Global patterns of nitrate isotope composition in rivers and adjacent aquifers reveal reactive nitrogen cascading

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Remediation of nitrate pollution of Earth’s rivers and aquifers is hampered by cumulative biogeochemical processes and nitrogen sources. Isotopes (δ15N, δ18O) help unravel spatio-temporal nitrogen(N)-cycling of aquatic nitrate (NO3−). We synthesized nitrate isotope data (n = ~5200) for global rivers and shallow aquifers for common patterns and processes. Rivers had lower median NO3− (0.3 ± 0.2 mg L−1, n = 2902) compared to aquifers (5.5 ± 5.1 mg L−1, n = 2291) and slightly lower δ15N values (+7.1 ± 3.8‰, n = 2902 vs +7.7 ± 4.5‰, n = 2291), but were indistinguishable in δ18O (+2.3 ± 6.2‰, n = 2790 vs +2.3 ± 5.4‰, n = 2235). The isotope composition of NO3− was correlated with water temperature revealing enhanced N-cascading in warmer climates. Seasonal analyses revealed higher δ15N and δ18O values in wintertime, suggesting waste-related N-source signals are better preserved in the cold seasons. Isotopic assays of nitrate biogeochemical transformations are key to understanding nitrate pollution and to inform beneficial agricultural and land management strategies.
the exponential growth of the human population con-
comitant with intensive development of fertilized agri-
cultural and industrial activity since the 1950s caused sharp 
increases in nitrogen loadings to rivers and surficial aquifers 
worldwide. Dissolved reactive nitrogenous species (e.g., NO3−, 
NH4+) are prevailing pollutants in many rivers and aquifers, 
stemming primarily from agricultural activities, municipal waste 
sources, and combustion derived nitrogen (N) deposition. The 
impact of reactive nitrogenous species on water and ecosystems 
(e.g., eutrophication, hypertrophication) and human health (e.g., 
methemoglobinemia, cancer, thyroid disease) occurs in series, 
known as N-cascading, and is of existential concern. The 
global nitrogen cycle is a subject of considerable debate 
and concern that anthropogenic loadings are driving it beyond 
Earth’s natural resilience boundaries. Reactive nitrogen cascades 
through aquatic ecosystems differentially, since some systems 
accumulate N, whereas others transform it through diverse bio-
geochemical N-cycling processes like nitrification, denitrification, 
N2-fixation, dissimilatory nitrate reduction (DNRA), ammonifi-
cation, and biological assimilation at rates dependent on the 
environmental conditions. These complexities make it difficult to 
unravel what point-based NO3− concentrations in rivers or their 
connected aquifers embody at any point in time, apart from 
regulatory pollutant exceedances, and quantitative knowledge of 
the roles of N-cycling processes in the terrestrial aquatic envi-
ronments remains deficient.

Stable isotopes of NO3− (δ15N, δ18O) are used, particularly in 
the last decades, to help identify the sources of high-N-pollution 
in aquatic systems. Contemporary preparative and isotope 
techniques enable fast low-cost isotopic analysis of 15N/14N and 
18O/16O ratios to ppb concentrations to incorporate comparative 
isotopic information from pristine aquatic environments and 
references therein. Distinctive δ15N and δ18O biplot clusters 
were initially proposed to “assign” organic and inorganic NO3− 
origins, however, NO3− sources rarely exhibit unique combinations of nitrogen (N) and oxygen (O) isotopic values as 
shown by ambiguous or widely overlapping clusters (e.g., soil, 
manure/sewage, NH4+ fertilizer), which make straightforward 
assignments problematic. Accordingly, others recommend 
including additional isotopes (e.g., δ13C), other chemicals (e.g., 
pharmaceuticals, food additives), and/or biological markers (e.g., 
chlorophyll-a, fecal coliforms) to better differentiate sources of 
nitrogen pollution in freshwaters, particularly as related to 
local sewage, animal waste, industrial and atmospheric sources.

One common misperception in the interpretation of NO3− 
isotopes in aquatic systems is that its isotopic composition is a 
conservative tracer of N source(s), despite isotope fractiona-
tions during numerous biogeochemical transformations that 
significantly alter the δ15N and δ18O values of nitrate. Instead, 
the δ15N and δ18O of NO3− in aquatic systems are more 
frequently a mixture of time and seasonally dependent N-source 
(s) and cumulative isotopic fractionations that occur during 
transport and biogeochemical transformation of dissolved N-
species. It is well-known that biological nitrate uptake (assimilation) and denitrification processes preferentially process 
the light isotopes, resulting in heavy isotope enrichment of 
residual nitrate. Despite uncertainties in the assessment of 
sources of nitrate pollution using δ15N and δ18O, these stable 
isotopes retain fundamental source and process information to 
help decipher the complexity of N-biogeochemistry in aquatic 
environments.

Common patterns of N-cycling processes using global spatial 
data sets of NO3− and its stable isotopes are established for soils 
and plant matter, but larger-scale spatiotemporal studies of 
nitrate in rivers and adjacent aquifers are lacking. Here we 
synthesized global river and adjacent shallow aquifer nitrate 
iso-14, 15O of NO3−) data sets from the scientific litera-
ture as well as new data (n = 5200) to (a) provide a first-
order global assessment of spatiotemporal patterns of NO3− and 
its isotopes in rivers and aquifers and (b) evaluate whether NO3− 
transformations are impacted by key environmental factors, such 
as temperature, climate, and seasonality. The investigation of the 
origin of nitrate pollution in rivers and groundwater at the local 
scale usually requires a deeper consideration of additional 
information such as N flux information from homogenized land-
use types; however, the detailed local evaluation was beyond the 
scope of this work. This synthesis provides a global foundational 
perspective to evaluate the potential and limitations of nitrate 
stable isotopes to track nitrogen pollution sources in aquatic 
systems, especially rivers, and to promote a data-based framework 
for further improving our understanding of the transformation 
mechanisms of nitrogen for sustainable management and reme-
diation of N-contaminated waters.

Results

Overall assessment of global river and groundwater nitrate 
data sets. Concentrations of nitrate in rivers and adjacent aquifers 
deviated from normal distributions and were highly skewed 
(skewness index > 2), hence median values were retained. Of all 
samples, rivers had substantially lower median NO3− concen-
trations (0.3 ± 0.2 mg L−1, n = 2902) compared to ground-
water (5.5 ± 5.1 mg L−1, n = 2291) (Supplementary Table 1). 
Around 2% of river water samples exceeded the WHO (World 
Health Organization) threshold of 50 mg L−1 (as NO3−), whereas 
in groundwater the exceedance was far higher (~34%). The 
Kruskal–Wallis test indicated that nitrate concentration differ-
ences between rivers and groundwater subsets were significant 
(p-value < 0.05) (Supplementary Table 2).

The distribution patterns for δ15N and δ18O of NO3− in 
rivers and groundwater deviated from normal distributions (Supple-
mentary Fig. 1). The isotope data were not as skewed as NO3− 
concentrations, whereas average values were retained to compare 
these isotope variables among them. Outliers were mostly extreme values 
falling within the upper and lower tails of the distributions of 
the Quantile–Quantile (Q–Q) plots (Supplementary Fig. 1). The 
δ15N values of nitrate in rivers were slightly lower (+7.1 ± 3.8‰, n = 2902) than aquifers (+7.7 ± 4.5‰, n = 2291, p-value < 0.05). 
Kendall presented an early compilation of nitrate isotope data in 
aquatic systems and reported groundwater nitrate has considerably higher δ15N values than rivers, however, others found 
more positive δ15N nitrate values in surface waters than nearby 
groundwater. Other case studies affirmed higher mean δ15N 
values for NO3− in shallow groundwater compared to local rivers 
in urban waters (Manila, Bangkok, and Jakarta) or under 
different land-use-based catchments. Overall, the δ18O values of 
nitrate in rivers and groundwater globally were indistinguish-
able from each other (+2.3 ± 6.2‰, n = 2790, vs +2.3 ± 5.4‰, n = 2235, p-value > 0.05, respectively) (Supplementary Tables 1 and 2).

Isotope variation with latitude. The global data set included river 
and groundwater samples spanning from 67°N to 38°S LAT and 
from 145°E to 123°W LON (Fig. 1). There were more samples 
between 30°N to 60°N than other latitudinal ranges, reflecting the 
preponderance of data from North America, Europe, and East 
Asia. However, Central Asia, Central, and South America, Oce-
ania, and Africa are poorly represented for nitrate isotope data 
due to a lack of water N pollution studies in those areas. The δ15N 
and δ18O of nitrate were negatively correlated with latitude for 
rivers (δ15N-N-O3− = −0.07 ± 0.01 × Latitude + 10.14 ± 0.46,
and 2013.

retrieved from literature or the IAEA Coordinated Research Project (CRP), respectively. Most data were from 30° N and 60° N LAT and between 2010 and 2013.

Relationship between 15N and 18O of NO3−. Both river and groundwater nitrate revealed positive correlations between the δ15N and δ18O of NO3− and latitude yielded a similar regression slope and intercept.

Variation of nitrate isotopic composition with water temperature. Water temperature (Twater) is often used as a proxy for potential rates of the microbial activity or primary productivity and was used to consider its relationship to the isotopic composition of NO3− in rivers and aquifers. In this assessment, the temperature was the river or groundwater temperature at the time of sampling without further considering overall climatic or seasonal aspects. Detailed results of least-square linear regression analyses of the nitrate isotopic variables for river water and groundwater subsets are summarized in Supplementary Table 5.

Fig. 1 Geographical distribution of δ15N and δ18O data for nitrate in river water and groundwater by number (#) of measurements by 0.1° latitude and year of sample collection. The size of each circle corresponds to the concentration of NO3-N in mg L−1. The light and dark gray bars indicate the data were retrieved from literature or the IAEA Coordinated Research Project (CRP), respectively. Most data were from 30° N and 60° N LAT and between 2010 and 2013.

p-value < 0.05, n = 521, R² = 0.07; δ18O-NO3− = −0.12 ± 0.02 × Latitude + 9.47 ± 0.79, p-value < 0.05, n = 506, R² = 0.06) and groundwater (δ15N-NO3− = −0.15 ± 0.03 × Latitude + 14.59 ± 1.16, p-value < 0.05, n = 301, R² = 0.08; δ18O-NO3− = −0.30 ± 0.03 × Latitude + 16.05 ± 1.43, p-value < 0.05, n = 299, R² = 0.18) (Supplementary Fig. 2 and Supplementary Table 3). When data from the Northern Hemisphere were considered separately, the correlation between δ15N and δ18O of NO3− and latitude yielded a similar regression slope.

The influence of δ18O from local water contributing to nitrate during its transformations, we normalized the δ18O-NO3− to the δ18O-H2O values from the same water sample wherever possible (i.e., “relatively to in situ H2O” instead of “relative to SMOW”)31. The normalization to in situ water 18O resulted in an improved but weak correlation between δ18O-NO3− vs H2O and δ15N-NO3− for river waters (δ18O-NO3− vs H2O = −0.21 ± 0.02 × δ15N-NO3− − 0.33 ± 0.19, R² = 0.09, p-value < 0.05, n = 946) and groundwater (δ18O-NO3− vs H2O = −0.10 ± 0.02 × δ15N-NO3− − 1.17 ± 0.06, R² = 0.15, p-value < 0.05, n = 1138). The slopes of the “normalized” regression became negative and decreased from ~0.6 to ~−0.2 for rivers and from ~−0.4 to ~−0.1 for aquifers. The river water samples from warmer tropical areas (e.g., Kenya, Ghana, Thailand) drove slopes to more negative δ18O and higher δ15N values. Results of the least-square linear regression analyses of the isotopic variables for river water and groundwater subsets are summarized in Supplementary Table 4.

The influence of 18O in water and molecular O2 on the 18O of NO3− produced during nitrification is typically described using a simple isotope mass balance14:

\[ \delta^{18}O-NO_3^- = 2/3 \times \delta^{18}O-H_2O + 1/3 \times \delta^{18}O-O_2 \]  

Eq. (1). Studies investigating nitrification in aquatic systems found the δ18O of NO3− does not necessarily conform to this model, even when the δ18O of O2 is assumed constant or in equilibrium with air35 (+23.5 = +24.2‰). To reduce the
with $T_w (\delta^{15}N-NO_3^- = 0.12 \pm 0.04 \times T_w + 6.55 \pm 0.82, R^2 = 0.03, p\text{-value} < 0.05, n = 253)$, but for groundwater, there was no correlation ($\delta^{15}N-NO_3^- = 0.05 \pm 0.05 \times T_w + 8.45 \pm 1.03, R^2 = 0.01, p\text{-value} > 0.05, n = 189$) (Fig. 3). The $\delta^{18}O$ of $NO_3^-$ in rivers had a weak but significant positive correlation with $T_w (\delta^{18}O-NO_3^- = 0.10 \pm 0.04 \times T_w + 5.19 \pm 0.90, R^2 = 0.02, p\text{-value} < 0.05, n = 275)$, whereas in groundwater the correlation was very strong ($\delta^{18}O-NO_3^- = 0.48 \pm 0.04 \times T_w - 3.79 \pm 0.95, R^2 = 0.41, p\text{-value} < 0.05, n = 170$). Regressions against $T_w$ using in situ normalized $\delta^{15}N$ values were conducted and resulted in less variable and slightly negative non-zero slopes like Fig. 2.

**Seasonal variation of stable isotopes of NO$_3^-$**. To investigate the effect of seasonality on the isotopic composition of nitrate in rivers and groundwater, we used data from temperate climates (C), given that seasonality is strong in this climatic zone compared to the others (see “Methods” section, Fig. 4). The mean $\delta^{15}N$ values of nitrate in rivers showed no difference between spring and fall ($+6.9 \pm 3.6\%o, n = 583$ to $+6.7 \pm 2.6\%o, n = 606, p\text{-value} > 0.05$), but a $-1\%$ increase in $\delta^{15}N$ in winter compared to all other seasons ($+7.7 \pm 3.8\%o, n = 200, p\text{-value} < 0.05$). The average $\delta^{18}O$ values of nitrate in rivers revealed a $\sim 2\%$ decrease from spring to autumn ($+2.2 \pm 5.9\%o, n = 530$ to $-0.1 \pm 4.5\%o, n = 562, p\text{-value} < 0.05$), and a $\sim 4.5\%$ increase from autumn to winter ($-0.1 \pm 4.5\%o, n = 562$ to $+4.4 \pm 4.5\%o, n = 149, p\text{-value} < 0.05$).

Shallow aquifers showed no $\delta^{15}N$ seasonality ($p\text{-value} > 0.05$) in mean $NO_3^-$ between spring, summer, autumn, and winter ($+7.8 \pm 4.4\%o, +7.7 \pm 3.9\%o, +8.1 \pm 3.8\%o$, and $+8.1 \pm 3.5\%o$, respectively). The average $\delta^{18}O$ of $NO_3^-$ in groundwater was similar in summer, autumn, and winter ($+0.6 \pm 4.8\%o, +0.6 \pm 4.6\%o$, and $+0.1 \pm 3.3\%o$, respectively) but was $\sim 2\%$ higher in spring ($+2.1 \pm 6.0\%o, n = 268, p\text{-value} < 0.05$) compared to all other seasons. However, considering that groundwater replenishment usually takes place in the wet period (October–April) in the northern hemisphere, we examined the difference in the isotopic values between the high recharge period and the low recharge period (May–September). There was no significant difference in $\delta^{15}N$ and $\delta^{18}O$ of $NO_3^-$ between the two periods, although the $\delta^{18}O$ of $NO_3^-$ was slightly lower in the high recharge period (Supplementary Fig. 3). No clear-cut pattern was observed for the relationship between $\delta^{15}N$ and $NO_3^-$ in rivers or groundwater by season to confirm single predominant biogeochemical processes, including denitrification$^{14}$ (Supplementary Fig. 4).

**Fig. 2 $\delta^{18}O$ vs $\delta^{15}N$ of NO$_3^-$ in rivers and groundwater.** Symbol size denotes NO$_3^-$ concentration in mg L$^{-1}$. The $\delta^{18}O$ values are relative to SMOW and normalized to in situ H$_2$O, as described in Venkiteswaran et al.$^{31}$. The results of the linear regression models of panels a-d are in Supplementary Table 4.

**Varying of stable isotopes of NO$_3^-$ with climate.** The average $\delta^{15}N$ of nitrate in rivers was significantly higher ($-3.0\%o$) in tropical (A) ($+10.2 \pm 5.1\%o, n = 137$) vs temperate (C) ($+7.5 \pm 3.7\%o, n = 834$) or cold (D) ($+7.4 \pm 4.0\%o, n = 252$) climates (Fig. 5). The same pattern was observed for $\delta^{18}O$ of NO$_3^-$ for rivers, having $\sim 2$–$3\%$ higher values in tropical (A) ($+6.6 \pm 5.9\%o, n = 137$) vs temperate (C) ($+4.4 \pm 5.4\%o, n = 722$) and cold (D) ($+3.8 \pm 5.6\%o, n = 275$) climates. Arid climates (B) showed significantly higher $\delta^{15}N$ values for nitrate ($+8.9 \pm 6.0\%o, n = 74$) compared to temperate (C) or cold climate (D), but similar $\delta^{18}O$ values for nitrate ($+5.5 \pm 8.1\%o, n = 74$) compared to other climate types. However, the arid climate zone had fewer data compared to the other climate types due to a limited number of river water nitrate studies in those climates.

The shallow aquifers had the lowest $\delta^{15}N$ values in temperate (C) ($+7.8 \pm 4.1\%o, n = 986$) and arid (B) climates ($+7.3 \pm 4.1\%o, n = 522$) and the highest in tropical/equatorial (A) ($+8.9 \pm 5.5\%o, n = 44$) and cold (D) climates ($+9.0 \pm 5.2\%o, n = 24$). However, these differences were indistinguishable ($p\text{-value} > 0.05$), hence no
Fig. 3 Variation of $\delta^{18}O$ and $\delta^{15}N$ of NO$_3^-$ in river waters and groundwater with water temperature ($T_w$) in °C. The linear regression of $\delta^{18}O$-NO$_3^-$ (‰) vs $T_w$ (°C) was more significant for groundwater ($R^2 = 0.41$) than river waters ($R^2 < 0.02$). The results of the linear regression models for panels a-d are in Supplementary Table 5.

Fig. 4 Variation of $\delta^{15}N$ and $\delta^{18}O$ of nitrate in rivers and groundwater classified by season. The symbol size expresses the NO$_3^-$-N in mg L$^{-1}$. Higher average $\delta^{15}N$ and $\delta^{18}O$ values in river water nitrate are found in winter. Error bars show average (gray line), median (light to dark gray division), 1st and 3rd quartile, and whiskers extend to 1.5 interquartile range.
N-isotopic impact solely from climate could be found. Groundwater nitrate $\delta^{18}O$ values exhibited the same pattern as rivers, with lower $\delta^{18}O$ values in cold (D) climates ($-0.1 \pm 6.4\%o$, $n = 24$), that differed significantly from those in tropical (A) ($+4.6 \pm 6.0\%o$, $n = 44$) climates. The average $\delta^{18}O$ of NO$_3^-$ values in arid (B) ($+3.4 \pm 4.0\%o$, $n = 505$) and temperate (C) ($+2.5 \pm 4.9\%o$, $n = 505$) climates were significantly different from the other two climate types. Cold climates had lower $\delta^{15}N$ and $\delta^{18}O$ records for groundwater compared to the other three climate types, which could bias this result. Rivers and groundwater showed no clear-cut relationship between $\delta^{15}N$ and NO$_3^-$ by climate type (Supplementary Fig. 5).

**Discussion**

The lower median NO$_3^-$ concentrations in rivers compared to adjacent shallow aquifers can be explained by the decrease of agricultural N inputs at the industrialized countries of the Northern hemisphere due to more severe fertilization regulations, whereas many aquifers do not yet exhibit decreasing nitrate contents. This is due to much longer mean residence times for N in the unsaturated zone and the aquifer38, which is controlled by several factors such as the thickness and hydro-lithological characteristics of the unsaturated zone, the spatiotemporal variations in recharge rate, and the diffusive and dispersion processes through the nitrogen stock in the soil and the unsaturated zones. The lower median NO$_3^-$ concentrations in rivers can also be explained by a complex combination of riparian and in situ attenuation and uptake by biological productivity, seasonal dilution (e.g., runoff, snowmelt), differential baseflow connectivity, and long-term dispersion processes, which lowers the ambient NO$_3^-$ concentration in rivers. Given that NO$_3^-$ sources are difficult to “fingerprint” using only stable isotopes, we suggest that the notable differences in NO$_3^-$ concentrations between river waters and adjacent aquifers are attributable to unique locational differences in situ N-cycling processes. As microbes have low biomass in most groundwater systems, NO$_3^-$ attenuation, and cycling is achieved by closed-system nitrification and denitrification and uncommonly by DNRA or anammox.39,40 Shallow aquifers are the recipients of NO$_3^-$ transport from residual leaching from soils and the unsaturated zone, where NH$_4^+$ and other nitrogenous compounds are favorably oxidized but can also be adsorbed or reduced depending on soil retention capacity46-49. Pronounced denitrification effects have been reported for some unconfined aquifers40,44,45.

Regression analysis of $\delta^{18}O$ vs $\delta^{15}N$ for river waters and groundwater generated slopes suggesting an “apparent” denitrification trend ($-0.5$)3,31. However, denitrification is unlikely to be the only process for the observed $\delta^{18}O$: $\delta^{15}N$ slope, which is expected to be canonical under closed-system conditions.46 The lack of a clear pattern between the isotopes and NO$_3^-$ concentration indicates that multiple processes that recycle nitrogen occur in river water systems. Granger and Wankel19 proposed that the deviations from the canonical slope of 1 for denitrification can be due to concurrent NO$_3^-$ production catalyzed by nitrification and/or anammox.

A non-zero normalized slope suggests that H$_2$O was not the only factor controlling the oxygen isotopic composition of nitrate (Eq.1). The large decrease in the slopes of the regression after normalization for in situ water confirmed a strong dependence of $\delta^{18}O$ of NO$_3^-$ on the $\delta^{18}O$ of water when the NO$_3^-$ is formed by oxidation of nitrite or ammonium.42 However, in situ water normalizations do not consider potential large daily or seasonal variations in the $\delta^{18}O$ of in situ gaseous or dissolved O$_2$35 involved in the N-cycling processes. Thus, the observed negative deviations of the slopes from “zero” suggest an influence of $\delta^{18}O$-O$_2$ on the isotopic composition of nitrate. The latter may be more profound for rivers than aquifers. Investigations of $\delta^{18}O$-O$_2$ in productive rivers show a preponderance of lower $\delta^{18}O$ values (as low as +3.4%), particularly during the growing season due to photosynthetic processes35,47,48. Tropical rivers typically show high rates of primary production, all-year-round growth, and less seasonal variation in solar irradiance than temperate latitudes49.

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**Fig. 5 Variation of $\delta^{15}N$ and $\delta^{18}O$ of NO$_3^-$ in rivers and groundwater by climate.** A: Tropical/Equatorial, B: Arid, C: Temperate, D: Cold. Symbol size expresses the NO$_3$N concentration (mg L$^{-1}$). Tropical climates had the highest $\delta^{15}N$ and $\delta^{18}O$ values for nitrate in rivers. Error bars show average (gray line), median (light to dark gray division), 1st and 3rd quartile, and whiskers extend to 1.5 interquartile range.
On a diel basis, $\delta^{18}O-O_2$ is lowest during the day due to the production of DO by photosynthesis from substrate water, whereas during the night $\delta^{18}O-O_2$ can be higher than the atmospheric equilibrium value due to preferential consumption of light $O_2$ by respiring organisms with inward diffusion of isotopically heavy $O_2$ from the atmosphere. Thus, the negative in situ normalized slopes of the $\delta^{18}O$ regression line for rivers suggest some influence of $^{18}O$ depleted $O_2$, especially for highly productive rivers where oxygen supersaturation and $^{18}O_2$ depletion and nitrification coexist. But even if the $\delta^{18}O$-nutrient contribution was considered as a fixed atmospheric constant, any kinetic and equilibrium isotope fractionation during oxygen exchange between $H_2O$ and $NO_2^-$ before final oxidation to $NO_3^-$ may also result in deviations from a 2:1 mixing ratio line ($\delta^{18}O-H_2O,\delta^{18}O-O_2$). Therefore, the relationship between $\delta^{18}O$-NO$_3^-$ and $\delta^{18}O$-$H_2O$ can be used to detect isotope fractionation effects related to the incorporation of oxygen atoms during ammonia oxidation to nitrite ($NO_2^-$), which readily exchanges oxygen atoms with in situ water during the oxidation of the latter to nitrate. Conversely, for aquifers the $\delta^{18}O$-$O_2$ values in recharge zone gas often deviate from the air ($+23.5\%o$), but mostly in a positive direction up to $>39\%$ as soil $O_2$ is consumed by microbial respiration. The substantial variability in the $\delta^{18}O$ of NO$_3^-$ associated with nitrification is also dependent on soil and water properties and is also a function of residence time.

The difficulty to elaborate clear relationships between $^{18}O$ of $H_2O$, $O_2$, and NO$_3^-$ is also due to the temporal variability of the oxygen variables and underscores the need for further investigations to better understand N-dynamics related to cycling. Unfortunately, oxygen isotope normalization processes that consider all potential reactive oxygen pools (e.g., in situ $H_2O$ and $^{18}O_2$) involved in N-cycling processes have not been fully investigated to our knowledge. New studies measuring $\delta^{18}O$-NO$_3^-$ and $\delta^{18}O$-$H_2O$ and $\delta^{18}O$-$O_2$ in the same water are therefore required to better explain the patterns and variations of the oxygen isotope composition of nitrate in rivers and groundwater.

The $\delta^{15}N$ and $\delta^{18}O$ of NO$_3^-$ in rivers showed a positive correlation with $T_w$, likely reflecting the influence of higher microbial activity and productivity with temperature and thus the biogeochemical N processing rates. Higher $\delta^{15}N$ values in rivers are likely driven by enhanced N-cascading. This means that due to increased temperature, metabolic rates accelerate, which progressively enriches $^{15}N$ in NO$_3^-$ compared to its source signature. Average $\delta^{15}N$ values of nitrate $>7\%o$ typically suggest that nitrogen has undergone multiple recycling after its initial fixation from $N_2$ (either biological or Haber–Bosch), as these processes bring reactive $N$ in the plant–soil–water continuum with a $^{15}N$ value of $\approx 0\%o$. An observed weak positive correlation of $\delta^{18}O$-NO$_3^-$ with $T_w$ and a negative correlation with latitude suggests that ambient $\delta^{18}O$-$H_2O$ has some impact on nitrate isotope values.

The pattern of decreasing $\delta^{15}N$-NO$_3^-$ in rivers with latitude agrees with the $N$ isotopic climatic patterns observed for soil organic matter (SOM). The pattern of higher $\delta^{15}N$-NO$_3^-$ values with the water temperature in rivers is observed from cold (D) to tropical (A) climates. Amundson et al. observed a weakly positive correlation between soil organic matter (SOM) $\delta^{15}N$ and Mean Annual Temperature (MAT) (slope = 0.13, $R^2 = 0.11$, $p$-value $< 0.1$, $n = 85$) for arid and tropical zones. A decrease in SOM $\delta^{15}N$ with lower MAT was seen at sites between about 10–40 degrees North or South albeit with a smaller data set (slope = $-0.08$, $R^2 = 0.09$, $p$-value $< 0.04$, $n = 49$). Craine et al. observed a positive trend of SOM $\delta^{15}N$ with MAT (slope = 0.18, $p$-value $< 0.001$, MAT > 9.8 °C) and a negative correlation with Mean Annual Precipitation (MAP) at a global scale that was attributed to soil and organic matter properties. Martinelli et al. reported higher values for $\delta^{15}N$ in N-rich soils and foliage from tropical forests compared to temperate forests. Brookshire et al. suggests that higher MAT enhances soil N-cycling as seen by $^{15}N$ enrichment in soil organic matter and the NO$_3^-$ concentration of tropical rivers.

The seasonality of $\delta^{15}N$ and $\delta^{18}O$ of nitrate in temperate rivers showed a pattern of isotopic enrichment of nitrate in spring and summer compared to autumn, which could be attributed to higher biological assimilation in surface water systems in the warmer seasons. The $\delta^{15}N$ values during winter compared to other seasons appeared to more closely reflect mixing of multiple organic-pollutant N-sources (e.g., sewage wastes, urban, and livestock wastewaters) as a result of reduced application and leaching of fertilizers, and restricted microbial and primary productivity during the cold season. Additionally, in the cold season, the relative fraction of NO$_3^-$ denitrified vs assimilated is higher, since increased $N$-loadings in winter or early spring runoff are often associated with high discharge events and snowmelt. Elevated $\delta^{15}N$ of NO$_3^-$ during winter can also be attributed to $\delta^{18}O$-$H_2O$ and $\delta^{18}O$-$O_2$ variation combined with variable soil/water and land-use properties of the different catchments. For example, high $\delta^{18}O$ values in nitrate can be attributed to the influence of $\delta^{18}O$-$O_2$ (Eq.1), especially in respiration-influenced ecosystems, increased atmospheric NO$_3^-$ deposition ($> +30\%o$), especially during high discharge events, or artificial NO$_3^-$ fertilizer ($> +20\%o$) mixed with nitrification-derived NO$_3^-$ in soils and infiltrating pore waters, especially in agricultural areas. Full equilibration or O-isotope exchange of NO$_3^-$ with $H_2O$ before being oxidized to NO$_3^-$ in soils, local respiration, and rapid redox cycling between NO$_3^-$ and NO$_2^-$ can also potentially increase $\delta^{18}O$-$NO_3^-$ values.

Shallow phreatic aquifers showed strongly dampened seasonal nitrate and isotopic variability compared to river waters. Given NO$_3^-$ concentrations are generally buffered in aquifers over longer timeframes (e.g., years), the range of $\delta^{15}N$ values in groundwater reflect longer-term climatic and land-use changes or established processes taking place in the soil and the unsaturated zone before NO$_3^-$ leaches into the groundwater system. The lack of a clear seasonal variation of $\delta^{15}N$ and $\delta^{18}O$ of NO$_3^-$ in groundwater is attributed to nitrifying and mixing of nitrate sources introduced at different points or periods in time due to variable residence times (from months to years) in the unsaturated zone. There was no significant difference in $\delta^{15}N$ and $\delta^{18}O$ of NO$_3^-$ between the high and low recharge periods, although the $\delta^{18}O$ of NO$_3^-$ was slightly lower in the high recharge period. This could imply a change in recharge source water (e.g., snowmelt vs rains) or the $^{18}O$ of O$_2$, which combined affect the $^{18}O$ values of microbially mediated nitrate. Because groundwater temperatures are usually stable and generally reflect MATs and in situ $\delta^{18}O$-$H_2O$ correlates with air temperature), it was unsurprising to see a correlation of $\delta^{18}O$ of NO$_3^-$ with $T_w$. This suggests we cannot discount the possibility that $^{18}O$ variance of groundwater nitrate is controlled by additional processes or isotope fractionations affecting O$_2$ involved in soil and vadose zone nitrification. However, the relationship between $\delta^{18}O$ in nitrate with latitude showed the highest oxygen isotope values tended to be at the warmer tropical/equatorial climates and can be attributed to a systematic change in the $\delta^{18}O$ of associated H$_2O$ (Eq.1) with decreasing latitude, continentality or altitude.

In summary, although rivers and shallow aquifers are primary and often well connected receiving environments for land-based anthropogenic N-pollution, shallow aquifers have five times higher median NO$_3^-$ concentration than rivers, indicating that...
\( \text{NO}_3^- \) is more persistent in aquifers and has long residence times, given lower microbial activity and a limited number of possible N-removal process mechanisms (e.g., denitrification). River waters and groundwater receive nitrate with \( \delta^{15}N \) and \( \delta^{18}O \) values spanning the expected natural and anthropogenic ranges (from \(-10\%\) to \(+26\%\) for \( \delta^{15}N \) and \(-17\%\) to \(+25\%\) for \( \delta^{18}O \)), indicating a globally consistent suite of N-sources. We identified several global drivers of the isotopic variability of nitrate in rivers and groundwater, recognizing that this simplification required aggregating many possible diverse nitrate sources typically observed at the catchment or complicated by seasonal variability. A deeper analysis of latitude and climatic factors is recommended to better distinguish the role of soil properties, precipitation patterns, land-use, and agricultural practices. The \( \delta^{18}O \) vs \( \delta^{15}N \) relationships observed suggested combinations of multiple biogeochemical (e.g., nitrification, anammox) and mixing processes taking place in rivers and aquifers in different seasons, which are not easily distinguishable with infrequent or synoptic samplings. River water nitrate had higher \( \delta^{15}N \) and \( \delta^{18}O \) values in winter, suggesting a change in N-processing due to lower temperatures and productivity and perhaps less alteration of the original isotopic composition of multiple organic-related N-sources. Across a wide range of global climate types, river waters and groundwater showed systematically higher \( \delta^{15}N \) and \( \delta^{18}O \) values in the arid and tropical/equatorial climates, mirroring an increase in MAT. This relationship was also seen in the increase in the \( \delta^{15}N \) of nitrate in rivers and groundwater with decreasing latitude and the positive correlation between \( \delta^{15}N \) and \( \delta^{18}O \) with \( T_w \).

We have provided a first-order attempt to identify global patterns in the isotopic composition of nitrate related to common environmental factors, such as climate and season impacting N-cascade-related processes in rivers and associated shallow groundwater. Our findings highlight the importance of water temperature as a driving force of biogeochemical microbial activity and productivity in rivers, however, the impact on the \( \delta^{15}N \) and \( \delta^{18}O \) of nitrate differed when examined on a seasonal or on a climate basis. Our findings suggest that higher frequency or seasonal sampling of river water systems is of critical importance to better and more deeply understand annual N-pollution dynamics, and winter sampling appears to better reflect the N-isotopic signals of the anthropogenic sources of nitrate. Further investigations to study N-cycling in river waters at a higher frequency are urgently needed (e.g., seasonal, diel) and systematic data collections of nitrate, particularly from the southern hemisphere climate types, such as arid and polar, for which data are limited or absent, are strongly recommended. Recent technological advances in isotopic assays of nitrate have vastly reduced the analytical and cost barriers to conduct high-frequency \( \delta^{15}N \) and \( \delta^{18}O \) of nitrate. Other key parameters, such as DO, redox potential, labile organic material, and solar radiation should also be systematically measured or considered to help better understand the controlling factors of the N-transformations in river waters, especially on a seasonal basis. Measuring \( \delta^{18}O \) of the water and dissolved oxygen from the same samples as \( \delta^{18}O \) of \( \text{NO}_3^- \) will allow a better assessment of N-cycling processes in rivers and groundwater. Understanding the fate of nitrogen in aquifers and the connectivity to rivers requires a fuller local understanding of hydrogeological processes, such as water source, flow paths, and water residence times (e.g., \( ^3 \)H tracers) that control nutrient transit times and contribution to baseflow (e.g., retardation of N due to the recycling of N in organic matter and subsequent mineralization processes). Our synthesis affirms nitrate isotope techniques are useful to assess the origin of nitrogen pollution in rivers and groundwater. However, we caution that interpretations of nitrate isopes may be complicated where nitrogen undergoes many biogeochemical transformations, especially in productive systems, that can mask the original isotopic source signal. Examination of N-cascading aspects are critically important to inform the implementation of beneficial land and agricultural management strategies aiming at mitigating increasingly serious nitrogen pollution in Earth’s rivers and aquifer systems. The clear isotopic linkage of the N-cascading in rivers to key climate parameters may have implications for future water quality management under changing climatic conditions.

**Methods**

**Literature data collection.** A literature search was conducted using online tools and bibliographic databases (e.g., Web of Science, Google Scholar, Scopus, etc.) for papers containing nitrate and isotope data from rivers and adjacent aquifers. We collected shallow aquifer data because many of the river studies contained data for adjacent aquifers owing to the possible connectivity of groundwater to riverine baseflow, although in most papers any connectivity of adjacent aquifers to the rivers was unconfirmed. Data compilations were restricted to journal articles where both \( \delta^{15}N \) and \( \delta^{18}O \) values of nitrate are available, and samples were then classified as River or Groundwater. Scientific papers focusing on precipitation, deep aquifers, soils, tap water, seawater, wastewater effluent, or artificial isotopic tracers were omitted. The list of cited data used is found in Supplementary References.

**Data preparation.** Established data preparation procedures resulted in a curated data set suitable for statistical analysis. Data were extracted from tables and supplementary materials, maps, and text (Supplementary Fig. 6). If the original work did not provide location coordinates, approximate latitude and longitude were obtained using Google Earth (https://www.google.com/earth/). The data assessment affirmed that \( \text{NO}_3^- \) was the species linked to the reported isotope ratios \( (\text{\^{15}N}/\text{\^{16}N} \text{ and } \text{\^{18}O}/\text{\^{16}O}) \) along with verification of units (as \( \text{NO}_3^- \) or N) and concentration (e.g., mg L\(^{-1} \), mmol L\(^{-1} \), µmol L\(^{-1} \), etc., etc.). Nitrate concentration data were converted to a common unit (mg L\(^{-1} \) or summer) and sampling date format (e.g., 2016/06/01) (Supplementary Table 1). The data set comprised 5194 sample points. Additional categorical and site information associated with the nitrate and isotopic data was obtained from the publications and is included in Supplementary Table 6. Important variables, such as pH, dissolved O\(_2\) (DO), dissolved organic nitrogen (DON), precipitation per catchment, Total Nitrogen (TN), that might contribute to a better understanding of the global N isotope variations of nitrate were either lacking or unavailable. For example, DO values were only available in 7.7% of the river waters data set and thus were excluded from further processing.

**Data curation.** Excessively high or suspicious values of \( \text{NO}_3^- \) and other N-species concentrations were verified by a careful review of the original paper. In some cases, authors were contacted to provide clarification. The \( \delta^{15}N \) and \( \delta^{18}O \) values of nitrate were sorted to avoid duplication from publications reusing the same data sets. Almost all groundwater data (~95%) was from adjacent shallow phreatic aquifers (<20 m depth below ground surface).

**Final data set and analytical aspects.** The data set comprised 5194 sample points from 70 papers (~86% of the whole data set, Supplementary References) plus new isotopic results. There were 2902 nitrate and associated isotope analyses from rivers and 2291 analyses from nearby shallow groundwater. Most samples were from between 2002 and 2016; fewer data exist from 1968 to 1990s due to analytical barriers particularly for \( \delta^{15}N \) and \( \delta^{18}O \). A total of 374 new river water nitrate and isotope samples from 10 countries were analyzed in the AIAE Isotope Hydrology Laboratory using the Ti(III) reduction method\(^{23} \) to reduce nitrate to N\(_2\)O gas in septum vials. The N\(_2\)O in the headspace was measured for \( \text{N}_2 \) and \( \text{NO}_3^- \) using a CF-IRMS (Isoprime-100 Trace Gas Analyser). The analytical uncertainties were ±0.2% and ±0.4% for \( \delta^{15}N \) and \( \delta^{18}O \), respectively. For Quality Controlling the water samples that do not give the expected N\(_2\)O yield on the IRMS based on the determined \( \text{NO}_3^- \) concentration, were repeated until they agree within 95% of the expected target before results are accepted. Oxygen isotopes \( (\delta^{18}O/\delta^{16}O) \) of the same water samples were analyzed by laser spectrosopy at the AIAE using a Los Gatos Research Liquid Water Isotope Analyser (TwIA-912). The Analytical uncertainty was ±1.5% (2SD). Chemical analysis of new samples included nitrate following standard discrete analyzer methods\(^{27} \).
Statistical evaluation. To evaluate the data, graphical plots by sample type (river or groundwater) were employed along with frequency distribution diagrams (histogram, Q-Q plot, P-P plot, boxplot) and metrics such as Pearson skewness. To check for normality, Shapiro–Wilk, and Kolmogorov–Smirnov tests were applied. The mean and standard deviation (SD) were used to describe the variation of variables with a skewness index of <2, whereas median and the median absolute deviation (MAD) were used otherwise. A single-factor analysis of variance (ANOVA) and Kruskal–Wallis non-parametric analysis of variance were performed to determine if differences between the means or the medians of the groups were significant. A conventional probability value (p-value) of 0.05 was used to indicate significance for differences between river and groundwater and seasonal subsets.

A least-square linear regression model of type I was applied to investigate relationships between $\delta^{15}$N and $\delta^{18}$O of nitrate to other variables. The least-square linear regression model was used to model the intercept and to determine if differences between the means or the medians of the groups were significant. To check for normality, Shapiro–Wilk, and Kolmogorov–Smirnov tests were performed. The mean and standard deviation (SD) were used to describe the variation of variables with a skewness index of $<$2, whereas median and the median absolute deviation (MAD) were used otherwise. A single-factor analysis of variance (ANOVA) and Kruskal–Wallis non-parametric analysis of variance were performed to determine if differences between the means or the medians of the groups were significant.

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Author contributions

LM. and L.I.W. conceived and wrote the paper. I.M. and L.R.M. undertook the formal analysis and visualization of the work. J.Y., D.C.G., P.B., and E.S. discussed the results and provided critical reviews and edits of the final manuscript. F.-Y.Y., G.M., C.A.H., C.B., L.B., N.V.E., W.F., J.R.F., A.G., N.K., S.-L.L., M.T.N.L., S.P., V.R., D.S.R., A.R., P.S., F.T., D.A.T., W.W., and N.W. contributed to the review and editing of the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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