Bias in calculating gross nitrification rates in forested catchments using the triple oxygen isotopic composition ($\Delta^{17}O$) of stream nitrate

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Abstract

A novel method has been proposed and applied in recent studies to quantify gross nitrification rate (GNR) in forested catchments using the triple oxygen isotopic composition (Δ^{17} O) of stream nitrate. However, the equations used in these calculations assume that the Δ^{17} O value of nitrate consumed through assimilation or denitrification in forest soils is equal to the Δ^{17} O value of stream nitrate. The GNR estimated from the Δ^{17} O value of stream nitrate was significantly higher than the GNRs in our simulated calculations for a forested catchment where the soil nitrate had Δ^{17} O values higher than those the stream nitrate. Given that most reported soil nitrate in forested catchments showed Δ^{17} O values higher than those of the stream nitrate, we concluded that the GNR estimated from the $\Delta^{17}O$ value of stream nitrate was, to an extent, an overestimate of the actual GNR. 1 Introduction Nitrate (NO₃⁻) is an important nitrogen nutrient for primary production in soils.

Nitrification is the microbial process that produces NO₃⁻ in forested ecosystems.

Thus, quantifying the nitrification rate can assist in the evaluation of the present and future states of forested ecosystems. The net nitrification rate can be estimated from an increase in NO₃⁻ concentration during a certain period. However, the gross nitrification rate (GNR), which includes the net nitrification rate plus the consumption rate of NO₃⁻ (e.g., through plant assimilation or denitrification), reflects the internal N cycling better than the net nitrification rate (Bengtsson et al., 2003), especially in

- forested ecosystems. Although the net nitrification rate is often negligible (Stark and
- 24 Hart, 1997), the consumption rate is significant in forested ecosystems, such that the
- 25 GNR often exceeds the net nitrification rate by several orders of magnitude (Verchot
- 26 et al., 2001).
- 27 Recent studies have successfully estimated the GNR in aquatic environments, such
- as lakes, using the Δ^{17} O values of NO₃⁻ as a conservative tracer to determine the
- 29 mixing ratio between atmospheric nitrate (NO₃ atm) and biologically produced nitrate
- 30 (NO₃-bio) (Tsunogai et al., 2011, 2018). The NO₃-atm is deposited in the water
- environment, while NO₃ bio is produced through nitrification. The NO₃ bio always
- shows the Δ^{17} O value close to 0 ‰ because its oxygen atoms are derived from either
- terrestrial O₂ or H₂O through nitrification. Contrarily, the NO₃⁻ atm always displays an
- anomalous enrichment in ¹⁷O with Δ^{17} O value being approximately +26 ± 3 % in
- 35 Japan (Tsunogai et al., 2010, 2016; Ding et al., 2022, 2023) because of oxygen
- transfers from atmospheric ozone (Michalski et al., 2003; Nelson et al., 2018).
- Additionally, Δ^{17} O is almost stable during "mass-dependent" isotope fractionation
- processes (Michalski et al., 2004; Tsunogai et al., 2016). This is because possible
- variations in the δ^{17} O and δ^{18} O values during the processes of biogeochemical isotope
- fractionation follow the relation of $\delta^{17}O \approx 0.5 \, \delta^{18}O$, which cancels out the variations
- in the Δ^{17} O value. Thus, regardless of the partial consumption through denitrification
- or assimilation after deposition in a water column, the Δ^{17} O can be used as a

- conservative tracer of NO₃ atm to calculate the mixing ratio of NO₃ atm to total NO₃
- $(NO_3^- atm/NO_3^- total)$ in a water column using the following equation:
- 45 $[NO_3^-]/[NO_3^-]/[NO_3^-] = [NO_3^-]/([NO_3^-]) + [NO_3^-] = \Delta^{17}O/\Delta^{17}O_{atm}$ (1)
- where the Δ^{17} O_{atm} and Δ^{17} O denote the Δ^{17} O values of NO₃⁻_{atm} and NO₃⁻ dissolved in
- 47 the water environment, respectively. Using the NO₃⁻atm/NO₃⁻total ratio estimated from
- the Δ^{17} O value of NO₃⁻ in a lake water column and the deposition rate of NO₃⁻ atm into
- 49 the lake, the GNR (i.e., production rate of NO₃-bio) can be successfully estimated. This
- approach works because the NO₃ atm/NO₃ total ratios are homogeneous in the water
- column due to the active vertical mixing; thus, we can constrain the NO₃⁻_{atm}/NO₃⁻_{total}
- ratios of NO₃⁻ consumed in the lake water column (Tsunogai et al., 2011, 2018).
- In addition to applications in water environments, the Δ^{17} O method has been
- applied to forested catchments to determine GNR (Fang et al., 2015; Hattori et al.,
- 55 2019; Huang et al., 2020). Using the deposition flux of NO₃ atm into the catchment
- and the leaching flux of unprocessed NO₃ atm and NO₃ bio via streams, the GNR in a
- 57 forested catchment was estimated similarly to the estimation for water environments
- (Fang et al., 2015). However, unlike in water environments, where the
- 59 NO_{3 atm}/NO_{3 total} ratio of nitrate consumed in the water column can be easily
- measured, it is often difficult to determine the NO₃⁻atm/NO₃⁻total ratio of NO₃⁻
- consumed in soil layers. Consequently, past studies have approximated these values as
- equal to those of stream NO₃⁻ leached from forested catchments without actual
- observation (Fang et al., 2015, Hattori et al., 2019, Huang et al., 2020). Such an

approximation should be used with extreme caution, as the NO₃ atm/NO₃ total ratio 64 $(\Delta^{17}\text{O values})$ of soil NO₃⁻ are not always equal to those of stream NO₃⁻ (Hattori et 65 al., 2019, Rose, 2014, Nakagawa et al., 2018). To clarify the details of the 66 approximation and its impact on the final estimated GNR, we present an accurate 67 relationship between the Δ^{17} O of soil NO₃⁻ and the GNR, using basic isotope mass 68 balance equations. Thereafter, we present possible range of variation in the GNRs 69 estimated for a forested catchment, using parameters such as Δ^{17} O values of stream 70 NO₃⁻ reported in a past study. Finally, we compared the GNRs estimated in this study 71 with those obtained from the Δ^{17} O values of stream NO₃⁻. 72

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2 Calculation

The total mass balance equation of NO₃⁻ including the GNR in catchments can be expressed as follows:

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$$NO_3^-$$
 deposition + $GNR = NO_3^-$ leaching + NO_3^- uptake + GDR (2)

- where NO₃⁻deposition, GNR, NO₃⁻leaching, NO₃⁻uptake, and GDR denote the deposition flux of NO₃⁻ into the catchment, GNR in the catchment, leaching flux of NO₃⁻ from the catchment, uptake rate of NO₃⁻ in the catchment, and gross denitrification rate in the catchment, respectively.
- The isotope mass balance for each Δ^{17} O value of NO₃⁻ in the catchment can be expressed using a similar equation:

- 84 $NO_3^-_{deposition} \times \Delta^{17}O(NO_3^-)_{atm} + GNR \times \Delta^{17}O(NO_3^-)_{nitrification} = NO_3^-_{leaching} \times \Delta^{17}O(NO_3^-)_{nitrification}$
- 85 3^{-})stream + NO_3^{-} uptake $\times \Delta^{17}O(NO_3^{-})$ uptake + $GDR \times \Delta^{17}O(NO_3^{-})$ denitrification (3)
- where $\Delta^{17}O(NO_3^-)$ atm, $\Delta^{17}O(NO_3^-)$ nitrification, $\Delta^{17}O(NO_3^-)$ stream, $\Delta^{17}O(NO_3^-)$ uptake, and
- 87 $\Delta^{17}O(NO_3^-)_{denitrification}$ denote the $\Delta^{17}O$ value of NO_3^- _{atm} deposited into the catchment,
- that of the NO₃-bio produced through nitrification, that of the NO₃-leached from the
- catchment, that of the NO₃⁻ assimilated by plants and other organisms in the
- catchment, and that of the NO₃⁻ decomposed through denitrification in the catchment,
- 91 respectively.
- If the Δ^{17} O values of the NO₃⁻ in the forested soil layers, where the NO₃⁻ was
- consumed through assimilation or denitrification, are equal to the Δ^{17} O value of NO₃⁻
- in the stream, we could obtain Eq. 4:

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$$\Delta^{17}\text{O(NO}_3^-)_{\text{uptake}} = \Delta^{17}\text{O(NO}_3^-)_{\text{denitrification}} = \Delta^{17}\text{O(NO}_3^-)_{\text{stream}}$$
 (4)

- Consequently, by combining Eqs. 3 and 4, we could obtain Eq. 5:
- 97 NO_3^- deposition $\times \Delta^{17}O(NO_3^-)$ atm $+ GNR \times \Delta^{17}O(NO_3^-)$ nitrification $= (NO_3^-$ leaching $+ NO_3^-$ uptak

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$$e + GDR$$
) $\times \Delta^{17}O(NO_3^-)_{stream}$ (5)

- We could estimate the GNR using Eq. 6 obtained from Eqs. 2 and 5 because we can
- approximate the Δ^{17} O values of NO₃-bio produced through nitrification
- 101 $(\Delta^{17}O(NO_3^-)_{nitrification})$ to 0 (Michalski et al., 2003; Tsunogai et al., 2010):

$$GNR = NO_3^{-}_{\text{deposition}} \times (\Delta^{17}O(NO_3^{-})_{\text{atm}} - \Delta^{17}O(NO_3^{-})_{\text{stream}})/\Delta^{17}O(NO_3^{-})_{\text{stream}}$$
(6)

Eq. 6 corresponds to the equations used in previous studies to quantify the GNR in the forested catchments (Eq. 4 in Fang et al., 2015; Eq. 8 in Hattori et al., 2019; Eq. 4 in Huang et al., 2020).

3 Results and Discussion

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The Δ^{17} O values of NO₃⁻ in forested soil layers should be equal to those of stream NO₃⁻ in Eq. 6, as presented in Eq. 4 to obtain Eq. 6. While the number of simultaneous observations of the oxygen isotopes of NO₃ in soil and stream in a given forested catchment is limited (Hattori et al., 2019, Osaka et al., 2010, Rose, 2014, Nakagawa et al., 2018), the observations showed that the oxygen isotopic ratios of soil NO₃⁻ are often heterogeneous. In addition, the oxygen isotopic ratios of soil NO₃⁻ mostly exceeded those of stream NO₃⁻. Different from water environments, vertical mixing of water/soil is limited in forested soil, so the Δ^{17} O values of soil NO₃⁻ are often heterogeneous. For example, Hattori et al. (2019) found a decreasing Δ^{17} O trend in soil NO₃⁻ with depth, ranging from over +20 % at the surface to less than +3 % at depths of 25–90 cm. Additionally, more than 60 % of the soil samples exhibited Δ^{17} O values significantly higher than those of stream NO₃⁻ determined simultaneously ($\Delta^{17}O(NO_3^-)_{stream} = +1$ to +3 %). A similar trend in the vertical distribution was observed in the δ^{18} O values of NO₃⁻ in another forested catchment, from above +35 ‰ at the surface soil to less than +10 ‰ at depths of 30–50 cm from the soil surface (Osaka et al., 2010). In addition, most of the soil NO₃⁻ also exhibited δ^{18} O values higher than those of the stream NO₃⁻ (Osaka et al., 2010). Rose (2014)

monitored the horizontal distribution of the Δ^{17} O of soil NO₃⁻ by randomly setting 15 tension-free lysimeters at depths of 0–10 cm in a 39-ha forested catchment. They reported significantly higher Δ^{17} O values in soil NO₃⁻ (+9.1 ± 5.8 ‰ on average) than those of stream NO₃⁻ (+0.5 ‰ on average) leached from the forested catchment. As most fine roots and root biomass are concentrated in the top 10 cm of the soil in forested catchments (Jackson et al., 1996; Li et al., 2020), most assimilation (uptake reactions) of NO₃⁻ should occur in that top 10 cm of soil. Consequently, the significant difference in the Δ^{17} O values between soil NO₃⁻ and stream NO₃⁻, particularly in surface soil layers, implies that the estimated GNRs in forested catchments obtained from Eq. 6 were inaccurate. To demonstrate the impact of this approximation on GNR estimation, we simulated GNR for two different forest soils within the same catchment. In the first scenario, soil NO₃⁻ exhibited a Δ^{17} O value close to that of Δ^{17} O(NO₃⁻)_{atm} at the surface, which decreased to the Δ^{17} O of stream NO₃⁻ at depth (heterogeneous soil) (Figs. 1a and 1b). In the second scenario, soil NO₃⁻ had Δ^{17} O values equal to those of stream NO₃⁻ throughout the soil profile (homogeneous soil) (Figs. 2a and 2b). To simulate the forested catchment studied by Hattori et al. (2019), we used the same parameters values for the current calculation, including 7.0 kg N ha⁻¹ y⁻¹ for NO_{3}^{-} deposition, 2.6 kg N ha⁻¹ y⁻¹ for NO_{3}^{-} leaching, +28.0 % for $\Delta^{17}O(NO_{3}^{-})$ and +2.2 % for $\Delta^{17}O(NO_3^-)_{stream}$. All symbols (e.g., GNR) are consistent with those used by Hattori et al. (2019).

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To estimate GNR in each forest soil type, we divided the soils into 10 vertical layers (i.e., 10 steps). In the heterogeneous soil, the Δ^{17} O values of NO₃⁻ gradually decreased with depth, from +28.0% to +2.2%, at a rate of -2.58% per step (Fig. 1b). In the homogeneous soil, Δ^{17} O values of NO₃⁻ were constant at +2.2% across all layers (Fig. 2b). Note that the y-axes in the models were layers, not depths (Tables S1, S2, and S3). While the Δ^{17} O values of soil NO₃⁻ always showed decreasing trends with depths irrespective to the seasons, Δ^{17} O values of soil NO₃⁻ showed significant temporal variation at each depth (Hattori et al., 2019). This was the reason why the layers were adopted for the y-axes in our models, instead of depths. As a result, the specific depth of each layer varies over time. In addition, the relation between depth and layer is not always linear. The temporal variation found in the vertical distributions of Δ^{17} O values in the forested catchment (Hattori et al., 2019) can be explained by our model as well without contradiction because the Δ^{17} O values of soil NO₃⁻, while showing large temporal variation at each depth, always showed decreasing trend with depth throughout their observation (Hattori et al., 2019). To estimate GNR in each layer, both the Δ^{17} O value and the NO₃⁻ leaching flux in soil are required. While Hattori et al. (2019) reported soil NO₃⁻ concentrations for each layer, indicating little vertical variation within the forested catchment, they did not measure the catchment water flux. Consequently, it is difficult to constrain the NO₃⁻ leaching flux for each layer of forest soil. Nevertheless, NO₃⁻_{deposition} was 7.0 kg $N\ ha^{-1}\ y^{-1}$ and $NO_3^-_{leaching}$ was 2.6 kg N $ha^{-1}\ y^{-1}$ in the catchment (Hattori et al.,

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2019). Additionally, because water fluxes decrease gradually with depth in various 166 forest settings (e.g., Christiansen et al., 2006), we assumed a gradual decrease in 167 NO_3^- , leaching flux from 7.0 to 2.6 kg N ha⁻¹ y⁻¹ at a rate of -0.44 kg N ha⁻¹ y⁻¹ per 168 layer (Figs. 1c and 2c). Similar trends in the NO₃⁻ leaching flux of soil have been 169 observed in other forested catchments (Callesen et al., 1999; Inoue et al., 2021). 170 Applying the total mass balance and isotope mass balance equations (Eqs. 2 and 3) 171 to each layer, we estimated GNR (Figs. 1e and 2e) and the total consumption rate of 172 NO₃⁻ (GDR + uptake) (Figs. 1d and 2d) in each layer. In this calculation, we assumed 173 the following: (1) Δ^{17} O values of NO₃⁻ were constant in each layer; (2) vertical flow 174 of NO₃⁻ in soil layers proceeds downward from the surface to the final layer (No. 10); 175 and (3) GNR and the NO₃⁻ consumption rate (GDR + uptake) are 0 in layers beyond 176 the final layer. By summing the GNR determined for each layer, we estimated the 177 178 total GNR in the forested catchment. The total GNR estimated for the catchment with the homogeneous Δ^{17} O values in 179 soil NO_3^- (homogeneous soil) was 83.6 kg of N ha⁻¹ y⁻¹ (Fig. 2e), exactly equal to 180 that estimated by Hattori et al. (2019) using Eq. 6. This result allows us to further 181 verify that past studies estimating GNR using Eq. 6 implicitly approximated that Δ^{17} O 182 values of soil NO₃⁻ consumed in forested catchments were homogeneous and always 183 equal to those of stream NO₃⁻. However, the total GNR estimated for the catchment 184 with heterogeneous Δ^{17} O values in soil NO₃⁻ (heterogeneous soil) was considerably 185

lower (13.0 kg of N ha⁻¹ y⁻¹; Fig. 1e), while the same parameters were used for NO₃⁻deposition, NO₃⁻leaching, $\Delta^{17}O(NO_3^-)_{atm}$, and $\Delta^{17}O(NO_3^-)_{stream}$.

As we increased the number of layers in the forest soils to 20, 30, 50, 100, and 188 1000, the estimated GNR for the heterogeneous soil decreased to 11.4, 11.0, 10.5, 189 10.3, and 10.1 kg N ha⁻¹ y⁻¹, respectively. Moreover, when we changed the 190 191 calculation method from stepwise summation to integration, the estimated GNR was 11.2 kg N ha⁻¹ y⁻¹. Furthermore, even if we assumed non-linear variation for the 192 leaching flux of soil NO₃⁻, in which the leaching flux of soil NO₃⁻ increased with soil 193 depth from layers 1 to 5 with an increasing rate of 0.44 kg of N ha⁻¹ y⁻¹ layer⁻¹, while 194 the leaching flux decreased with soil depth from layers 6 to 10 with a decreasing rate 195 of 1.32 kg of N ha⁻¹ y⁻¹ layer⁻¹ (Table S3), the newly estimated total GNR (19.1 kg of 196 N ha⁻¹ y⁻¹) was still comparable with that estimated for the forested catchment with 197 the heterogeneous soil shown by Figure 1 (13.0 kg of N ha⁻¹ y⁻¹). As a result, we 198 concluded that the differences in the Δ^{17} O values of the soil NO₃⁻ consumed in a 199 forested catchment from that of stream NO₃⁻ resulted in a significant deviation in the 200 201 GNR estimated using Eq. 6 from the actual GNR. In addition, the most important parameter to determine GNR was the Δ^{17} O values of NO₃⁻ consumed in soil layers. 202 That is, the other parameters such as the number of layers and the vertical changes in 203 the leaching flux of soil NO₃⁻ had little impact on total GNR. 204

By combining the total mass balance and isotope mass balance shown in Eqs. 2 and 3, Eq. 7 was obtained to accurately estimate the total GNR:

 $GNR = NO_3^-_{leaching} - NO_3^-_{deposition} + (NO_3^-_{deposition} \times \Delta^{17}O(NO_3^-)_{atm} -$

 $NO_3^{-}_{leaching} \times \Delta^{17}O(NO_3^{-})_{stream} / \Delta^{17}O(NO_3^{-})_{soil}$ (7)

where $\Delta^{17}O(NO_3^-)_{soil}$ denotes the "average" $\Delta^{17}O$ of NO_3^- consumed through assimilation or denitrification in the forested catchment. Most of the soil NO_3^- measured to date exhibited $\Delta^{17}O$ values higher than those of stream NO_3^- leached from the catchments (Hattori et al., 2019, Rose, 2014). Consequently, the total GNR estimated from stream NO_3^- using Eq. 6 exceeded the total GNR estimated from soil NO_3^- using Eq. 7, to an extent. Therefore, the total GNR estimated from Eq. 6 was

If we can estimate the downward water flux at each soil layer, along with the NO_3^- concentration and $\Delta^{17}O$ value of NO_3^- in each soil layer using, e.g., a tension-free lysimeter (Inoue et al., 2021), we could estimate the vertical change in the NO_3^- leaching flux for each soil layer, along with the $\Delta^{17}O$ values of soil NO_3^- . Thereafter, applying Eq. 7 to each layer, we can more estimate the total GNR for the forested catchment accurately, by integrating the GNR estimated for each soil layer, together with the NO_3^- consumption rate in the forested catchment.

4 Conclusion

overestimated to an extent.

Past studies have proposed the $\Delta^{17}O$ method for determining the GNR in forested catchments. The equations used in the calculation implicitly assumed that the $\Delta^{17}O$ values of NO_3^- consumed in forested soils are homogeneous and equal to those of the

stream NO_3^- . However, the values are often heterogeneous and do not always equal those of the stream in forested soils. It is essential to clarify/verify the $\Delta^{17}O$ values of NO_3^- in forested soils and streams before applying the $\Delta^{17}O$ values of stream NO_3^- to estimate the total GNR.

Data availability. All data are presented in the Supplement.

Author contributions. WD, UT, and FN designed the study. WD and UT performed data analysis and wrote the paper.

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

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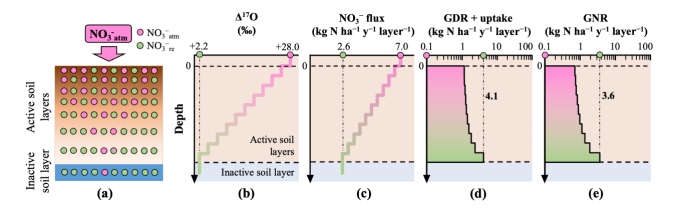


Figure. 1. Distribution of NO_3^- atm in the simulated forested soil with heterogeneous distribution of $\Delta^{17}O$ values of NO_3^- (a). Vertical distribution of the following parameters in the forested soil: assumed $\Delta^{17}O$ values of NO_3^- (b), assumed leaching flux of NO_3^- (c), estimated NO_3^- consumption rate (GDR + uptake) (d), and estimated GNR (e).

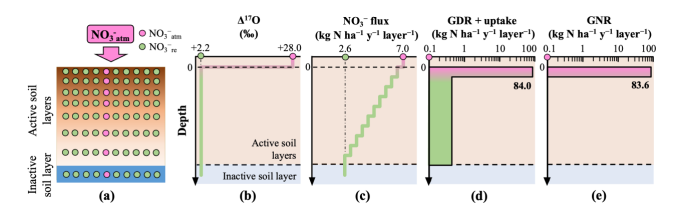


Figure. 2. Distribution of NO_3^- atm in the simulated forested soil with homogeneous distribution of $\Delta^{17}O$ values of NO_3^- (a). Vertical distribution of the following parameters in the forested soil: assumed $\Delta^{17}O$ values of NO_3^- (b), assumed leaching flux of NO_3^- (c), estimated NO_3^- consumption rate (GDR + uptake) (d), and estimated GNR (e).