Seasonal variation in effects of herbivory on foliar nitrogen of a threatened conifer

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Abstract. Invasive herbivores can dramatically impact the nitrogen (N) economy of native hosts. In deciduous species, most N is stored in stem tissues, while in evergreen conifer species N is stored in needles, making them potentially more vulnerable to herbivory. In eastern forests of the USA, the long-lived, foundational conifer eastern hemlock (Tsuga canadensis) is under the threat of extirpation by the invasive hemlock woolly adelgid (HWA: Adelges tsugae). We assessed the impact of HWA infestation on the patterns of seasonal foliar N availability in hemlock planted in a deciduous forest understory. Over the course of a year, we sampled needles and twigs and measured N, carbon (C), C:N ratio, and total protein concentrations. Tissue sampling events were timed to coincide with key life-history transitions for HWA to determine the association between HWA development and feeding with these foliar nutrients. In uninfested trees, needle and twig N concentrations fluctuated across seasons, indicating the potential importance of N storage and remobilization for the N economy of eastern hemlock. Although N levels in HWA-infested trees also cycled annually, the degree to which N concentrations fluctuated seasonally in tissues was significantly affected by HWA feeding. These fluctuations exceeded N levels observed in control trees and coincided with HWA feeding. HWA feeding generally increased N concentrations but did not affect protein levels, suggesting that changes in N do not occur via adelgid-induced protein breakdown. Herbivore-induced mobilization of N to feeding sites and its rapid depletion may be a significant contributor to eastern hemlock mortality in US forests.

Keywords: Adelges tsugae; nitrogen remobilization; nitrogen storage; protein; Tsuga canadensis.

Introduction
Nitrogen (N) utilized by trees for growth can be derived from both external and internal sources. Through fertilization (Tripler et al. 2006, LeBauer and Treseder 2008), mineralization of soil organic matter (Reich et al. 1997), atmospheric deposition (Nadelhoffer et al. 1999, Thomas et al. 2010), or internal resorption (Millard and Grelet 2010), increases in N availability can have a profound effect on tree growth, highlighting its limiting nature (Rennenberg et al. 2009). Internal storage and seasonal redistribution...
can also be a critical component of the N economy of trees (Millard and Grelet 2010). While deciduous trees store N in woody tissues, often in the form of bark storage proteins (Cooke and Weih 2005), evergreen conifers store N in their foliage primarily in the form of photosynthetic proteins such as ribulose bisphosphate carboxylase (RuBisCo) (Cammarano et al. 1996). For evergreen conifers, seasonal cycling of N among needles is a critical component of their N economy (Nambiar and Fife 1987, Millard and Proe 1992; Millard and Grelet 2010). For instance, remobilized N has been estimated to provide anywhere from 30 to 60% of the N incorporated into new growth (Rapp et al. 1979, Miller 1984, Nambiar and Fife 1987), highlighting the importance of N recycling. High levels of herbivory can often compromise future plant growth; in conifers, this impact may be partially due to the loss of valuable N stored in needles (Honkanen et al. 1999, Millard et al. 2001) or because of host manipulation by herbivores; many insects cause the accumulation of N at the site of feeding (Girouesse et al. 2005, Goggin 2007).

Evergreen conifers primarily store N in their youngest needles, and the high nutritional value of newly-produced plant tissues makes them especially prone to attack by both chewing and sucking herbivores (Coley 1980, Crawley 1983, Raupp and Denno 1983). For example, simulated browse of 1-year old Pinus sylvestris needles reduced new needle growth in spring (Honkanen et al. 1999) and N remobilization by 50% (Millard et al. 2001). While eastern hemlock is browsed by white-tailed deer (Odocoileus virginianus), whose impact can reduce growth and increase mortality (Stockeler et al. 1957, Freligh and Lorimer 1985, Mladenoff and Stearns 1993), this tree faces a more serious threat to its future. Throughout the eastern USA, eastern hemlock is experiencing significant declines in the wake of invasion by the hemlock wooly adelgid (HWA; Adelges tsugae). Introduced to the USA from Japan in 1950 (Souto et al. 1996), this exotic hemipteran feeds by inserting its stylet into xylem cells at the base of hemlock needles (Young et al. 1996), this exotic hemipteran feeds by inserting its stylet into xylem cells at the base of hemlock needles (Young et al. 1996), this exotic hemipteran feeds by inserting its stylet into xylem cells at the base of hemlock needles (Young et al. 1996). Lacking natural predators, HWA has spread rapidly throughout the eastern USA and can cause hemlock mortality in as little as 4 years (McClure 1991). Adelgid numbers can rapidly increase in an invaded area, as they undergo multiple generations a year, and easily disperse through wind, wildlife, and human activity such as logging (McClure 1989, 1990). Their lifecycle in the invaded range consists of two parthenogenetic stages on hemlock per year, a rapidly-developing spring generation and an overwintering generation with a longer development period (Ward et al. 2004). The April–June progrediens generation emerges in early spring and settles on hemlock foliage produced in the previous growing season. Females in this generation can produce ~75 eggs per individual and their offspring (sistens generation; July–April) settle on newly produced foliage. Once settled, they enter aestival until late autumn, after which individuals feed throughout the winter and are capable of producing anywhere from 70 to 200 eggs per female (Paradis 2011). Understanding the seasonal dynamics of nitrogen availability and herbivore activity may elucidate the mechanisms driving hemlock decline and mortality.

Relatively, few studies have investigated the effects of sap-feeding herbivores on woody species (reviewed in Zvereva et al. 2010). Conifers may be especially susceptible to sap-feeders because they allocate more storage compounds to foliage than do deciduous trees (Chapin et al. 1990). Nutrient storage in the foliage may make conifers more likely to succumb to intense sap-feeding events (Fernandezes 1990, Furuta and Aloe 1994, Paine 2000), as temporary nitrogen depletion has the potential to adversely affect tree health over short timescales, particularly during the periods of rapid growth. Indeed, the limited research on the effects of sap-feeders on woody species has revealed that they can negatively affect photosynthesis, growth and reproduction (Zvereva et al. 2010), and in certain contexts may even surpass defoliators in their impact on photoassimilates (Llewellyn 1972, Schowalter et al. 1981). Given their sessile nature, the survival and fecundity of specialist sap-feeding herbivores depends on local resource availability in their plant hosts. As a result, the timing of herbivore development relative to fluctuations in host nitrogen is critical for insect survival and fecundity (Mattson 1980). As a sessile hemipteran, HWA is also dependent on high-quality local resources during active feeding periods; elevated N levels are found at HWA feeding sites (Gómez et al. 2012, Soltis et al. 2015).

Previous common garden studies suggest that the impacts of HWA feeding on N dynamics are dependent upon factors that vary over time. Foliage from artificially-infested trees had a lower N concentration than uninsected foliage 5 to 10 months after initial infestation (Miller-Pierce et al. 2010). One year later, however, no HWA effect on foliar N was observed. In a separate study using experimental samples collected in the spring, newly-produced HWA-infested foliage had a higher amino acid content than uninfested foliage after one year of infestation (Gómez et al. 2012). The amino acid content of foliage from trees infested for 3 years, however, did not differ from control foliage. The changes to N dynamics across multiple years of data collection suggest that the effect of HWA on foliar N varies throughout the progression of infestation. While free amino acids are one of many components making up total N, these studies together suggest a drop in N or a spike in amino acids...
acids in the first year of infestation, followed by recovery to levels comparable to uninfested trees. Further, the contrasting results between total N and amino acid dynamics may be due to seasonal variation in the effects of infestation. It is thus still unclear how and when HWA may be causing N resource depletion.

We examined seasonal variation in foliar N availability in eastern hemlocks grown in an understory common garden to 1) assess baseline N dynamics in healthy trees; and 2) understand how these dynamics are affected by adelgid feeding. We examined tissue-level concentrations of N, carbon (C), C:N, and protein as simple indicators of bulk N dynamics. By measuring these indicators over the course of a year, we aimed to clarify previous results indicating a time-dependent HWA effect.

Methods

Experimental design

We collected a one-year time course of samples from seedlings in an understory common garden established at the Kingston Wildlife Research Station (Kingston, RI). One-year-old seedlings were purchased from Van Pines Nursery (West Olive, MI, USA) and planted in a grid in April 2011. The seedlings were inoculated with HWA-infested foliage following standard protocols (Butin et al. 2007) over the course of three growing seasons, in April 2011, 2012, and 2013. Control trees received a sham treatment using uninfested hemlock foliage to control for tree handling. Seedlings were artificially infested in a randomized complete block design, such that 15 trees each were allocated to HWA and control treatments. All seedlings were caged in wire and netting to prevent deer herbivory and HWA dispersal (Agribon-15, Johnny’s Selected Seeds, Waterville, ME, USA; 90% light transmission). On our first sample date, the seedlings were < 1 m in height, approximately 2-years old, with 1 year of HWA infestation in the common garden. Complete details of the common garden are available elsewhere (Gonda-King et al. 2014).

Adelgid abundance

To assess infestation levels, HWA densities were monitored in late fall (October 2012) and early spring (April 2013). Briefly, in each season, two 5 cm branchlets were selected and HWA individuals were counted.

Tissue sampling and analysis

Our time course included a total of seven sampling events from fall 2012 to summer 2013, in an effort to characterize N dynamics over 1 year in the phenology and life history of three HWA generations (G) (Fig. 1).

Figure 1. Life cycle of Adelges tsugae in the eastern USA in relation to Tsuga canadensis seasonal growth. Vertical arrows indicate dates of tissue sampling to assess effects of A. tsugae on the nitrogen economy of eastern hemlock. Adapted from Gonda-King et al. (2014).

These included: HWA G1 mid-diapause (September 24), HWA G1 immediately preceding diapause break (October 8), HWA G1 newly active (October 31), HWA G1 maturing and active (November 26), HWA G2 newly hatched (April 6), HWA G1 of year 2 newly hatched and preparing to enter diapause (July 8) (Fig. 1). We sampled young (2012) and mature (2011) foliage (twigs and attached needles) for all analyses. In July, we sampled only the newest (2013) elongated twigs and young (2012) twigs; mature (2011) twigs had already lost most needles. Sampled branches were haphazardly selected from the top third of the tree, and the same branches were subsampled for each foliar age class. We collected one pooled sample (>10 g of foliage from one to three branch segments with attached needles) per tree per age class at each sampling event. Immediately following these measurements, samples were oven-dried at 75 °C to constant mass. Needle samples were finely ground in a KLECO ball mill (Garcia Machine, Visalia, CA, USA), and twigs were finely ground in a Wiley mill (Thomas Scientific, Swedesboro, NJ, USA). Ground samples were stored desiccated in sealed vials prior to analysis. We determined %C and %N of 5 ± 0.5 mg subsamples by dry combustion on a CHNOS analyzer (vario Micro cube, Elementar Americas, Mt. Laurel, NJ, USA). Finally, protein concentration of tissues was analyzed using 10 ± 0.5 mg subsamples by Bio-Rad Bradford assay (Bradford 1976).

Statistical analyses

We performed all analyses in R v. 3.2.2 (R Core Team 2015). Linear mixed-effects models (LMMs) were fit using packages lme4 (Bates et al. 2014) and lmerTest (Kuznetsova et al. 2013) for analyses of N(%), C(%), C:N, and protein data. The package lmerTest uses algorithms of SAS Proc Mixed (Littell et al. 1996) to both assess the significance of fixed effects using Type III sums-of-squares and estimate denominator degrees of freedom using Satterthwaite’s approximation. For measures on mature (2011) and young (2012) foliage from September 24 to April 6, we fit LMMs with sample date (quantitative), insect treatment (HWA or control), tissue age (mature or young), tissue type (needles or twig), as well as all possible interactions as factors. Tree identity was
included as a random effect. We then proceeded with a backward model selection procedure using the step function, where terms are removed sequentially based on having the lowest statistical significance ($P > 0.1$). Protein concentration data for Oct 31 were not included in the statistical analysis due to sample loss.

For analyses of N(%), C(%), C:N, and protein in newest (2013) foliage (sampled only on July 8), we fit separate LMMs with factors insect treatment and tissue type, and their interaction, with tree identity as a random effect. The same resource measures taken on young (2012) foliage on July 8 were included in this analysis for comparison. When appropriate, data were transformed to meet the assumptions of normality and homoscedasticity. Results from these best-fit models [see Supporting Information—Tables S1 and S2] are presented.

To compare HWA densities in the fall and spring, we used a Wilcoxon Signed-Rank test.

**Results**

**Seasonal N availability in eastern hemlock**

Nitrogen levels in needles of uninfested trees cycled annually (Fig. 2A). In the fall (September 24), N concentrations in young (2012) and mature (2011) needles averaged 1.94% (±0.05%) and 1.63% (±0.06%) of needle biomass respectively. These levels however decreased by 31% and 20% over the winter, reaching averages of 1.34% (±0.03%) and 1.3% (±0.05%) as these tissues aged leading into spring (April 6). New growth the following year was similar to September 24 levels. Mirroring N concentrations, protein levels of needles and twigs exhibited significant seasonal cycling (Fig. 2E and F: $F_{3,390.10} = 18.48, P < 0.001$). Twig N also cycled in an annual fashion (Fig. 2B), though to a less pronounced degree and in an opposite direction.

**Effects of hemlock woolly adelgid on seasonal N availability**

Adelgid densities significantly differed between the fall and spring of 2011 and 2012 respectively (Wilcoxon Signed-Rank test: $P = 0.001$). Sistens-generation HWA were five times as dense in the fall (Mean ± SE: 1.67 ± 0.38 insects/cm) compared to the spring (0.32 ± 0.16 insects/cm).

Adelgid feeding affected N concentrations of needle and twig tissues produced in 2011 and 2012 (Fig. 2A).
and B); however, these effects depended on both the age of tissue examined ($F_{1,523.09} = 16.63, P < 0.001$), as well as the season in which tissues were sampled ($F_{1,522.09} = 4.29, P = 0.04$). For instance, HWA infestation elevated the N concentration of young (2012) needles sampled September 24 by 13% relative to controls. Though initially elevated, N in these needles decreased over winter as sistens emerged from aestivation in late fall and began to feed, dropping 36% to an average 1.4% (±0.06%) in April: a level no different from controls (post hoc Tukey HSD test: $P > 0.05$). Moreover, new foliage produced in 2013 following feeding by progrediens on young (2012) and mature (2011) tissues was found to be 11% higher in N concentration than foliage produced by controls, although this effect was only marginally significant ($F_{1,28} = 3.80, P = 0.06$). Nitrogen dynamics in twigs mirrored that of needles. Young (2012) twigs sampled in September had N levels 28% higher than controls. As sistens began to feed, these levels dropped 19%, to a level 11% lower than controls at the end of November. Interestingly, as sistens continued feeding over winter, N levels increased steadily in HWA-infested twigs to a level 9% higher than controls in April. Finally, despite affecting N, HWA infestation had no effect on protein levels of tissues between September and April (HWA main effect: $F_{1,29,26} = 0.04, P = 0.83$) or in July (HWA main effect: $F_{1,27,52} = 0.07, P = 0.80$).

Adelgid effects on C:N ratios from fall to spring varied according to tissue type and when tissues were examined (Treatment x Tissue x Time interaction: $F_{1,519.74} = 9.27, P = 0.002$). This effect was largely driven by sistens feeding during winter and resulting shifts in C:N ratios of mature (2011) needles and twigs (Fig. 2C and D). After spring, C:N ratios of tissues produced in 2013 were significantly affected following progrediens feeding ($F_{1,28} = 4.89, P = 0.04$). Carbon-to-nitrogen ratios were lower for HWA-infested tissues, driven by aforementioned increases in N, rather than changes in C (HWA main effect: $F_{1,28} = 0.0002, P = 0.99$).

**Discussion**

Nitrogen concentrations of tissues fluctuated across seasons in a dynamic manner. The nature of these fluctuations varied across tissue types however, and were significantly affected by HWA feeding. Specifically, we observed high N levels at HWA feeding sites and then a rapid depletion of this resource across seasons as feeding commenced. This suggests that HWA may act as sinks for remobilized N and that changes in N economy is likely a significant contributor to eastern hemlock mortality in North American forests.

**Seasonal variation in N availability in eastern hemlock**

Nitrogen concentrations in needles, primary tissues for N storage and utilization in conifers, fluctuated over the course of the year by up to 36%. These fluctuations exhibited an annual cycle, indicative of potential movement of N between storage and actively growing tissues across seasons. Overall, N levels were relatively high in needles in the fall, and then steadily declined until April the following year, as N likely turned over and translocated to new, expanding needles in spring. Though N concentrations are elevated in needles, differences between tissues of different age were observed. The highest levels of N were observed in current year needles (2012) in the fall, while older needles produced in 2011 were comparatively lower. This pattern is consistent with findings for other conifers (Millard and Grelet 2010), highlighting how the bulk of a host’s N is stored in the youngest age class of needle. Our data on patterns of protein concentrations suggest that protein storage and remobilization likely features as part of the hemlock’s N economy as well. Protein levels decreased during fall and winter in current year (2012) needles, with protein levels spiking again in July in maturing 2013 foliage. These losses have been shown in another conifer Pinus sylvestris, and also correlate with reduced Rubisco activity over this period (Gezelius and Hallén 1980, Gezelius et al. 1981), as Rubisco is likely hydrolyzed and resulting N is translocated to other parts of the tree during this period of shoot elongation (Fife and Nambiar 1982, Nåsholm and Ericsson 1990).

Nitrogen concentrations in twigs fluctuated in an annual cycle similar to needles, highlighting a likely important role for these tissues in N storage and mobilization. Though both cycled, the patterns and levels of N were in stark contrast, reflecting the inherent differences between these tissues in their chemistry, structure and functionality. This turnover, by upwards of 22%, is driven by slight, but steady increases in N concentration in twigs over winter until spring, when N concentration declines steeply, as observed in young (2012) twigs. Concurrent with losses of N in needles, this pattern may capture N remobilization and translocation, if N from previous-year growth is allocted to branch elongation and new needle expansion in the early summer (Millard and Proe 1992).

**Effects of hemlock woolly adelgid on seasonal N availability**

Although adelgid feeding altered N concentrations, the degree and direction of the impact varied according to tissue age, sampling date, and coincided with key life-history transitions over the course of the HWA life cycle.

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In early fall (September), HWA sistens produced in the summer are settled on young (2012) and mature (2011) tissues. These tissues are significantly elevated in N relative to control trees, following feeding by the progresiens generation the previous spring. Once sistens emerge from aestivation (late October and November) and begin to actively feed, N concentrations in young (2012) and mature (2011) needles and twigs drops considerably as sistens develop and invest in production of the next generation (progresiens) that will emerge the following spring (April). Though sistens depress N as they feed, once hatched in April, spring-generation HWA (progresiens) commence feeding immediately on young (2012) twigs, enhancing N local to feeding sites. Moreover, this enhancement of N locally can affect N content of the newly produced foliage (2013 growth). Adelgid feeding marginally increased N levels in the newest (2013) tissues relative to controls. This finding supports previous work showing elevated concentrations of free amino acids in uninfested, newly-produced foliage (Gómez et al. 2012). Finally, in July, summer-generation larvae (sistens) produced by progresiens settle onto the newest (2013) twigs and enter aestivation, repeating the cycle.

There are several potential explanations for elevated N concentrations in HWA-attacked tissues. First, increased N concentration in HWA-infested tissues may indicate that HWA can act as a strong sink for N. Previous research using measures of N and stable isotopes has indicated the potential for HWA to increase N local to feeding sites (Gonda-King et al. 2014, Soltis et al. 2015). Aphids and other sap-feeding hemipterans have been similarly shown to increase N transport to feeding sites (Girousse et al. 2005, Goggin 2007). Second, following infestation by HWA, eastern hemlock trees often exhibit a reduction in new growth in spring (Preißer and Elkinton 2008; Miller-Pierce et al. 2010). Relative to controls, elevated N in new foliage produced in 2013 may merely reflect a concentration effect if both uninfested and infested trees allocate the same amount of N to new growth. Lower levels of N in controls would reflect a dilution effect as foliage expands.

Adelgid feeding had no effect on protein concentrations in eastern hemlock foliage. These results are in support of previous findings that utilized the same assay (Soltis et al. 2015), suggesting that HWA feeding does not accelerate protein breakdown in infested tissues. However, our methods for measuring proteins may limit our ability to detect effects or draw conclusions on how proteins are fully impacted by HWA. The Bradford assay is highly specific to proteins with molecular weights in excess of 3 kDa, particularly with arginine, basic, or aromatic amino acid residues. Shorter polypeptides, other nitrogen-containing molecules, and less soluble proteins are not detected (Jones et al. 1989). Other forms of N in the needles, including free amino acids, short polypeptides, nucleic acids, secondary metabolites, and inorganic N, were detected in our total-N combustion, but not by the Bradford assay. If HWA are primarily feeding on free amino acids or short, soluble polypeptides available in the xylem ray parenchyma, the lack of effect on proteins as detected by the Bradford assay is perhaps not surprising.

In summary, our data demonstrate seasonal variation in impacts of an invasive herbivore on foliar N of a threatened host. Akin to other evergreen conifers, internal cycling of N for growth is likely an important component of the overall N economy for eastern hemlock. However, winter and spring feeding by HWA sistens and progresiens generations respectively may limit remobilization and transport to new tissues. These findings highlight the significant, but understudied impact that sap-feeders can exert on N of hosts. Future work should clarify how HWA impacts on total N pools and fluxes scale to affect entire forested systems.

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Contributions by the Authors

N.S., S.G., E.P. and C.O. conceived the idea for the study. N.S., S.G., J.M. and A.B. performed the research. N.S. and R.S. analyzed data, with R.S. leading the writing of the manuscript. All authors contributed to revisions.

Conflict of Interest Statement

None declared.

Supporting Information

The following additional information is available in the online version of this article —

Table S1. Analysis of variance (ANOVA) tables for effects of Adelges tsugae (HWA) on N(%), C:N ratio, and protein concentrations of mature (2011 growth) and
young (2012 growth) Tsuga canadensis tissues from September to April.

Table S2. Analysis of variance (ANOVA) tables for effects of Adelges tsugae (HWA) on N(%), C:N ratio, and protein concentrations of new Tsuga canadensis tissues produced during the 2013 season.

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