{FK1+FK2} * F_{FK1+FK2} - \underline{\delta_{FK1}} * [NO_3^-]_{FK1} * F_{FK1}) / ([NO_3^-]_{FK2} * C_{FK1+FK2} + C_{FK1}) / ([NO_3^-]_{FK2} * C_{FK1+FK2} + $
17	(10)

318	where δ_{FK1} , δ_{FK2} , and $\delta_{FK1+FK2}$ denote the $\delta^{15}N$ (or $\delta^{18}O$ or $\Delta^{17}O$) of stream nitrate eluted
319	from the FK1, FK2, and FK1+FK2 catchment, respectively. The $\delta^{15}N$ (or $\delta^{18}O$ or $\Delta^{17}O$)
320	values of stream nitrate measured at stations A and B were used for δ_{FK1} and $\delta_{FK1+FK2}$,
321	respectively.
322	
323	3 Results
324	3.1 Deposition rate of atmospheric nitrate
325	The mean annual precipitation (P) from 2009 to 2021 was 1777 mm and 3981 mm
326	for FK and MY catchments, respectively (Chiwa, 2020; Chiwa, personal
327	communication, September 21, 2022). The mean annual temperature (T_{avg}) was
328	reported to be 15.9 °C and 10.8 °C for FK and MY catchments, respectively (Chiwa,
329	2020). Chiwa (2020) estimated the mean annual precipitation (P) and mean annual
330	temperature (T_{arg}) to be 1769 mm and 15.9 °C, respectively, at FK catchments, and
331	3837 mm and 10.8 °C, respectively, at MY catchment. Based on these data, the annual
332	flux of stream water (F _{stream}) was estimated to be $902.0893.7 \pm 162.3$ mm at FK
333	catchments and $3266.13121.9 \pm 162.3$ mm at MY catchment, respectively, using Eq.
334	(65), corresponding to 1.25×10^5 m ³ /year in FK1, 5.54×10^5 m ³ /year in FK2, and 1.34
335	$\times 10^6 \text{ m}^3/\text{year in MY}.$
336	Chiwa (2020) also-reported the annual bulk deposition rates of atmospheric nitrate
337	(D_{bulk}) to be 34.0 mmol m ⁻² year ⁻¹ at FK catchments and 24.2 mmol m ⁻² year ⁻¹ at MY
338	catchment. On the other hand, the annual dry deposition rate of atmospheric nitrate

339	(D_{dry}) deposited <u>onin</u> the forest $(D_{dry}(F))$ and on the water surface $(D_{dry}(W))$ were
340	estimated to be 39.98 mmol m ⁻² year ⁻¹ and 4.1 mmol m ⁻² year ⁻¹ , respectively, at FK
341	catchments, and 18.4 mmol m ⁻² year ⁻¹ and 2.4 mmol m ⁻² year ⁻¹ , respectively, at MY
342	catchment. As a result, D_{atm} was estimated to be 69.3 mmol m ⁻² year ⁻¹ at FK catchments
343	and 40.1 mmol m ⁻² year ⁻¹ at MY catchments, using Eq. (32).

345 3.2 Concentration and isotopic composition of stream nitrate

The concentrations of stream nitrate at eluted from the FK1, FK2 (only), and MY 346 catchments ranged from 97.5 μ M to 121.3 μ M, from 73.965.7 μ M to 148.5142.6 μ M, 347 348 and from 3.5 μ M to 15.3 μ M, respectively, with the average concentrations of 109.5 349 μ M, 90.994.2 μ M, and 7.3 μ M, respectively, and the standard deviations (SD) of 6.3 μ M, 18.5 μ M, and 3.0 μ M, respectively, which corresponds to the coefficients of 350 variation (CV) of 5.7 %, 20.4 %, and 40.7 %, respectively (Fig. 2a). All catchments 351 showed <u>no clear</u><u>little</u> seasonal variation during the observation periods. The variation 352 ranges and the average concentrations of stream nitrate in eluted from the three 353 354 catchments agreed well with the past observations performed in the same catchments 355 (Chiwa, 2021).-356 The stable isotopic compositions of stream nitrate at eluted from the FK1, FK2 (only),

and MY catchments ranged from -0.9 % to +1.5 %, from -1.42 % to +5.84.5 %, and from -0.8 % to $\pm 2.4 \%$, respectively, for $\delta^{15}N$ (Fig. 2b), from $\pm 3.9 \%$ to $\pm 8.5 \%$, from =0.7-2.2 % to $\pm 2.8 \%$, and from -5.6 % to $\pm 1.7 \%$, respectively, for $\delta^{18}O$ (Fig. 2c), 360 and from +2.0 ‰ to +3.3 ‰, from +0.68 ‰ to +2.24 ‰, and from +0.2 ‰ to +1.0 ‰, respectively, for $\Delta^{17}O$ (Fig. 2d), with <u>little</u> no clear seasonal variation during the 361 362 observation periods. The concentration-weighted averages for the δ^{15} N, δ^{18} O, and Δ^{17} O values of stream nitrate were +0.2 ‰, +6.4 ‰, and +2.6 ‰, respectively, at FK1, 363 364 +1.00.9 ‰, +0.51.7 ‰, and +1.51.7 ‰, respectively, at FK2, +0.7 ‰, -2.5 ‰, and +0.6 ‰, respectively, at MY. These values were typical for stream nitrate eluted from 365 forested catchments (Hattori et al., 2019; Huang et al., 2020; Nakagawa et al., 2013, 366 2018; Riha et al., 2014; Sabo et al., 2016; Tsunogai et al., 2014, 2016). 367 368

369 3.3 Concentration of unprocessed atmospheric nitrate and the M_{atm}/D_{atm} ratio in each
 370 catchment

The concentration of unprocessed atmospheric nitrate ($[NO_3^-]_{atm}$) in the streams of 371 372 eluted from the FK1, FK2 (only), and MY catchments ranged from 8.64 to 14.30 μ M, from 3.882.27 to 11.1610.71 µM, and from 0.03 to 0.46 µM with the average 373 374 concentration of 10.80 ± 1.65 , $\frac{6.095.06}{5.06} \pm \frac{1.050.92}{0.92}$, and $0.16 \pm 0.05 \mu$ M, respectively, 375 even though these studiedy catchments showed little seasonal variations during the observation periods (Fig. 2e). The annual export flux of nitrate (M_{total}), the annual 376 export flux of NO_{3⁻atm} (M_{atm}), and the M_{atm}/D_{atm} ratio were $\frac{98.897.9}{100} \pm 17.8 \text{ mmol m}^{-2}$ 377 year⁻¹, 9.7 \pm 2.3 mmol m⁻² year⁻¹, and 14.113.9 \pm 4.43 % at FK1 catchment, 378 respectively, $\frac{84.282.0 \pm 15.314.8}{15.314.8}$ mmol m⁻² year⁻¹, $\frac{5.44.6 \pm 1.24}{1.24}$ mmol m⁻² year⁻¹, and 379 380 $6.67.9 \pm 2.15$ % at FK2 catchment, respectively, $23.722.6 \pm 1.2$ mmol m⁻² year⁻¹, 0.5 ± 1.2

381 $0.\underline{2}4$ mmol m⁻² year⁻¹, and $1.\underline{3}2 \pm 0.\underline{5}4$ % at MY catchment, respectively (Table 2).

382

383 4 Discussion

384 4.1 Deposition rate of atmospheric nitrate at the study catchments

Based on the air monitoring data determined at the stations of Fukuoka (33°51'N, 385 130°50'E) and Miyazaki (31°83'N, 131°42'E) from 2011 to 2017, the Environmental 386 Laboratories Association of Japan (2017) reported D_{atm} to be 57.8 mmol m⁻² year⁻¹ at 387 Fukuoka and 49.1 mmol m⁻² year⁻¹ at Miyazaki. Those values are consistent with the 388 D_{atm} estimated in this study (69.3 and 40.1 mmol m⁻² year⁻¹ at the FK and MY 389 390 catchments, respectively), within a difference of approximately 20_%. Thus, we 391 concluded that the D_{atm} estimated in this study was reliable within the error margin of 392 20_% (Table 2). Because the D_{atm} determined at the FK catchments was the highest 393 among the forested catchments in Table 3, we further compared the D_{atm} of the FK 394 catchments with those from the other air monitoring stations in Japan reported in past studies, along with that of the MY catchment (Table S1). While the Datm of the MY 395 396 catchment corresponded to the average level among the sites compiled in Table S1, the D_{atm} of the FK catchments exceeded the average level significantly. In addition, the D_{atm} 397 of the FK catchments corresponded to one of the highest among the Japanese forested 398 399 areas (Table S1). While aAll the catchments in Japanthis study can be suffered from the long-range transport of air pollutants derived from megacities in East Asian region 400 401 (Chiwa, 2021; Chiwa et al., 2012 and 2013). In addition, the shorter transport distance

402	from the Fukuoka metropolitan area (total population: 1.62 million people; population
403	density: 4715 people/km ²) may be mainly responsible for the D_{atm} higher in FK than in
404	MY, because the FK catchments are only 15 km west of the Fukuoka metropolitan area.
405	As a result, the local emission in the Fukuoka metropolitan area should be responsible
406	for the high D _{atm} at the FK catchments.

408 4.2 Excess leaching of unprocessed atmospheric nitrate from FK catchments

409 The isotopic compositions (δ^{15} N, δ^{18} O, and Δ^{17} O) of stream nitrate eluted from the

410 FK and MY catchments were typical for those eluted from forested catchments (Hattori

411 et al., 2019; Huang et al., 2020; Nakagawa et al., 2013, 2018; Riha et al., 2014; Sabo et

412 <u>al., 2016; Tsunogai et al., 2014, 2016).</u> The striking features found in the FK catchments

413 were that, in addition to the high $[NO_3^-]$ and high M_{total} that had been clarified prior to

414 this in a past study (Chiwa, 2021), both $[NO_3^-]_{atm}$ and M_{atm} in FK were higher than those

415 in <u>eluted from</u> MY (Table 2). Especially, the average [NO_{3⁻atm}] in the stream eluted ______in

416 the<u>from the</u> FK1 stream catchment was the highest ever reported in forested streams

417 determined through continuous monitoring for more than 1 year (Bostic et al., 2021;

418 Bourgeois et al., 2018b, 2018a; Hattori et al., 2019; Huang et al., 2020; Nakagawa et

419 al., 2018; Rose et al., 2015; Sabo et al., 2016; Tsunogai et al., 2014, 2016).

420 The observed high $[NO_3^-_{atm}]$ in the FK1-stream <u>eluted from the FK1 catchment</u> could 421 be caused just by the high $[NO_3^-_{atm}]$ in deposition in the catchment D_{atm} . Thus, we 422 compiled all past data ever reported in forested streams through continuous monitoring 423 in Table 3, where the data of average $[NO_3^-]$, average $[NO_3^-]$, M_{atm} , M_{total} , D_{atm} , and 424 M_{atm}/D_{atm} ratio were included for comparison.<u>,</u> and <u>t</u> he result showed that the FK1 425 catchment has the highest M_{atm}/D_{atm} ratio, along with M_{atm} , was the highest as well in 426 <u>the FK1 catchment</u> among the forested catchments (Table 3).

Elevated loading of nitrogen through atmospheric deposition was responsible for the 427 occurrence of nitrogen saturation in forest ecosystems, from which elevated levels of 428 nitrate are exported (Aber et al., 1989). Nakagawa et al. (2018) proposed that the 429 Matm/Datm ratio can be an index for evaluating the nitrogen saturation in each forested 430 catchment, because the Matm/Datm ratio directly reflects the present demand for 431 432 atmospheric nitrate deposited in each forested catchment, and thus reflects the nitrogen saturation in each forested catchment. The high M_{atm}/D_{atm} ratios observed in the FK 433 434 catchments implied that the demand for atmospheric nitrate was low in the FK 435 catchments and <u>thethat the stages of nitrogen saturation at the FK catchments were</u> higher than those at other forested catchments., and thus That is, the nitrogen saturation 436 437 at the FK catchments was responsible for the observed higher observed [NO₃⁻] and high 438 M_{total} at the FK catchments than at MY and any other catchment ever studied (Table 3). 439 The stand age of forests can affect the retention or loss of N (Fukushima et al., 2011; Ohrui and Mitchell, 1997). Fukushima et al. (2011) evaluated N uptake rates of 440 Japanese cedars at different ages (5-89 years old) and demonstrated that the N uptake 441 rates of Japanese cedars were higher in younger stands (53 kg N ha⁻¹ year⁻¹ in 16 years 442 old) than in older stands (29 kg N ha⁻¹ year⁻¹ in 31 years old; 24 kg N ha⁻¹ year⁻¹ in 42 443

444	years old; 34 kg N ha ⁻¹ year ⁻¹ in 89 years old). In addition, Yang and Chiwa (2021)
445	found that the nitrate concentration in the soil water taken beneath the rooting zone of
446	matured artificial Japanese cedar plantations (607 \pm 59 $\mu M;$ 64-69 years old) was
447	significantly higher than that of normal Japanese oak plantations (8.7 \pm 8.1 $\mu M;$ 24
448	years old). Moreover, by adding ammonium nitrate (50 kg N ha ⁻¹ year ⁻¹) to the forest
449	floor directly, Yang and Chiwa (2021) found that the nitrate concentration in the soil
450	water of the matured artificial Japanese cedar plantations increased significantly faster
451	than that of the normal Japanese oak plantations, probably because of the lower N
452	uptake rates in the matured artificial Japanese cedar plantations. Because most of the
453	artificial Japanese cedar/cypress plantations in the FK and MY catchments have reached
454	their maturity (> 50 years; Yang and Chiwa, 2021), the higher proportion of matured
455	artificial Japanese cedar/cypress plantations in the FK1 catchment (Table 1) was highly
456	responsible for the observed elevated leaching of nitrate, caused by the reduction in N
457	uptake rates.

As a result, we concluded that the FK forested catchments were under the high nitrogen saturation stage, FK1 catchment especially, and the high-nitrogen saturation stage of in the FK1 catchments was responsible for the elevated M_{total} , M_{atm} , $[NO_3^-]$, $[NO_3^-]$ found in the stream eluted from the catchment (Figs. 3a, 3b, and 3c).

463 4.3 The M_{atm}/D_{atm} ratio as an index of nitrogen saturation

464 Past studies have used the concentration of stream nitrate as one of the important

465	indexes to evaluate the stage of nitrogen saturation in each forest (Aber, 1992; Huang
466	et al., 2020; Rose et al., 2015; Stoddard, 1994). The strong linear relationship (R^2 =
467	0. <u>7681</u> ; $P < 0.0001$) between the stream nitrate concentration and the M _{atm} /D _{atm} ratio,
468	except for the Qingyuan forested catchment (Fig. 3d), further supported that the
469	M_{atm}/D_{atm} ratio can be used as an alternative index of nitrogen saturation, as pointed out
470	in Nakagawa et al. (2018).
471	The differences in the number of storm and/or snowmelt events could affect the
472	<u>M_{atm}/D_{atm} ratio as well, because NO₃⁻_{atm} could be injected into the stream water directly,</u>
473	along with the storm / snowmelt water (Tsunogai et al., 2014; Ding et al., 2022; Inamdar
474	and Mitchell, 2006). In recent study, however, we found that the storm events have little
475	impacts on the M _{atm} /D _{atm} ratio, based on monitoring temporal variation of [NO ₃ - a_{tm}] in
476	a stream water during storm events (Ding et al., 2022). In addition, the low Matm/Datm
477	ratio found in Uryu forested catchment (0.7 %; Table 3) implied that the snowmelt has
478	little impacts on the M_{atm}/D_{atm} ratio as well, because 30% of the annual mean
479	precipitation was snow in Uryu forested catchment (Tsunogai et al., 2014).
480	The differences in the amount of precipitation, temperature, and the flux of stream
481	water could affect the M_{atm}/D_{atm} ratio as well. As a result, the annual amount of
482	precipitation, mean temperature, and the annual mean flux of stream water (F _{stream}) in
483	the forested catchments were compiled in Table S2. While the stream nitrate
484	concentration showed the strong linear relationship ($R^2 = 0.76$; $P < 0.0001$) with the
485	M_{atm}/D_{atm} ratio (Fig. 3d), the precipitation, temperature, and F_{stream} did not show

486 <u>significant relationship with the M_{atm}/D_{atm} ratio (P > 0.14; Fig. 4). As a result, we</u> 487 <u>concluded that the M_{atm}/D_{atm} ratio was mainly controlled by the progress of nitrogen</u> 488 <u>saturation, rather than the differences in the number of storm and/or snowmelt events,</u> 489 <u>the amount of precipitation, temperature, and the flux of stream water.</u>

490 Moreover, tThe M_{atm}/D_{atm} ratio is a more reliable and robust index than the stream nitrate concentration, as explained below. The Qingyuan forested catchment can be 491 classified into the highest nitrogen saturation stage based only on the highest stream 492 nitrate concentration of 150 µM (Table 3). However, based on the leaching flux of 493 nitrogen via stream water monitored by Huang et al. (2020) for 4 years in the Qingyuan 494 495 forested catchment, along with the deposition flux of nitrogen, we can obtain the M_{atm}/D_{atm} ratio in the catchment to be a medium level of 5.8 ± 1.3 %, implying that the 496 497 nitrogen saturation stage was not sovery high (Table 3). Huang et al. (2020) also 498 concluded that the input of nitrogen exceeded the output in the catchment, and thus, the 499 catchment was at stage 2 of nitrogen saturation. The Matm/Datm ratio in the Qingyuan forested catchment with a medium level among all forested catchments (Fig. 3d) should 500 501 be a more reliable index of nitrogen saturation.

Compared with those in the other forested catchments in Table 3, the annual amount of precipitation (P) has the lowest value of 709 mm in the Qingyuan forested catchment. The flux of stream water (F_{stream}) has the lowest value of 309 mm as well. Thus, we concluded that nitrate was relatively concentrated in the catchment because of the small precipitation, resulting in relative enrichment in the concentrations of both nitrate (150 507 μ M) and unprocessed atmospheric nitrate (8.9 μ M) in the stream.

508	While the concentration of stream nitrate, as an index of nitrogen saturation
509	traditionally, can be influenced by the amount of precipitation, as demonstrated in the
510	Qingyuan forested catchment, the $M_{\text{atm}}/D_{\text{atm}}$ ratio is independent of the amount of
511	precipitation (Fig. 4). Therefore, we concluded that the M_{atm}/D_{atm} ratio can be used as
512	a more robust index for evaluating nitrogen saturation in each forested catchment.
513	The only concern on using the M_{atm}/D_{atm} ratio as the index of nitrogen saturation is
514	the impact of the differences in the residence time of water in each catchment. The
515	residence time of water varies from 1 month to more than 1 year in forested catchments
516	(Asano et al., 2002; Farrick and Branfireun, 2015; Kabeya et al., 2008; Rodgers et al.,
517	2005; Soulsby et al., 2006; Tetzlaff et al., 2007). The M _{atm} /D _{atm} ratio could be lower in
518	catchments with longer residence time of water. We would like to clarify this in future
519	studies by adding much more data of stream nitrate eluted from various forested
520	catchments.

521

522 **5 Conclusions**

Both the concentrations and Δ^{17} O of stream nitrate were determined for more than 2 years in the forested catchments of FK (FK1 and FK2) and MY to determine the M_{atm}/D_{atm} ratio for each catchment. The FK catchments exhibited higher M_{atm}/D_{atm} ratio than the MY catchment and other forested catchments reported in past studies, implying that the progress of nitrogen saturation in the FK catchments was severe. Both age and

528	proportion of artificial plantation in the FK catchments were responsible for the
529	progress of nitrogen saturation. In addition, although past studies have commonly used
530	the concentration of stream nitrate as an index to evaluate the progress of nitrogen
531	saturation in forested catchments, it can be influenced by the amount of precipitation.
532	As a result, we concluded that the $M_{\text{atm}}/D_{\text{atm}}$ ratio should be used as a more reliable
533	index for evaluating the progress of nitrogen saturation because the $M_{\text{atm}}/D_{\text{atm}}$ ratio is
534	independent from the amount of precipitation.
535	
536	Data availability. All the primary data are presented in the Supplement. The other data
537	are available upon request to the corresponding author (Weitian Ding).
538	
539	Author contributions. UT, FN, KS, and MC designed the study. MC and TK performed
540	the field observations. WD, UT, and FN determined the concentrations and isotopic
541	compositions of the samples. WD, TS, FN, and UT performed data analysis, and WD
542	and UT wrote the paper with input from MC, TK, and KS.
543	
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545	
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562 **Reference**

- 563 Aber, J. D.: Nitrogen cycling and nitrogen saturation in temperate forest ecosystems,
- 564 Trends Ecol. Evol., 7(7), 220–224, doi:10.1016/0169-5347(92)90048-G, 1992.
- 565 Aber, J. D., Nadelhoffer, K. J., Steudler, P. and Melillo, J. M.: Nitrogen Saturation in
- 566 Northern Forest Ecosystems, Bioscience, 39(6), 378–386, doi:10.2307/1311067,
- 567 1989.
- 568 Aikawa, M., Hiraki, T., Tamaki, M. and Shoga, M.: Difference between filtering-type
- 569 bulk and wet-only data sets based on site classification, Atmos. Environ., 37(19),

- 570 2597–2603, doi:10.1016/S1352-2310(03)00214-0, 2003.
- 571 Alexander, B., Hastings, M. G., Allman, D. J., Dachs, J., Thornton, J. A. and
- 572 Kunasek, S. A.: Quantifying atmospheric nitrate formation pathways based on a
- 573 global model of the oxygen isotopic composition (δ^{17} O) of atmospheric nitrate,
- 574 Atmos. Chem. Phys., 9(14), 5043–5056, doi:10.5194/acp-9-5043-2009, 2009.
- 575 Asano, Y., Uchida, T. and Ohte, N.: Residence times and flow paths of water in steep
- 576 <u>unchannelled catchments, Tanakami, Japan, J. Hydrol., 261(1–4), 173–192</u>,
- 577 <u>doi:10.1016/S0022-1694(02)00005-7, 2002.</u>
- 578 Environmental Laboratories Association of Japan: Acid Rain National Survey Report
- 579 2017, https://tenbou.nies.go.jp/envgis_explain/acid_rain/content.html.
- 580 Bostic, J. T., Nelson, D. M., Sabo, R. D. and Eshleman, K. N.: Terrestrial Nitrogen
- 581 Inputs Affect the Export of Unprocessed Atmospheric Nitrate to Surface Waters:
- 582 Insights from Triple Oxygen Isotopes of Nitrate, Ecosystems, doi:10.1007/s10021-
- 583 021-00722-9, 2021.
- 584 Bourgeois, I., Savarino, J., Némery, J., Caillon, N., Albertin, S., Delbart, F., Voisin,
- 585 D. and Clément, J. C.: Atmospheric nitrate export in streams along a montane to
- urban gradient, Sci. Total Environ., 633, 329–340,
- 587 doi:10.1016/j.scitotenv.2018.03.141, 2018a.
- 588 Bourgeois, I., Savarino, J., Caillon, N., Angot, H., Barbero, A., Delbart, F., Voisin, D.
- and Clément, J. C.: Tracing the Fate of Atmospheric Nitrate in a Subalpine Watershed
- 590 Using Δ¹⁷O, Environ. Sci. Technol., 52(10), 5561–5570, doi:10.1021/acs.est.7b02395,

591 2018b.

- 592 Chiwa, M.: Ten-year determination of atmospheric phosphorus deposition at three
- 593 forested sites in Japan, Atmos. Environ., 223(May 2019), 1–7,
- 594 doi:10.1016/j.atmosenv.2019.117247, 2020.
- 595 Chiwa, M.: Long-term changes in atmospheric nitrogen deposition and stream water
- 596 nitrate leaching from forested watersheds in western Japan, Environ. Pollut.,
- 597 287(November 2020), 117634, doi:10.1016/j.envpol.2021.117634, 2021.
- 598 Chiwa, M., Enoki, T., Higashi, N., Kumagai, T. and Otsuki, K.: The Increased
- 599 Contribution of Atmospheric Nitrogen Deposition to Nitrogen Cycling in a Rural
- 600 Forested Area of Kyushu, Japan, Water, Air, Soil Pollut., 224(11), 1763,
- 601 doi:10.1007/s11270-013-1763-2, 2013.
- 602 Chiwa, M., Onikura, N., Ide, J. and Kume, A.: Impact of N-Saturated Upland Forests
- on Downstream N Pollution in the Tatara River Basin, Japan, Ecosystems, 15(2),
- 604 230–241, doi:10.1007/s10021-011-9505-z, 2012.
- 605 Chiwa, M.: Characteristics of atmospheric nitrogen and sulfur containing compounds
- in an inland suburban-forested site in northern Kyushu, western Japan, Atmos. Res.,
- 607 96(4), 531–543, doi:10.1016/j.atmosres.2010.01.001, 2010.
- Ding, W., Tsunogai, U., Nakagawa, F., Sambuichi, T., Sase, H., Morohashi, M., and
- 609 Yotsuyanagi, H.: Tracing the source of nitrate in a forested stream showing elevated
- 610 concentrations during storm events, Biogeosciences, 19, 3247–3261,
- 611 https://doi.org/10.5194/bg-19-3247-2022, 2022.

- 612 Endo, T., Yagoh, H., Sato, K., Matsuda, K., Hayashi, K., Noguchi, I. and Sawada, K.:
- 613 Regional characteristics of dry deposition of sulfur and nitrogen compounds at
- 614 EANET sites in Japan from 2003 to 2008, Atmos. Environ., 45(6), 1259–1267,
- 615 doi:10.1016/j.atmosenv.2010.12.003, 2011.
- 616 Farrick, K. K. and Branfireun, B. A.: Flowpaths, source water contributions and water
- 617 residence times in a Mexican tropical dry forest catchment, J. Hydrol., 529, 854–865,
- 618 <u>doi:10.1016/j.jhydrol.2015.08.059, 2015.</u>
- 619 Kabeya, N., Shimizu, A., Nobuhiro, T. and Tamai, K.: Preliminary study of flow
- 620 regimes and stream water residence times in multi-scale forested watersheds of central
- 621 Cambodia, Paddy Water Environ., 6(1), 25–35, doi:10.1007/s10333-008-0104-3,
 622 <u>2008.</u>
- 623 Fukushima, K., Tateno, R. and Tokuchi, N.: Soil nitrogen dynamics during stand
- 624 development after clear-cutting of Japanese cedar (Cryptomeria japonica) plantations,
- 625 J. For. Res., 16(5), 394–404, doi:10.1007/s10310-011-0286-1, 2011.
- 626 Galloway, J. N., Aber, J. D., Erisman, J. W., Seitzinger, S. P., Howarth, R. W.,
- 627 Cowling, E. B. and Cosby, B. J.: The nitrogen cascade, Bioscience, 53(4), 341–356,
- 628 doi:10.1641/0006-3568(2003)053[0341:TNC]2.0.CO;2, 2003.
- 629 Hattori, S., Nuñez Palma, Y., Itoh, Y., Kawasaki, M., Fujihara, Y., Takase, K. and
- 630 Yoshida, N.: Isotopic evidence for seasonality of microbial internal nitrogen cycles in
- a temperate forested catchment with heavy snowfall, Sci. Total Environ., 690, 290-
- 632 299, doi:10.1016/j.scitotenv.2019.06.507, 2019.

- 633 Hirota, A., Tsunogai, U., Komatsu, D. D. and Nakagawa, F.: Simultaneous
- 634 determination of δ^{15} N and δ^{18} O of N₂O and δ^{13} C of CH₄ in nanomolar quantities from
- a single water sample, Rapid Commun. Mass Spectrom., 24, 1085–1092,
- 636 doi:10.1002/rcm.4483, 2010.
- Huang, S., Wang, F., Elliott, E. M., Zhu, F., Zhu, W., Koba, K., Yu, Z., Hobbie, E.
- A., Michalski, G., Kang, R., Wang, A., Zhu, J., Fu, S. and Fang, Y.: Multiyear
- 639 Measurements on Δ^{17} O of Stream Nitrate Indicate High Nitrate Production in a
- 640 Temperate Forest, Environ. Sci. Technol., 54(7), 4231–4239,
- 641 doi:10.1021/acs.est.9b07839, 2020.
- 642 Inoue, T., Nakagawa, F., Shibata, H. and Tsunogai, U.: Vertical Changes in the Flux
- of Atmospheric Nitrate From a Forest Canopy to the Surface Soil Based on Δ^{17} O
- 644 Values, J. Geophys. Res. Biogeosciences, 126(4), 1–18, doi:10.1029/2020JG005876,
 645 2021.
- 646 Inamdar, S. P. and Mitchell, M. J.: Hydrologic and topographic controls on storm-
- 647 event exports of dissolved organic carbon (BOC) and nitrate across catchment scales,
- 648 <u>Water Resour. Res., 42(3), 1–16, doi:10.1029/2005WR004212, 2006.</u>
- 649 Kaiser, J., Hastings, M. G., Houlton, B. Z., Röckmann, T. and Sigman, D. M.: Triple
- oxygen isotope analysis of nitrate using the denitrifier method and thermal
- decomposition of N₂O, Anal. Chem., 79(2), 599–607, doi:10.1021/ac061022s, 2007.
- 652 Komatsu, D. D., Ishimura, T., Nakagawa, F. and Tsunogai, U.: Determination of the
- $^{15}N/^{14}N$, $^{17}O/^{16}O$, and $^{18}O/^{16}O$ ratios of nitrous oxide by using continuous-flow

- 654 isotope-ratio mass spectrometry Daisuke, Rapid Commun. Mass Spectrom., 22, 1587–
- 655 1596, doi:10.1002/rcm.3493, 2008a.
- 656 Komatsu, H., Maita, E. and Otsuki, K.: A model to estimate annual forest
- evapotranspiration in Japan from mean annual temperature, , 330–340,
- 658 doi:10.1016/j.jhydrol.2007.10.006, 2008b.
- 659 Konno, U., Tsunogai, U., Komatsu, D. D., Daita, S., Nakagawa, F., Tsuda, A.,
- 660 Matsui, T., Eum, Y. J. and Suzuki, K.: Determination of total N₂ fixation rates in the
- ocean taking into account both the particulate and filtrate fractions, Biogeosciences,
- 662 7(8), 2369–2377, doi:10.5194/bg-7-2369-2010, 2010.
- 663 Matsuda, K.: Estimation of dry deposition for sulfur and nitrogen compounds in the
- atmosphere : Updated parameterization of deposition velocity, J. Japan Soc. Atmos.
- 665 Environ., 43(6), 332–339, doi:10.11298/taiki1995.43.332, 2008.
- 666 McIlvin, M. R. and Altabet, M. A.: Chemical conversion of nitrate and nitrite to
- nitrous oxide for nitrogen and oxygen isotopic analysis in freshwater and seawater,
- 668 Anal. Chem., 77(17), 5589–5595, doi:10.1021/ac050528s, 2005.
- 669 Michalski, G., Scott, Z., Kabiling, M. and Thiemens, M. H.: First measurements and
- 670 modeling of Δ^{17} O in atmospheric nitrate, Geophys. Res. Lett., 30(16), 3–6,
- doi:10.1029/2003GL017015, 2003.
- 672 Michalski, G., Meixner, T., Fenn, M., Hernandez, L., Sirulnik, A., Allen, E. and
- 673 Thiemens, M.: Tracing Atmospheric Nitrate Deposition in a Complex Semiarid
- 674 Ecosystem Using Δ^{17} O, Environ. Sci. Technol., 38(7), 2175–2181,

- 675 doi:10.1021/es034980+, 2004.
- 676 Mitchell, M. J., Iwatsubo, G., Ohrui, K. and Nakagawa, Y.: Nitrogen saturation in
- Japanese forests: An evaluation, For. Ecol. Manage., 97(1), 39–51,
- 678 doi:10.1016/S0378-1127(97)00047-9, 1997.
- 679 Morin, S., Sander, R. and Savarino, J.: Simulation of the diurnal variations of the
- 680 oxygen isotope anomaly (Δ^{17} O) of reactive atmospheric species, Atmos. Chem. Phys.,
- 681 11(8), 3653–3671, doi:10.5194/acp-11-3653-2011, 2011.
- 682 Nakagawa, F., Suzuki, A., Daita, S., Ohyama, T., Komatsu, D. D. and Tsunogai, U.:
- 683 Tracing atmospheric nitrate in groundwater using triple oxygen isotopes: Evaluation
- based on bottled drinking water, Biogeosciences, 10(6), 3547–3558, doi:10.5194/bg-
- 685 10-3547-2013, 2013.
- Nakagawa, F., Tsunogai, U., Obata, Y., Ando, K., Yamashita, N., Saito, T.,
- 687 Uchiyama, S., Morohashi, M. and Sase, H.: Export flux of unprocessed atmospheric
- nitrate from temperate forested catchments: A possible new index for nitrogen
- 689 saturation, Biogeosciences, 15(22), 7025–7042, doi:10.5194/bg-15-7025-2018, 2018.
- 690 Nelson, D. M., Tsunogai, U., Ding, D., Ohyama, T., Komatsu, D. D., Nakagawa, F.,
- 691 Noguchi, I. and Yamaguchi, T.: Triple oxygen isotopes indicate urbanization affects
- 692 sources of nitrate in wet and dry atmospheric deposition, Atmos. Chem. Phys., 18(9),
- 693 6381–6392, doi:10.5194/acp-18-6381-2018, 2018.
- 694 Ohrui, K. and Mitchell, M. J.: Nitrogen Saturation in Japanese Forested Watersheds
- 695 Author (s): Kiyokazu Ohrui and Myron J. Mitchell Published by : Wiley Stable

- 696 URL : http://www.jstor.org/stable/2269507 Accessed : 05-07-2016 04 : 51 UTC Your
- use of the JSTOR archive indicates your ac, 7(2), 391-401, 1997.
- 698 Paerl, H. W. and Huisman, J.: Climate change: A catalyst for global expansion of
- harmful cyanobacterial blooms, Environ. Microbiol. Rep., 1(1), 27–37,
- 700 doi:10.1111/j.1758-2229.2008.00004.x, 2009.
- 701 Peterjohn, W. T., Adams, M. B. and Gilliam, F. S.: Symptoms of nitrogen saturation
- in two central Appalachian hardwood forest ecosystems, Biogeochemistry, 35(3),
- 703 507–522, doi:10.1007/BF02183038, 1996.
- Riha, K. M., Michalski, G., Gallo, E. L., Lohse, K. A., Brooks, P. D. and Meixner, T.:
- 705 High Atmospheric Nitrate Inputs and Nitrogen Turnover in Semi-arid Urban
- 706 Catchments, Ecosystems, 17(8), 1309–1325, doi:10.1007/s10021-014-9797-x, 2014.
- 707 Rodgers, P., Soulsby, C., Waldron, S. and Tetzlaff, D.: Using stable isotope tracers to
- 708 <u>identify hydrological flow paths, residence times and landscape controls in a</u>
- 709 mesoscale catchment, Hydrol. Earth Syst. Sci. Discuss., 9, 139–155, 2005.
- 710 Rose, L. A., Elliott, E. M. and Adams, M. B.: Triple Nitrate Isotopes Indicate
- 711 Differing Nitrate Source Contributions to Streams Across a Nitrogen Saturation
- 712 Gradient, Ecosystems, 18(7), 1209–1223, doi:10.1007/s10021-015-9891-8, 2015.
- 713 Sabo, R. D., Nelson, D. M. and Eshleman, K. N.: Episodic, seasonal, and annual
- r14 export of atmospheric and microbial nitrate from a temperate forest, Geophys. Res.
- 715 Lett., 43(2), 683–691, doi:10.1002/2015GL066758, 2016.
- 716 Sappa, G., Ferranti, F. and Pecchia, G. M.: Validation Of Salt Dilution Method For

- 717 Discharge Measurements In The Upper Valley Of Aniene River (Central Italy), Recent
- 718 Adv. Environ. Ecosyst. Dev., (October 2015), 42–48, 2015.
- 719 Soulsby, C., Tetzlaff, D., Rodgers, P., Dunn, S. and Waldron, S.: Runoff processes,
- 720 stream water residence times and controlling landscape characteristics in a mesoscale
- 721 catchment: An initial evaluation, J. Hydrol., 325(1–4), 197–221,
- 722 <u>doi:10.1016/j.jhydrol.2005.10.024, 2006.</u>
- 723 Stoddard, J. L.: Long-Term Changes in Watershed Retention of Nitrogen, , 223–284,
- 724 doi:10.1021/ba-1994-0237.ch008, 1994.
- 725 Tetzlaff, D., Malcolm, I. A. and Soulsby, C.: Influence of forestry, environmental
- 726 <u>change and climatic variability on the hydrology, hydrochemistry and residence times</u>
- 727 of upland catchments, J. Hydrol., 346(3–4), 93–111,
- 728 <u>doi:10.1016/j.jhydrol.2007.08.016, 2007.</u>
- 729 Tsunogai, U., Komatsu, D. D., Daita, S., Kazemi, G. A., Nakagawa, F., Noguchi, I.
- and Zhang, J.: Tracing the fate of atmospheric nitrate deposited onto a forest
- row ecosystem in Eastern Asia using Δ^{17} O, Atmos. Chem. Phys., 10(4), 1809–1820,
- 732 doi:10.5194/acp-10-1809-2010, 2010.
- 733 Tsunogai, U., Daita, S., Komatsu, D. D., Nakagawa, F. and Tanaka, A.: Quantifying
- nitrate dynamics in an oligotrophic lake using Δ^{17} O, Biogeosciences, 8(3), 687–702,
- 735 doi:10.5194/bg-8-687-2011, 2011.
- 736 Tsunogai, U., Komatsu, D. D., Ohyama, T., Suzuki, A., Nakagawa, F., Noguchi, I.,
- 737 Takagi, K. and Nomura, M.: Quantifying the effects of clear-cutting and strip-cutting

- on nitrate dynamics in a forested watershed using triple oxygen isotopes as tracers, ,
- 739 (1), 5411–5424, doi:10.5194/bg-11-5411-2014, 2014.
- 740 Tsunogai, U., Miyauchi, T., Ohyama, T., Komatsu, D. D., Nakagawa, F., Obata, Y.,
- 741 Sato, K. and Ohizumi, T.: Accurate and precise quantification of atmospheric nitrate
- in streams draining land of various uses by using triple oxygen isotopes as tracers,
- 743 Biogeosciences, 13(11), 3441–3459, doi:10.5194/bg-13-3441-2016, 2016.
- 744 Tsunogai, U., Miyauchi, T., Ohyama, T., Komatsu, D. D., Ito, M. and Nakagawa, F.:
- 745 Quantifying nitrate dynamics in a mesotrophic lake using triple oxygen isotopes as
- 746 tracers, Limnol. Oceanogr., 63, S458–S476, doi:10.1002/lno.10775, 2018.
- 747 Vitousek, P. M. and Howarth, R. W.: Nitrogen limitation on land and in the sea: How
- can it occur?, Biogeochemistry, 13(2), 87–115, doi:10.1007/BF00002772, 1991.
- 749 Watanabe, M., Miura, S., Hasegawa, S., Koshikawa, M. K., Takamatsu, T., Kohzu,
- A., Imai, A. and Hayashi, S.: Coniferous coverage as well as catchment steepness
- influences local stream nitrate concentrations within a nitrogen-saturated forest in
- 752 central Japan, Sci. Total Environ., 636, 539–546, doi:10.1016/j.scitotenv.2018.04.307,
- 753 2018.
- 754 Yamazaki, A., Watanabe, T. and Tsunogai, U.: Nitrogen isotopes of organic nitrogen
- in reef coral skeletons as a proxy of tropical nutrient dynamics, Geophys. Res. Lett.,
- 756 38(19), 1–5, doi:10.1029/2011GL049053, 2011.
- 757 Yang, R. and Chiwa, M.: Low nitrogen retention in a Japanese cedar plantation in a
- 758 suburban area, western Japan, Sci. Rep., 11(1), 1–7, doi:10.1038/s41598-021-84753-

759 1, 2021.

Overstory vegetation (%) FK1 FK2 MY Artificial Japanese cedar/cypress plantation 74 40 16 Other artificial coniferous plantations <1 <1 7 Natural trees 10 54 75 Others 16 5 2

760 **Table 1.** Plant information for each forested catchment (Chiwa, 2021).

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Table 2. Average concentrations of stream nitrate $([NO_3^-]_{avg})$, the average concentrations of unprocessed $NO_3^-_{atm}$ in streams $([NO_3^-]_{avg})$, the annual export flux of NO_3^- per unit area of catchments (M_{total}) , the annual export flux of $NO_3^-_{atm}$ per unit area of catchments (M_{atm}) , the deposition flux of $NO_3^-_{atm}$ per unit area of catchment (D_{atm}) , and the M_{atm}/D_{atm} ratios in the study catchments.

	FK1	FK2	MY
$[NO_3^-]_{avg}(\mu M)$	109.5	<u>90.9</u> 94.2	7.3
$[NO_{3}^{-}_{atm}]_{avg} (\mu M)$	10.80 ± 1.65	<u>6.095.06</u> ± <u>0.92</u> 1.05	0.16 ± 0.05
$M = (mm \circ 1 m^{-2} m^{-1})$	<u>98.8</u> 97.9 ±	<u>82.0</u> 84.2 ±	<u>23.7</u> 22.6 ±
$M_{total} \ (mmol \ m^{-2} \ yr^{-1})$	17.8	<u>14.8</u> 15.3	1.2
$M_{atm} \ (mmol \ m^{-2} \ yr^{-1})$	9.7 ± 2.3	<u>4.6</u> 5.4 ± <u>1.2</u> 1.4	$0.5\pm0.\underline{2}1$
$D_{atm} \ (mmol \ m^{-2} \ yr^{-1})$	69.3 ± 13.9	69.3 ± 13.9	40.1 ± 8.0
M_{atm}/D_{atm} (%)	<u>14.1</u> 13.9 ± 4. <u>4</u> 3	<u>6.6</u> 7.9 ± <u>2.1</u> 2.5	1.32 ± 0.54

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Table 3. The annual amount of precipitation (P), the average concentration of stream nitrate ($[NO_3^-]_{avg}$), the nitrogen saturation stage, the average concentration of unprocessed $NO_3^-_{atm}$ in streams ($[NO_3^-_{atm}]_{avg}$), the annual export flux of NO_3^- per unit area of catchment (M_{total}), the annual export flux of $NO_3^-_{atm}$ per unit area of catchment (M_{atm}), the deposition flux of $NO_3^-_{atm}$ per unit area of catchment (D_{atm}), and the

 M_{atm}/D_{atm} ratio in the FK1, FK2, and MY, along with those in the catchments studied in

past studies using Δ^{17} O of nitrate as a tracer.

	Р	[NO ₃ ⁻] _{avg}	N ata aa*	[NO3 ⁻ atm]avg	Matm	M _{total}	Datm	$M_{atm}\!/D_{atm}$
	mm	μΜ	N stage*	μM	m	mol m ⁻² yr	-1	%
FK1 ^a	<u>1777</u> 1769	109.5	-	10.8	9.7	<u>98.8</u> 97.9	69.3	<u>14.1</u> 13.9
FK2 ^a	<u>1777</u> 1769	<u>90.9</u> 94.2	-	<u>5.06</u> 6.1	<u>4.6</u> 5.4	<u>82.0</u> 84.2	69.3	<u>6.6</u> 7.9
MY ^a	<u>3981</u> 3837	7.3	-	0.2	0.5	<u>23.7</u> 22.6	40.1	1. <u>3</u> 2
KJ ^b	2500	58.4	-	3.3	4.3	76.4	45.6	9.4
IJ1 ^b	3300	24.4	2	1.4	2.9	50.1	44.5	6.5
IJ2 ^b	3300	17.1	-	0.6	1.2	35.1	44.5	2.6
Fernow1 ^c	1450	17.9	1	1.6	0.8	9.3	23.4	3.6
Fernow2 ^c	1450	34.3	2	3.4	1.5	14.8	23.4	6.3
Fernow3 ^c	1450	60.0	3	4.2	2.4	34.5	23.4	10.3
Uryu ^d	1170	0.7	-	0.1	0.1	1.0	18.6	0.7
Qingyuan ^e	709	150.0	2	8.9	2.9	49.3	50.0	5.8

780 a: This study

b: Nakagawa et al., 2018; Nakahara et al., 2010

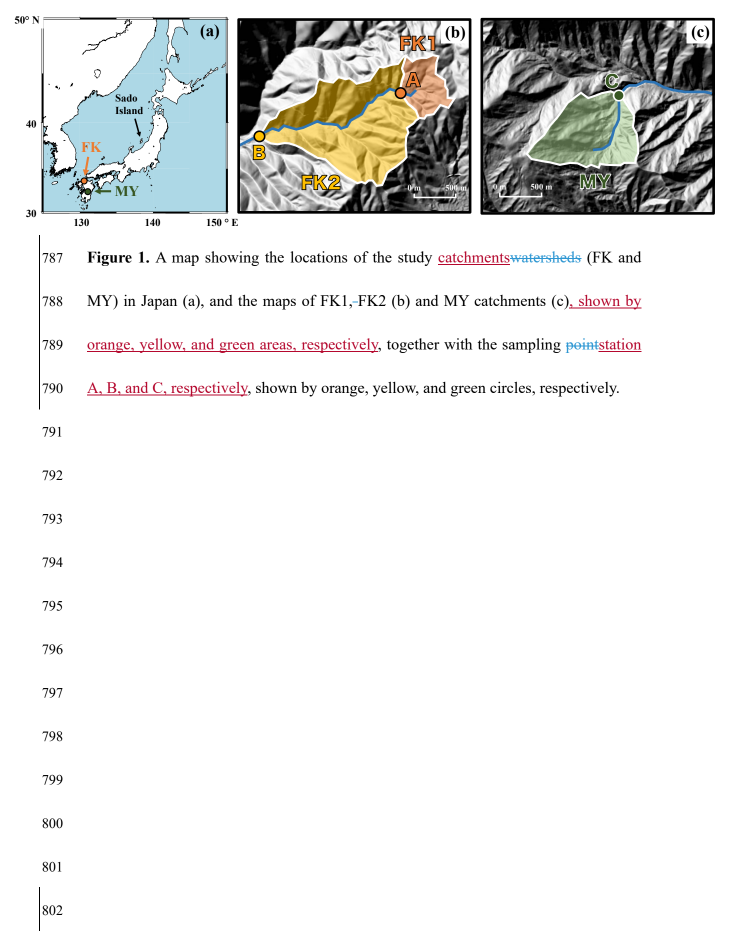
782 c: Rose et al., 2015

783 d: Tsunogai et al., 2014

784 e: Huang et al., 2020

785 *: N saturation stage estimated in past studies

786 -: No data



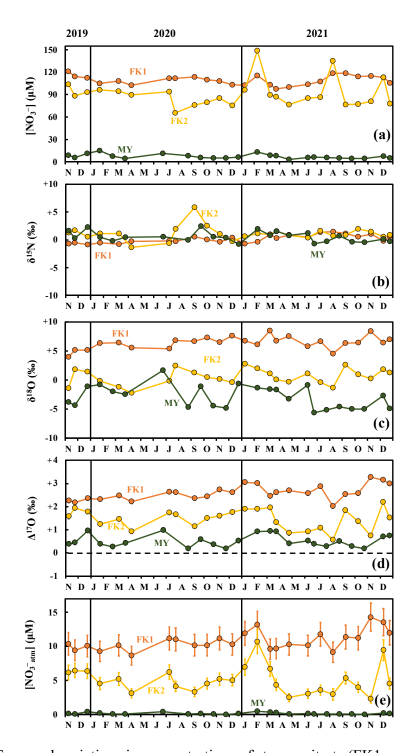
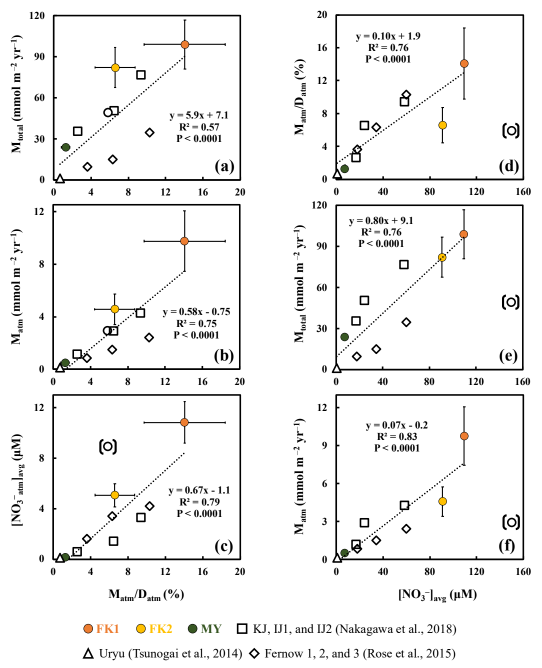


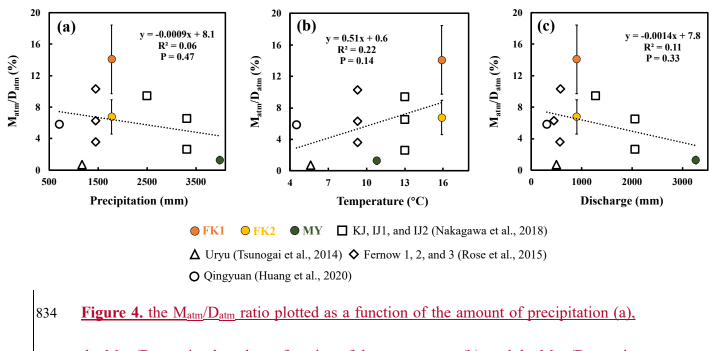
Figure 2. Temporal variations in concentrations of stream nitrate (FK1: orange circles; FK2: yellow circles; MY: green circles) (a), together with those in δ^{15} N (b), δ^{18} O (c), and Δ^{17} O (d) of nitrate, and the concentration of unprocessed NO_{3⁻atm} ([NO_{3⁻atm}]) (e) in the stream water of the FK1, FK2, and MY forested catchments. Error bars smaller than the sizes of the symbols are not presented.

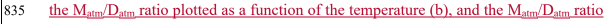


O Qingyuan (Huang et al., 2020)

Figure 3. Annual export flux of nitrate per unit area (M_{total}) plotted as a function of the M_{atm}/D_{atm} ratio in each forested catchment (a); the annual export flux of unprocessed atmospheric nitrate per unit area (M_{atm}) plotted as a function of the M_{atm}/D_{atm} ratio (b); the average concentration of NO_{3⁻atm} ([NO_{3⁻atm}]_{avg}) plotted as a function of the M_{atm}/D_{atm} ratio (c); the M_{atm}/D_{atm} ratio plotted as a function of the average concentration

813	of nitrate ([NO ₃ ⁻] _{avg}) (d); the M_{total} plotted as a function of [NO ₃ ⁻] _{avg} (e); the M_{atm}
814	plotted as a function of $[NO_3^-]_{avg}$ (f) (FK1: orange circles; FK2: yellow circles; MY:
815	green circles). Those determined for the forested catchments in past studies are plotted
816	as well (Qingyuan: white circle (Huang et al., 2020); KJ, IJ1, and IJ2: white squares
817	(Nakagawa et al., 2018); Fernow 1, 2, and 3: white diamonds (Lucy et al., 2015); Uryu:
818	white triangle (Tsunogai., 2014)). The data obtained in the Qingyuan forested
819	catchment are shown in parentheses and excluded from the calculation to estimate
820	correlation coefficients (see text for the reason).
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836 plotted as a function of flux of stream water (c) (FK1: orange circles; FK2: yellow

837 <u>circles; MY: green circles). Those determined for the forested catchments in past studies</u>

838 <u>are plotted as well.</u>

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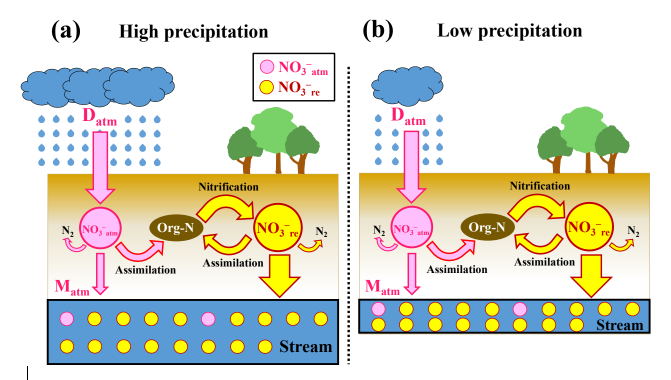


Figure 54. Schematic diagram showing the biogeochemical processing of nitrate in 842 843 forested catchments under high precipitation (a) and low precipitation (b), where NO_{3⁻atm} (unprocessed atmospheric nitrate) is represented by pink circles, NO_{3⁻re} by 844 yellow circles, the flows of NO_{3-atm}^{-} by pink arrows, and those of NO_{3-re}^{-} (remineralized 845 846 nitrate) by yellow arrows (modified after Nakagawa., 2018). Although the deposition rates of NO_{3-atm}^{-} (D_{atm}) and the biogeochemical reaction rates between (a) and (b) are 847 the same, we can expect high $[NO_3^-]$ in (b). On the other hand, the M_{atm}/D_{atm} ratio 848 between (a) and (b) are the same. 849