Soil warming alters nitrogen cycling in a New England forest: implications for ecosystem function and structure

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Abstract Global climate change is expected to affect terrestrial ecosystems in a variety of ways. Some of the more well-studied effects include the biogeochemical feedbacks to the climate system that can either increase or decrease the atmospheric load of greenhouse gases such as carbon dioxide and nitrous oxide. Less well-studied are the effects of climate change on the linkages between soil and plant processes. Here, we report the effects of soil warming on these linkages observed in a large field manipulation of a deciduous forest in southern New England, USA, where soil was continuously warmed 5°C above ambient for 7 years. Over this period, we have observed significant changes to the nitrogen cycle that have the potential to affect tree species composition in the long term. Since the start of the experiment, we have documented a 45% average annual increase in net nitrogen mineralization and a three-fold increase in nitrification such that in years 5 through 7, 25% of the nitrogen mineralized is then nitrified. The warming-induced increase of available nitrogen resulted in increases in the foliar nitrogen content and the relative growth rate of trees in the warmed area. Acer rubrum (red maple) trees have responded the most after 7 years of warming, with the greatest increases in both foliar nitrogen content and relative growth rates. Our study suggests that considering species-specific responses to increases in nitrogen availability and changes in nitrogen form is important in predicting future forest composition and feedbacks to the climate system.

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Keywords  Soil warming · Net nitrogen mineralization · Nitrification · Species-specific nutrient acquisition strategies

Introduction

Climate change is predicted to play a significant role in altering the function and structure of forest ecosystems (Melillo et al. 1990). Many studies have addressed how changes in major processes such as photosynthesis and plant and microbial respiration are projected to affect ecosystem carbon (C) balance (e.g., Melillo et al. 1993, 2002, 2011; Ryan 1991; Ainsworth and Long 2005), water balance (Zavaleta et al. 2003), and cycling rates of key nutrients including nitrogen (N) and phosphorus (P) (Van Cleve et al. 1990; Peterjohn et al. 1994; Rustad et al. 2001; Melillo et al. 2002). A growing number of studies have examined how these functional changes within ecosystems can feedback to the climate system (Peterjohn et al. 1994; Shaver et al. 2000; Field et al. 2007; Melillo et al. 2011).

Direct and indirect feedbacks to the climate system as a result of warming are predicted to occur, with the direction (positive or negative) and magnitude being dependent on specific ecosystem conditions such as nutrient availability, moisture regime and species composition (Shaver et al. 2000; Field et al. 2007). In addition, ecosystem responses can vary over time, as organisms acclimate to changes in the environment or as community shifts in composition. For example, increased microbial activity, a characteristic response to warming, can further drive climate change by accelerating the decay of soil organic matter and thereby releasing carbon dioxide (CO₂), a greenhouse gas, to the atmosphere (Peterjohn et al. 1994; Rustad and Fernandez 1998; Shaver et al. 2000; Luo et al. 2001; Rustad et al. 2001; Melillo et al. 2002, 2011; Eliasson et al. 2005). This response to warming is reported to decrease over time as labile carbon is depleted or as the microbial communities adapt to warmer conditions (Melillo et al. 2002, 2011; Bradford et al. 2008). However, over the long term, it is unclear if warming will cause additional changes in ecosystems, further altering the magnitude or direction of the microbial community’s response to climate change. Varying and interacting responses of ecosystem processes to warming make it difficult to predict the overall scale and direction of long-term feedbacks to the climate system.

A number of climate change experiments in forests (e.g., Peterjohn et al. 1994; Rustad et al. 2001) have also shown that soil warming increases net N mineralization, the transformation of organically bound N to ammonium (NH₄⁺) and nitrification, the transformation of NH₄⁺ to nitrate (NO₃⁻). With increases in nitrification, ecosystems could experience gaseous and solution losses of N, potentially affecting water quality and creating a positive feedback to the climate system (Aber et al. 1989).

In an ecosystem where plant growth is limited by N availability, an increase in N has the potential to enhance photosynthetic rates and carbon storage in trees (Melillo et al. 2002, 2011). This can happen through increases N deposition in precipitation (Melillo and Gosz 1983; Thomas et al. 2009). Increased N availability to plants can also occur in response to soil warming (Melillo et al. 1995, 2002, 2011) as N is moved from the soil where the C:N mass ratio is often less than 30:1, to the plants where the C:N mass ratio in woody tissue is 200–300:1 (Melillo et al. 2002, 2011). Warming responses that cause enhanced carbon storage in woody tissue produce negative feedbacks to the climate system, which consequently slow the rate of atmospheric CO₂ accumulations (Melillo et al. 2011).

Changes in N cycling in response to warming are also likely to have long-term consequences for plant community structure. Tree species vary in their abilities to acquire and retain N based on characteristics such as root morphology and physiology (Marschner and Dell 1994), fungal associations (Lambers et al. 2008), preferred form of N uptake (NO₃ vs. NH₄ vs. amino acids; Schimel and Bennett 2004; Finzi and Berthrong 2005) and N resorption efficiency (Killingbeck 1996; Kobe et al. 2005). As human-induced environmental changes continue to affect forest ecosystems, species-specific strategies and responses to changes in the N cycle may become increasingly important in determining plant–soil interactions, forest species composition and the associated long-term feedbacks to the climate system.

Models that project the redistribution of tree species in response to climate change have often used a “climate envelope” approach, with species tracking shifts in key climate parameters including temperature and moisture over space and time (e.g., VEMAP Members 1995; Iverson and Prasad 1998; Iverson et al. 2008; Mohan et al. 2009). Climate-driven changes in the nitrogen cycle have not routinely been considered in vegetation redistribution models. A recent review by Ostle et al. (2009) argues that this may be a serious omission and urges that this potential shortcoming be addressed in new modeling efforts.

Here, we report the results of a large soil warming study at the Harvard Forest in central Massachusetts, USA, designed to explore the complex links among climate change, ecosystem biogeochemistry and plant responses. In a mixed deciduous stand, we have increased soil temperature 5°C using buried resistance cables (Melillo et al. 2002, 2011). For 7 years, we measured biogeochemical and plant responses in 900-m² heated and control (unwarmed) areas to examine how a temperate forest ecosystem is affected by warming-induced changes in the N cycle. Our measurements have included in situ net N
mineralization and nitrification, soil–water N concentrations, nitrous oxide (N₂O) emissions from the soil surface to the atmosphere, N concentration in green leaves and leaf litter, nitrate reductase activity in leaves of dominant species, and growth responses of the canopy trees. By examining how warming influences various fluxes and pools of N within the ecosystem, we document some short- and longer-term effects on ecosystem structure and function.

Materials and methods

Study site

The research site was an even-aged, mixed deciduous forest in central Massachusetts, USA (42°28′N, 72°10′W). Historical records, stonewalls and soil-horizon characteristics indicate that the area was used for either pastureland or low-intensity agriculture prior to 1908. White pines dominated the site by the early 1900s, but were destroyed in the 1938 hurricane. Blow downs were salvaged and the area was left to regrow naturally to its current state: a mid-aged stand of mixed hardwoods dominated by red oaks (Quercus rubra). Soils are mainly Canton series (coarse-loamy over sandy or sandy-skeletal, mixed, semi-active, mesic Typic Dystrudepts) with a surface pH of 5.2, and subsurface pH of 5.5. The average bulk density is 0.37 g cm⁻³ in the organic layer and 0.78 g cm⁻³ in the mineral layer. The climate is cool, temperate and humid. The mean weekly air temperature varies from a high of about 20°C in July to a low of about −6°C in January. Precipitation is distributed evenly throughout the year and averages about 108 cm annually.

During the summer and fall of 2001, about 5 km of heating cable were installed by hand to minimize disturbance in a 30 × 30 m area (Melillo et al. 2011). Cables were buried at 10 cm depth, spaced 20 cm apart. An adjacent 900-m² area was delineated to serve as the control area, with a 5-m buffer inbetween the areas. Results from our previous nearby soil-warming experiment using 5-m² plots confirmed that the soil disturbance associated with the installation of heating cables had no effect on soil temperatures and only minor and variable impacts on soil moisture and growth of small-statured woody vegetation (Melillo et al. 2002). Pre-treatment biogeochemical data were collected during the 2002 growing season and heating commenced in May 2003.

When supplied with 240 VAC, the heating cables have a power output of 13.6 W m⁻¹ and produce a power density of about 77 W m⁻². Within the heated area, there were 160 resistance cables, each of which is approximately 30 m long. A typical resistance per cable was 96 Ω. The heated area was monitored with 80 thermistors installed at 5 cm depth and 6 moisture probes at 10 cm depth, while the control area had 12 thermistors and 6 moisture probes at analogous depths. Each minute, heating cables were turned on or off automatically to maintain a 5°C temperature differential between heated and control areas.

Due to the scale and cost of the experiment, plot-level replication was not possible. Pre-treatment data were collected during the 2002 growing season to compare the baseline biogeochemistry for the two large areas, control and heated. Our pre-treatment measurements indicated that the control and heated areas were structurally and functionally similar prior to initiating treatment in the heated area (Melillo et al. 2011). In 2002, the tree biomass in the two areas was similar (202 Mt ha⁻¹ in the control area and 192 Mt ha⁻¹ in the heated area). Woody increment in the pre-treatment year was 1.73 Mt C ha⁻¹ in the control area and 1.69 Mt C ha⁻¹ in the heated area. N mineralization in the two areas was also similar during the pre-treatment year as determined by in situ incubations (method described below).

Net N mineralization and nitrification

Using the in situ buried bag incubation method (Eno 1960; Melillo 1977), we measured the rates of net N mineralization and nitrification in plots in the heated and control areas. We divided each area into 36 plots, each 5 × 5 m, and randomly chose 10 plots per treatment for the N mineralization and nitrification study. For each sampling, we took two 10 cm-deep soil cores from each plot in each treatment. One core, designated as the “initial” sample, was split into organic and mineral horizons and each horizon was placed into a gas-permeable polyethylene bag and transported to the laboratory for analysis. The second core, designated as the “final” sample, was placed intact inside another gas-permeable polyethylene bag and positioned back in the ground. The final bags were incubated for 5 weeks April–November and for 5 months over the winter and then harvested and transported to the laboratory for analysis.

In the laboratory, soils were separated into organic and mineral horizons and sieved through a 5.6-mm screen to remove rocks and roots. A subsample of the soil was weighed and dried at 105°C for 24 h for soil moisture analyses. Approximately 10 g of soil were placed in 100 mL of 2 M KCl, extracted for 36 h and filtered. The extracts were analyzed for nitrite/nitrate (NO₂⁻ + NO₃⁻ - N) and ammonium (NH₄⁺ - N) using a Lachat QuikChem FIA + 8000 Series Flow Injection Analyzer with Omnion 3.0 software. Using these techniques, the detection limit for NO₂⁻ + NO₃⁻ - N was 0.02 mg L⁻¹ and the detection limit for NH₄⁺ - N was 0.005 mg L⁻¹.
For each sample, extractable \( \text{NO}_2^- + \text{NO}_3^- - \text{N} \) and \( \text{NH}_4^+ - \text{N} \) were scaled to kg N ha\(^{-1}\) using sample mass, moisture content and bulk density. Net mineralization rates were calculated as final minus initial extractable \( \text{NO}_2^- + \text{NO}_3^- - \text{N} + \text{NH}_4^+ - \text{N} \) divided by incubation length. Net nitrification rates were calculated as final minus initial extractable \( \text{NO}_2^- + \text{NO}_3^- - \text{N} \) divided by incubation length.

Soil solution chemistry

In the fall of 2001, a total of 14 porous cup tension lysimeters were installed at a 50 cm depth (7 control, 7 heated). Lysimeter collections began in June 2002, approximately 8 months after installation, in order to minimize the effects of soil disturbance caused by lysimeter installation. Soil solution samples were collected once a month by draining any existing water from lysimeters using a syringe, and then applying a tension of 0.05 MPa with a hand pump. Approximately 24 h later, samples were collected, filtered, placed into scintillation vials and frozen for later analysis of \( \text{NO}_2^- + \text{NO}_3^- - \text{N} \) and \( \text{NH}_4^+ - \text{N} \).

Filtered samples were analyzed for \( \text{NO}_2^- + \text{NO}_3^- - \text{N} \) and \( \text{NH}_4^+ - \text{N} \) using a Lachat QuikChem FIA + 8000 Series Flow Injection Analyzer with Omnion 3.0 software. Using these techniques, the detection limit for \( \text{NO}_2^- + \text{NO}_3^- - \text{N} \) was 0.02 mg L\(^{-1}\) and the detection limit for \( \text{NH}_4^+ - \text{N} \) was 0.005 mg L\(^{-1}\).

\( \text{N}_2\text{O} \) soil efflux

Nitrous oxide was measured monthly April through November 2002–2007 using head-space air extracted with syringes from closed chambers of permanently installed gas collars and analyzed using electron capture gas chromatography similar to the procedure used by Bowden et al. (1990). On each sampling date, fluxes were measured in the afternoon and then analyzed using a Shimadzu model GC8A gas chromatograph (GC) equipped with a 3-m \( \times \) 3.2-mm-diameter stainless steel column filled with Poropak Q (80/100 mesh) and a 63Ni detector. Carrier gas was \( \text{N}_2 \) at 20 mL min\(^{-1}\). Column and detector temperatures were 65 and 310°C, respectively. Standards and samples were transferred into the GC using a 14-port valve (Valco Instruments Incorporated) equipped with a 1-mL sample loop. A 20-mm \( \times \) 4-mm-ID Silica gel (0.5 g, 80/100 mesh) water trap attached to the inlet of the injection valve was used to reduce water vapor carried into the GC (Bowden et al. 1990). Certified \( \text{N}_2\text{O} \) standards of 310 and 999 ppbv (Scott Specialty Gases, Plumsteadville, PA, USA) were used for calibration. Bowden et al. (1990) showed that standard curves are linear over this range and we found little variation throughout the day.

\( \text{Foliar N} \)

Green foliage was collected once per year from 2003 to 2009 during July in the heated and control areas. Sun leaves were collected from red oak (\textit{Quercus rubra}), red maple (\textit{Acer rubrum}), white ash (\textit{Fraxinus Americana}), sugar maple (\textit{Acer saccharum}), hemlock (\textit{Tsuga canadensis}) and birch (\textit{Betula lenta}) were shot down from the canopy and dried at 60°C for 24 h and ground using a Wiley Mill. A subsample of the leaf litter collected each fall was ground using a Wiley Mill in preparation for analysis. Both green and brown leaf samples were dried again at 60°C prior to analysis and analyzed for %N using Perkin-Elmer CHN analyzer. Average percent resorption was calculated as:

\[
\% \text{ Resorption} = 100 \times \left( \frac{\text{green leaf } \% \text{ N} - \text{ brown leaf } \% \text{ N}}{\text{green leaf } \% \text{ N}} \right)
\]

\( \text{Nitrate reductase enzyme activity} \)

We sampled nitrate reductase activity (NRA) during an expected peak in seasonal \( \text{NO}_3^- \) availability. Our methods were derived primarily from Downs et al. (1993). Green leaf samples were collected from red oak, red maple and white ash canopy trees July 6, 2009 within 2 h of solar noon on a sunny day. The samples were washed and cut into 10- to 15-mm\(^2\) sections, excluding midribs. A buffered solution (pH 7.0) of 40 \( \mu \)M \( \text{KNO}_3 \) with 1.5% propanol solution was added to the leaf tissues, and a 100-kPa vacuum was applied to each sample for 3 min and released. The tissues were incubated in the dark for 2 h at 30°C. A 1.0-mL aliquot of each sample was then diluted 1:2 with deionized water. After adding 0.5 mL each of 1% sulfanilamide in 1:5 \( \text{HCl} \) and 0.1% \( n \)-naphtylethylene diamine dihydrochloride, the samples were colorimetrically analyzed for nitrite using a Shimadzu UV 1201 spectrophotometer at 535 nm. Percent moisture was calculated gravimetrically using a subsample of each leaf. To normalize for changes in canopy mass with warming, NRA was multiplied by leaf mass for each species.

\( \text{Growth responses of trees by species} \)

Allometric equations were applied to monthly measurements of dendrometer bands on all trees >5 cm DBH in order to calculate changes in above- and below-ground woody biomass and vegetation carbon storage over time (Jenkins et al. 2003). Relative growth rates were calculated as:

\[
\text{Relative growth rate} = \left[ \frac{(\text{biomass}_{t+1})}{(\text{biomass}_{t})} - 1 \right] \times 100
\]

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where $t$ equals time, $\text{Biomass}$, is the woody biomass at time $t$ and $\text{Biomass}_{t+1}$ is the biomass at time $t+1$.

Litterfall was collected throughout the fall in permanent litter baskets (0.32 m$^2$ in size from 2003 to 2009 and 0.21 m$^2$ in size from 2006 to 2010). Litter was sorted by species, dried at 60°C for 24 h, and weighed. Annual litterfall by species was used as a proxy for canopy mass in subsequent analyses.

Statistical analyses

Annual net N mineralization rates were compared between the heated and control areas using Friedman’s test, a non-parametric repeated measures analysis in SAS (v.9.1.3). Similarly, relative growth rates were compared between heated and control areas using a Friedman’s test. In the cases where our samples were not collected from the same individual through time, including NRA, foliar %N, N resorption and litterfall, we compared data in the heated versus control areas using non-parametric (Wilcoxon) rank-sum tests. Results are reported on a “warming year” basis beginning in May, to coincide with heating cable activation (e.g., May 2002–April 2003 is the “pre-treatment” year, May 2003–April 2004 is “year 1,” and so on).

Results

Based on in situ soil incubations, we found that warming significantly increased annual rates of net N mineralization ($P < 0.01$) (Fig. 1a). The average increase of net N mineralization in the heated area relative to the control area was about 45% over 7 years of warming; a mean annual increase in the net N mineralization of 27.4 kg N ha$^{-1}$ in the organic and mineral soil (0–10 cm) horizons. We observed a progressive increase in NO$_3^-$ production over the 7-year study period (Fig. 1b). By the 5th year of warming, the production of NO$_3^-$ had increased by a factor of 3 in the heated area, with 25% of the NH$_4^+$ produced during the mineralization process transformed to NO$_3^-$.

Net N mineralization and nitrification varied both annually and seasonally in the heated and control areas. We defined spring as April–May, summer as June–August, autumn as September–October and winter as November–March. Peaks in net N mineralization occurred in the summer months, averaging 45 and 37% of the annual totals in the control and heated areas, respectively (Fig. 2). In the spring, net nitrification averaged 25% of the annual total in the control area and 32% in the heated area. In the fall, nitrification averaged 18% in the control area and 11% in the heated area. In the winter, nitrification averaged 3% in the control area and 8% in the heated area. While the overall trends were the same for both areas, the proportion of N mineralized in the summer decreased in response to warming, while the proportion of N mineralized in the winter increased with warming. The proportion of net nitrification occurring in the spring and winter increased in response to warming and decreased in the fall and summer.

Over 7 years of warming, we saw no significant solution or gaseous losses of N. While we saw occasional spikes in NH$_4^+$ and NO$_3^-$ solution losses in the heated and control areas, there was no significant effect of warming on solution losses through time. Gaseous N$_2$O fluxes from both heated and control areas were near detection limits, and no discernable pattern emerged over years of warming (Online Resource 1).

Foliar percent N has been significantly higher in the heated area since the 2nd year of warming ($P < 0.05$ for each year) (Fig. 3a). While the dominant species (oak, red
maple and white ash) all showed higher percent N with warming in most years, red maple had the largest and most consistent response to warming. When foliar N was expressed as a percent increase over the controls, red maple shows a 25% increase in leaf N by year 6 (Fig. 3b).

Overall, we saw no significant stand-level responses of leaf N resorption efficiency with warming. However, we did see significant species-level differences (Fig. 4a–c).

Red maple had the highest leaf N resorption compared to red oak and white ash in the warmed area (P < 0.01 for each year), and its leaf N resorption efficiency increased relative to the control by year 3 (P < 0.05 for each year). By year 6, oaks had a significant decrease in N resorption in response to warming (P = 0.04). White ash had a variable response in leaf resorption, with a positive response to warming in year 2 and a negative response to warming in year 3 (P = 0.05 and 0.01, respectively).
We sampled NRA during an expected peak in seasonal NO$_3^-$ availability in 2009. Overall, oaks had the highest NRA in the heated and control areas, followed by white ash and red maple (Table 1).

Warming had significant effects on plant C cycling. Overall, there was an increase in C storage with warming (Melillo et al. 2011). In addition, after correcting for pre-treatment differences, we saw a significant increase in the relative growth rate in trees on the heated area through time ($P = 0.007$) (Fig. 5a). We saw varied species-specific responses to warming. Red maple showed the largest and most consistent response to warming, doubling its relative growth rate in the warmed areas ($P = 0.01$) (Fig. 5b). While red oak had the greatest increase in biomass and trends toward increased relative growth rate in the heated area, the difference was not significant ($P = 0.20$) (Fig. 5c). White ash showed no consistent pattern of increased relative growth rates with warming ($P = 0.47$) (Fig. 5d).

In addition to increases in woody biomass, we saw overall increases in litterfall with warming (60 kg C ha$^{-1}$). Litterfall was used here as a proxy for canopy mass. Oak litterfall showed the greatest increase in response to warming by year 5 ($P = 0.006$), with an average annual increase in litterfall of 35% relative to the control over the past 7 years. The other dominant species showed more variation through time, with no clear pattern emerging.

### Discussion

Over 7 years of treatment, we have seen significant warming-induced changes in the N cycle that include increases in net mineralization and nitrification and increases in N concentrations in leaves of the canopy trees. We saw no evidence of increases in gaseous or solution N losses from the heated area relative to the control; the system has maintained a closed N cycle in spite of warming. The increase in N availability in the warmed area has led to increases in leaf N and plant C storage relative to the control (Melillo et al. 2011) and to an increase in relative growth rates, especially for red maples. Relative to other tree species, red oak exhibited high rates of nitrate reductase activity, although the relatively high reductase activity was observed in trees growing in both the control and heated areas.

#### Soil nitrogen

The increase in N mineralization in response to warming that we documented in this study has also been observed in other studies; some in forests (Peterjohn et al. 1994; Hartley et al. 1999; Rustad et al. 2001; Melillo et al. 2002), some in grasslands (Shaw and Harte 2001) and some in tundra (Chapin et al. 1995). In our study, net N mineralized in the heated area remained elevated relative to the control for the full 7 years. This sustained increase in net N mineralization with warming has been accompanied by increases in net nitrification, which has also been reported by others (Hartley et al. 1999; Barnard et al. 2004; Emmett et al. 2004).

### Table 1

| Species      | Heated        | Control       |
|--------------|---------------|---------------|
| Red maple    | 10.3 ± 25.8   | 95.7 ± 107    |
| White ash    | 35.0 ± 7.8    | 96.7 ± 22.7   |
| Red oak      | 2,072.4 ± 271.4 | 2,524.1 ± 303.8 |

Fig. 5  

- **a** Average relative growth rate for heated (filled circle) ($n = 74$) and control (open circles) areas ±1SE. 
- **b** Relative growth rate of oak species (red and black oaks) in control (open circles) ($n = 10$) and heated (filled circles) ($n = 22$) areas. 
- **c** Relative growth rate of red maple in control (open circles) ($n = 43$) and heated (filled circles) ($n = 33$) areas. 
- **d** Relative growth rate of white ash in control (open circles) ($n = 7$) and heated (filled circles) ($n = 10$) areas as a percent of initial growing season biomass ±1SE.
The increase in nitrification with soil warming is likely due to a combination of multiple interacting soil and plant responses, many of which we have not measured in this study. We have found that soil warming leads to decreases in labile soil organic matter and microbial biomass (Bradford et al. 2008; Frey et al. 2008). One possible implication of these changes in the labile C pool and microbial biomass is a decrease in microbial immobilization of N. This would cause an increase in NH₄⁺ available to nitrifying bacteria and an increase in net nitrification.

Tree responses to warming

Soil warming has led to changes in plant N cycling and relative growth rates. As soil N availability has increased with warming, the trees in the heated area have exhibited increased leaf N. By year 2, significant warming-induced changes in green leaf N content were apparent, especially for red maples. Leaf N is positively correlated to photosynthetic rate and carbon storage in many plants growing across the globe (Field and Mooney 1986; Reich et al. 1994, 1995, 1997; Ollinger et al. 2008). This relationship, observed in ecosystems around the world, suggests that the increases in leaf N we see with warming likely corresponds to increases in C assimilation in the heated area, which has implications for C cycling and storage.

Coupled with increases in leaf N, warming leads to general increases in the relative growth rate of trees, particularly for red maple. Recent studies have shown that red maple saplings increase in relative growth rate with increasing N availability (Finzi and Canham 2000; Zuccherio and Finzi 2007). Red maple’s strategy of taking up and storing more N, resulting in an increase in relative growth rate, suggests that it may have a competitive advantage in a warmer world.

Red oaks too may have a competitive advantage in a warmer world. This tree species had the greatest NRA, demonstrating its potential to take advantage of increases in NO₃⁻ in a warmer world. However, the higher levels of NRA have not yet resulted in increased relative growth rate for oaks in the warmed areas.

Multiple factors

Interactions between increasing temperatures and other factors predicted to change in the future, such as CO₂ concentrations, could reinforce the ecosystem responses to warming alone. Bazazz and Miao (1993) found that, when N was added to seedlings in growth chambers at Harvard Forest with elevated CO₂ (700 μL/L CO₂), biomass of red maple and red oak increased significantly relative to controls. Therefore, as CO₂ concentrations increase and warming stimulates increases in N availability, it is possible that we may see further increases in growth rates of these species. Results from Duke University’s Free-Air Carbon Dioxide Enrichment experiment (FACE) in North Carolina show red maples also benefiting from CO₂ enrichment (Mohan et al. 2007).

N cycling and forest community composition in a warmer world

Model-based predictions of future species composition in the northeastern USA in response to climate change have relied predominantly on variables such as temperature, precipitation, elevation, soil type and properties, and land use and fragmentation (Iverson and Prasad 2001, 2002; Hickler et al. 2004; Iverson et al. 2008). As suggested by Ostle et al. (2009), species’ responses to both form and abundance of N should be incorporated into species distribution models to better predict future forest composition. We have seen that the amount and form of N available may play a role in the growth rate and potential dominance of tree species in these temperate forests. We may be able to better predict the composition of our future forests by incorporating N availability and species-specific N form preference into species distribution models. By examining the N concentration in leaves, resorption of N during senescence, and the NRA in canopy leaves, in addition to the growth response of trees to warming, we can better predict how forested ecosystems will respond to climate change.

Conclusions

We have seen significant increases in net N mineralization and net nitrification with soil warming. The absence of solution and gaseous N losses suggest that the additional N mineralized as a result of warming is available for plant and microbial uptake. Warming significantly increases leaf N and growth of trees in the warmed area. The increases in available N have a significant effect on the relative growth rate of red maples, suggesting that this species will have a competitive advantage in taking up the additional N in a warmer world if the N economy is predominantly NH₄⁺ based. Interestingly, the red maples do not take advantage of the growing NO₃⁻ resource in the warmed area. Of the canopy dominants, oaks show the highest levels of NRA, suggesting that they are using much of the NO₃⁻. If the future N economy has a greater proportion of NO₃⁻, we would expect oaks to have a competitive advantage. By distinguishing species-specific nutrient acquisition strategies, we can begin to understand how canopy dominance and species composition may shift as the form and abundance of N changes in a warmer world.
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