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

# nitrate from forested catchments under high nitrogen saturation

Weitian Ding<sup>1</sup>, Urumu Tsunogai<sup>1</sup>, Fumiko Nakagawa<sup>1</sup>, Takashi Sambuichi<sup>1</sup>, Masaaki Chiwa<sup>2</sup>, Tamao Kasahara<sup>3</sup>, Ken'ichi Shinozuka<sup>4</sup>

<sup>1</sup>Graduate School of Environmental Studies, Nagoya University, Furo-cho, Chikusa-ku,

Nagoya 464-8601, Japan

<sup>2</sup>Kyushu University Forest, Kyushu University, Japan

<sup>3</sup>Faculty of Agriculture, Kyushu University, Japan

<sup>4</sup>River Basin Research Center, Gifu University, 1-1 Yanagido, Gifu, 501-1193, Japan

Corresponding author: Weitian Ding

Email: ding.weitian.v2@s.mail.nagoya-u.ac.jp

#### 3 Abstract

Owing to the elevated loading of nitrogen through atmospheric deposition, some 4 forested ecosystems become nitrogen saturated, from which elevated levels of nitrate 5 are exported. The average concentration of stream nitrate eluted from upstream and 6 downstream of the Kasuya Research forested catchments (FK1 and FK2 catchments) 7 in Japan were more than 90 µM, implying that these forested catchments were under 8 nitrogen saturation. To verify that these forested catchments were under the nitrogen 9 saturation, we determined the export flux of unprocessed atmospheric nitrate relative to 10 the entire deposition flux (Matm/Datm ratio) in these catchments, because the Matm/Datm 11 ratio has recently been proposed as a reliable index to evaluate nitrogen saturation in 12 forested catchments. Specifically, we determined the temporal variation in the 13 concentrations and stable isotopic compositions, including  $\Delta^{17}$ O, of stream nitrate in 14 15 the FK catchments for more than 2 years. In addition, for comparison, the same parameters were also monitored in the Shiiba Research forested catchment (MY 16 catchment) in Japan during the same period, where the average stream nitrate 17 18 concentration was low, less than 10 µM. While showing the average nitrate concentrations of 109.5, 90.9, and 7.3 µM in FK1, FK2, and MY, respectively, the 19 catchments showed average  $\Delta^{17}$ O values of +2.6, +1.5, and +0.6 ‰ in FK1, FK2, and 20 21 MY, respectively. Thus, the average concentration of unprocessed atmospheric nitrate ( $[NO_3_{atm}]$ ) was estimated to be 10.8, 5.1, and 0.2 µM in FK1, FK2, and MY, 22 respectively, and the M<sub>atm</sub>/D<sub>atm</sub> ratio was estimated to be 14.1, 6.6, and 1.3 % in FK1, 23

24	FK2, and MY, respectively. The estimated $M_{atm}/D_{atm}$ ratio in FK1 (14.1 %) was the
25	highest ever reported from temperate forested catchments monitored for more than 1
26	year. Thus, we concluded that nitrogen saturation was responsible for the enrichment
27	of stream nitrate in the FK catchments, together with the elevated NO3 <sup>-</sup> <sub>atm</sub> leaching
28	from the catchments. While the stream nitrate concentration ( $[NO_3^-]$ ) can be affected
29	by the amount of precipitation, the $M_{\text{atm}}/D_{\text{atm}}$ ratio is independent of the amount of
30	precipitation; thus, the $M_{atm}/D_{atm}$ ratio can be used as a robust index for evaluating
31	nitrogen saturation in forested catchments.

## 33 **1 Introduction**

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

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

3

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  $\mu$ 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 soil nitrate can be buffered 53 by groundwater with long residence time, so that the seasonal variation is unclear in 54 stream nitrate concentration in Japan, even in normal forests under the nitrogen 55 saturation stage of 0 (Mitchell et al., 1997); and (2) the stream nitrate concentration can 56 57 be enriched or diluted depending on the volume of rainfall, so the concentration level can be high in low precipitation area irrespective of the stage of nitrogen saturation. 58

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  $M_{atm}/D_{atm}$  <sup>66</sup> ratios in forested catchments with nitrogen deficiency.

To estimate the M<sub>atm</sub>/D<sub>atm</sub> ratio accurately and precisely in each forested catchment, 67 the fraction of unprocessed atmospheric nitrate (NO<sub>3<sup>-</sup>atm</sub>) in the stream needs to be 68 estimated accurately and precisely. Triple oxygen isotopic compositions of nitrate 69 ( $\Delta^{17}$ O) have recently been used as a conservative tracer of NO<sub>3<sup>-</sup>atm</sub> deposited onto each 70 forested catchment (Inoue et al., 2021; Michalski et al., 2004; Nakagawa et al., 2018; 71 Tsunogai et al., 2014; Ding et al., 2022), showing distinctively different  $\Delta^{17}$ O from that 72 of remineralized nitrate (NO3<sup>-</sup>re), derived from organic nitrogen through general 73 chemical reactions, including microbial N mineralization and microbial nitrification. 74 While NO<sub>3<sup>-</sup>re</sub>, the oxygen atoms of which are derived from either terrestrial O<sub>2</sub> or H<sub>2</sub>O 75 through microbial processing (i.e., nitrification), always shows the relation close to the 76 "mass-dependent" relative relation between <sup>17</sup>O/<sup>16</sup>O ratios and <sup>18</sup>O/<sup>16</sup>O ratios; NO<sub>3<sup>-</sup>atm</sub> 77 displays an anomalous enrichment in <sup>17</sup>O reflecting oxygen atom transfers from 78 atmospheric ozone (O<sub>3</sub>) during the conversion of NO<sub>X</sub> to NO<sub>3<sup>-</sup>atm</sub> (Alexander et al., 79 80 2009; Michalski et al., 2003; Morin et al., 2011; Nelson et al., 2018). As a result, the 81  $\Delta^{17}$ O signature defined by the following equation (Kaiser et al., 2007) enables us to distinguish NO<sub>3<sup>-</sup>atm</sub> ( $\Delta^{17}$ O > 0) from NO<sub>3<sup>-</sup>re</sub> ( $\Delta^{17}$ O = 0): 82  $\Delta^{17} O = \frac{1 + \delta^{17} O}{(1 + \delta^{18} O)^{\beta}} - 1$ 83 (1)

84 where the constant 
$$\beta$$
 is 0.5279 (Kaiser et al., 2007),  $\delta^{18}O = R_{\text{sample}}/R_{\text{standard}} - 1$  and  $R$  is  
85 the <sup>18</sup>O/<sup>16</sup>O ratio (or the <sup>17</sup>O/<sup>16</sup>O ratio in the case of  $\delta^{17}O$  or the <sup>15</sup>N/<sup>14</sup>N ratio in the case  
86 of  $\delta^{15}N$ ) of the sample and each standard reference material. In addition,  $\Delta^{17}O$  is almost

87	stable during "mass-dependent" isotope fractionation processes within terrestrial
88	ecosystems. Therefore, while the $\delta^{15}N$ or $\delta^{18}O$ signature of $NO_3^-$ atm can be overprinted
89	by the biological processes subsequent to deposition, $\Delta^{17}O$ can be used as a robust tracer
90	of unprocessed $NO_3^{atm}$ to reflect its accurate mole fraction within total $NO_3^-$ ,
91	regardless of the progress of the partial metabolism (partial removal of nitrate through
92	denitrification and assimilation) subsequent to deposition (Michalski et al., 2004;
93	Nakagawa et al., 2013, 2018; Tsunogai et al., 2011, 2014, 2018).
94	Past studies reported that the maximum concentration of stream nitrate was 58.4 $\mu$ M
95	in the KJ forested catchment in Japan, with the maximum value of the $M_{\text{atm}}/D_{\text{atm}}$ ratio
96	was 9.4 % (Nakagawa et al., 2018; Sase et al., 2022). Whether the index of the $M_{atm}/D_{atm}$
97	ratio can be applied to forested catchments, where the leaching of stream nitrate is much
98	higher than the KJ forested catchment, remained unclarified. Besides, the advantages
99	of the $M_{\text{atm}}/D_{\text{atm}}$ ratio within the past indexes of nitrogen saturation have not been
100	discussed.

101 Chiwa (2021) has recently reported the enrichment of nitrate of more than 90  $\mu$ M on 102 the annual average in forested streams eluted from the FK catchments (FK1 and FK2) 103 in Kasuya Research Forest, Kyushu University, Japan (Figs. 1a and 1b). The observed 104 enrichment of stream nitrate implied that these forested catchments were under nitrogen 105 saturation. Thus, in this study, we determined the M<sub>atm</sub>/D<sub>atm</sub> ratio in the FK1 and FK2 106 forested catchments by monitoring both the concentration and  $\Delta^{17}$ O of stream nitrate 107 for more than 2 years to verify that these forested catchments were under nitrogen

108 saturation. For comparison, the MY forested catchment in Shiiba Research Forest, 109 Kyushu University, Japan (Figs. 1a and 1c), was also monitored during the same period, 110 where the average stream nitrate concentration was low (less than 10  $\mu$ M). Furthermore, 111 the M<sub>atm</sub>/D<sub>atm</sub> ratios in these forested catchments were compared with those reported in 112 past studies to verify the reliability of the M<sub>atm</sub>/D<sub>atm</sub> ratio as an index of nitrogen 113 saturation.

114

115 2 Methods

116 2.1 Study sites

The FK forested catchments (33°38'N, 130°31'E) are located in a suburban area, 117 about 15 km west of the Fukuoka metropolitan area (the fourth largest metropolitan 118 area in Japan). The main plantation in these catchments was Japanese cedar/cypress 119 120 (Table 1). The MY forested catchment (32°22'N, 131°09'E) is located in a rural area at the village of Shiiba in southern Japan's Central Kyushu Mountain range. This 121 catchment is a mixed forest consisting of coniferous trees such as Abies firma Sieb. et 122 123 Zucc., and Tsuga sieboldii Carr., and deciduous broadleaved trees such as Quercus crispula Blume, Fagus crenata Blume, and Acer sieboldianum Miq. Details on the 124 studied forested catchments have been described in the past studies (Chiwa, 2020, 2021). 125 126

127 2.2 Sampling

128 The stream water eluted from the FK1 (14 ha), FK2 (62 ha), and MY (43 ha)

7

129	forested catchments were collected about once every month in principle from 2019/11
130	to 2021/12 (Fig. 1). At the FK catchments, stream water was collected at upstream
131	(station A) and downstream (station B) locations (Fig. 1b). At the MY catchment,
132	stream water was collected at station C (Fig. 1c). Samples of stream water to determine
133	the concentration and stable isotopic compositions ( $\delta^{15}N$ , $\delta^{18}O$ , and $\Delta^{17}O$ ) of stream
134	nitrate were collected manually in bottles washed with deionized water before sampling
135	and then rinsed at least twice with the sample before sampling at each sampling site.

137 2.3 Analysis

All the stream water samples were passed through a membrane filter (pore size 0.45 138 μm) within two days after sampling and stored in a refrigerator (4 °C) until analysis. 139 140 The concentrations of nitrate were measured by ion chromatography (Prominence HIC-141 SP, Shimadzu, Japan). To determine the stable isotopic compositions of nitrate in the stream water samples, nitrate in each sample was chemically converted to N<sub>2</sub>O using a 142 method originally developed to determine the <sup>15</sup>N/<sup>14</sup>N and <sup>18</sup>O/<sup>16</sup>O ratios of seawater 143 144 and freshwater nitrate (McIlvin and Altabet, 2005) that was later modified (Konno et al., 2010; Tsunogai et al., 2011; Yamazaki et al., 2011). In brief, 11 mL of each sample 145 solution was pipetted into a vial with a septum cap. Then, 0.5 g of spongy cadmium 146 was added, followed by 150 µL of a 1 M NaHCO<sub>3</sub> solution. The sample was then shaken 147 for 18-24 h at a rate of 2 cycles s<sup>-1</sup>. Then, the sample solution (10 mL) was decanted 148 into a different vial with a septum cap. After purging the solution using high-purity 149

150	helium, 0.4 mL of an azide-acetic acid buffer, which had also been purged using high-
151	purity helium, was added. After 45 min, the solution was alkalinized by adding 0.2 mL
152	of 6 M NaOH. Then, the stable isotopic compositions ( $\delta^{15}N$ , $\delta^{18}O$ , and $\Delta^{17}O$ ) of the
153	N <sub>2</sub> O in each vial were determined using the continuous-flow isotope ratio mass
154	spectrometry (CF-IRMS) system at Nagoya University. The analytical procedures
155	performed using the CF-IRMS system were the same as those detailed in previous
156	studies (Hirota et al., 2010; Komatsu et al., 2008a). The obtained values of $\delta^{15}$ N, $\delta^{18}$ O,
157	and $\Delta^{17}O$ for the N <sub>2</sub> O derived from the nitrate in each sample were compared with those
158	derived from our local laboratory nitrate standards to calibrate the values of the sample
159	nitrate to an international scale and to correct for both isotope fractionation during the
160	chemical conversion to $N_2O$ and the progress of oxygen isotope exchange between the
161	nitrate derived reaction intermediate and water (ca. 20 %). In this study, we adopted the
162	internal standard method to calibrate the stable isotopic compositions of sample nitrate.
163	Specifically, three kinds of the local laboratory nitrate standards were used in this study,
164	which were named to be GG01 ( $\delta^{15}N = -3.07 \%$ , $\delta^{18}O = +1.10 \%$ , and $\Delta^{17}O = 0 \%$ ),
165	HDLW02 ( $\delta^{15}N = +8.94$ ‰, $\delta^{18}O = +24.07$ ‰), and NF ( $\Delta^{17}O = +19.16$ ‰), which the
166	GG01 and the HDLW02 were used to determine the $\delta^{15}N$ and $\delta^{18}O$ of stream nitrate,
167	and the GG01 and the NF was used to determine the $\Delta^{17}$ O of stream nitrate. The GG01,
168	HDLW02, and NF had been calibrated using the internationally distributed isotope
169	reference materials (USGS 34 and USGS 35). The oxygen exchange rate between
170	nitrate and water during the chemical conversion was calculated through Eq. (2):

171 Oxygen exchange rate (%) = 
$$\Delta^{17}O(N_2O)_{NF} / \Delta^{17}O(NO_3)_{NF}$$
 (2)

where the  $\Delta^{17}O(N_2O)_{NF}$  denote the  $\Delta^{17}O$  value of N<sub>2</sub>O that convert from the NF

172

)NF / $\Delta^{1}$	$O(NO_3)$	)NF		

nitrate, the  $\Delta^{17}O(NO_3)_{NF}$  denote the  $\Delta^{17}O$  value of NF nitrate ( $\Delta^{17}O = +19.16$  ‰) 173 (Tsunogai et al., 2016; Nakagawa et al., 2013, 2018; Ding et al., 2022). 174 The  $\delta^2$ H and  $\delta^{18}$ O values of H<sub>2</sub>O of the stream water samples were analyzed using 175 the cavity ring-down spectroscopy method by employing an L2120-i instrument 176 (Picarro Inc., Santa Clara, CA, USA) equipped with an A0211 vaporizer and 177 autosampler. The errors (standard errors of the mean) in this method were  $\pm 0.5\%$  for 178  $\delta^2$ H and  $\pm 0.1\%$  for  $\delta^{18}$ O. Both the VSMOW and standard light Antarctic precipitation 179 (SLAP) were used to calibrate the values to the international scale. The  $\delta^{18}$ O values of 180  $H_2O$  were used to calibrate the differences in  $\delta^{18}O$  of  $H_2O$  between the samples and 181 those our local laboratory nitrate standard samples (Tsunogai et al., 2010, 2011, 2014). 182 183 To determine whether the conversion rate from nitrate to N<sub>2</sub>O was sufficient, the concentration of nitrate in the samples was determined each time we analyzed the 184 isotopic composition using CF-IRMS based on the  $N_2O^+$  or  $O_2^+$  outputs. We adopted 185 the  $\delta^{15}$ N,  $\delta^{18}$ O, and  $\Delta^{17}$ O values only when the concentration measured via CF-IRMS 186 correlated with the concentration measured via ion chromatography prior to isotope 187 analysis within a difference of 10 %. We repeated the analysis of  $\delta^{15}$ N,  $\delta^{18}$ O, and  $\Delta^{17}$ O 188 values for each sample at least three times to attain high precision. All samples had a 189 nitrate concentration of greater than 3.5 µM, which corresponded to a nitrate quantity 190 greater than 35 nmol in a 10 mL sample. Thus, all isotope values presented in this study 191

have an error (standard error of the mean) better than  $\pm 0.2$  ‰ for  $\delta^{15}$ N,  $\pm 0.3$  ‰ for  $\delta^{18}$ O, and  $\pm 0.1$  ‰ for  $\Delta^{17}$ O.

Nitrite  $(NO_2^{-})$  in the samples interferes with the final N<sub>2</sub>O produced from nitrate because the chemical method also converts  $NO_2^{-}$  to N<sub>2</sub>O (McIlvin and Altabet, 2005). Therefore, it is sometimes necessary to remove  $NO_2^{-}$  prior to converting nitrate to N<sub>2</sub>O. In this study, however, we skipped the processes for removing  $NO_2^{-}$  because all the stream samples analyzed for stable isotopic composition had  $NO_2^{-}$  concentrations lower than the detection limit (0.05  $\mu$ M).

200

201 2.4 Deposition rate of atmospheric nitrate

The annual deposition rate of atmospheric nitrate (D<sub>atm</sub>; total dry and wet deposition 202 rate of atmospheric nitrate) in each catchment was estimated using the annual "bulk" 203 204 deposition rate of atmospheric nitrate (D<sub>bulk</sub>) calculated in Chiwa (2020) at each catchment by multiplying the volume-weighted mean concentration of nitrate in the 205 bulk deposition samples collected every 2 weeks at each catchment for 10 years (from 206 207 2009/1 to 2018/12) by the annual amount of precipitation. The bulk deposition samples were those accumulated in a plastic bucket installed in an open site of each catchment 208 55 cm above the ground. The distances between the monitoring sites of bulk deposition 209 in the FK1, FK2, and MY forested catchments and the stations of stream water sampling 210 (stations A, B, and C) were 3.9, 2.9, and 4.5 km, respectively. The concentrations of 211 nitrate in the bulk deposition samples were measured by ion chromatography. 212

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. (3) to estimate  $D_{atm}$  from  $D_{bulk}$ :

217 
$$D_{atm} = D_{bulk} - D_{dry}(W) + D_{dry}(F)$$
(3)

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

219 respectively.

The D<sub>dry</sub>(W) and D<sub>dry</sub>(F) at each catchment were determined using an inferential method (Endo et al., 2011) through Eqs. (4) and (5), respectively:

222 
$$D_{dry}(W) = [NO_3^{-}_{atm}]_{gas} \times V_{gas}(W) + [NO_3^{-}_{atm}]_p \times V_p(W)$$
(4)

223 
$$D_{dry}(F) = [NO_3^-_{atm}]_{gas} \times V_{gas}(F) + [NO_3^-_{atm}]_p \times V_p(F)$$
(5)

224 where [NO<sub>3<sup>-</sup>atm]gas</sub> denotes the concentration of gaseous nitrate in air; [NO<sub>3<sup>-</sup>atm]p</sub> 225 denotes the concentration of particle nitrate in air; Vgas(W) and Vgas(F) denote the deposition velocities of gaseous nitrate on the water surface and forest, respectively; 226 and  $V_p(W)$  and  $V_p(F)$  denote the deposition velocities of particulate nitrate on the water 227 228 surface and forest, respectively. Those determined by Chiwa (2010) using the annular denuder method from 2006/5 to 2007/4 were used for the  $[NO_3^-]_{gas}$  and  $[NO_3^-]_p$  in the 229 FK catchments. Those determined by the National Institute for Environmental Studies 230 (Environmental Laboratories Association of Japan, 2017) using the filter-pack method 231 at Miyazaki (31°83'N, 131°42'E) from 2011 to 2017 were used for the [NO<sub>3</sub><sup>-</sup>]<sub>gas</sub> and 232  $[NO_3^-]_p$  in the MY catchment. The  $V_{gas}(F)$ ,  $V_{gas}(W)$ ,  $V_p(F)$ , and  $V_p(W)$  of each 233

catchment were determined by applying the estimation file for dry deposition (Matsuda,235 2008;

236	http://www.hro.or.jp/list/environmental/research/ies/katsudo/acid_rain/kanseichinchak
237	u/kanseichinchaku.html), where $V_{\text{gas}}$ and $V_{\text{p}}$ were calculated using the meteorological
238	data of wind speed, temperature, humidity, radiation, and cloud amount and land use.
239	The meteorological data monitored by Japan Meteorological Agency at the nearest
240	Fukuoka station (33°34'N, 130°22'E) and Miyazaki station (31°56'N, 131°24'E) from
241	2009 to 2021 were used for the FK and MY catchments, respectively. The forested land
242	use of 100 % was chosen for each area.

243

# 244 2.5 Flux of stream water

The flux of stream water ( $F_{stream}$ ) in each catchment was not measured fully 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:

$$F_{\text{stream}} = P - E \tag{6}$$

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

255 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 256 of E (mm) =  $31.4T_{avg}$  (°C) + 376 to estimate E in each forested catchment in Japan, 257 where the standard error of 162.3 mm was included in the estimated evapotranspiration 258 flux (E). They also confirmed that the estimated F<sub>stream</sub> using the model corresponded 259 well with the observed F<sub>stream</sub> in three forested catchments, with estimated errors of less 260 than 6 %. As a result, we utilized the water balance method proposed by Komatsu et al. 261 (2008) to quantify the F<sub>stream</sub> in each catchment. 262

263

264 2.6 Concentration of unprocessed NO<sub>3<sup>-</sup>atm</sub> in each water sample

265 The  $\Delta^{17}$ O data of nitrate in each sample was used to estimate the concentration of 266 NO<sub>3<sup>-</sup>atm</sub> ([NO<sub>3<sup>-</sup>atm</sub>]) in each water sample by applying Eq. (7):

267 
$$[NO_3^-_{atm}]/[NO_3^-] = \Delta^{17}O/\Delta^{17}O_{atm}$$
 (7)

where [NO<sub>3<sup>-</sup>atm</sub>] and [NO<sub>3<sup>-</sup></sub>] denote the concentrations of NO<sub>3<sup>-</sup>atm</sub> and nitrate (total) in 268 each water sample, respectively, and  $\Delta^{17}O_{atm}$  and  $\Delta^{17}O$  denote the  $\Delta^{17}O$  values of 269 270 NO<sub>3<sup>-</sup>atm</sub> and nitrate (total) in the stream water sample, respectively. In this study, we used the annual average  $\Delta^{17}$ O value of NO<sub>3<sup>-</sup>atm</sub> determined at the Sado-Seki monitoring 271 station in Japan (Sado Island; Fig. 1a) from April 2009 to March 2012 ( $\Delta^{17}O_{atm} =$ 272 +26.3 %; Tsunogai et al., 2016) for  $\Delta^{17}O_{atm}$  in Eq. (7) to estimate [NO<sub>3</sub><sup>-</sup><sub>atm</sub>] in the stream. 273 We allow for an error range of 3 % in  $\Delta^{17}O_{atm}$ , where the factor changes in  $\Delta^{17}O_{atm}$ 274 from +26.3 ‰ caused by both areal and seasonal variations in the  $\Delta^{17}$ O values of 275

NO<sub>3<sup>-</sup>atm</sub> have been considered (Nakagawa et al., 2018; Tsunogai et al., 2016; Ding et
al., 2022).

The annual export flux of unprocessed  $NO_{3}^{-}_{atm}$  per unit area of the catchment (M<sub>atm</sub>) was determined by applying Eq. (8):

280 
$$M_{\text{atm}} = [NO_3^{-}_{\text{atm}}]_{\text{avg}} \times F_{\text{stream}}$$
(8)

where  $[NO_3^-atm]_{avg}$  denotes the annual average  $[NO_3^-atm]$  in each stream. The index of nitrogen saturation (M<sub>atm</sub>/D<sub>atm</sub> ratio) was calculated by dividing M<sub>atm</sub> with D<sub>atm</sub> in each catchment.

284

285 2.7 Concentration and isotopic compositions of stream nitrate eluted only from the FK2286 catchment

The concentration and isotopic compositions ( $\delta^{15}N$ ,  $\delta^{18}O$ , and  $\Delta^{17}O$ ) of stream nitrate determined at station B were the mixtures of those eluted from FK1 and FK2 catchments (Fig. 1b). Assuming that the stream nitrate eluted from FK1 catchment was stable during the flow path from station A to station B. The concentration of stream nitrate eluted from the FK2 catchment was determined by applying Eq. (9):

292 
$$[NO_3^-]_{FK2} = ([NO_3^-]_{FK1+FK2} * F_{FK1+FK2} - [NO_3^-]_{FK1} * F_{FK1}) / F_{FK2}$$
(9)

where  $F_{FK1}$ ,  $F_{FK2}$ , and  $F_{FK1+FK2}$  denote the flux of stream water eluted from the FK1, FK2 (only), and FK1+FK2 catchment, respectively.  $[NO_3^-]_{FK1}$ ,  $[NO_3^-]_{FK2}$ , and  $[NO_3^-]_{FK1+FK2}$  denote the concentration of stream nitrate eluted from the FK1, FK2 (only), and FK1+FK2 catchment, respectively. In this study, the flow rates measured at

297	stations A and B on 2021/01/15 by using the salt dilution method (Sappa et al., 2015)
298	was used for $F_{FK1}$ (0.85 L/s) and $F_{FK1+FK2}$ (4.75 L/s), respectively, and the measured
299	$[NO_3^-]$ at stations A and B was used for $[NO_3^-]_{FK1}$ and $[NO_3^-]_{FK1+FK2}$ , respectively.
300	Because the relation between the measured flow rates was comparable with the relation
301	between the catchment area of FK1 (14 ha) and that of FK1+FK2 (76 ha), we concluded
302	that the measured flow rates of 0.85 L/s and 4.75 L/s were reasonable as for those
303	representing the $F_{FK1}$ and $F_{FK1+FK2}$ , respectively. According to the mass balance of water,
304	we can estimate the $F_{FK2}$ eluted from the FK2 catchment only to be 3.90 L/s.
305	Assuming that the stream nitrate eluted from FK1 catchment was stable during the
306	flow path from station A to station B, the $\delta^{15}$ N, $\delta^{18}$ O, and $\Delta^{17}$ O values of stream nitrate
307	eluted from the FK2 catchment only were determined by applying Eq. (10):
308	$\delta_{FK2} = (\delta_{FK1+FK2} * [NO_3^{-}]_{FK1+FK2} * F_{FK1+FK2} - \delta_{FK1} * [NO_3^{-}]_{FK1} * F_{FK1}) / ([NO_3^{-}]_{FK2} * F_{FK1}) / ([NO_3^{-}]_{FK2} * F_{FK1}) / ([NO_3^{-}]_{FK2} * F_{FK1}) / ([NO_3^{-}]_{FK1} * F_{FK1} * F_{FK1} * F_{FK1}) / ([NO_3^{-}]_{FK1} * F_{FK1} $
309	(10)
310	where $\delta_{FK1}$ , $\delta_{FK2}$ , and $\delta_{FK1+FK2}$ denote the $\delta^{15}N$ (or $\delta^{18}O$ or $\Delta^{17}O$ ) of stream nitrate eluted
311	from the FK1, FK2, and FK1+FK2 catchment, respectively. The $\delta^{15}N$ (or $\delta^{18}O$ or $\Delta^{17}O$ )
312	values of stream nitrate measured at stations A and B were used for $\delta_{FK1}$ and $\delta_{FK1+FK2}$ ,
313	respectively.
314	
215	2 D 14-

- 315 **3 Results**
- 316 3.1 Deposition rate of atmospheric nitrate

The mean annual precipitation (P) from 2009 to 2021 was 1777 mm and 3981 mm

for FK and MY catchments, respectively (Chiwa, 2020; Chiwa, personal communication, September 21, 2022). The mean annual temperature ( $T_{avg}$ ) was reported to be 15.9 °C and 10.8 °C for FK and MY catchments, respectively (Chiwa, 2020). Based on these data, the annual flux of stream water ( $F_{stream}$ ) was estimated to be 902.0 ± 162.3 mm at FK catchments and 3266.1 ± 162.3 mm at MY catchment, respectively, using Eq. (6).

Chiwa (2020) reported the annual bulk deposition rates of atmospheric nitrate (D<sub>bulk</sub>) 324 to be 34.0 mmol m<sup>-2</sup> year<sup>-1</sup> at FK catchments and 24.2 mmol m<sup>-2</sup> year<sup>-1</sup> at MY 325 catchment. On the other hand, the annual dry deposition rate of atmospheric nitrate 326  $(D_{dry})$  deposited on the forest  $(D_{dry}(F))$  and on the water surface  $(D_{dry}(W))$  were 327 estimated to be 39.9 mmol m<sup>-2</sup> year<sup>-1</sup> and 4.1 mmol m<sup>-2</sup> year<sup>-1</sup>, respectively, at FK 328 catchments, and 18.4 mmol m<sup>-2</sup> year<sup>-1</sup> and 2.4 mmol m<sup>-2</sup> year<sup>-1</sup>, respectively, at MY 329 330 catchment. As a result, D<sub>atm</sub> was estimated to be 69.3 mmol m<sup>-2</sup> year<sup>-1</sup> at FK catchments and 40.1 mmol  $m^{-2}$  year<sup>-1</sup> at MY catchment, using Eq. (3). 331

332

## 333 3.2 Concentration and isotopic composition of stream nitrate

The concentrations of stream nitrate eluted from the FK1, FK2 (only), and MY catchments ranged from 97.5  $\mu$ M to 121.3  $\mu$ M, from 65.7  $\mu$ M to 148.5  $\mu$ M, and from 3.5  $\mu$ M to 15.3  $\mu$ M, respectively, with the average concentrations of 109.5  $\mu$ M, 90.9  $\mu$ M, and 7.3  $\mu$ M, respectively, and the standard deviations (SD) of 6.3  $\mu$ M, 18.5  $\mu$ M, and 3.0  $\mu$ M, respectively, which corresponds to the coefficients of variation (CV) of

339	5.7 %, 20.4 %, and 40.7 %, respectively (Fig. 2a). All catchments showed no clear
340	seasonal variation during the observation periods. The variation ranges and the average
341	concentrations of stream nitrate eluted from the three catchments agreed well with the
342	past observations performed in the same catchments (Chiwa, 2021).
343	The stable isotopic compositions of stream nitrate eluted from the FK1, FK2 (only),
344	and MY catchments ranged from –0.9 ‰ to +1.5 ‰, from –1.4 ‰ to +5.8 ‰, and from
345	$-0.8$ ‰ to +2.4 ‰, respectively, for $\delta^{15}N$ (Fig. 2b), from +3.9 ‰ to +8.5 ‰, from -2.2 ‰
346	to +2.8 ‰, and from –5.6 ‰ to +1.7 ‰, respectively, for $\delta^{18}$ O (Fig. 2c), and from +2.0 ‰
347	to +3.3 ‰, from +0.6 ‰ to +2.2 ‰, and from +0.2 ‰ to +1.0 ‰, respectively, for $\Delta^{17}O$
348	(Fig. 2d), with no clear seasonal variation during the observation periods. The
349	concentration-weighted averages for the $\delta^{15}$ N, $\delta^{18}$ O, and $\Delta^{17}$ O values of stream nitrate
350	were +0.2 ‰, +6.4 ‰, and +2.6 ‰, respectively, at FK1, +1.0 ‰, +0.5 ‰, and +1.5 ‰,
351	respectively, at FK2, $+0.7 \%$ , $-2.5 \%$ , and $+0.6 \%$ , respectively, at MY.
352	
353	3.3 Concentration of unprocessed atmospheric nitrate and the $M_{\text{atm}}/D_{\text{atm}}$ ratio in each
354	catchment
355	The concentration of unprocessed atmospheric nitrate ( $[NO_3^-]_{atm}$ ) in the streams eluted
356	from the FK1, FK2 (only), and MY catchments ranged from 8.64 to 14.30 $\mu M,$ from
357	2.27 to 10.71 $\mu M,$ and from 0.03 to 0.46 $\mu M$ with the average concentration of 10.80 $\pm$
358	1.30, 5.06 $\pm$ 0.67, and 0.16 $\pm$ 0.03 $\mu M,$ respectively, even though these studied
359	catchments showed little seasonal variations during the observation periods (Fig. 2e).

360	The annual export flux of nitrate ( $M_{total}$ ), the annual export flux of $NO_3^-$ <sub>atm</sub> ( $M_{atm}$ ), and
361	the $M_{atm}/D_{atm}$ ratio were 98.8 $\pm$ 17.8 mmol m <sup>-2</sup> year <sup>-1</sup> , 9.7 $\pm$ 2.1 mmol m <sup>-2</sup> year <sup>-1</sup> , and
362	14.1 $\pm$ 4.1 % at FK1 catchment, respectively, 82.0 $\pm$ 14.8 mmol m^{-2} year^{-1}, 4.6 $\pm$ 1.0
363	mmol m <sup>-2</sup> year <sup>-1</sup> , and 6.6 $\pm$ 2.0 % at FK2 catchment, respectively, 23.7 $\pm$ 1.2 mmol m <sup>-</sup>
364	$^2$ year $^{-1},0.5\pm0.1$ mmol m $^{-2}$ year $^{-1},$ and 1.3 $\pm$ 0.4 % at MY catchment, respectively
365	(Table 2). The uncertainties of $[NO_3^-atm]$ , $M_{atm}$ , and $M_{atm}/D_{atm}$ ratio in each catchment
366	were determined from the uncertainties of $\Delta^{17}O$ , $\Delta^{17}O_{atm}$ , $F_{stream}$ , and $D_{atm}$ according to
367	the equations of error propagation. The details were described in Appendix A.
368	
369	4 Discussion
370	4.1 Deposition rate of atmospheric nitrate at the study catchments
371	Based on the air monitoring data determined at the stations of Fukuoka (33°51'N,
372	130°50'E) and Miyazaki (31°83'N, 131°42'E) from 2011 to 2017, the Environmental
373	Laboratories Association of Japan (2017) reported $D_{atm}$ to be 57.8 mmol m <sup>-2</sup> year <sup>-1</sup> at
374	Fukuoka and 49.1 mmol $m^{-2}$ year <sup>-1</sup> at Miyazaki. Those values are consistent with the
375	$D_{atm}$ estimated in this study (69.3 and 40.1 mmol $m^{-2}\ year^{-1}$ at the FK and MY
376	catchments, respectively), within a difference of approximately 20 %. Thus, we
377	concluded that the $D_{\text{atm}}$ estimated in this study was reliable within the error margin of
378	20 % (Table 2). Because the $D_{\text{atm}}$ determined at the FK catchments was the highest
379	among the forested catchments in Table 3, we further compared the $D_{\text{atm}}$ of the FK

studies, along with that of the MY catchment (Table S1). While the Datm of the MY 381 catchment corresponded to the average level among the sites compiled in Table S1, the 382 Datm of the FK catchments exceeded the average level significantly. In addition, the Datm 383 of the FK catchments corresponded to one of the highest among the Japanese forested 384 areas (Table S1). All the catchments in Japan can be suffered from the long-range 385 transport of air pollutants derived from megacities in East Asian region (Chiwa, 2021; 386 Chiwa et al., 2012 and 2013). In addition, the shorter transport distance from the 387 Fukuoka metropolitan area (total population: 1.62 million people; population density: 388 4715 people/km<sup>2</sup>) may be mainly responsible for the  $D_{atm}$  higher in FK than in MY, 389 390 because the FK catchments are only 15 km west of the Fukuoka metropolitan area.

391

#### 392 4.2 Excess leaching of unprocessed atmospheric nitrate from FK catchments

The isotopic compositions ( $\delta^{15}$ N,  $\delta^{18}$ O, and  $\Delta^{17}$ O) of stream nitrate eluted from the 393 FK and MY catchments were typical for those eluted from forested catchments (Hattori 394 et al., 2019; Huang et al., 2020; Nakagawa et al., 2013, 2018; Riha et al., 2014; Sabo et 395 396 al., 2016; Tsunogai et al., 2014, 2016). The striking features found in the FK catchments 397 were that, in addition to the high  $[NO_3^-]$  and high  $M_{total}$  that had been clarified in a past study (Chiwa, 2021), both [NO<sub>3<sup>-</sup>atm</sub>] and M<sub>atm</sub> in FK were higher than those eluted from 398 MY (Table 2). Especially, the average  $[NO_3^{-}_{atm}]$  in the stream eluted from the FK1 399 catchment was the highest ever reported in forested streams determined through 400 continuous monitoring for more than 1 year (Bostic et al., 2021; Bourgeois et al., 2018b, 401

402 2018a; Hattori et al., 2019; Huang et al., 2020; Nakagawa et al., 2018; Rose et al., 2015;

403 Sabo et al., 2016; Tsunogai et al., 2014, 2016).

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

Elevated loading of nitrogen through atmospheric deposition was responsible for the 411 occurrence of nitrogen saturation in forest ecosystems, from which elevated levels of 412 nitrate are exported (Aber et al., 1989). Nakagawa et al. (2018) proposed that the 413 M<sub>atm</sub>/D<sub>atm</sub> ratio can be an index for evaluating the nitrogen saturation in each forested 414 415 catchment, because the Matm/Datm ratio directly reflects the present demand for atmospheric nitrate deposited in each forested catchment, and thus reflects the nitrogen 416 saturation in each forested catchment. The high Matm/Datm ratios observed in the FK 417 418 catchments implied that the demand for atmospheric nitrate was low in the FK catchments and that the stages of nitrogen saturation at the FK catchments were higher 419 than those at other forested catchments. That is, the nitrogen saturation at the FK 420 catchments was responsible for the observed high [NO<sub>3</sub><sup>-</sup>] and high M<sub>total</sub> at the FK 421 catchments than at MY and any other catchment ever studied (Table 3). 422

423	The stand age of forests can affect the retention or loss of N (Fukushima et al., 2011;
424	Ohrui and Mitchell, 1997). Fukushima et al. (2011) evaluated N uptake rates of
425	Japanese cedars at different ages (5-89 years old) and demonstrated that the N uptake
426	rates of Japanese cedars were higher in younger stands (53 kg N ha <sup>-1</sup> year <sup>-1</sup> in 16 years
427	old) than in older stands (29 kg N ha <sup>-1</sup> year <sup>-1</sup> in 31 years old; 24 kg N ha <sup>-1</sup> year <sup>-1</sup> in 42
428	years old; 34 kg N ha <sup>-1</sup> year <sup>-1</sup> in 89 years old). In addition, Yang and Chiwa (2021)
429	found that the nitrate concentration in the soil water taken beneath the rooting zone of
430	matured artificial Japanese cedar plantations (607 $\pm$ 59 $\mu M;$ 64-69 years old) was
431	significantly higher than that of normal Japanese oak plantations (8.7 $\pm$ 8.1 $\mu M;$ 24
432	years old). Moreover, by adding ammonium nitrate (50 kg N ha <sup>-1</sup> year <sup>-1</sup> ) to the forest
433	floor directly, Yang and Chiwa (2021) found that the nitrate concentration in the soil
434	water of the matured artificial Japanese cedar plantations increased significantly faster
435	than that of the normal Japanese oak plantations, probably because of the lower N
436	uptake rates in the matured artificial Japanese cedar plantations. Because most of the
437	artificial Japanese cedar/cypress plantations in the FK and MY catchments have reached
438	their maturity (> 50 years; Yang and Chiwa, 2021), the higher proportion of matured
439	artificial Japanese cedar/cypress plantations in the FK1 catchment (Table 1) was highly
440	responsible for the observed elevated leaching of nitrate, caused by the reduction in N
441	uptake rates.

442 As a result, we concluded that the FK forested catchments were under the high 443 nitrogen saturation stage, FK1 catchment especially, and the nitrogen saturation in the FK1 catchment was responsible for the elevated M<sub>total</sub>, M<sub>atm</sub>, [NO<sub>3</sub><sup>-</sup>], [NO<sub>3</sub><sup>-</sup><sub>atm</sub>] found
in the stream eluted from the catchment (Figs. 3a, 3b, 3c and 3d).

446

447 4.3 The  $M_{atm}/D_{atm}$  ratio as an index of nitrogen saturation

Past studies have used the concentration of stream nitrate as one of the important indexes to evaluate the stage of nitrogen saturation in each forest (Aber, 1992; Huang et al., 2020; Rose et al., 2015; Stoddard, 1994). The strong linear relationship ( $R^2 = 0.76$ ; P < 0.0001) between the stream nitrate concentration and the M<sub>atm</sub>/D<sub>atm</sub> ratio, except for the Qingyuan forested catchment (Fig. 3d), further supported that the M<sub>atm</sub>/D<sub>atm</sub> ratio can be used as an alternative index of nitrogen saturation, as pointed out in Nakagawa et al. (2018).

455 The differences in the number of storm and/or snowmelt events could affect the  $M_{atm}/D_{atm}$  ratio as well, because NO<sub>3<sup>-</sup>atm</sub> could be injected into the stream water directly, 456 along with the storm/snowmelt water (Tsunogai et al., 2014; Ding et al., 2022; Inamdar 457 and Mitchell, 2006). In recent study, however, we found that storm events have little 458 459 impact on the  $M_{atm}/D_{atm}$  ratio, based on monitoring temporal variation of  $[NO_3^-]_{atm}$  in 460 stream water during storm events (Ding et al., 2022). In addition, the low M<sub>atm</sub>/D<sub>atm</sub> ratio found in Uryu forested catchment (0.7 %; Table 3) implied that the snowmelt has 461 little impact on the Matm/Datm ratio as well, because 30% of the annual mean 462 precipitation was snow in Uryu forested catchment (Tsunogai et al., 2014). 463 The differences in the amount of precipitation, temperature, and the flux of stream 464

465 water could affect the  $M_{atm}/D_{atm}$  ratio as well. As a result, the annual amount of

precipitation, mean temperature, and the annual mean flux of stream water (F<sub>stream</sub>) in 466 467 the forested catchments were compiled in Table S2. While the stream nitrate concentration showed a strong linear relationship ( $R^2 = 0.76$ ; P < 0.0001) with the 468 469  $M_{atm}/D_{atm}$  ratio (Fig. 3d), the precipitation, temperature, and  $F_{stream}$  did not show a significant relationship with the  $M_{atm}/D_{atm}$  ratio (P > 0.14; Fig. 4). As a result, we 470 471 concluded that the Matm/Datm ratio was mainly controlled by the progress of nitrogen saturation, rather than the differences in the number of storm and/or snowmelt events, 472 473 the amount of precipitation, temperature, and the flux of stream water.

The differences in the residence time of water in each catchment could also impact 474 the Matm/Datm ratio, as the residence time of water in forested catchments ranges from 475 476 one month to more than one year (Asano et al., 2002; Farrick and Branfireun, 2015; Kabeya et al., 2008; Rodgers et al., 2005; Soulsby et al., 2006; Tetzlaff et al., 2007). It 477 is difficult to explain high [NO<sub>3</sub><sup>-</sup>] and high M<sub>total</sub> eluted from the catchment by the 478 479 residence time of water alone, while the Matm/Datm ratio could be higher in catchments with a shorter water residence time, as the majority of nitrate eluted from the catchment 480 with a high M<sub>atm</sub>/D<sub>atm</sub> ratio was NO<sub>3</sub><sup>-</sup><sub>re</sub> produced by microbial nitrification. The 481 significant correlation between  $M_{total}$  and  $M_{atm}/D_{atm}$  ratios (P < 0.0001; Fig. 3a) 482 supported nitrogen saturation as the leading cause of high M<sub>total</sub> in catchments with a 483 high Matm/Datm ratio. Additionally, the high loading of atmospheric nitrogen, the type of 484 plantation, and the old age of plantation in the FK1 catchment all supported the 485 conclusion that the FK1 catchment was under nitrogen saturation. 486

487 The Matm/Datm ratio is a more reliable and robust index than the stream nitrate

488 concentration, as explained below. The Qingyuan forested catchment can be classified into the highest nitrogen saturation stage based only on the highest stream nitrate 489 concentration of 150  $\mu$ M (Table 3). However, based on the leaching flux of nitrogen via 490 stream water monitored by Huang et al. (2020) for 4 years in the Qingyuan forested 491 catchment, along with the deposition flux of nitrogen, we can obtain the Matm/Datm ratio 492 in the catchment to be a medium level of  $5.8 \pm 1.3$  %, implying that the nitrogen 493 saturation stage was not so high (Table 3). Huang et al. (2020) also concluded that the 494 input of nitrogen exceeded the output in the catchment, and thus, the catchment was at 495 stage 2 of nitrogen saturation. The M<sub>atm</sub>/D<sub>atm</sub> ratio in the Qingyuan forested catchment 496 with a medium level among all forested catchments (Fig. 3d) should be a more reliable 497 index of nitrogen saturation. 498 Compared with those in the other forested catchments in Table 3, the annual amount 499 500 of precipitation (P) has the lowest value of 709 mm in the Qingyuan forested catchment. The flux of stream water (F<sub>stream</sub>) has the lowest value of 309 mm as well. Thus, we 501 concluded that nitrate was relatively concentrated in the catchment because of the small 502

503 precipitation, resulting in relative enrichment in the concentrations of both nitrate (150

504  $\mu$ M) and unprocessed atmospheric nitrate (8.9  $\mu$ M) in the stream.

505 While the concentration of stream nitrate, as an index of nitrogen saturation 506 traditionally, can be influenced by the amount of precipitation, as demonstrated in the 507 Qingyuan forested catchment, the  $M_{atm}/D_{atm}$  ratio is independent of the amount of 508 precipitation (Fig. 4). Therefore, the  $M_{atm}/D_{atm}$  ratio can be used as a more robust index 509 for evaluating nitrogen saturation in each forested catchment.

510

## 511 **5 Conclusions**

Both the concentrations and  $\Delta^{17}$ O of stream nitrate were determined for more than 2 512 years in the forested catchments of FK (FK1 and FK2) and MY to determine the 513 Matm/Datm ratio for each catchment. The FK catchments exhibited higher Matm/Datm ratio 514 than the MY catchment and other forested catchments reported in past studies, implying 515 that the progress of nitrogen saturation in the FK catchments was severe. Both age and 516 proportion of artificial plantation in the FK catchments were responsible for the 517 progress of nitrogen saturation. In addition, although past studies have commonly used 518 the concentration of stream nitrate as an index to evaluate the progress of nitrogen 519 saturation in forested catchments, it can be influenced by the amount of precipitation. 520 521 As a result, we concluded that the Matm/Datm ratio should be used as a more reliable index for evaluating the progress of nitrogen saturation because the Matm/Datm ratio is 522 independent from the amount of precipitation. 523

524

Appendix A: Calculating of uncertainties in the values of [NO<sub>3<sup>-</sup>atm</sub>], M<sub>atm</sub>, and M<sub>atm</sub>/D<sub>atm</sub>
ratio

527 The uncertainty in the values of  $[NO_3^-a_{tm}]$  was estimated from the uncertainties in 528 the  $\Delta^{17}O$  values of stream nitrate ( $\Delta^{17}O$ ) and  $NO_3^-a_{tm}$  ( $\Delta^{17}O_{atm}$ ) according to the 529 divisive equation of error propagation (A1):

530 
$$\sigma_{[NO_{3}]_{atm}} = [NO_{3}] * \sqrt{(\frac{1}{\Delta^{17}O_{atm}} * \sigma_{\Delta^{17}O})^{2} + (\frac{\Delta^{17}O}{\Delta^{17}O_{atm}} * \sigma_{\Delta^{17}O_{atm}})^{2}}$$
(A1)

where  $\sigma_{[NO_3^-am]}$ ,  $\sigma_{\Delta^{17}O}$ , and  $\sigma_{\Delta^{17}O_{atm}}$  denote the uncertainties in  $[NO_3^-atm]$ ,  $\Delta^{17}O$  values of stream nitrate, and  $\Delta^{17}O$  values of  $NO_3^-atm$ , respectively. The standard error of the mean (SE) of ±0.1 ‰ and the areal/seasonal variations of ±3 ‰ was used in calculating  $\sigma_{\Delta^{17}O}$  and  $\sigma_{\Delta^{17}O_{atm}}$ , respectively. As a result, the uncertainty in  $[NO_3^-atm]$ ( $\sigma_{[NO_3^-atm]}$ ) was ±1.30, ±0.67, and ±0.03 µM at FK1, FK2, and MY catchments, respectively. The uncertainty in the values of M<sub>atm</sub> was estimated from the uncertainties in

538  $[NO_3_{atm}]$  and in F<sub>stream</sub> according to the multiplicative equation of error propagation 539 (A2):

540 
$$\sigma_{M_{atm}} = \sqrt{(F_{stream} * \sigma_{[NO_3]_{atm}})^2 + ([NO_3]_{atm}] * \sigma_{F_{stream}})^2}$$
 (A2)

541 where  $\sigma_{M_{atm}}$ ,  $\sigma_{[NO_{3}]}$ , and  $\sigma_{F_{stream}}$  denote the uncertainties in  $M_{atm}$ ,  $[NO_{3}]$  and

542 F<sub>stream</sub>, respectively. Komatsu et al. (2008) proposed the uncertainty in F<sub>stream</sub> to be

 $\pm 162.3$  mm when using the water balance method in estimating F<sub>stream</sub>. Here, the

544 uncertainty in  $M_{atm}$  ( $\sigma_{M_{atm}}$ ) was  $\pm 2.1, \pm 1.0$ , and  $\pm 0.1 \text{ mmol m}^{-2} \text{ yr}^{-1}$  at FK1, FK2, and 545 MY catchments, respectively.

546 The uncertainty in  $M_{atm}/D_{atm}$  ratio was estimated from the uncertainties in  $M_{atm}$  and 547 in  $D_{atm}$  according to the divisive equation of error propagation (A3):

548 
$$\sigma_{M_{atm}/D_{atm} ratio} = \sqrt{\left(\frac{1}{D_{atm}} * \sigma_{M_{atm}}\right)^2 + \left(\frac{M_{atm}}{D_{atm}^2} * \sigma_{D_{atm}}\right)^2}$$
 (A3)

where  $\sigma_{M_{atm}/D_{atm} ratio}$ ,  $\sigma_{M_{atm}}$ , and  $\sigma_{D_{atm}}$  denote the uncertainty in  $M_{atm}/D_{atm}$  ratio, M<sub>atm</sub>, and D<sub>atm</sub>, respectively. Comparing the deposition rate of NO<sub>3</sub><sup>--</sup><sub>atm</sub> obtained at the other atmospheric monitoring stations nearby, the uncertainty of 20 % was adopted for those of D<sub>atm</sub> in each catchment, which corresponds to the uncertainty in D<sub>atm</sub> of ±13.9, ±13.9, ±8.0 mmol m<sup>-2</sup> yr<sup>-1</sup> at FK1, FK2, and MY catchments, respectively. As

554	a result, the uncertainty in $M_{atm}/D_{atm}$ ratio was ±4.1 %, ±2.0 %, and ±0.4 % at FK1,
555	FK2, and MY catchments, respectively.

557 *Data availability.* All the primary data are presented in the Supplement. The other data 558 are available upon request to the corresponding author (Weitian Ding).

559

560	Author contributions.	UT, FN,	KS, and	MC designed	l the study. MC	and TK performed
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the field observations. WD, UT, and FN determined the concentrations and isotopic

562 compositions of the samples. WD, TS, FN, and UT performed data analysis, and WD

and UT wrote the paper with input from MC, TK, and KS.

564

565 *Competing interests.* The authors declare that they have no conflict of interest.

566

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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

786 **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  $([NO_3^-_{atm}]_{avg})$ , 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	90.9	7.3
$[NO_{3}^{-}_{atm}]_{avg} (\mu M)$	$10.80\pm1.30$	$5.06\pm0.67$	$0.16\pm0.03$
$M_{total} \ (mmol \ m^{-2} \ yr^{-1})$	$98.8 \pm 17.8$	$82.0\pm14.8$	$23.7\pm1.2$
$M_{atm} \ (mmol \ m^{-2} \ yr^{-1})$	$9.7\pm2.1$	$4.6\pm1.0$	$0.5\pm0.1$
$D_{atm} \ (mmol \ m^{-2} \ yr^{-1})$	$69.3\pm13.9$	$69.3 \pm 13.9$	$40.1\pm\!\!8.0$
$M_{atm}/D_{atm}$ (%)	$14.1\pm4.1$	$6.6\pm2.0$	$1.3\pm0.4$

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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

	Р	[NO <sub>3</sub> <sup>-</sup> ] <sub>avg</sub>	N stage*	[NO3 <sup>-</sup> atm]avg	M <sub>atm</sub>	M <sub>total</sub>	D <sub>atm</sub>	$M_{atm}/D_{atm}$
	mm	μΜ	IN stage	μΜ	mn	nol m <sup>-2</sup> y	$r^{-1}$	%
FK1 <sup>a</sup>	1777	109.5	-	10.8	9.7	98.8	69.3	14.1
FK2 <sup>a</sup>	1777	90.9	-	5.06	4.6	82.0	69.3	6.6
MY <sup>a</sup>	3981	7.3	-	0.2	0.5	23.7	40.1	1.3
KJ <sup>b</sup>	2500	58.4	-	3.3	4.3	76.4	45.6	9.4
IJ1 <sup>b</sup>	3300	24.4	2	1.4	2.9	50.1	44.5	6.5
IJ2 <sup>b</sup>	3300	17.1	-	0.6	1.2	35.1	44.5	2.6
Fernow1 <sup>c</sup>	1450	17.9	1	1.6	0.8	9.3	23.4	3.6
Fernow2 <sup>c</sup>	1450	34.3	2	3.4	1.5	14.8	23.4	6.3
Fernow3 <sup>c</sup>	1450	60.0	3	4.2	2.4	34.5	23.4	10.3
Uryu <sup>d</sup>	1170	0.7	-	0.1	0.1	1.0	18.6	0.7
Qingyuan <sup>e</sup>	709	150.0	2	8.9	2.9	49.3	50.0	5.8

802 past studies using  $\Delta^{17}$ O of nitrate as a tracer.

a: This study

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

805 c: Rose et al., 2015

806 d: Tsunogai et al., 2014

807 e: Huang et al., 2020

- 808 \*: N saturation stage estimated in past studies
- 809 -: No data

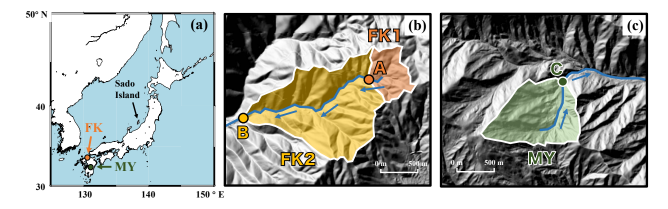


Figure 1. A map showing the locations of the study catchments (FK and MY) in Japan (a), and the maps of FK1, FK2 (b) and MY catchments (c), shown by orange, yellow, and green areas, respectively, together with the sampling station A, B, and C, respectively, shown by orange, yellow, and green circles, respectively. The blue arrows indicate the flow direction of stream water.

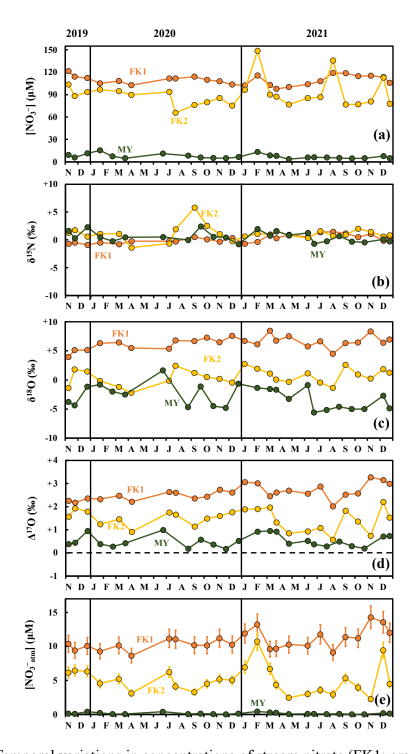
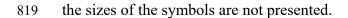
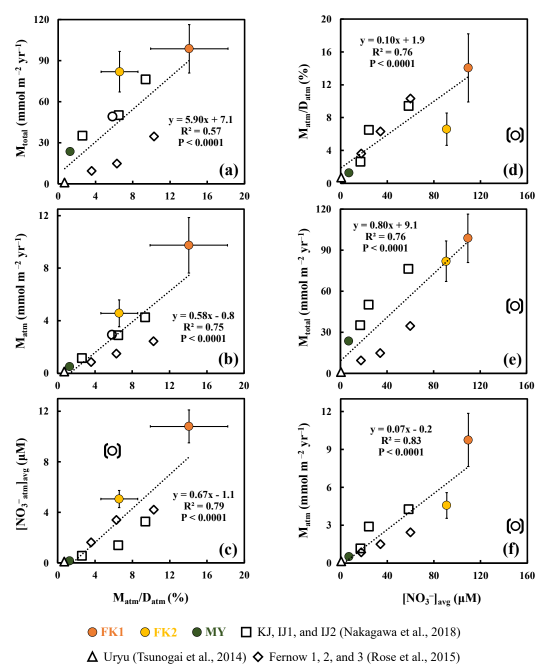


Figure 2. Temporal variations in concentrations of stream nitrate (FK1: orange circles;
FK2: yellow circles; MY: green circles) (a), together with those in δ<sup>15</sup>N (b), δ<sup>18</sup>O (c),

and  $\Delta^{17}O(d)$  of nitrate, and the concentration of unprocessed NO<sub>3<sup>-</sup>atm</sub> ([NO<sub>3<sup>-</sup>atm</sub>]) (e) in

818 the stream water of the FK1, FK2, and MY forested catchments. Error bars smaller than





**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<sub>atm</sub>/D<sub>atm</sub> 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  $NO_{3^-atm}$  ([ $NO_{3^-atm}$ ]<sub>avg</sub>) plotted as a function of the M<sub>atm</sub>/D<sub>atm</sub> ratio (c); the  $M_{atm}/D_{atm}$  ratio plotted as a function of the average concentration

825	of nitrate ([NO <sub>3</sub> <sup>-</sup> ] <sub>avg</sub> ) (d); the $M_{total}$ plotted as a function of [NO <sub>3</sub> <sup>-</sup> ] <sub>avg</sub> (e); the $M_{atm}$
826	plotted as a function of $[NO_3^-]_{avg}$ (f) (FK1: orange circles; FK2: yellow circles; MY:
827	green circles). Those determined for the forested catchments in past studies are plotted
828	as well (Qingyuan: white circle (Huang et al., 2020); KJ, IJ1, and IJ2: white squares
829	(Nakagawa et al., 2018); Fernow 1, 2, and 3: white diamonds (Lucy et al., 2015); Uryu:
830	white triangle (Tsunogai., 2014)). The data obtained in the Qingyuan forested
831	catchment are shown in parentheses and excluded from the calculation to estimate
832	correlation coefficients (see text for the reason).

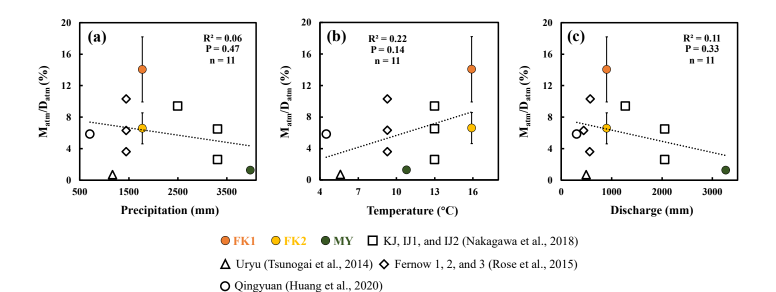


Figure 4. the  $M_{atm}/D_{atm}$  ratio plotted as a function of the amount of precipitation (a), the  $M_{atm}/D_{atm}$  ratio plotted as a function of the temperature (b), and the  $M_{atm}/D_{atm}$  ratio plotted as a function of flux of stream water (c) (FK1: orange circles; FK2: yellow circles; MY: green circles). Those determined for the forested catchments in past studies

are plotted as well.

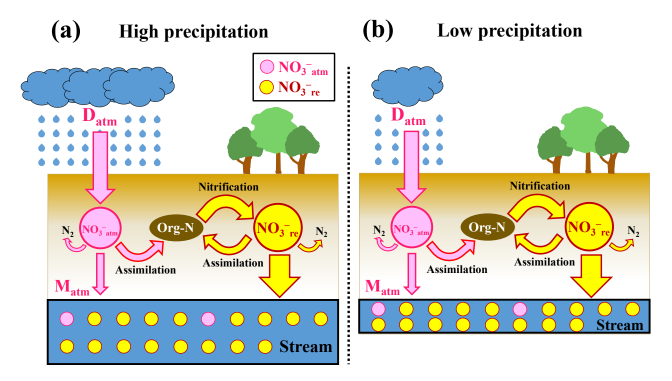


Figure 5. Schematic diagram showing the biogeochemical processing of nitrate in 839 840 forested catchments under high precipitation (a) and low precipitation (b), where NO<sub>3<sup>-</sup>atm</sub> (unprocessed atmospheric nitrate) is represented by pink circles, NO<sub>3<sup>-</sup>re</sub> by 841 yellow circles, the flows of  $NO_{3-atm}^{-}$  by pink arrows, and those of  $NO_{3-re}^{-}$  (remineralized 842 843 nitrate) by yellow arrows (modified after Nakagawa., 2018). Although the deposition rates of  $NO_{3-atm}^{-}$  (D<sub>atm</sub>) and the biogeochemical reaction rates between (a) and (b) are 844 the same, we can expect high  $[NO_3^-]$  in (b). On the other hand, the  $M_{atm}/D_{atm}$  ratio 845 between (a) and (b) are the same. 846