HIGHLIGHTED STUDENT RESEARCH



Soluble soil aluminum alters the relative uptake of mineral nitrogen forms by six mature temperate broadleaf tree species: possible implications for watershed nitrate retention

Mark B. Burnham¹ \bigcirc · Jonathan R. Cumming¹ · Mary Beth Adams² · William T. Peterjohn¹

Received: 22 March 2017 / Accepted: 3 September 2017 © Springer-Verlag GmbH Germany 2017

Abstract Increased availability of monomeric aluminum (Al^{3+}) in forest soils is an important adverse effect of acidic deposition that reduces root growth and inhibits nutrient uptake. There is evidence that Al^{3+} exposure interferes with NO₃⁻ uptake. If true for overstory trees, the reduction in stand demand for NO_3^- could increase NO_3^- discharge in stream water. These effects may also differ between species that tolerate different levels of soil acidity. To examine these ideas, we measured changes in relative uptake of NO_3^{-} and NH_4^{+} by six tree species in situ under increased soil Al³⁺ using a ¹⁵N-labeling technique, and measured soluble soil Al levels in a separate whole-watershed acidification experiment in the Fernow Experimental Forest (WV). When exposed to added Al^{3+} , the proportion of inorganic N acquired as NO₃⁻ dropped 14% across species, but we did not detect a reduction in overall N uptake, nor did tree species differ in this response. In the long-term acidification experiment, we found that soluble soil Al was mostly in the free Al³⁺ form, and the concentration of Al³⁺ was ~65 μ M higher (~250%) in the mineral soil of the acidified watershed vs. an untreated watershed. Thus, increased levels of soil

We found that soil solution Al reduces the uptake of NO_3 by mature trees. This effect and the impact on watershed NO_3 export are novel findings important in areas impacted by acid deposition.

Communicated by Jason P. Kaye.

Mark B. Burnham mburnha1@mix.wvu.edu

² USDA Forest Service Northern Research Station, Morgantown, WV, USA Al^{3+} under acidic deposition cause a reduction in uptake of NO_3^- by mature trees. When our ¹⁵N uptake results were applied to the watershed acidification experiment, they suggest that increased Al^{3+} exposure could reduce tree uptake of NO_3^- by 7.73 kg N ha⁻¹ year⁻¹, and thus increase watershed NO_3^- discharge.

Keywords Acid deposition \cdot Nitrogen cycle \cdot Nitrogen export \cdot Tree nutrition \cdot ¹⁵N tracer

Introduction

The eastern US has a history of elevated acid deposition. Emissions of SO₂ and NO_r from the combustion of fossil fuels in power plants in the Ohio River Basin and automobiles throughout the region have caused acidic deposition and elevated inputs of nitrogen (N) and sulfur (S) during the late-20th and early 21st centuries (Driscoll et al. 2001; Galloway et al. 2004). The increased deposition of these materials onto downwind ecosystems can increase soil acidity, especially in poorly buffered soils, and lead to a variety of adverse effects (Lovett et al. 2009). These effects include loss of base cations (i.e., Ca, Mg, etc.), altered plant mineral nutrition, reduced root growth, and reduced forest productivity. Through time, elevated supply of N could also exceed forest N demand and cause N saturation (Aber et al. 1998). Thus, acid deposition has the potential to significantly impact the biogeochemistry of temperate forest ecosystems through soil acidification and N saturation.

An increase in soil acidity typically causes higher solubility of monomeric aluminum (Al^{3+}) (de Vries et al. 2003). We define monomeric aluminum as Al^{3+} , but other studies sometimes include different inorganic complexes in surface soils, such as various oxides of Al. Some discrepancy in

¹ Department of Biology, West Virginia University, 53 Campus Drive, Morgantown, WV 26506, USA

plant responses to Al between studies could be caused, in part, by measurements of different forms of aluminum. We focused on Al³⁺ because of its increase in concentration at low pH and severe impact on plant roots. Root growth is severely reduced when exposed to Al³⁺ in solution (Delhaize and Ryan 1995; Poschenrieder et al. 2009), and while this alone can inhibit plant development, Al³⁺ also has a number of secondary effects on plant roots, including reduced water and nutrient uptake (Kochian 1995). The effects of Al^{3+} on plants have been studied extensively in the lab, and particularly on herbaceous plants and tree seedlings. However, its impact on plant growth in field conditions can be much more variable than in the lab. Al has relatively complex dissolution reactions in the soil that are dependent on the soil composition. Buffering by base cation release (i.e., calcium) (Monterroso et al. 1999; de Vries et al. 2003) and the formation of Al complexes with organic acids (Mulder and Stein 1994; Brumme et al. 2009) may lead to varying levels of free Al^{3+} species, and diverse effects, across a landscape (de Vries et al. 2003; Li and Johnson 2016). For example, Rosenberg and Butcher (2010) found no correlation between foliar and BaCl₂-extractable soil Al concentration for red spruce in acid forest soils. In addition, de Wit et al. (2010) found that 7 years of AlCl₃ addition to a Norway spruce forest did not impede root growth as seen in lab studies with seedlings of other tree species (e.g., Lux and Cumming 1999), but the additions did reduce foliar magnesium (Mg) concentration. Therefore, while soluble Al³⁺ in soil may not affect the growth of mature trees in the field to the degree suggested by laboratory studies, other aspects of their function may be altered, such as mineral nutrition.

Because of these potential negative effects, many plants reduce their exposure to Al^{3+} by altering the Al species present in the rhizosphere. When soil Al^{3+} increases, plant roots exude organic acids, such as citrate and malate, which chelate free Al^{3+} and reduce negative growth and nutritional effects (Delhaize and Ryan 1995; Kochian 1995). Thus, while bulk soil Al^{3+} may increase under acid deposition, its effect would be lower in the rhizosphere of Al-tolerant plants that exude chelating organic acids. This further complicates the potential biogeochemical effect of acid deposition-induced Al^{3+} solubility, and therefore, it is necessary to measure the chelation of Al^{3+} in rhizosphere and bulk soil to adequately assess its impact on stand-scale growth and nutrient cycling.

When soluble AI^{3+} increases in the soil, several negative effects on plants could translate to changes in N demand and thus an impact on the N biogeochemistry of forest catchments. Should soil AI^{3+} rise to a level that reduces plant growth, overall N uptake by vegetation would be reduced, leading to elevated stream water N export. Even in the absence of a reduction in growth, the presence of soluble AI^{3+} can impede other aspects of tree nutrition that may alter N demand (de Wit et al. 2010). In particular, Al^{3+} exposure can reduce NO₃⁻ uptake by plants (Jarvis and Hatch 1986; Durieux et al. 1993; Calba and Jaillard 1997; Watanabe et al. 1998; Jerzykiewicz 2001; Pal'ove-Balang and Mistrík 2007; Zhou et al. 2016). While the exact mechanism is not established, soluble soil Al^{3+} can interfere with cell membrane H⁺-ATPase activity, reducing the cell's capacity to pump out H^+ (Zhou et al. 2016). This would strongly reduce the cell's ability to transport NO₃⁻ across the cell membrane, since NO₃⁻ cotransporters require 2H⁺ per NO₃⁻ moved (Britto and Kronzucker 2005). Thus, the result could be a shift in relative uptake of mineral N forms, towards greater uptake of NH₄⁺ and reduced uptake of NO_3^- (Cumming 1990). Since NO_3^- is highly mobile in soils, any reduction in the uptake of NO_3^{-1} induced by higher levels of Al^{3+} has the potential to increase stream water NO₃⁻ export.

Increased N supply by acid deposition could cause elevated NO₃⁻ in stream water due to N saturation, and an Al³⁺-mediated decrease in stand NO₃⁻ demand would compound this effect. In a long-term, whole-watershed fertilization/acidification experiment at the Fernow Experimental Forest, N added as (NH₄)₂SO₄ has caused a persistent reduction in the pH, and increase in the stream water NO₃⁻ concentration and discharge (Fig. 1; Adams et al. 1997; Edwards et al. 2006). While an initial increase in net nitrification was measured in the fertilized watershed relative to the reference watershed, more recent in situ and lab estimates of net nitrification rates in the upper 10 cm of mineral soil, collected at 100 points within each watershed, were unable to detect any difference in the net rate of NO₃⁻ production (Gilliam and Peterjohn, unpublished data), despite the persistence of elevated NO_3^{-} concentration in stream water leaving the fertilized watershed (Fig. 1). This suggests that elevated NO_3^{-1} loss from the fertilized/acidified watershed may be influenced by a decrease in NO_3^- demand, potentially due to elevated Al^{3+} in the soil under acidified conditions. Therefore, the main objectives of this study were to determine if tree roots are exposed to higher levels of free, unchelated Al³⁺ under experimental soil acidification, if this exposure could change the relative uptake of different forms of mineral N by important tree species in situ, and to provide an initial assessment of the potential impact that any such change might have on stream water NO₃⁻ export from a forested watershed. We hypothesized that (1) an increase in tree root exposure to soluble Al³⁺ would shift the relative uptake of mineral N away from NO_3^- and towards NH_4^+ due to the hindrance of NO_3^- uptake pathways, (2) that species would vary in their sensitivity, with species that are more tolerant of acidic soils, such as Acer rubrum and Quercus rubra, being less affected by increased levels of soluble $A1^{3+}$, and (3) that soil acidification causes levels of Fig. 1 Annual stream water NO_3^- discharge from the acidified (WS3) and reference (WS7) watersheds. Vertical dashed line indicates the start of the annual addition of 35 kg N ha⁻¹ year⁻¹ as $(NH_4)_2SO_4$ to the acidified watershed. Three-year running averages are displayed to better visualize temporal trends in the data. Only years with values for all months were included for a given watershed



soluble Al^{3+} that have the potential to elevate stream water NO_3^{-} discharge from a watershed if N uptake by most of the species present were Al sensitive.

Materials and methods

Site description

This research was conducted in the Fernow Experimental Forest (FEF) in Tucker County, WV, USA. This site is a mixed hardwood forest, and the soil is primarily a Calvin channery silt loam (loamy-skeletal, mixed, mesic Typic Dystrochrept). Elevation ranges from 762 to 854 m, and average annual precipitation totals ~145 cm (Kochenderfer 2006). To test if Al^{3+} affects the relative uptake of NO_3^{-} vs. NH_4^{+} , we used an area of the FEF with no assigned long-term treatment, to avoid affecting the $\delta^{15}N$ of the experimental areas. The area was last used in the 1980s when 0.2 ha plots were harvested to varying levels of basal area. However, we selected mature canopy trees that were similarly sized to those in the nearby acidified watershed (<1 km away), and we avoided areas with signs of harvest. To assess the potential effects of acidification on plant available Al in the soil, we used the long-term watershed acidification experiment at the FEF. This is a paired watershed experiment consisting of two adjacent watersheds-an acidified 34-ha watershed (WS 3, 1883 tree stems ha^{-1}) and a similarly aged, 24-ha reference watershed (WS 7, 1473 tree stems ha⁻¹) (Kochenderfer 2006). The forest on the acidified watershed is currently dominated by *Prunus serotina* (52% of the total basal area), A. rubrum (10.9%), Betula lenta (7.2%), and Liriodendron

tulipifera (6.4%). In 1969–1970, the watershed was clearcut, and then allowed to naturally regrow thereafter. To experimentally acidify the soils in WS 3, 35 kg N ha⁻¹ year⁻¹ of $(NH_4)_2SO_4$ have been aerially applied in three doses per year since 1989. The reference watershed (WS 7) is currently dominated by P. serotina (29.4% of the total basal area), B. lenta (19.1%), L. tulipifera (17.9%), Acer saccharum (11.3%), A. rubrum (6%), and Q. rubra (4%). This watershed was clearcut between 1963-1964 and 1966-1967 (lower half, then upper half) (Patric and Reinhart 1971). The reference watershed has never received additions of (NH₄)₂SO₄. In 2011, after 21 years of treatment, the pH of the top 10 cm of mineral soil was significantly lower in the acidified watershed than in the reference watershed (pH 4.2 vs. 4.6), and the extractable soil Al (extracted with 1 N ammonium acetate) was significantly higher in the acidified watershed than in the reference watershed (0.45 ± 0.03 vs. $0.32 \pm 0.01 \text{ meq } 100 \text{ g}^{-1}$; Peterjohn, unpublished data).

Relative uptake of NO₃⁻ and NH₄⁺

In the early July of 2014, we used an in situ ¹⁵N-labeling method to determine the relative uptake of NO_3^- and NH_4^+ by mature overstory trees (McKane et al. 2002; Andresen and Michelsen 2005). NO_3^- and NH_4^+ pools under canopy trees were labeled with sub-fertilization amounts of either ¹⁵NH₄Cl or K¹⁵NO₃. Five canopy trees of six important species found in WS 3 and WS 7 (*A. saccharum, A. rubrum, B. lenta, L. tulipifera, P. serotina,* and *Q. rubra*) were selected from an area adjacent to the experimental watersheds in the FEF to avoid labeling the natural ¹⁵N pool in the soils of the long-term experimental areas. Under each

tree's canopy, and within 4 m of the stem, four 625-cm² plots were used for the injection of labeled N solutions. One of the four solutions was applied to each plot: (1) $^{15}NH_4Cl$; (2) $^{15}NH_4Cl + Al^{3+}$; (3) K $^{15}NO_3$; and (4) K $^{15}NO_3 + Al^{3+}$. The N concentrations in each treatment solution were 3.5 mM. The past measurements of lysimeter soil water Al³⁺ from the acidified watershed yielded concentrations from zero to nearly 600 μ M (Lux 1999). We used 600 μ M Al³⁺ in our treatment solutions assess the potential of $A1^{3+}$ to impact tree N form uptake. Since some added Al³⁺ would rapidly associate with exchange sites on soil particles, the resulting Al³⁺ concentration in solution was in the range of measured lysimeter values, up to 600 μ M. Al³⁺ was added as $Al_2(SO_4)_3$, and all solutions were acidified to pH 4.0-4.5 using HCl, to best match the soil pH. Each plot consisted of a 100-hole grid frame (10×10 hole commercial pegboard and 2.54 cm between holes) laid on the ground to guide the injection of labeled N solutions. At each hole, 1 mL of N solution was injected at a depth of 3 cm (approximately the top of the A horizon) using a side-port syringe needle for a total of 52.5 mg ¹⁵N added to each plot.

After 3 h, a sample of fine roots (<2 mm diameter) of the nearby canopy tree was removed from a depth of ~3 cm. In addition to the ¹⁵N-labeled plots, we collected roots from an unlabeled area around each tree for measurement of root ¹⁵N natural abundance. To maximize our confidence that the roots were from the intended tree, the roots were traced as far as possible towards the canopy tree. In addition, we compared the morphology of the collected roots to the fine roots of nearby seedlings of the same species. Four of the species had distinct root characteristics; however, the roots of the two Acer spp. were very similar. Thus, we selected A. saccharum trees that had no nearby A. rubrum trees within ~15 m, and vice versa. We placed all collected roots on ice and transported them to the lab, where they were immediately placed in 1 M CaSO₄ for 1 min to remove unassimilated nutrients from the Donnan free space (Thornton et al. 2007). This was done to isolate the signal to N that had been transported across a cell membrane, and remove N that was passively present in the root apoplast. This may be a low amount of N, but even a small amount could greatly influence the results when working with highly δ^{15} N-enriched solutions. Root samples were then dried at 65 °C for 48 h, and then ground in a dental amalgamator (Henry Schein, Inc., Melville, NY, USA). From each plot, powdered root samples (~5 mg each) were wrapped in tin capsules and analyzed for tissue δ^{15} N and N content (% N) by the Central Appalachian Stable Isotope Facility at the Appalachian Laboratory of the University of Maryland (Frostburg, MD, USA).

Since the δ values of the labeled samples were highly enriched, we converted δ^{15} N values to R_{sample} , the ratio of

 15 N to 14 N in the root sample, and calculated the value of *F*, the fraction of the heavy isotope in the sample (Fry 2006):

$$R_{\text{sample}} = \left(\left(\frac{\delta^{15} N}{1000} \right) \times R_{\text{std}} \right) + R_{\text{std}},$$
$$F = \frac{R_{\text{sample}}}{1 + R_{\text{sample}}},$$

where $R_{\rm std} = {}^{15}{\rm N}/{}^{14}{\rm N}$ ratio in atmospheric N₂ (0.0036764). We then used the tissue N content, and *F* values to determine the µmol ${}^{15}{\rm N}$ g⁻¹ in root tissue. Finally, we estimated the rate of ${}^{15}{\rm N}$ taken up by roots from the labeled N pools by dividing the ${}^{15}{\rm N}$ excess (${}^{15}{\rm N}$ content of labeled–unlabeled roots from the same tree) by the exposure time (3 h). The total uptake rate of inorganic N from the labeled pools was the sum of our estimate of NO₃⁻ and NH₄⁺ uptake rates.

Soil Al determination

To determine the effect of whole-watershed acidification on both chelated and free monomeric soluble soil Al, we measured aqueous Al in organic and mineral soils from the two watersheds in the whole-watershed acidification study. We collected organic and mineral soil (top 15 cm) from ten plots in each watershed, combining four separate subsamples collected within each ~10-m radius plot into two composite samples-one for the organic and one for the mineral soil. In the lab, the mineral soils were further separated into mineral bulk soil and mineral rhizosphere soil. Any roots in the mineral soil were gently shaken to remove excess soil, and any soil remaining attached to the root was considered mineral rhizosphere soil. Due to the high density of roots in the organic horizon, this fraction was considered all rhizosphere soil. We sieved all soils through a 2-mm mesh and stored them at 4 °C. Soil moisture content was measured on a subsample from each soil by mass loss after drying for 48 h in a 65 °C oven. To measure total (chelated + monomeric) aqueous Al in soil solution, we used undried, fresh, sieved soil samples, combining 10 mL of distilled H₂O with 10 g of mineral soil, and 20 mL of H₂O with 10 g of organic soil. The goal of this procedure was to collect Al that is currently present in soil water close to the soil surface (top 15 cm). This region of soil has high fine root density (~57 g m⁻² in the O-horizon and ~ 230 g m⁻² in the top 15 cm of mineral soil in the acidified watershed; Carrara unpublished data). Our water addition diluted the existing soil water ~3:1 and allowed us to collect now-diluted soil solution after centrifugation. We chose to measure only the Al in soil water rather than using an ionic extractant to best estimate the Al that is delivered to the root surface via the soil solution. Thus, the Al values that we present are concentrations (μM) in

aqueous soil solution after accounting for the dilution factor using the initial soil moisture content, which is intended to be similar to what would be measured in lysimeters (Lux 1999; Edwards et al. 2002). All soils were shaken for 30 min and centrifuged for 5 min at $4400 \times g$, and then, the supernatant passed through a 0.45 µm filter. To separate free monomeric Al³⁺ from chelated Al in solution, we passed each sample through a Cleanert SCX cation exchange column (Bonna-Agela Technologies, Inc., Wilmington, DE, USA) to remove Al³⁺ from solution. The concentration of Al in the filtered and deionized (after the exchange column) solutions was then analyzed using a Varian SpectrAA 220FS graphite tube atomic absorption spectrometer (Varian, Inc., Palo Alto, CA, USA). The amount of chelated Al was subtracted from the total water-soluble Al to obtain the monomeric Al³⁺ content of each extract. Using soil moisture measurements for each sample, we adjusted the diluted Al values to the concentrations of the original soil water in each sample.

Statistical analyses

We used a complete three-way ANOVA and a Tukey's HSD post hoc analysis ($\alpha = 0.05$) to test for differences in soil Al between watersheds, soil fractions (organic, mineral bulk, and mineral rhizosphere), forms of Al (chelated vs.

unchelated), and to test all interactions between the three factors. We focused on the differences in unchelated Al^{3+} between watersheds in the mineral rhizosphere and organic horizon, since these soil fractions should best characterize the exposure of tree roots to potentially damaging Al^{3+} .

To determine if A1³⁺ addition affected total uptake of N from the 15 N-labeled pool (15 NH₄ uptake + 15 NO₃ uptake), we used a two-way ANOVA with ¹⁵N uptake as the response variable and species and Al³⁺ addition as factors. The residuals for the rates of N uptake from the labeled pool were nonnormal, so we natural log-transformed these data to fulfill the normality assumption of ANOVA. Thus, the reported rates of uptake of N from the ¹⁵N labeled pools are backtransformed means (\pm SE). To test for an effect of Al³⁺ on NO_3^- uptake, we used the NH_4^+ and NO_3^- uptake rates from the labeled pools for each tree to calculate the total ¹⁵N uptake from the labeled pools, as well as the percentage taken up as NO_3^{-} , both in the presence and absence of added Al³⁺. We then used a two-way ANOVA with a Tukey's HSD post hoc analysis ($\alpha = 0.05$) to determine the effects of A1³⁺ and species on the percentage of ¹⁵N uptake that was NO_3^{-} , and to test if the effect of Al^{3+} depended on species. The model included the effect of tree nested within species. To determine if any species took up significantly more NO_3^- than NH_4^+ without added Al^{3+} , or significantly



Fig. 2 Percent of ¹⁵N taken up from the labeled pools as NO_3^- in the presence or absence of added Al^{3+} for the six temperate broadleaf tree species, and averaged across all species (far right). Bars that do not share a like letter are significantly different (Tukey's HSD post

hoc analysis, $\alpha = 0.05$). No individual species percent ¹⁵N uptake as NO₃⁻ was significantly affected by Al³⁺ addition. Dotted line shows 50% threshold of ¹⁵N uptake as NO₃⁻ for visual comparison

less NO₃⁻ than NH₄⁺ with added Al³⁺, we performed onetailed *t* tests to determine if the contribution of NO₃⁻ to total uptake of N from the labeled pool was greater (no added Al³⁺) or less (added Al³⁺) than 50%.

Results

Relative uptake of NO₃⁻ and NH₄⁺

Across tree species, the total N uptake rate from the labeled pool (${}^{15}NH_4^+ + {}^{15}NO_3^-$) was 0.120 µmol ${}^{15}N$ g⁻¹ h⁻¹, which is similar to rates measured in prior studies from the ${}^{15}N$ pool (McKane et al. 1990). There was no significant effect

Table 1 Total soil solution Al (μ M) (monomeric Al³⁺ + chelated Al) in three soil fractions within the acidified and reference watersheds, and the percent of total Al that was chelated

	Acidified Mean (SE)	Reference Mean (SE)
Organic soil		
Total Al	46.0 (7.3) ^{bc}	76.9 (12.2) ^{ab}
Percent chelated	38% (5.1)	32% (3.1)
Mineral bulk soil		
Total Al	103.3 (11.1) ^a	29.9 (6.3) ^c
Percent chelated	16% (2.8)	23% (9.5)
Mineral rhizosphere soil		
Total Al	113.2 (9.4) ^a	41.8 (5.5) ^{bc}
Percent chelated	11% (2.2)	20% (4.0)

Total Al values that do not share a like letter are significantly different (Tukey's HSD post hoc analysis, $\alpha = 0.05$)

Fig. 3 Monomeric soil solution Al^{3+} (μM) in three soil fractions of the acidified and reference watersheds. Bars that do not share a like letter are significantly different (Tukey's HSD post hoc analysis, $\alpha = 0.05$)

of species or Al treatment on total uptake of N from the labeled pool, and the effect of Al did not depend on species. Among all species, 59% (±5.2%) of N from the labeled pool was taken up as NO_3^- (0.074 ± 0.02 µmol ¹⁵N g⁻¹ h⁻¹), and 41% as NH₄⁺ (0.046 \pm 0.05 µmol ¹⁵N g⁻¹ h⁻¹), in the absence of added Al³⁺, and these proportions were not significantly different between species. However, under added Al^{3+} , NO_3^{-} uptake from the labeled pool decreased to 44.6% (±5.0%) of total N uptake (0.065 \pm 0.03 µmol ¹⁵N g⁻¹ h⁻¹) (F = 4.38, P = 0.047) (Fig. 2), and NH₄ accounted for 55.4% of total N uptake from the labeled pool (0.094 \pm 0.03 μ m ol ${}^{15}N$ g⁻¹ h⁻¹). While the mean percent of N uptake as NO_3^- declined from >50% for all species without added Al^{3+} to <50% under added Al^{3+} , no individual species decline was significant. For A. rubrum, there was a trend towards NO_3^{-} uptake contributing >50% to total uptake of N from the labeled pool (t = 2.03, P = 0.056), but no other species' NO₃⁻ uptake significantly differed from 50% of total uptake of N, regardless of Al treatment.

Soil Al determination

The total soil solution Al (across all soil forms and fractions) was 77% higher in the acidified watershed than the reference, an increase of 37.9 μ M Al (SE 7.3, *F* = 5.19, *P* < 0.001). Total Al was higher in the fertilized watershed in both the mineral bulk (245%) and mineral rhizosphere (171%) soil fractions, whereas there was no significant difference in total Al in the organic horizon (Table 1). Within each soil fraction (organic and mineral), <50% of the total soil solution Al was chelated in both watersheds



(Table 1), and the percent chelated did not significantly differ between watersheds.

Monomeric soil solution Al^{3+} was 103% higher (36.9 µM) in the acidified watershed than the reference (Tukey's t = 6.12, P < 0.001), and within the different soil fractions, it was 64.1 µM higher (283%) (Tukey's t = 6.14, P < 0.001) in mineral bulk soil and 67.5 µM higher (203%) (Tukey's t = 6.47, P < 0.001) in mineral rhizosphere soil in the acidified watershed compared to the reference watershed (Fig. 3). In the organic soil, there was no significant difference in monomeric soil solution Al^{3+} between the watersheds, despite a high statistical power (>0.98) to detect a similar difference in this soil horizon as the bulk and rhizosphere mineral soils.

Discussion

In the absence of added Al³⁺ from the ¹⁵N label addition, we found little difference in relative uptake of NO₃⁻ vs. NH₄⁺ for six temperate tree species under field conditions, whereas many prior studies found that NH₄ is the dominant mineral N form utilized by tree species (Buchmann et al. 1995; Gessler et al. 1998; Kronzucker et al. 1997; Lovett and Mitchell 2004; Malagoli et al. 2000; McFarlane and Yanai 2006; Min et al. 2000; Rothstein et al. 1996; Socci and Templer 2011; Templer and Dawson 2004). Our study differs from most of these in two important ways. First, the studies that tend to show the highest relative uptake of NH_4^+ over NO_3^- used coniferous species, whereas we studied temperate deciduous species that have been exposed to decades of elevated atmospheric N deposition. For example, Buchmann et al. (1995) labeled the soil of a *Picea abies* plantation and estimated that uptake of ${}^{15}NH_4^+$ was between two and four times higher than ¹⁵NO₃⁻. Second, many previous studies placed live or excised roots directly into nutrient solutions containing one or both mineral N forms. While this is valuable when studying the physiology of N uptake at the root surface, the higher diffusional resistance of NH_4^+ vs. NO_3^- in soil results in a greater delivery of NO₃⁻ to the root surface under natural conditions (Chapman et al. 2012). As a result, nutrient solution studies may underestimate the relative contribution of NO₃⁻ to tree N nutrition under field conditions. Similarly, the use of excised roots severs the transpiration stream, which drives mass flow to the root surface and is an important factor in plant NO₃⁻ uptake (Oyewole et al. 2014). Under more natural conditions, NO₃⁻ is more mobile than NH_4^+ , and the movement of NO_3^- via diffusion to the root surface may lead to greater relative uptake of NO₃⁻ than can be measured using nutrient solutions (Fahey and Yavitt 2005). Indeed, the keystone species A. saccharum (sugar maple) may be a good example of how N uptake assessments under artificial conditions may be misleading. Sugar maples are typically thought to utilize NH_4^+ as the primary mineral N source (Lovett and Mitchell 2004), a conclusion supported by excised root (Rothstein et al. 1996; Eddy et al. 2008; Socci and Templer 2011), nutrient solution depletion (McFarlane and Yanai 2006; Socci and Templer 2011), and greenhouse seedling studies (Templer and Dawson 2004). However, when we measured the relative importance of NO₃⁻ uptake in situ, we found a much higher relative contribution of NO₃⁻ to total uptake of N for mature trees than was indicated by many previous studies. Furthermore, the only other study that measured uptake of NO₃⁻ by mature A. saccharum trees under nearly in situ conditions found significant uptake of NO₃⁻, and also found that A. saccharum took up NO_3^{-} at a higher rate than three other temperate broadleaf species (Fahey and Yavitt 2005). Therefore, we suggest that in situ ¹⁵N-labeling techniques may provide meaningful insight into the mineral N uptake dynamics of mature trees under natural conditions.

Our in situ findings of a reduction in the relative amount of NO₃⁻ uptake under Al exposure in our ¹⁵N labeling experiment support our first hypothesis, and these results generally agree with prior greenhouse- and laboratorybased studies on herbaceous and woody plants. NO₃⁻ uptake reductions in plants exposed to Al have been found in maize (Durieux et al. 1993; Calba and Jaillard 1997), cucumber (Jerzykiewicz 2001), barley (Watanabe et al. 1998), and Lotus (Pal'ove-Balang and Mistrík 2007), as well as in the tropical tree Melaleuca cajuputi (Watanabe et al. 1998) and coniferous tree Pinus rigida (Cumming 1990). In addition, Al had a greater impact on growth when P. rigida seedlings were grown with primarily NO_3^- vs. NH_4^+ or mixed N sources (Cumming and Weinstein 1990). Thus, our in situ measurement of this pulse effect on six important tree species suggests that acidic deposition has the potential to reduce stand NO₃⁻ demand in a temperate deciduous forest, at least short term, as Al³⁺ becomes soluble in the soil under field conditions. Should the Al³⁺ effect on NO₃⁻ uptake persist, reduced stand NO3⁻ demand would be sustained and impact longer term discharge of N.

Our experiment of Al^{3+} addition to ¹⁵N-labeled solutions was a pulse addition of Al^{3+} , which contrasts somewhat with the long-term effects of whole-watershed acidification. We altered the Al^{3+} concentration at the interface of the organic and mineral soil (3 cm depth), but we only measured a longterm acidification effect on Al^{3+} concentration in the mineral soil. Therefore, one assumption of our method was that Al^{3+} would similarly impact fine root uptake of NO_3^{-} in deeper mineral soil (up to 15 cm) as at the interface between mineral and organic soil. Pulses of Al^{3+} exposure could result from rain storms that increase soil moisture and mobilize Al^{3+} in acidic soils, leading to greater movement of Al^{3+} to the root surface via mass flow. The spike of Al^{3+} in soil solution caused by our experimental addition was similar to what we observed in the soil of the long-term acidified watershed, so the physiological responses of the trees may also be similar. However, it is unknown if trees acclimate to long-term Al³⁺ exposure, thus recovering their uptake of NO₃⁻ under more natural conditions. Some evidence from herbaceous plants suggests that the effect of Al³⁺ persists. Maize plants showed no signs of short-term acclimation to Al³⁺ after 8 h of exposure (Durieux et al. 1993), although they recovered rapidly once they were removed from Al³⁺ solutions. The uptake of NO_3^- by white clover was also affected by Al³⁺ over a period of 5 weeks (Jarvis and Hatch 1986). In trees, there is also evidence of long-term effects on growth and tissue Ca:Al ratios (Vanguelova et al. 2007), although the effects vary between methods and species. Phillips and Yanai (2004) added AlCl₃ to A. saccharum trees in the field for 2 years, and found that Al content in the rhizosphere was reduced relative to bulk soil, suggesting that Al leached from the rhizosphere due to increased organic acid efflux from tree roots. However, we did not find a decrease in soluble Al³⁺ in the rhizosphere soils of our paired watershed study, so the effects of Al³⁺ would not be relieved in this manner. While a reduction in the uptake of NO_3^- may lessen over time, prior evidence suggests that at least some effect of Al³⁺ persists while it remains in soil solution.

Surprisingly, contrary to our second hypothesis, our ¹⁵N labeling results suggest that the tree species we studied did not differ in the impact of Al³⁺ on percent of N uptake as NO_3^{-} . This contrasts with prior evidence of variable Al^{3+} sensitivity between species (Kochian 1995; Watanabe et al. 1998), including temperate deciduous trees (Halman et al. 2015). Since we collected roots 3 h after treatment application, we measured their initial response to added Al^{3+} . It is possible that some species would increase Al³⁺-resistance over a longer time period by, for example, increasing root efflux of organic acids to chelate rhizosphere Al^{3+} (Kochian 1995). In addition, our treatment levels of Al^{3+} were relatively low to mimic the measured increase in the soil of the acidified watershed. We estimate that our levels of added Al³⁺ (~50–100 μ M exposed to plant roots, or 2.16–4.32 mg Al m^{-2}) were approximately 2% of the treatment level of Halman et al. (2015) (182 mg Al m^{-2} year⁻¹), who also studied temperate forest trees. It is possible that the species reacted similarly because these levels were lower than the threshold for Al^{3+} response by sensitive species (Vanguelova et al. 2007). Thus, low levels of Al^{3+} in acidified soils can rapidly affect uptake of NO₃⁻ across dominant temperate tree species.

Measuring uptake of N in situ by isotopically labeling the available pool presents some significant challenges. First, the use of a labor intensive and higher cost ¹⁵N labeling method limited our sample size to five trees of each species. As a result, our ability to detect differences between species was likewise limited. We conducted an iterative post hoc power

analysis, following the methods of Sokal and Rohlf (1981), using α of 0.05. This revealed that our sample size led to a relatively low statistical power $(1 - \beta)$ of ~0.1 to detect a similar effect of Al^{3+} on uptake of NO_3^{-} as a percent of total uptake of N within species as we found across species. The sample size would need to be increased to 66 or greater, depending on species, to reach a statistical power of 0.8. Our results can still be applied to stands given the Al³⁺ effect among species, and further studies on the effects within species could vield interesting results. Second, assimilated N is moved away from the roots into the tree, and the rate at which this happens is difficult to estimate in situ. Given our relatively short time from ¹⁵N addition to root excavation (3 h), our estimated uptake rates should be close approximations of the actual uptake of ¹⁵N from the labeled pools. The movement of N from the roots into the tree could affect the measured proportion of uptake as NO_3^- vs. NH_4^+ if they have different residence times in the root tissue. The reduction of NO₃⁻ occurs mostly in leaves in temperate deciduous tree species (Tang et al. 2012), potentially minimizing this effect. However, differential movement of the two N forms out of root tissue could result in an underestimation of the relative contribution of NH_4^+ to overall N uptake if reduced NO₃⁻ is stored in roots. Finally, it is also difficult to measure total N uptake using an in situ labeling method in undisturbed soil. To do so, an accurate measurement of the ¹⁵N atom percent in the soil at the root surface after the label is added would be necessary. As such, we have presented our results as uptake of ¹⁵N from the labeled pool, rather than total uptake of N, and focused on the proportions taken up as the two different mineral N forms. With efforts to minimize these methodological concerns, our measurements of root uptake from undisturbed soil provide important advantages that should be considered when conducting research in situ.

Not surprisingly, in the whole-watershed acidification experiment, we found that soluble soil Al³⁺ increased under long-term treatment (since 1989), indicating that soil acidification causes an increase in monomeric Al^{3+} in the upper mineral soil. We found comparable levels of soluble soil Al³⁺ that we measured (87–101 μ M in bulk soil, Table 1) to prior measurements in lysimeter-collected soil water in the same watershed (107 µM; Lux 1999), which suggests that our aqueous extraction method yielded accurate measurements of actual soil solution Al³⁺. Monomeric Al³⁺ was elevated in both the bulk and rhizosphere mineral soils, so Al³⁺ directly impacts tree roots in the mineral soil. There was no significant difference in organic soil Al³⁺ between watersheds, yet the measured levels may still be high enough $(30-50 \ \mu\text{M})$ to affect root uptake of NO₃⁻ in this soil horizon. It is possible that long-term acidic deposition in the region caused these levels of Al³⁺ even in the reference watershed, as was seen by Lux (1999). In addition, additional soluble soil Al³⁺ under experimental acidification of

the treated watershed could readily associate with exchange sites on organic material, reducing the treatment's effect on Al^{3+} in soil solution. Since the organic horizon is an area of high root density, the fact that Al³⁺ did not increase in this horizon under experimental acidification could relieve some of the effect of Al^{3+} on root uptake of NO_3^{-} at the stand level. However, there were actually more roots m^{-2} in the top 15 cm of mineral soil than in the organic horizon in these watersheds (~57 g m⁻² in the O-horizon vs. ~230 g m⁻² in the mineral soil in the acidified watershed; Carrara unpublished data). As a result, our results still support the hypothesis that acidification increases Al³⁺ to levels that diminish the relative uptake of NO₃⁻, potentially impacting watershed NO₃⁻ dynamics. Furthermore, we did not detect a decrease in overall N uptake from the labeled pool under Al³⁺ treatment; instead, uptake remained stable, but the proportion of N taken up as NO₃⁻ decreased. This emphasizes that soluble soil Al³⁺ can impact the pool of mineral N used by overstory trees under long-term acidic deposition, and thus potentially increase NO_3^{-} discharge from the watershed.

As an initial assessment of the potential impact of Al on the export of NO₃⁻ in stream water at the scale of a small watershed, we applied the results of our ¹⁵N-labeling experiment to estimates of total N uptake by the trees growing in the acidified watershed at the FEF (WS 3). Under wholewatershed fertilization and acidification, the discharge of NO₃-N increased from 4.17 kg N ha⁻¹ year⁻¹ pre-fertilization (1982-1989) to 13.82 kg N ha⁻¹ year⁻¹ post-fertilization (1990-2009), an increase of 9.65 kg N ha⁻¹ year⁻¹ (Fig. 1). However, at the same time, there was no detectable difference in mineral soil net nitrification rate between the two watersheds (Gilliam and Peterjohn unpublished data). Therefore, reduced stand NO₃⁻ demand due to soil Al may contribute to the higher NO_3^- discharge in stream water in the acidified watershed. We estimated tree uptake of NO₃⁻ in the acidified watershed by multiplying an estimate of total N uptake (N return in leaf litter + aboveground woody N storage) (50.95 kg ha⁻¹ year⁻¹) by our ¹⁵N-label measurement of percent of uptake as NO₃⁻ both without and with added Al³⁺ (59 and 44.6% of total N uptake as NO₃⁻, respectively). Aboveground woody N storage was calculated by multiplying bole wood N content in the outer 1 cm by the annual stand woody biomass increase reported by DeWalle et al. (2006). The resulting difference between NO_3^- uptake without and with added Al^{3+} , an estimate of unassimilated, excess soil NO₃⁻ available for leaching due to the impact of Al³⁺, is 7.73 kg N ha⁻¹ year⁻¹. If this amount was completely discharged in stream water, the effect of increased Al³⁺ would account for up to 76% of the 9.65 kg N ha⁻¹ year⁻¹ increase in stream water NO_3^- due to whole-watershed acidification. Perhaps, more realistically, if ~70% of this unassimilated NO₃⁻ were retained in the watershed, as measured by Adams et al. (2006), then elevated Al^{3+}

would still cause 23% of the increase in stream water NO_3^- . While this initial estimate is specific to our study site, the potential magnitude of the effect of elevated soil Al^{3+} on watershed NO_3^- discharge is large enough to warrant more detailed assessments at a variety of locations.

Acknowledgements The authors thank Chris Walter, Joe Carara, Katie Sesa, Hannah Hedrick, Leah Baldinger, Rachel Arrick, Jessica Graham, and Hoff Lindberg for their field and lab work on this project. We also thank Dr. Robert McKane for his aid in developing 15 N labeling methods. We greatly appreciate Dr. James McGraw's input on statistical analyses. Finally, we acknowledge the contributions of the USDA Forest Service Fernow Experimental Forest staff for the long-term management and support of this experiment. This work was supported by the Long-Term Research in Environmental Biology (LTREB) program at the National Science Foundation (Grant Nos. DEB-0417678 and DEB-1019522) and the WVU Department of Biology and Eberly College of Arts and Sciences.

Author contributions MBB, JRC, and WTP conceived and designed the ¹⁵N-labeling experiment, MBA aided with site selection and soil analysis methodology for both the whole-watershed acidification study and the labeling experiment. MBB and WTP analyzed the data and wrote the manuscript; MBA and JRC reviewed the manuscript and provided editorial advice and comments.

References

- Aber J, McDowell W, Nadelhoffer K, Magill A, Bernston G, Kamakea M, McNulty S, Currie W, Rustad L, Fernandez I (1998) Nitrogen saturation in temperate forest ecosystems. Bioscience 48:921–934
- Adams MB, Angradi TR, Kochenderfer JN (1997) Stream water and soil solution responses to 5 years of nitrogen and sulfur additions at the Fernow Experimental Forest, West Virginia. For Ecol Manag 95:79–91
- Adams M, Peterjohn W, Gilliam F (2006) Acidification and Nutrient cycling. In: Adams M, DeWalle D, Hom J (eds) The Fernow watershed acidification study. Springer, Dordrecht, pp 207–236
- Andresen L, Michelsen A (2005) Off-season uptake of nitrogen in temperate heath vegetation. Oecologia 144:585–597
- Britto DT, Kronzucker HJ (2005) Nitrogen acquisition, PEP carboxylase, and cellular pH homeostasis: new views on old paradigms. Plant Cell Environ 28:1396–1409. doi:10.1111/j.1365-3040.2005.01372.x
- Brumme R, Meesenburg H, Bredemeier M, Jacobsen C, Schonfelder E, Meiwes K, Eichhorn J (2009) Changes in soil solution chemistry, seepage, losses, and input–output budgets at three beech forests in response to atmospheric depositions. In: Brumme R, Khanna P (eds) Functioning and management of European Beech Ecosystems. Berlin, p 499
- Buchmann N, Schultz E, Gebauer G (1995) ¹⁵N-ammonium and ¹⁵N-nitrate uptake of a 15-year-old *Picea abies* plantation. Oecologia 102:361–370
- Calba H, Jaillard B (1997) Effect of aluminum on ion uptake and H⁺ release by maize. New Phytol 137:607–616

- Chapman N, Miller AJ, Lindsey K, Whalley WR (2012) Roots, water, and nutrient acquisition: let's get physical. Trends Plant Sci 17:701–710. doi:10.1016/j.tplants.2012.08.001
- Cumming JR (1990) Nitrogen source effects on Al toxicity in nonmycorrhizal and mycorrhizal pitch pine (*Pinus rigida*) seedlings. II. Nitrate reduction and NO₃⁻ uptake. Can J Bot 68:2653–2659
- Cumming J, Weinstein L (1990) Nitrogen source effects on Al toxicity in nonmycorrhizal and mycorrhizal pitch pine (*Pinus rigida*) seedlings. I. Growth and nutrition. Can J Bot 68:2644–2652
- de Vries W, Reinds G, Vel E (2003) Intensive monitoring of forest ecosystems in Europe 2: atmospheric deposition and its impacts on soil solution chemistry. For Ecol Manag 174:97–115. doi:10.1016/ S0378-1127(02)00030-0
- de Wit HA, Eldhuset TD, Mulder J (2010) Dissolved Al reduces Mg uptake in Norway spruce forest: results from a long-term field manipulation experiment in Norway. For Ecol Manag 259:2072– 2082. doi:10.1016/j.foreco.2010.02.018
- Delhaize E, Ryan P (1995) Aluminum toxicity and tolerance in plants. Plant Physiol 107:315–321
- DeWalle D, Kochenderfer J, Adams M, Miller G, Gilliam F, Wood F, Odenwald-Clemens S, Sharpe W (2006) Vegetation and acidification. In: Adams M, DeWalle D, Hom J (eds) The Fernow watershed acidification study. Springer, Dordrecht, pp 137–188
- Driscoll CT, Lawrence GB, Bulger AJ, Butler TJ, Cronan CS, Eagar C, Lambert KF, Likens GE, Stoddard JL, Weathers KC (2001) Acidic deposition in the northeastern US: sources and inputs, ecosystems effects, and management strategies. Bioscience 51:180–198. doi:10.1641/0006-3568(2001)051[0180:ADITNU]2.0.CO;2
- Durieux R, Jackson W, Kamprath E, Moll R (1993) Inhibition of nitrate uptake by aluminum in maize. Plant Soil 151:97–104
- Eddy WC, Zak DR, Holmes WE, Pregitzer KS (2008) Chronic atmospheric NO₃⁻ deposition does not induce NO₃⁻ use by *Acer saccharum* Marsh. Ecosystems 11:469–477. doi:10.1007/ s10021-008-9134-3
- Edwards PJ, Kochenderfer JN, Coble DW, Adams MB (2002) Soil leachate responses during 10 years of induced whole-watershed acidification. Water Air Soil Pollut 140:99–118. doi:10.102 3/A:1020181800320
- Edwards PJ, Williard KWJ, Wood F, Sharpe WE (2006) Soil water and stream water chemical responses. In: Adams M, DeWalle D, Hom J (eds) The Fernow watershed acidification study. Springer, Dordrecht, pp 71–136
- Fahey TJ, Yavitt JB (2005) An in situ approach for measuring rootassociated respiration and nitrate uptake of forest trees. Plant Soil 272:125–131. doi:10.1007/s11104-004-4212-6

Fry B (2006) Stable isotope ecology, 1st edn. Springer, New York

- Galloway J, Dentener F, Capone D, Boyer E, Howarth R, Seitzinger S, Asner G, Cleveland C, Green P, Holland E, Karl D, Michaels A, Porter J, Townsend A, Vorosmarty C (2004) Nitrogen cycles: past, present, and future. Biogeochemistry 70:153–226
- Gessler A, Schneider S, Von Sengbusch D, Weber P, Haneman U, Huber C, Rothe A, Kreutzer K, Rennenberg H (1998) Field and laboratory experiments on net uptake of nitrate and ammonium by the roots of spruce (*Picea abies*) and beech (*Fagus sylvatica*) trees. New Phytol 138:275–285
- Halman JM, Schaberg PG, Hawley GJ, Hansen CF, Fahey TJ (2015) Differential impacts of calcium and aluminum treatments on sugar maple and American beech growth dynamics. Can J For Res 45:52–59. doi:10.1139/cjfr-2014-0250
- Jarvis SC, Hatch DJ (1986) The effects of low concentrations of aluminum on the growth and uptake of nitrate-N by white clover. Plant Soil 95:43–55
- Jerzykiewicz J (2001) Aluminum effect on nitrate assimilation in cucumber (*Cucumis sativus* L.) roots. Acta Physioligiae Plant 23:213–219

- Kochenderfer JN (2006) Fernow and the Appalachian Hardwood region. In: Adams M, DeWalle D, Hom J (eds) The Fernow watershed acidification study. Springer, Dordrecht, pp 17–39
- Kochian LV (1995) Cellular mechanisms of aluminum toxicity and resistance in plants. Annu Rev Plant Physiol Plant Mol Biol 46:237–260. doi:10.1146/annurev.pp.46.060195.001321
- Kronzucker HJ, Siddiqi MY, Glass ADM (1997) Conifer root discrimination against soil nitrate and the ecology of forest succession. Nature 385:59–61
- Li W, Johnson CE (2016) Relationships among pH, aluminum solubility and aluminum complexation with organic matter in acid forest soils of the Northeastern US. Geoderma 271:234–242. doi:10.1016/j.geoderma.2016.02.030
- Lovett G, Mitchell M (2004) Sugar maple and nitrogen cycling in the forests of Eastern North America. Front Ecol Environ 2:81–88
- Lovett GM, Tear TH, Evers DC, Findlay SEG, Cosby BJ, Dunscomb JK, Driscoll CT, Weathers KC (2009) Effects of air pollution on ecosystems and biological diversity in the eastern US. Ann N Y Acad Sci 1162:99–135. doi:10.1111/j.1749-6632.2009.04153.x
- Lux HB (1999) The effects of aluminum and nitrogen on mycorrhizal and non-mycorrhizal tulip poplar. MS Thesis, West Virginia University, Morgantown, WV, USA
- Lux HB, Cumming JR (1999) Effect of aluminum on the growth and nutrition of tulip-poplar seedlings. Can J For Res 29:2003–2007. doi:10.1139/cjfr-29-12-2003
- Malagoli M, Canal AD, Quaggiotti S, Pegoraro P, Bottacin A (2000) Differences in nitrate and ammonium uptake between Scots pine and European larch. Plant Soil 221:1–3
- McFarlane KJ, Yanai RD (2006) Measuring nitrogen and phosphorus uptake by intact roots of mature *Acer saccharum* Marsh., *Pinus resinosa* Ait., and *Picea abies* (L.) Karst. Plant Soil 279:163–172. doi:10.1007/s11104-005-0838-2
- McKane R, Grigal D, Russelle M (1990) Spatiotemporal differences in ¹⁵N uptake and the organization of an old-field plant community. Ecology 71:1126–1132
- McKane RB, Johnson LC, Shaver GR, Nadelhoffer KJ, Rastetter EB, Fry B, Giblin AE, Kielland K, Kwiatkowski BL, Laundre JA, Murray G (2002) Resource-based niches provide a basis for plant species diversity and dominance in arctic tundra. Nature 415:68– 71. doi:10.1038/415068a
- Min X, Siddiqi MY, Guy RD, Glass ADM, Kronzucker HJ (2000) A comparative kinetic analysis of nitrate and ammonium influx in two early-successional tree species of temperate and boreal forest ecosystems. Plant Cell Environ 23:321–328. doi:10.1046/j.1365-3040.2000.00546.x
- Monterroso C, Alvarez E, Fernandez-Marcos M, Macias F (1999) Geochemistry of aluminium and iron in mine soils from As Pontes, Galicia (N.W. Spain). Water Air Soil Pollut 110:81–102. doi:10. 1023/A:1005055704392
- Mulder J, Stein A (1994) The solubility of aluminum in acidic forest soils: long-term changes due to acid deposition. Geochim Cosmochim Acta 58:85–94. doi:10.1016/0016-7037(94)90448-0
- Oyewole OA, Inselsbacher E, Nasholm T (2014) Direct estimation of mass flow and diffusion of nitrogen compounds in solution and soil. New Phytol 201:1056–1064. doi:10.1111/nph.12553
- Pal'ove-Balang P, Mistrík I (2007) Impact of low pH and aluminium on nitrogen uptake and metabolism in roots of *Lotus japonicus*. Biologia (Bratisl) 62:715–719. doi:10.2478/s11756-007-0133-1
- Patric J, Reinhart K (1971) Hydrologic effects of deforesting two mountain watersheds in West Virginia. Water Resour Res 7:1182–1188
- Phillips RP, Yanai RD (2004) The effect of AlCl₃ additions on rhizosphere soil and fine root chemistry of sugar maple (*Acer saccharum*). Water Air Soil Pollut 159:339–356. doi:10.1023/B:W ATE.0000049187.35869.7d

- Poschenrieder C, Amenos M, Corrales I, Doncheva S, Barcelo J (2009) Root behavior in response to aluminum toxicity. In: Baluska F (ed) Plant–environment interactions. Springer, Berlin, p 313
- Rosenberg MB, Butcher DJ (2010) Investigation of acid deposition effects on southern Appalachian red spruce (*Picea rubens*) by determination of calcium, magnesium, and aluminum in foliage and surrounding soil using ICP-OES. Instrum Sci Technol 38:341–358. doi:10.1080/10739149.2010.508968
- Rothstein DE, Zak DR, Pregitzer KS, Url S, Zak R (1996) Nitrate deposition in northern hardwood forests and the nitrogen metabolism of *Acer saccharum* marsh. Oecologia 108:338–344
- Socci AM, Templer PH (2011) Temporal patterns of inorganic nitrogen uptake by mature sugar maple (*Acer saccharum* Marsh.) and red spruce (*Picea rubens* Sarg.) trees using two common approaches. Plant Ecol Divers 4:141–152. doi:10.1080/17550874.2011.6245 57
- Sokal RR, Rohlf FJ (1981) Biometry, 2nd edn. W. H. Freeman and Company, New York
- Tang MH, Porder S, Lovett GM (2012) Species differences in nitrate reductase activity are unaffected by nitrogen enrichment in northeastern US forests. For Ecol Manag 275:52–59. doi:10.1016/j. foreco.2012.03.006
- Templer PH, Dawson TE (2004) Nitrogen uptake by four tree species of the Catskill Mountains, New York: implications for

forest N dynamics. Plant Soil 262:251–261. doi:10.1023/B:P LSO.0000037047.16616.98

- Thornton B, Osborne SM, Paterson E, Cash P (2007) A proteomic and targeted metabolomic approach to investigate change in *Lolium perenne* roots when challenged with glycine. J Exp Bot 58:1581–1590. doi:10.1093/jxb/erl294
- Vanguelova EI, Hirano Y, Eldhuset TD, Sas-Paszt L, Bakker MR, Puttsepp U, Brunner I, Lohmus K, Godbold D (2007) Tree fine root Ca/Al molar ratio—indicator of Al and acidity stress. Plant Biosyst 141:460–480. doi:10.1080/11263500701626192
- Watanabe T, Osaki M, Tadano T (1998) Effects of nitrogen source and aluminum on growth of tropical tree seedlings adapted to low pH soils. Soil Sci Plant Nutr 44:655–666. doi:10.1080/00380768.19 98.10414489
- Zhou X, Gu Z, Xu H, Chen L, Tao G, Yu Y, Li K (2016) The effects of exogenous ascorbic acid on the mechanism of physiological and biochemical responses to nitrate uptake in two rice cultivars (*Oryza sativa* L.) under aluminum stress. J Plant Growth Regul 35:1013–1024. doi:10.1007/s00344-016-9599-9