Evaluating Regional Patterns in Nitrate Sources to Watersheds in National Parks of the Rocky Mountains using Nitrate Isotopes

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In the Rocky Mountains, there is uncertainty about the source areas and emission types that contribute to nitrate (NO_3) deposition, which can adversely affect sensitive aquatic habitats of high-elevation watersheds. Regional patterns in NO₃ deposition sources were evaluated using NO₃ isotopes in five National Parks, including 37 lakes and 7 precipitation sites. Results indicate that lake NO₃ ranged from detection limit to 38 μ eq/L, δ^{18} O (NO₃) ranged from -5.7 to +21.3‰, and δ^{15} N (NO₃) ranged from -6.6 to +4.6‰. δ^{18} O (NO₃) in precipitation ranged from +71 to +78‰. $\delta^{15}N$ (NO₃) in precipitation and lakes overlap; however, $\delta^{15}N(NO_3)$ in precipitation is more depleted than $\delta^{15}N$ (NO₃) in lakes, ranging from -5.5to -2.0%. $\delta^{15}N$ (NO₃) values are significantly related (p < 0.05) to wet deposition of inorganic N, sulfate, and acidity, suggesting that spatial variability of δ^{15} N (NO₃) over the Rocky Mountains may be related to source areas of these solutes. Regional patterns show that NO₃ and δ^{15} N (NO₃) are more enriched in lakes and precipitation from the southern Rockies and at higher elevations compared to the northern Rockies. The correspondence of high NO₃ and enriched δ^{15} N (NO₃) in precipitation with high NO₃ and enriched δ^{15} N (NO₃) in lakes, suggests that deposition of inorganic N in wetfall may affect the amount of NO₃ in lakes through a combination of direct and indirect processes such as enhanced nitrification.

Introduction

In the western United States, anthropogenic emissions of NO_x (nitrogen oxides) and NH_3 (ammonia) from energy generation activities, transportation, industry, and agricultural activities contribute to deposition of dissolved inorganic

nitrogen (DIN = $NO_3 + NH_4$) in high-elevation watersheds (1, 2). There is considerable uncertainty about the source areas and emission types that contribute to deposition of DIN, which can adversely affect sensitive aquatic habitats of high-elevation lake basins (3). Deposition data indicate that DIN in wetfall has increased steadily over much of the Rocky Mountains in recent years for a variety of reasons (4-6), including increases in motor vehicle emissions which have offset reductions in NO_x emissions from fossil-fuel burning industries (7) and regional increases in ammonia emissions (8). The percent of DIN in wet deposition contributed by NH₄ has increased from 1992-1996 to 2002-2006, and is now approximately 50% of measured DIN in over half of the National Atmospheric Deposition Program/National Trends Network (NADP/NTN) sites in the Rocky Mountains (8), including sites located near national parks. A previous study that evaluated all major emission sources (including both stationary and mobile sources) across the Rocky Mountains found that the Colorado Front Range, which is located near large urban centers, has the highest N emissions (2). Spatial trends in deposition of DIN in wetfall over the Rocky Mountains show that deposition is greatest near Rocky Mountain National Park and Great Sand Dunes National Park and Preserve near the Colorado Front Range in the southern Rocky Mountains, compared to National Parks in the northern Rocky Mountains that are also located further west (9) (Supporting Information Figures SI-1 and SI-2).

Federal and state resource managers are investigating policy options to alleviate this problem by reducing anthropogenic emissions of NO_x and NH_3 . However, identifying source areas and emission types is complicated (10, 11). Isotopic tracers of N measured in precipitation and water samples show promise in helping to identify these emission sources (10, 11). Previously published studies (12, 13) indicate δ^{15} N (NO₃) values in NO_x emissions from coal-fired power plants have isotopic values ranging from +6 to +13% (12, 13). δ^{15} N (NO₃) values from motor vehicle NO_x emissions in tailpipe exhaust range from -13 to -2% (12). The following δ^{15} N (NO₃) values have also been reported for vehicle NO_x emissions in tailpipe exhaust (+3.7‰) and roadside vegetation (+3.8%) (14, 15). The use of these NO₃ isotopes has been limited in part because analytical techniques for NO3 isotopes required large sample volumes that made it logistically difficult to sample in areas with topographically complex terrain.

Researchers have shown that elevated levels of atmospheric N deposition in the Front Range of Colorado have caused substantial changes in the state and function of terrestrial and aquatic ecosystems at high elevations (1, 16-20). In one study, inorganic N retention of DIN in wetfall averaged 72% in high-elevation ecosystems (21). N deposition in excess of the total combined plant and microbial demand can cause watershed N saturation and increased rates of N leaching from soils to aquatic ecosystems (22), which is occurring in the Colorado Rockies (20, 23). This excess N can result in a cascade of ecological effects in surface waters that includes acidification, eutrophication, and increased emissions of N₂O, a greenhouse gas. Eutrophication increases primary productivity in lakes and streams and alters diatom species distributions that form the base of the food web in many high-elevation lakes (24). The combined effects of increasing N deposition and drought have sharply increased streamwater concentrations of NO3 in Rocky Mountain National Park (Rocky) in recent years (20). Changes in the water quality of the headwater systems affects not only fish, wildlife, and

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ecosystem integrity, but also downstream ecosystems and water users.

Increased aquatic productivity resulting from eutrophication accelerates the accumulation of organic matter in the water column and in lake sediments. Decomposition of this organic matter promotes hypoxia in lakes when they are ice-covered during winter, and may adversely affect fish populations (*25*). In Rocky, mineralization of organic N in pond sediments has caused concentrations of dissolved ammonia in vernal ponds to reach levels that may be harmful to threatened amphibians that breed there (*26*). Although N deposition is greatest in the Front Range of Colorado, other high-elevation sites in the Rocky Mountains also show symptoms of early stage N saturation. The progression toward N saturation is expected to continue if N deposition continues at current or higher levels in the future (*27*).

The denitrifier method to determine the dual isotopic composition (δ^{15} N and δ^{18} O) of NO₃ is well suited for studies of NO₃ contributions to streams and lakes (*28*). This method requires only 20–60 nmol of NO₃ and enables high throughput of samples (*29, 30*). The development of a new analytical technique for analyzing δ^{15} N (NH₄) holds promise in terms of tracing other sources of N deposition. However, it requires large sample volumes and is beyond the scope of this study.

To evaluate NO₃ sources, we analyzed both δ^{15} N and δ^{18} O of NO₃ using the denitrifier method in samples collected from lakes in the Rocky Mountains that span a range of NO₃ deposition (*31*). The objectives of this study were to (1) evaluate the spatial distribution of δ^{18} O (NO₃) and δ^{15} N (NO₃) in lake samples from five National Parks collected during baseflow conditions; and (2) compare the isotopic composition of the lake water from these watersheds with that of precipitation collected at nearby NADP/NTN sites. This study is the first comprehensive evaluation using NO₃ isotopes to investigate the possible relationship between atmospheric deposition of NO₃ in wetfall and the NO₃ in lakes of the Rocky Mountains.

Experimental Section

Study Area and Field Methods. The five National Parks in the Rocky Mountains included in this study are Glacier National Park (Glacier), Yellowstone National Park (Yellowstone), Grand Teton National Park (Grand Teton), Rocky Mountain National Park (Rocky), and Great Sand Dunes National Park and Preserve (Great Sand Dunes) (Supporting Information Figure SI-2). The areas under investigation are in the headwaters of most of the major rivers in the western United States, and their airsheds extend across state and national political boundaries. Precipitation chemistry was measured at 7 NADP/NTN sites located near National Parks (Supporting Information Figure SI-2). The precipitation samples collected weekly at NADP/NTN sites from 2000 were pooled into bimonthly, volume-weighted-mean composites, and analyzed for NO₃ concentrations and for δ^{18} O (NO₃) and δ^{15} N (NO₃) (10). For this study, bimonthly δ^{18} O (NO₃) and δ^{15} N (NO₃) values were aggregated into average annual wet deposition values. Sample duplicates had an average standard deviation of 0.6‰ for δ^{18} O (NO₃) and 0.4‰ for δ^{15} N (NO₃). Dry deposition was not included in this evaluation.

Lakes were randomly selected for sampling during late summer 2004 and are spatially distributed within each of the National Parks. The 56 sampled lakes range in elevation from 2000 to 3800 m and from 1 to 46 ha in area, with 65% of the lakes less than 5 ha in area. Surface waters were collected from the outflow of each lake as grab samples during the low-flow period from August to September. Samples were collected at baseflow when NO₃ concentrations in surface waters are generally near or at their annual minima as a result of biological assimilation (2). Surface waters that have elevated NO₃ concentrations (i.e., greater than about 5 μ eq/ L) during baseflow conditions may be approaching the initial stage of N saturation (*2*) and may be particularly susceptible to inputs of DIN in wetfall. Polyethylene bottles (250 mL) were soaked with deionized (DI) water overnight and then rinsed with DI water 5 times; bottles were further rinsed 3 times with sample water at the time of collection. Samples were frozen after collection and transported to the Kiowa Environmental Chemistry Laboratory (*32*) run by the Niwot Ridge Long-Term Ecological Research Program (University of Colorado, INSTAAR), which specializes in analysis of dilute waters such as those found in the study area (*33*).

Laboratory Analyses. All lake samples were analyzed for pH, acid neutralizing capacity (ANC), conductance, and major ions. ANC and pH were measured immediately after melting or after return to the laboratory using the Gran titration technique. Subsamples were immediately filtered through prerinsed (300 mL), 47-mm Gelman A/E glass fiber filters with ca. $1-\mu M$ pore size. Filtered samples were stored in the dark at 4 °C for subsequent analyses within 1-4 weeks. Anions were measured using ion chromatography (Dionex DX 500) employing chemical ion suppression and conductivity detection. Base cations were analyzed with a Varian AA6 atomic absorption spectrophotometer using an airacetylene flame. Quality assurance for this study was addressed with field duplicate samples separated by 10-15 samples in each run. Analytical precision for all solutes was less than 2% and detection limits were less than 1 μ eg/L.

Frozen aliquots were analyzed for δ^{18} O (NO₃) and δ^{15} N (NO₃) using the denitrifier method at the USGS Stable Isotope Laboratory in Menlo Park. In this method, denitrifying bacteria (*Pseudomonas aureofaciens*) quantitatively convert the N and O from NO₃ into gaseous nitrous oxide (N₂O) for isotopic analysis (*29, 30*). A minimum of 20 nmol NO₃ was required to analyze samples on a Micromass IsoPrime isotope ratio mass spectrometer (IRMS). Sample duplicates had an average standard deviation of 0.7‰ for δ^{18} O (NO₃) and 0.2‰ for δ^{15} N (NO₃).

To evaluate regional differences in spatial patterns, $\delta^{18}O$ (NO₃) and $\delta^{15}N$ (NO₃) values from lakes were compared to $\delta^{15}N$ (NO₃) values in precipitation from collocated NADP/ NTN sites. Also, isotopic values of NO₃ in lakes were compared with emissions within a specified buffer distance calculated using emissions inventories (*2*, *7*, *10*). For this study, stationary source NO_x emission inventory data that are readily available (*7*) were used as a surrogate for various anthropogenic N emission sources that are not as readily available including motor vehicles, agriculture, feedlots, power plants, and other industrial emission sources (*2*).

Basin characteristics were evaluated to determine their potential influence on spatial patterns in NO₃ concentrations, $\delta^{18}O$ (NO₃), and $\delta^{15}N$ (NO₃) values. Forty-eight basin characteristics were derived using Geographic Information Systems (GIS) software, and included bedrock type, slope, aspect, elevation, lake area, soil type, and vegetation type, following the protocols presented in ref *34*.

Results and Discussion

NO₃ concentrations in the 56 lakes sampled ranged from below the detection limit ($\sim 1 \mu eq/L$) to $38 \mu eq/L$ (Supporting Information Figure SI-3). Mean values were highest in Rocky ($20 \mu eq/L$) and lowest in Yellowstone ($0.2 \mu eq/L$). An analysis of variance test (ANOVA) shows that mean concentrations of NO₃ varied significantly among National Parks (n = 56, p< 0.001). A follow-up Tukey–Kramer HSD test shows that the mean value of $20 \mu eq/L$ for NO₃ at Rocky was significantly higher than that of the other 4 parks.

Of the 56 lakes sampled, 37 lakes had sufficient mass of NO₃ to analyze for δ^{15} N (NO₃) and δ^{18} O (NO₃) (Supporting Information Table SI-1). Lake samples with NO₃ mass greater

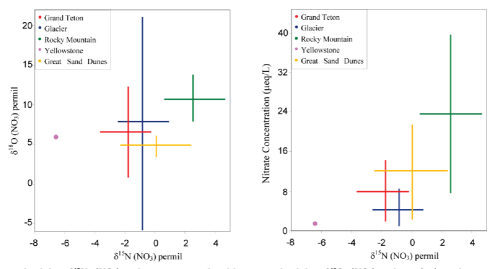


FIGURE 1. Range in lake $\delta^{15}N$ (NO₃) values compared with range in lake $\delta^{18}O$ (NO₃) values (1a) and range in lake NO₃ concentrations (1b).

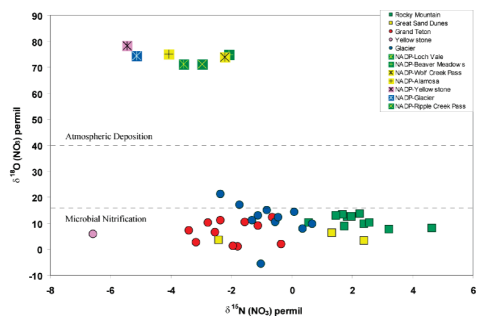


FIGURE 2. $\delta^{15}N$ (NO₃) values compared with $\delta^{18}O$ (NO₃) values at NADP/NTN sites and at lakes. Dashed lines represent estimated $\delta^{18}O$ (NO₃) end-members.

than 20 nmoles were distributed among the parks as follows: Glacier (n = 11), Grand Teton (n = 11), Great Sand Dunes (n = 3), Rocky (n = 11), and Yellowstone (n = 1).

 δ^{18} O (NO₃) values in lake samples ranged from -5.7 to +21.3%, with a median value of +10.1 (Figure 1a). Both of the lakes with elevated δ^{18} O (NO₃) values are located in Glacier and receive direct input from glacier outflow, which may explain the enriched δ^{18} O (NO₃) values. δ^{15} N (NO₃) values for the 37 lakes ranged from -6.6 to +4.6% (Figure 1a), with a median value of -0.6%. Somewhat surprisingly, this range in δ^{15} N (NO₃) values is similar to δ^{15} N (NO₃) values in wet NO3 deposition from NADP/NTN sites across the Northeastern and Mid-Atlantic U.S. (10). A plot of δ^{15} N (NO₃) values compared to NO₃ concentration, color coded by park, is shown in Figure 1b. In general, there was a trend for the lake δ^{15} N (NO₃) values to increase with increasing NO₃ concentrations ($r^2 = 0.5$; p < 0.05). Lake δ^{18} O (NO₃) values, however, did not increase with increasing NO₃ concentrations ($r^2 =$ 0.02; p > 0.1).

We compared the isotopic composition of NO_3 from these catchments to the isotopic composition of NO_3 in precipitation collected at the 7 nearby NADP/NTN sites in Figure 2 (Supporting Information Table SI-2). The δ^{18} O (NO₃) values in precipitation ranged from +71 to +78‰, significantly more enriched than the -5.7 to +21.3‰ of samples in lake water (p < 0.001). The δ^{18} O (NO₃) values in lake water are not indicative of a direct atmospheric source (Figure 2). The δ^{15} N (NO₃) values in precipitation tended to be significantly more depleted than values in lake waters (p < 0.001), ranging from -5.5 to -2.0‰ (Figure 2). δ^{15} N (NO₃) values in precipitation generally increased from north to south. For example, δ^{15} N (NO₃) values were -5.5‰ for Glacier, and increased to -2.0‰ continuing south toward Wolf Creek Pass, Colorado. There was a significant positive trend for the atmospheric δ^{15} N (NO₃) values to increase with increasing NO₃ concentrations ($r^2 = 0.7$; p < 0.01).

The occurrence of higher NO₃ concentrations and enriched δ^{15} N (NO₃) values in precipitation in National Parks characterized by higher NO₃ concentrations and enriched δ^{15} N (NO₃) values in lake waters suggests that atmospheric deposition of DIN in wetfall affects the amount of NO₃ in lakes. δ^{15} N (NO₃) values in both precipitation and lakes are more enriched in Colorado parks than in the northern parks in Wyoming and Montana. The correspondence of enriched

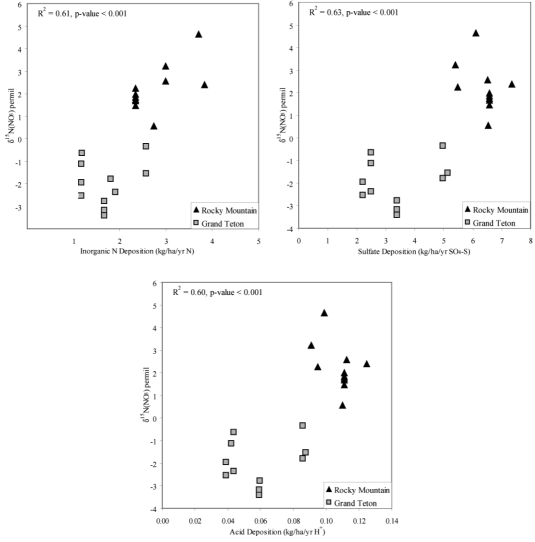


FIGURE 3. $\delta^{15}N$ (NO₃) values of lake water compared to average annual deposition of inorganic N (DIN), SO₄, and H⁺ in wetfall for two parks (estimates of wetfall from ref *31*).

 δ^{15} N with higher concentrations of NO₃ in precipitation and lakes suggests that areas with higher deposition are affected by a source of anthropogenic N emissions that is enriched in δ^{15} N. These results are similar to an earlier study in which significant correlations between δ^{15} N (NO₃) values in precipitation and stationary source NO_x emissions within source areas of 500–600 km in the eastern U.S. were attributed to regional transport of NO_x (10).

To evaluate this idea, isotopic values of δ^{15} N (NO₃) from lake water were compared with total stationary source NO_x emissions in USEPA Region 8 during 1990-1999, the longest recent period of record (7), as a proxy for total anthropogenic N emissions (2). Total stationary source NO_x emissions within a variety of buffer distances ranging from 50 to 600 km were tested, and 300 km was the most highly correlated with the data for all parks. Results for Colorado and Wyoming indicate that there is a significant positive correlation ($r^2 = 0.8$, p < 0.80.05) between δ^{15} N (NO₃) from lake waters and NO_x emissions within a 300 km buffer for Rocky, Grand Teton, Great Sand Dunes, and Yellowstone (Supporting Information Figure SI-4). For Glacier, there are no reported stationary sources of NO_x emissions within a 300 km buffer during 1990–1999 (7). As NO_x emissions increase, δ^{15} N (NO₃) values in NO₃ from lakes increase, suggesting a relationship between spatial variations in $\delta^{15}N$ across the Rocky Mountains and N emissions. This correlation suggests a contribution of regional anthropogenic N emission sources. This is particularly

apparent for National Parks in Colorado (Rocky and Great Sand Dunes) which are located near larger anthropogenic N emission sources compared to parks in northern Wyoming (Grand Teton and Yellowstone).

To further evaluate a potential connection between atmospheric deposition of pollutants and NO₃ in lake waters, lake δ^{15} N (NO₃) values were compared with average annual deposition estimates of DIN, SO₄, and H⁺ in wetfall (*31*) at Rocky and Grand Teton (Figure 3). Results indicate that lake water δ^{15} N (NO₃) values are significantly correlated (p < 0.001) with average annual deposition estimates of DIN ($r^2 = 0.61$), SO₄ ($r^2 = 0.63$), and H⁺($r^2 = 0.60$). This correlation is consistent with the spatial variability of δ^{15} N (NO₃) in lake waters being related to the atmospheric deposition of pollutants in wetfall (Figure 3).

Modeling results using 48 GIS attributes for each watershed show that there is a significant positive relation (p < 0.05) between elevation for lakes located at elevations greater than 2500 m and δ^{15} N (NO₃) values in lake water (Figure 4a). There was no significant relation between δ^{18} O (NO₃) values in lake waters and any GIS attribute. Geographic patterns of NO₃ concentrations of high-elevation lakes in the Rockies have previously been reported and results show particularly high NO₃ concentrations in Rocky (*35*). Here we show an increase in δ^{15} N (NO₃) values of lake waters with increasing elevation of the lakes (Figure 4a). There was also a trend for δ^{15} N (NO₃) values in precipitation to increase with increasing

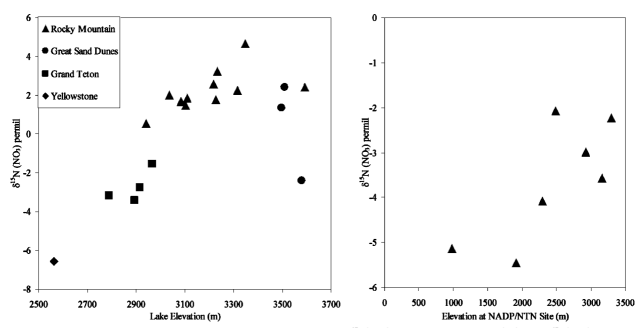


FIGURE 4. Lakes with elevation greater than 2500 m compared with $\delta^{15}N(NO_3)$ values in lake water (4a) and $\delta^{15}N(NO_3)$ values in NADP/NTN precipitation compared with elevation (4b).

elevation (p < 0.05) (Figure 4b). It is possible that the enriched δ^{15} N (NO₃) values in precipitation with increasing elevation may be influencing the NO₃ isotopic values collected from lake water.

The source of NO₃ in lake and stream waters from mountain catchments in the Sierra Nevada has previously been examined with a simple end-member mixing analysis using δ^{18} O (NO₃) values (*36*). Here we have direct measurements of the atmospheric end-member, where δ^{18} O (NO₃) values range from +71 to +78‰. The range in δ^{18} O (NO₃) values of lake water is from -5.7 to + 21.3 (Figure 2), with 95% of samples having δ^{18} O (NO₃) values less than +15‰ (Figure 2). However, the terrestrial end-member is difficult to parameterize without direct measurements of subsurface δ^{18} O (NO₃) values. The generally accepted upper limit for δ^{18} O of microbial NO₃ is +15‰ for the terrestrial end-member (*11*).

Several previous studies have evaluated N sources in streamwater draining undisturbed catchments using a dual NO₃ isotope approach ($\delta^{18}O$ (NO₃) and $\delta^{15}N$ (NO₃)) to differentiate watershed NO₃ sources (*18, 37, 38*). NO₃ in streamwater was found to be mainly derived from nitrification in the Catskill mountains of New York (*37*). Most of the NO₃ in streamflow was nitrified within two forested catchments in New Hampshire (*38*). Similarly, most of the streamwater NO₃ in Rocky had an isotope signature indicative of substantial biological cycling of atmospherically derived N prior to release from the ecosystem (*18*).

However, assuming a δ^{18} O (NO₃) value of less than +15‰ for a terrestrial source could underestimate the actual contribution of atmospheric NO₃ to the NO₃ in lake waters. Laboratory incubation experiments and field studies have shown that the δ^{18} O (NO₃) formed by microbial nitrification range between +2 and +14‰, assuming that soil–water δ^{18} O (NO₃) values vary between -15 and -5‰ (39). In the Catskill Mountains of New York, δ^{18} O (NO₃) values from +13.2 to +16.0‰ were measured for NO₃ derived by nitrification in incubated soil samples (37). In Sleepers River, a snowmeltdominated catchment in Vermont, δ^{18} O values of stream NO₃ ranged from -7.7 to +18.3‰ and generally were correlated with NO₃ concentrations (28). It was concluded that a significant amount of NO₃ during snowmelt was directly from atmospheric deposition of NO₃ (28). A Δ^{17} O in surface water did not undergo biologic processing before being exported from the system (40). Thus, it is possible that the direct atmospheric contribution to NO₃ may be underestimated in earlier reports. A quantitative source apportionment of atmospheric NO₃ to the NO₃ in lake waters is difficult because of the wide range of δ^{18} O values from microbial nitrification.

The trend toward increasing $\delta^{15}N$ (NO₃) values with increasing concentrations of NO₃ in lake waters, which in turn are associated with increasing elevation and increasing inputs of DIN in wetfall, is intriguing. One potential explanation that deserves additional study is that the more enriched values of δ^{15} N (NO₃) may result from increasing rates of net nitrification in the watersheds. Well-drained soils typically show an increase in total soil- δ^{15} N with increasing soil depth and age (11). This increase in δ^{15} N is attributed to fractionation during net mineralization and generally results from the metabolism of microbial heterotrophs that produce δ^{15} N-enriched biomass as a result of δ^{15} N-depleted waste (41, 42). DIN deposition in wetfall in the Rocky Mountains increases with increasing elevation (>2500 m) compared to lower elevations (<2500 m), due in part to orographically enhanced precipitation amounts at high elevations (2, 31). This increased DIN deposition in wetfall may lead to enhanced N cycling in high-elevation watersheds. The percent of DIN in wet deposition contributed by NH₄ is now approximately 50% of measured DIN in the Rocky Mountains (8). However, export of NH₄ in these watersheds is small, making up less than 6% of DIN compared to approximately 94% of DIN that is NO₃ in the baseflow lake outlet samples. This indicates that at least some of the NH4 in atmospheric deposition of DIN in wetfall is mineralized and nitrified to NO_3 that is exported to lake waters (17). Research in high-elevation areas of the Colorado Rockies show high rates of N-mineralization (43) and less NH₄ assimilation due to a lack of vegetation, particularly in talus areas (44). These areas tend to be carbon limited (45), driving systems toward net nitrification (43, 46). It is possible that high rates of DIN deposition in wetfall at high elevations in the Colorado Rockies (2, 3), characterized by enriched $\delta^{15}N$ (NO₃) values (10), may lead to enhanced nitrification and more enriched δ^{15} N values in the NO₃ exported to lake waters.

The role of denitrification in these systems was also considered, however, denitrification does not appear to substantially affect δ^{15} N (NO₃). If denitrification were important in these systems, a progression toward decreased NO₃ concentration would be expected. This trend does not exist in the data. Instead, the data show a significant correlation with increasing NO₃ concentrations and increasing δ^{15} N (NO₃) values in lake waters suggesting that enhanced nitrification may be important in these systems. Our results are consistent with previous work evaluating pathways for NO₃ release from an alpine watershed using δ^{18} O (NO₃) and δ^{15} N (NO₃) which found that denitrification does not affect fluxes of NO₃ from surface water or talus springs (*18*).

The results presented in this study suggest that relatively high anthropogenic emissions of NO_x may be contributing to high NO₃ concentrations in high-elevation lakes in the southern Rocky Mountains through a combination of direct and indirect processes such as enhanced nitrification. This study provides valuable information on spatial patterns in δ^{15} N (NO₃) in lakes and precipitation across the Rocky Mountains and has important implications as N emissions (stationary and mobile sources) and inorganic N deposition continue to increase into the future. Results of this study may be helpful to resource managers who are considering the best way to reduce N emissions to control inorganic N deposition in sensitive, protected areas.

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Supporting Information Available

Four additional figures including a national map of DIN deposition (Figure SI-1), a map of the study sites (Figure SI-2), nitrate concentrations at 56 lakes aggregated by National Park (Figure SI-3), and $\delta^{15}N$ (NO₃) values compared with NO_x emissions (Figure SI-4). Two additional tables including a table of NO₃ concentration, $\delta^{18}O$ (NO₃) and $\delta^{15}N$ (NO₃) values for National Park lakes (Table SI-1) and NADP/ NTN precipitation sites (Table SI-2). This information is available free of charge via the Internet at http://pubs.acs.org.

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