Phenology and allocation of belowground plant carbon at local to global scales

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Dissertation

PHENOLOGY AND ALLOCATION OF BELOWGROUND PLANT CARBON AT LOCAL TO GLOBAL SCALES

by

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ABSTRACT

Forests play an important role in mitigating climate change by removing carbon dioxide (CO₂) from the atmosphere via photosynthesis and storing it in plant tissues and soil organic matter (SOM). Plant roots are a major conduit for transporting recently fixed CO₂ belowground, where carbon (C) remains in SOM or returns to the atmosphere via respiration of soil microbes. Compared to aboveground plant processes related to the C cycle, there is little understanding of how belowground plant-C allocation to roots, symbiotic root fungi and secretions into the soil influence the gain or loss of C from the soil. Further, the uncertainty in the timing and amount of root growth that occurs in forests is a barrier to understanding how root activity responds to global change and feeds back to the C cycle. Therefore, the objective of my research is to quantify the timing and magnitude of C allocation to roots and soil via data compilation, field studies and modeling across broad spatial scales. Using data compilation at the global scale, I show that root and shoot phenology are often asynchronous and that evergreen trees commonly have later root growth compared to deciduous trees using meta-analysis across four
biomes. At the plot scale, field studies in a mid-latitude forest demonstrate that deciduous stands allocate more C belowground earlier in the growing season compared to a conifer stand. The difference in phenology between stands can be attributed to the timing of root growth. At the root scale, zymographic analysis demonstrates that microbial extracellular enzyme activity is concentrated near the surface of roots and that the rhizosphere can extend well beyond 2 mm from the root surface. Finally, I developed a new model of microbial physiology and extracellular enzyme activity to assess how climate change may affect plant – microbe interactions and soil organic matter decomposition. I show that increases in temperature and the quantity of C inputs substantially alter decomposition. Collectively, these results demonstrate the importance of belowground allocation to the C cycle of terrestrial ecosystems.
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<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>$\theta$</td>
<td>volumetric soil moisture</td>
</tr>
<tr>
<td>2-D</td>
<td>two dimensional</td>
</tr>
<tr>
<td>$a$</td>
<td>proportion of the enzyme pool acting on the soil organic carbon pool</td>
</tr>
<tr>
<td>$A$</td>
<td>pre-exponential constant</td>
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<td>AIC</td>
<td>Akaike information criterion</td>
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<tr>
<td>AM</td>
<td>arbuscular mycorrhizal</td>
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<tr>
<td>ANOVA</td>
<td>analysis of variance</td>
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<tr>
<td>AP</td>
<td>acid phosphatase</td>
</tr>
<tr>
<td>BD</td>
<td>bulk density</td>
</tr>
<tr>
<td>BG</td>
<td>beta-glucosidase</td>
</tr>
<tr>
<td>C</td>
<td>carbon</td>
</tr>
<tr>
<td>$C_{\text{roots}}$</td>
<td>carbon pool associated with fine roots</td>
</tr>
<tr>
<td>$C_{\text{soil}}$</td>
<td>carbon pool associated with soil</td>
</tr>
<tr>
<td>CMIP5</td>
<td>Fifth Coupled Model Intercomparison Project</td>
</tr>
<tr>
<td>C:N</td>
<td>carbon-to-nitrogen ratio</td>
</tr>
<tr>
<td>$CO_2$</td>
<td>carbon dioxide</td>
</tr>
<tr>
<td>CUE</td>
<td>carbon use efficiency</td>
</tr>
<tr>
<td>CV</td>
<td>coefficient of variation</td>
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<tr>
<td>DAMM</td>
<td>Dual Arrhenius and Michaelis-Menten</td>
</tr>
<tr>
<td>DEATH</td>
<td>microbial turnover</td>
</tr>
<tr>
<td>DEPOLY</td>
<td>depolymerization of soil organic matter</td>
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</table>
Dliq .......................................................... diffusion coefficient for unprotected soil organic matter in liquid
DOC ........................................................................................................... dissolved organic carbon
DOM ........................................................................................................... dissolved organic matter
DON ........................................................................................................... dissolved organic nitrogen
DOY .......................................................................................................... day of year
E\textsubscript{a} ................................................................................................ activation energy
EEA ............................................................................................................. extracellular enzyme activity
EPROMD ..................................................................................................... enzyme production
ELOSS ........................................................................................................ enzyme turnover rate
EU ............................................................................................................. enzyme units
F\textsubscript{efflux} ............................................................................................ soil respiration rate
F\textsubscript{exudates} ........................................................................................ root exudation rate
F\textsubscript{leaching} ..................................................................................... flux of dissolved organic carbon into streamwater
F\textsubscript{litterfall} ....................................................................................... litterfall
F\textsubscript{resp} ............................................................................................. fine and coarse root respiration rate
F\textsubscript{roots} ............................................................................................ gross fine root production rate
frac ................................................................. fraction of unprotected soil organic matter
GPP ............................................................................................................. gross primary production
GROWTH ............................................................................................... microbial biomass growth
H\textsubscript{2}O ................................................................................................. water
HWA .......................................................................................................... hemlock woolly adelgid
HSD ................................................................................................. Honestly Significant Difference

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inputDOM ............................................................. root input to the soil
Km ........................................................................... half-saturation constant
MCNiP .................................................................. Microbial Carbon and Nitrogen Physiology
MCP ......................................................................... Microbial Carbon Physiology
MIC ........................................................................... microbial biomass
MICtoSOM ........................................... fraction of dead microbial biomass returned to soil organic matter
MIMICS ................................................................. Microbial Mineral Carbon Stabilization
MMP ................................................................. mean monthly precipitation
MMT ................................................................. median monthly temperature
N ................................................................................... nitrogen
NAG ........................................................................... N-acetyl-beta-D-glucosaminidase
NEON ................................................................. National Ecological Observatory Network
NPP ............................................................... net primary production
NSC ................................................................. nonstructural carbohydrate
O ................................................................................... organic
O2 .................................................................................. oxygen
O2airfrac ................................................................ volume fraction of oxygen in air
p ........................................................................... proportion of assimilated C allocated to enzyme production
PAR .............................................................. photosynthetically active radiation
PD ........................................................................... particle density
Pg .................................................................................. petagram
PFT ................................................................... plant functional type
aminopeptidase

enzyme turnover rate

microbial turnover rate

proportion of assimilated N allocated to enzyme production

root-mean-square error

sum of squared residuals

soil organic carbon

volumetric water content

soil organic matter

temperature

total belowground carbon flux

rate of dissolved organic matter uptake

ultraviolet

maximum reaction rate

spatial weights matrix
CHAPTER ONE: INTRODUCTION

For over a century, researchers have recognized the link between atmospheric carbon dioxide (CO$_2$) concentrations and the re-radiation of thermal energy back to the Earth’s surface (Arrhenius & Sandström, 1903, Chamberlin, 1899). In 1957, Roger Revelle and Hans Seuss were among the first to suggest that the observed accumulation of fossil-fuel-derived CO$_2$ in the atmosphere would not only contribute to a warmer climate, but that the extent of climate change may depend on interactions between the atmosphere, biosphere, and geosphere that are not yet known (Revelle & Suess, 1957). “Thus,” they write, “human beings are now carrying out a large scale geophysical experiment of a kind that could not have happened in the past nor be reproduced in the future.”

From 1750 to the present, the terrestrial biosphere and ocean have mitigated climate change by sequestering 300 of the 550 Pg C released from the combustion of fossil fuels (Stocker et al., 2013). There is at present substantial uncertainty regarding the size of the terrestrial sink over the coming century with continued climate change (Ahlström et al., 2012). There is also debate as to whether the feedback between climate change and the land sink for atmospheric CO$_2$ will mitigate or exacerbate future climate change (Stocker et al., 2013). On the one hand, if the growing season lengthens as a result of warming temperatures, C uptake is expected to increase because of sustained photosynthesis (Richardson et al., 2009). On the other hand, extension of the growing
season may exacerbate drought, fire frequency, and stimulate decomposition resulting in a net loss of CO$_2$ to the atmosphere (Piao et al., 2008, Wu et al., 2013).

The terrestrial sink for atmospheric CO$_2$ is in plant biomass and soil. Typically 2- to 3-fold more C is stored in soil than biomass, increasing substantially with latitude (Ciais et al., 2014). The activity of soil microbes, often referred to as heterotrophs, is temperature sensitive, with greater respiration consuming soil C and releasing CO$_2$ to the atmosphere. Heterotrophic respiration is expected to increase with warming temperatures, but the magnitude of this effect depends on the extent to which microbes physiologically acclimate to warm temperatures, microbial community composition is changed, or the availability of soil C declines (Bradford et al., 2008, Hartley et al., 2007). As a result, the long-term effect of global warming on heterotrophic respiration is unclear (Frey et al., 2013, Melillo et al., 2002).

The research presented in this dissertation is a continuation of the effort to understand feedbacks between plants, microbes, soil and the atmosphere. Ultimately all soil C is derived from plant matter and thus plants are intimately tied to the activity of soil microbes. The main process by which this occurs is referred to as “allocation” of C to leaves, stems, and roots.

Plant C allocation is an inherently seasonal process. The timing of seasonal events, or phenology, of aboveground plant organs and the tools used to observe it are advanced compared to studies of root growth and mortality. For example, there are records from the 9th century of cherry tree flowering in Kyoto, Japan, as well as a myriad
of historical phenological records of various plant phenomena dating back to 18th century England and 19th century United States (Aono & Kazui, 2008, Margary, 1926, Miller-Rushing & Primack, 2008). Since the 1970s, “remote sensing” technology using satellites, unmanned aerial vehicles, and mounted cameras have been instrumental in estimating growing season length and C uptake over the globe (Sharma et al. 2013, Hufkens et al. 2012).

In contrast to aboveground phenology, root dynamics are hidden from view, and there is no non-invasive technique analogous to remote-sensing that can be used to accurately assess root biomass, growth, or turnover. Root biomass has been estimated using coring and excavation for over half of a century (Lyford, 1980, Lyford & Wilson, 1964). In these and similar studies, root growth was measured by taking the difference in root biomass between two sample intervals. Beginning in the early 1980s, Vogt, Persson, Gower, and others popularized “sequential” coring in plots over a finer resolution of time and ingrowth techniques to separate rates of root growth from those of mortality (Persson, 1980, Vogt et al., 1986). By the late 1980s, minirhizotron cameras were developed to capture magnified images of root growth and death along the surface of a plexiglass tube installed into the soil (Brown & Upchurch, 1987). Though root boxes and minirhizotrons have been in use for some time, only recently has the automation of image capture and annotation begun to develop (Roberti et al., 2014, Zeng et al., 2008).

Carbon allocated belowground is used for root production, respiration and exudation, but these fluxes are difficult to measure individually. In 1989, Raich and
Nadelhoffer proposed that calculating the difference between outputs (i.e., soil CO$_2$ efflux) from the soil and inputs (i.e., litterfall) to the soil provides a useful estimate of the total quantity of C allocated belowground (Davidson et al., 2002, Litton et al., 2007). This estimate assumes that soils are in steady state. Otherwise, it is necessary to measure the change in the C pool associated with roots and soil, as in:

$$\text{TBCF} = F_{\text{efflux}} - F_{\text{litter}} + \Delta(C_{\text{roots}} + C_{\text{soil}})$$  \[1\]

where TBCF is the total belowground carbon flux, $F_{\text{efflux}}$ is annual soil respiration, $F_{\text{litter}}$ is annual litterfall, and $\Delta(C_{\text{roots}} + C_{\text{soil}})$ is the annual change in the C pool associated with roots and soil. If $\Delta(C_{\text{roots}} + C_{\text{soil}})$ is equal to zero, then it is only necessary to measure $F_{\text{efflux}}$ and $F_{\text{litter}}$.

The assumption that the pool of C in roots and soil does not change over time is often violated. For example, Giardina & Ryan (2002) found that the change in the root and soil pool was about 14% of measured TBCF in a Hawaiian Eucalyptus plantation. In my study, I found that 0.2 - 40% of TBCF can be represented by the change in the root pool. The most accurate method to estimate belowground C allocation may be to compare multiple methods, such as using the difference between soil CO$_2$ efflux and litterfall in combination with measuring belowground allocation to roots [production, respiration, exudation] directly (Drake et al., 2011).

Once C enters the soil via root exudation or turnover, it is available to the microbial community for decomposition. Soil microbes are able to utilize a diverse array
of substrates for metabolism (Bai et al., 2013). Given the metabolic flexibility of microbes and the amount of energy stored in soil organic matter, it is paradoxical that there should be more than 1500 Pg of soil organic matter remaining on Earth (Ciais et al., 2014). Most soil organic matter has an irregular structure and is therefore difficult to characterize by chemical or spectral analysis. Indeed, for the past several decades, the prevailing paradigm to explain the persistence of SOM was that most organic matter cannot be degraded because of its irregular chemical structure (Oades, 1989, Skjemstad et al., 1996). Recent research has placed more emphasis on the physical separation of SOM from the microbial extracellular enzymes that would degrade them (Schmidt et al., 2011) and the physical protection of SOM bound to or within soil aggregates (Conant et al., 2011).

The wealth of research over the past century has clarified many feedbacks between the biosphere and atmosphere, by demonstrating that phenology has changed with warming, and that respiration and C allocation may also change (Bradford et al., 2008, Melillo et al., 2011, Richardson et al., 2009). As a result, we have a greater ability to monitor and simulate C pools and fluxes at global scales than we did only two decades ago (Stocker et al., 2013, Tang & Riley, 2015).

Despite these advances, many mechanisms of belowground C allocation are not yet clear. Individual studies have observed many different seasonal patterns of root growth (Bevington & Castle, 1985, Iivonen et al., 2001, McCormack et al., 2014), but it is not clear if particular biomes or plant types have synchronous or lagged phenology.
Because belowground C allocation data are few, we know little about the seasonal C partitioning to roots and how this varies between tree species, growth form (e.g., evergreen, deciduous), or mycorrhizal association. In part because of the paucity of data with which to parameterize models and infer process-level relationships, terrestrial biosphere models struggle to represent the soil C cycle mechanistically and often default to defining belowground C allocation as an instantaneous, fixed fraction of net C uptake at each time step (Medvigy et al., 2009, Oleson et al., 2010, Smith et al., 2013). There is little understanding of the priority of C allocation in both plants and microbes, though some promising frameworks for determining allocation to different plant organs are emerging from evolutionary theory based on competition for light and nutrients (Craine & Dybzinski, 2013, Dybzinski et al., 2011).

Microbial allocation to extracellular enzymes, biomass growth, and respiration is largely unknown, and is a major difficulty in parameterizing microbial physiology models (Averill, 2014). Fortunately, allometric relationships for microbial communities, as well as models of microbial metabolism are emerging frameworks for estimating microbial allocation to different processes (Harcombe et al., 2014, Sinsabaugh et al., 2014).

The interaction between plant roots and microbes is similarly data-limited. For example, including my findings, estimates of the size of the rhizosphere, defined as the zone of soil around a root where microbial exoenzyme activity is stimulated, ranges from 0.1 cm to over 7 cm (De Neergaard & Magid, 2001, Spohn & Kuzyakov, 2014). This
wide range reflects the few studies that have made these measurements, and suggests that there is either a large methodological uncertainty about how to quantify the rhizosphere extent, a large heterogeneity in the actual size of the rhizosphere, or both.

The research community is uncertain how rhizosphere and bulk soil microbes will respond to changes in temperature and substrate supply (Davidson & Janssens, 2006, Tang & Riley, 2015, Wieder et al., 2013). Long-term field studies of soil warming indicate that a large initial increase in heterotrophic respiration decreases over time, suggesting some combination of acclimation, microbial community shift, or substrate limitation (Frey et al., 2013, Melillo et al., 2002, Melillo et al., 2011). Regional climate projections also predict that some areas of North America, particularly the northeastern United States, will receive more precipitation, and others less (Melillo et al., 2014). It is unclear how changes in soil moisture in combination with warming will affect microbial activity.

**Dissertation Overview**

The objective of the research in this dissertation is to better understand the seasonality, magnitude, and partitioning of C to root and soil processes at global-to-local scales. To achieve this objective, I used a combination of data compilation, field studies, and computer simulation. In Chapter 2, my objective was to determine the relationship between above and belowground phenology. I performed a meta-analysis, compiling data that quantified the offset between the maximum in root and shoot growth in woody and herbaceous perennial plants. Using data from 40 studies, 63 species, and 4 biomes, I
found that root and shoot phenology are often asynchronous and that evergreen trees commonly have later root growth compared to deciduous trees. I also found broad differences in phenology across biomes, finding that boreal biomes had later root growth than did temperate biomes, possibly due to the predominance of evergreen trees in boreal ecosystems.

In Chapter 3, my objective was to determine the magnitude, timing and partitioning of seasonal belowground C allocation at the plot scale, using field studies at the Harvard Forest in Petersham, MA. I recorded the total amount of C allocated belowground monthly during the growing season in three mono-dominant stands, white ash (*Fraxinus americana*), red oak (*Quercus rubra*), and eastern hemlock (*Tsuga canadensis*). I found that the red oak stand allocated more C belowground earlier in the growing season compared to the eastern hemlock stand. This difference in the phenology of belowground C allocation can be attributed to the timing of root growth, making this finding consistent with the observation in Chapter 2 that deciduous trees have earlier root growth than evergreen trees. The magnitude of the belowground C flux was highest in the red oak stand, consistent with the observed increase in aboveground biomass of red oak at the Harvard Forest over the past 20 years (Keenan *et al*., 2012, Urbanski *et al*., 2007). The belowground C flux was lowest in the eastern hemlock stand due to a decline in allocation to root production over the study period. This decline is coincident with the arrival and spread of the hemlock woolly adelgid in this stand.

In Chapter 4, my objective was to estimate the spatial extent of the rhizosphere at the scale of individual roots, using a recently-developed enzyme imaging, or
zymography, technique. I incubated chromatography paper soaked in the substrate of four C-, N-, or P-releasing enzymes on rooted soil surfaces. When an extracellular enzyme decomposed substrate fixed to the paper, the cleavage resulted in a colorimetric or fluorometric tag that remained on the paper. These zymograph images were digitized along with photographs of roots in the sample area. I developed a quantitative framework for analyzing these image data. I used two regression models, a spatial error model that accounted for the autocorrelation between image pixels, and a “break-point” regression model that estimated the location of a change in the slope of the relationship between extracellular enzyme activity and distance from a root. I used this break point as a proxy for the rhizosphere extent. The rhizosphere extent varied depending on the resolution of image analysis, as a result of increased sample size, background staining, or misalignment of images. Nevertheless, I found that extracellular enzyme activity was concentrated near the surface of roots. The break points that I estimated suggest that the rhizosphere can extend beyond 2 mm from the root surface, suggesting that recent estimates of rhizosphere contributions to decomposition and nutrient mineralization are very conservative (Finzi et al., 2015).

In Chapter 5, my objective was to develop a combined model of microbial physiology and extracellular enzyme activity to assess how climate change may affect plant–microbe interactions, soil organic matter decomposition, and soil C storage. I merged two existing models: the Dual-Arrhenius Michaelis Menten (DAMM) model (Davidson et al., 2012) and the Microbial Carbon and Nitrogen Physiology (MCNiP) model (Finzi et al., 2015). The combined model reproduced heterotrophic respiration
measured in a trenched plot at the Harvard Forest in central Massachusetts. It also captured the seasonal pattern of C efflux better than MCNiP alone, and predicted C efflux during wet-up events better than both models alone. DAMM-MC NiP predicted that stands with a wide C:N ratio [litter, roots, microbial biomass] stored more C, consistent with recent empirical findings at the global scale (Averill et al., 2014). When subject to a series of climate change simulations, the model predicted that C mineralization increases with either +5ºC warming or a 50% increase in soil moisture. The response to added soil moisture was unexpected given that oxygen (O2) availability limits microbial activity at high soil moisture, and suggests that in this model O2 limitation has a small effect on microbial respiration relative to substrate diffusion.

This research demonstrates that belowground C allocation affects C balance on the plot-to-global scale. There are broad differences in root phenology across biomes and plant types, and in a mid-latitude temperate forest, there are stand-level differences in both the magnitude and partitioning of C belowground, driven primarily by allocation to root growth. A significant amount of root C is allocated to the microbial community, where it stimulates microbial activity. I demonstrate that the C:N ratio of root exudates influences both the magnitude of SOM decomposition and the mechanism by which decomposition is stimulated. These findings demonstrate the importance of plant roots to the terrestrial C cycle.
CHAPTER TWO: ARE ABOVE AND BELOWGROUND PHENOLOGY IN SYNC?

Abstract

Globally, root production accounts for 33-67% of terrestrial net primary productivity and influences decomposition via root production and turnover, carbon (C) allocation to mycorrhizal fungi and root exudation. As recognized aboveground, the timing of phenological events affects terrestrial C balance, yet there is no parallel understanding for belowground phenology. In this paper I examine the phenology of root production and its relationship to temperature, soil moisture, and aboveground phenology. Synthesizing 87 observations of whole plant phenology from 40 studies, I found that on average root growth occurs 25 ± 8 days after shoot growth but that the offset between the peak in root and shoot growth varies > 200 days across biomes (boreal, temperate, Mediterranean, and subtropical). Root and shoot growth are positively correlated with median monthly temperature and mean monthly precipitation in boreal, temperate, and subtropical biomes. However, a temperature hysteresis in these biomes leads to the hypothesis that internal controls over C allocation to roots are an equally, if not more, important driver of phenology. The specific mechanism(s) are as yet unclear but are likely mediated by some combination of photoassimilate supply, hormonal signaling, and growth form.
Introduction

It is widely acknowledged that roots play a fundamental role in terrestrial C cycling, consuming up to 70% of net primary production (Grier et al., 1981, Jackson et al., 1997), yet there is little understanding of the factors controlling patterns of root growth (Pregitzer et al., 2000). Fine root production can occur in a single flush but often occurs in multiple flushes throughout the growing season (Bevington & Castle, 1985, Harris et al., 1995, Reich et al., 1980, Steinaker et al., 2010). Soil temperature, moisture and nutrient availability affect the growth of roots (Fukuzawa et al., 2013, Noguchi et al., 2013), but there is often no temporal correlation between these abiotic factors and root growth apart from obvious growing-to-nongrowing season transitions (Hendrick & Pregitzer, 1996, Joslin et al., 2001; Table 2.S1). In these cases internal signaling such as photoassimilate transport may control root growth (Sloan & Jacobs, 2008), such that roots cannot grow when shoots are consuming the majority of photoassimilate. Supporting this, several studies have observed patterns of alternating root and shoot growth (Cardon et al., 2002, Drew & Ledig, 1980, Mickelbart et al., 2012, Reich et al., 1980).

Most conceptual and terrestrial biosphere models allocate C belowground as a fixed fraction of net C uptake, which by definition makes root phenology synchronous with aboveground growth (Table 2.1). The purpose of this review is to show that current data support asynchrony between above and belowground growth. I acknowledge that the available data are few and that inference regarding broad-scale patterns is subject to change as more data become available. I hope, however, that bringing currently available
data to light in this regard will generate the collection of new data, and refine current understanding of belowground phenology and its relevance at the ecosystem scale.

A compilation of available data

It is commonly assumed that root growth peaks early in the growing season and is therefore synchronous with aboveground growth (Medvigy et al., 2009). While this can be true (Misson et al., 2006, Scagel et al., 2007), there are many exceptions (Lahti et al., 2005, Palacio & Montserrat-Martí, 2007, Willaume & Pagès, 2006). Some studies report root growth lagging shoot growth by several weeks, an observation attributed to air temperature warming faster than soil temperature in the spring (Steinaker & Wilson, 2008). Others report root growth preceding shoot growth by several weeks to months (Broschat, 1998, Ploetz et al., 1992). In a common garden study in Pennsylvania, some species such as Acer negundo and Pinus strobus had large interannual variability in root phenology while others such as Liriodendron tulipfera did not (McCormack et al., 2014), suggesting that some trees may be environmentally cued while others are inflexible in their timing (i.e., phenological programming sensu Hendrick & Pregitzer, 1996, Joslin et al., 2001).

To address broad-scale patterns in phenology, I conducted a literature survey to quantify the offset between the maximum in root and shoot growth in woody and herbaceous perennial plants. Web of Science was searched using the following keywords: belowground phenology, root phenology, root allocation, and root growth in combination with shoot phenology, aboveground, stem growth, leaf out, budburst or greenness. I considered only studies that simultaneously measured both root and shoot production. Of
the 13,934 results from the keyword search, only 40 studies had suitable data. There were a total of 87 datasets containing shoot and root growth for 63 species (see Paper selection, Gross and net root production, Monthly root and shoot growth, and Temperature and precipitation data in Supplementary Information for a more detailed description of the data used in this survey). The data are compiled in Tables 2.S2 and 2.S3. Each observation was classified into one of four biomes (boreal, temperate, Mediterranean and subtropical) based on Whittaker’s biome classification system (1970).

In order to visualize findings from the literature, I quantified the difference between peak shoot and root growth using the equation,

\[ \text{Offset (days)} = \text{DOY}_{\text{maximum root}} - \text{DOY}_{\text{maximum shoot}} \]  \[1\]

where DOY is the day of year of maximum root or shoot growth as indicated. In plants with multiple root and shoot flushes, maximum root or shoot growth rate was used to calculate offset. Positive offset values therefore indicate peak shoot growth preceding the peak in root growth whereas negative values would indicate root growth preceding the peak in shoot growth. Differences in root and shoot data collection methods marginally affect offset \( (F_{6,80} = 2.17, P = 0.055) \), so they were included in stepwise model selection (see Offset in Supplementary Information). Soil coring methods tended to detect later root growth relative to shoot growth.

**Primary Data Findings**

There was wide variation in the timing of maximum shoot growth relative to root growth (Figure 2.1). In the majority of cases maximum shoot production occurred before root production (offset > 0 in 54 out of 87 observations) and the mean offset for all
studies (25 ± 8 days) was greater than zero (one sample t-test; \( t = 3.15, df = 86, P < 0.01 \)) indicating that root and shoot growth are not synchronous on a broad geographic scale.

There was no difference in offset between tree, shrub and herbaceous growth forms.

As the data are largely from extratropical, northern hemisphere localities, there is a significant correlation between growth and temperature (Table 2.2, Figure 2.51). At the biome scale, subtropical plants were significantly different from all other biomes, with the peak in root growth occurring 45 ± 19 (n = 11) days earlier than shoot growth, whereas offset in boreal, Mediterranean, and temperate biomes occurred 48 ± 8 (n = 20), 36 ± 19 (n = 11), and 28 ± 12 (n = 45) days after shoot production, respectively (Figure 2.2a). The generally late root relative to shoot growth in boreal biomes may, however, be confounded with tree growth form. Conifer (n = 14) root growth peaked 44 ± 12 days later than deciduous tree species (n = 20, Figure 2.2b).

Similar to temperature, there is a positive linear correlation between growth and mean monthly precipitation (MMP) in boreal and subtropical biomes (Table 2.2). In these biomes median monthly temperature (MMT) and MMP are highly correlated (\( \rho = 0.89 \) and \( \rho = 0.85 \) for boreal and subtropical climate variables respectively), so it is difficult to separate the precipitation from the temperature effect. By contrast, precipitation predicts growth poorly in temperate and Mediterranean biomes (Table 2.2). In the temperate data set precipitation did not have large seasonal variation (Figure 2.3b). In the Mediterranean data set shoot growth occurs in a large spring pulse following winter rain, whereas root growth appears to proceed at a steadily rising rate over the year (Figure 2.3c).
To visualize the phenology of root relative to shoot growth, I plotted the proportion of peak root growth as a function of peak shoot growth and generated a hysteresis plot for each biome. Deviations from the 1:1 line indicate dominance of root relative to shoot growth (or vice versa) across the year (Figure 2.3e-h). In boreal ecosystems root growth remains low throughout spring shoot expansion with the largest proportion of root growth observed in the summer through autumn (Figure 2.3e). In temperate ecosystems, however, root growth is entirely proportional to shoot growth, with all data plotting closely to the 1:1 line (Figure 2.3f). The Mediterranean observations are unlike the others. There is no clear hysteresis between root and shoot growth (Figure 2.3g). The subtropical biome is similar to the temperate biome in that root growth mirrors shoot growth not deviating from the 1:1 line (Figure 2.3h).

**Some Implications of the Data**

The available data suggest that root and shoot growth is largely asynchronous. At broad spatial scales temperature and precipitation influence this asynchrony, for example, the positive offset between peak shoot and root growth with decreasing annual temperatures (data not shown). As suggested by Steinaker & Wilson (2008), air temperature rises more rapidly than soil temperature in the spring and hence root growth is delayed later into the spring or summer (Figure 2.3a-d). The boreal dataset supports this hypothesis. A corollary to this observation is that thermal buffering allows soils to remain warm through the autumn and, as a result, the duration of root production can be 40% longer than shoot production (Steinaker & Wilson, 2008, Steinaker et al., 2010).
Once again, this observation is strongly supported by the boreal zone data set (Figure 2.3a).

There is, however, at least as much variation in the offset between peak root and shoot growth within and between biomes as that explained by climate variables (Table 2.2). This leads to the ecologically interesting hypothesis that endogenous control of plant C allocation is an important driver of root phenology. What evidence is there in support of this hypothesis?

For the purposes of this paper, I define endogenous cuing as any factor that influences the growth of roots other than direct effects of temperature and precipitation. One of the clearest examples of endogenous root cuing is the production, storage and transport of photoassimilate (Palmroth et al., 2006, Pregitzer et al., 2000). There are important stores of carbohydrates in plants that can fuel production (Richardson et al., 2013), and root growth depends on these stores as well as newly fixed C from aboveground organs. Isotopic labelling studies have confirmed that substantial C used in root biomass is newly fixed (Keel et al., 2006, Trumbore et al., 2006). Root growth stops or is greatly reduced in response to experimental manipulations such as girdling and stem chilling that cut off the supply of photoassimilates from the canopy (Högberg et al., 2001, Johnsen et al., 2006) and thus belowground phenology must be in part regulated by aboveground phenology (Litton et al., 2007).

Does the difference in root growth phenology implicate the supply of photoassimilate? It does appear to provide a parsimonious explanation. Leaf area and photosynthetic rates in temperate deciduous forests tend to be highest in the spring and
decline through late summer and rapidly in autumn as a consequence of canopy senescence (Wu et al., 2010). This seasonality is highly synchronous with the most active period of root growth, and the progressive summer decline in photosynthesis and autumnal leaf senescence correlate with reductions in root growth (Figure 2.3f). The prolonged autumnal root growth in the boreal zone may also reflect photoassimilate control in that the retention of live-needles in the canopy sustains the supply of photoassimilate even as temperatures cool and day length declines (but soils stay warm). Additional support for photoassimilate-regulated autumnal root growth is observed when separating the boreal dataset into evergreen trees vs. deciduous woody and herbaceous species. Doing so shows that autumnal root growth in evergreen trees dominates the hysteresis (Figure 2.S2a) in Figure 2.3f. Autumnal root growth in the deciduous plants follows that found in the temperate biome data (i.e., maximum root growth in the spring and very little in the autumn; Figure 2.S2b). Boreal evergreen trees may also utilize stored C during late season root growth (Nordgren et al., 2003).

The subtropical and Mediterranean data are challenging to interpret. The subtropical data come from evergreen species, largely palms, in a consistently warm environment, yet there is a distinct phenology in shoot and root production with the peak in both following the wettest months of the year (Figure 2.3d). I speculate that the distinct phenology may be analogous to observations from seasonally dry Amazonian rainforests where evapotranspiration rates are highest in the dry season (Hutyra et al., 2007) because of reductions in cloudiness and light-limitation of photosynthesis (Restrepo-Coupe et al., 2013). It is possible that the reduction in late summer and autumnal precipitation is
sufficiently large that PAR does not limit photosynthesis during this period of time and peak growth occurs.

The Mediterranean observations are the most challenging to interpret under the assumption that the majority of C used for root growth is newly fixed. Shoot and root growth are not correlated with MMT or MMP and root growth is apparently decoupled from shoot growth (Figure 2.3g). The one exception is a concentrated pulse of vernal shoot growth following winter precipitation and the delay in peak root relative to shoot growth (Figure 2.3c). The strong asynchrony between shoot and root growth that extends across the year, however, suggests that endogenous cuing and subsequent allocation of stored carbohydrates is a dominant driver of root growth in Mediterranean plants. Root and stem nonstructural carbohydrates generally decline during the growing season and re-accumulate in autumn (Loescher et al., 1990, Richardson et al., 2013), as stored carbohydrates are allocated to respiration and growth during the growing season. However, limited data availability in this biome may prevent any meaningful conclusions.

Finally, I note that the time scale of the data analysis here cannot address the occurrence of alternating above and belowground growth, for example, as found in Quercus spp. (Cardon et al., 2002, Reich et al., 1980). Whether this is common is not well known, but at least these data suggest coordination of C allocation across the growing season, which may be mediated by both above and belowground plant organs. For example, roots produce and transport several shoot regulating hormones acropetally, such as abscisic acid, cytokinin and strigolactone, that can affect stomatal closure, shoot
and bud growth respectively (Domagalska & Leyser, 2011, Sharp, 2002). Roots may also control photoassimilate transport by modifying the rate that photoassimilates in the phloem are taken up by root tissues (Patrick, 1997) that feedback to genetic regulatory networks (Koch, 1996). Numerous studies provide support for shared control of C allocation (Davidson & Holbrook, 2009, Farrar & Jones, 2008).

**Moving Forward**

At the extreme, root and shoot phenology can be offset by ±200 days, and both are mechanistically linked by temperature, water and C allocation. This dataset establishes possible generalizations regarding root and shoot phenology based on biome and growth form (i.e., evergreen, deciduous). Temperature and moisture are positively correlated with the phenology of both shoot and root growth in three of four biomes suggesting that abiotic factors both directly and indirectly affect root physiology. Endogenous factors (e.g., allocation of photoassimilate, source-sink dynamics, hormonal control) are also likely to be important drivers of phenology but as yet I am not able to make any broad conclusions with the possible exception that photoassimilate supply, storage and transport are key drivers of root growth phenology. The temperature hysteresis in Figure 2.3 provides support for endogenous controls—particularly the boreal and Mediterranean datasets. At the present time, there are few data available in the literature probably because of the difficulty in making measurements of seasonal root growth. Current datasets are derived from techniques that are relatively indirect (soil respiration), often destructive (coring), labor intensive (minirhizotrons) and hence expensive to implement. In addition to the paucity of data, many methods are difficult to
compare often with known biases (e.g., minirhizotron vs. isotopic methods; Guo et al., 2008), and suffer from chronic under-sampling (Taylor et al., 2013). However, for my purposes these methods capture temporal change in root length or biomass well enough to characterize its timing, though perhaps not its magnitude. I hope that improved scaling methods (Taylor et al., 2014) and standardization across large networks (Keller, 2010) will alleviate sampling difficulties and allow for more accurate and generalizable data to emerge over time. Root growth is an important conduit for photosynthetically-fixed C into the soil with well-established feedbacks on C and N cycling (Averill et al., 2014, Brzostek et al., 2013, Drake et al., 2011, Schmidt et al., 2011). Quantitative models assume the phenology of root growth is synchronous with that of aboveground phenology despite empirical evidence to the contrary. Whether the addition of belowground phenology will affect total C efflux in terrestrial biosphere models is presently unknown. In the temperate biome where phenology is largely in sync such a change may be unnecessary, but in boreal biomes late season root allocation may explain observed fall increases in soil respiration that are currently poorly explained by temperature and soil moisture (Davidson et al., 2006, Giasson et al., 2013). Since both root growth and decomposition are known to be temperature-sensitive, and the latter also substrate limited, understanding the phenology and drivers of above- vs. belowground-C allocation is important for estimating ecosystem C fluxes under global change.

Acknowledgements

I would like to thank Edward Brzostek, John Drake, Allison Gill, Luke McCormack, Andrew Richardson, Andrew Reinmann, Patrick Sorensen and Anthony
Walker for helpful comments and discussion. This research was supported by the American Association of University Women (AAUW) American Dissertation Fellowship, Office of Science (BER), U.S. Department of Energy (Grant No. 10-DOE-1053) and by the National Science Foundation (DEB-0743564).
<table>
<thead>
<tr>
<th>Model</th>
<th>Aboveground Phenology</th>
<th>Belowground Phenology</th>
<th>Allocation</th>
<th>Citation</th>
</tr>
</thead>
<tbody>
<tr>
<td>TRIFFID</td>
<td>Temperature</td>
<td>n/a</td>
<td>Allometric relationships (root:leaf, stem:leaf) and partitioning into ‘spreading’ and ‘growth’ based on LAI</td>
<td>Cox 2001</td>
</tr>
<tr>
<td>Hyland</td>
<td>n/a</td>
<td>n/a</td>
<td>Fixed coefficients</td>
<td>Friend et al. 1997, Friend and White 2000, Levy et al. 2004</td>
</tr>
<tr>
<td>PnET-BGC</td>
<td>GDD</td>
<td>Monthly turnover rate</td>
<td>Linear function of foliar production</td>
<td>Kram et al. 1999</td>
</tr>
<tr>
<td>ORCHIDEE</td>
<td>GDD, Soil moisture</td>
<td>Some root growth at leaf onset, stress can cause root death</td>
<td>Allocation fractions for leaves, stems, roots determined based on water, light and N availability</td>
<td>Krinner et al. 2005</td>
</tr>
<tr>
<td>3PG</td>
<td>PAR, number of frost days</td>
<td>Stress affects root growth</td>
<td>Allometric relationships</td>
<td>Landsberg and Waring 1997</td>
</tr>
<tr>
<td>IBIS</td>
<td>Temperature, Productivity threshold</td>
<td>n/a</td>
<td>Fixed fraction of C uptake allocated to leaves, stems, roots</td>
<td>McGuire et al. 2001</td>
</tr>
<tr>
<td>TEM</td>
<td>Evapotranspiration</td>
<td>n/a</td>
<td>Not explicit</td>
<td>McGuire et al. 2001</td>
</tr>
<tr>
<td>ED2</td>
<td>Logistic functions derived from MODIS data</td>
<td>n/a</td>
<td>PFT-dependent allocation relationships, root:leaf varies with water or N limitation</td>
<td>Medvigy et al. 2009</td>
</tr>
<tr>
<td>CLM 4.0</td>
<td>GDD, Soil moisture, Daylength</td>
<td>n/a</td>
<td>Fixed ratios of fine root:leaf and coarse root:stem</td>
<td>Oleson et al. 2010, Thornton and Zimmerman 2007</td>
</tr>
<tr>
<td>TREGRO</td>
<td>GDD, NSC status</td>
<td>GDD, NSC status</td>
<td>Allocated based on priority leaf&gt;branch&gt;stem&gt;coarse roots&gt;fine roots</td>
<td>Weinstein et al. 1991</td>
</tr>
<tr>
<td>BIOME-BGC</td>
<td>Temperature, Photoperiod, Climate-specific GDD</td>
<td>n/a</td>
<td>PFT-dependent allometric ratios such as root:leaf and stem:leaf</td>
<td>White et al. 1997</td>
</tr>
<tr>
<td>Sheffield-DGVM</td>
<td>Temperature and soil moisture</td>
<td>n/a</td>
<td>LAI optimized to maximize C uptake, wood and roots receive a constant PFT-dependent fraction of remaining C</td>
<td>Woodward et al. 1995, Woodward and Lomas 2004</td>
</tr>
</tbody>
</table>

n/a = not applicable
Table 2.2. Regression statistics for the proportion of maximum monthly root or shoot growth as a function of median monthly temperature (MMT) or mean monthly precipitation (MMP). The slope estimate (β) for MMT are in units of growth per °C and MMP growth per mm precipitation. Significant variables and overall model significance are indicated by: NS = not significant, * P < 0.05, ** P < 0.01 and *** P < 0.001.

<table>
<thead>
<tr>
<th>Biome</th>
<th>Organ</th>
<th>MMT</th>
<th>$R_{adj}^2$</th>
<th>MMP</th>
<th>$R_{adj}^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>All biomes</td>
<td>Root</td>
<td>0.02***</td>
<td>0.48</td>
<td>0.01*</td>
<td>0.09</td>
</tr>
<tr>
<td></td>
<td>Shoot</td>
<td>0.01**</td>
<td>0.21</td>
<td>0.01**</td>
<td>0.17</td>
</tr>
<tr>
<td>Boreal</td>
<td>Root</td>
<td>0.02***</td>
<td>0.79</td>
<td>0.01***</td>
<td>0.83</td>
</tr>
<tr>
<td></td>
<td>Shoot</td>
<td>0.02*</td>
<td>0.47</td>
<td>0.01**</td>
<td>0.67</td>
</tr>
<tr>
<td>Temperate</td>
<td>Root</td>
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<td>0.80</td>
<td>0.01**</td>
<td>0.01</td>
</tr>
<tr>
<td></td>
<td>Shoot</td>
<td>0.02***</td>
<td>0.75</td>
<td>0.011NS</td>
<td>0.01</td>
</tr>
<tr>
<td>Mediterranean</td>
<td>Root</td>
<td>0.01NS</td>
<td>0.01</td>
<td>0.01NS</td>
<td>0.18</td>
</tr>
<tr>
<td></td>
<td>Shoot</td>
<td>0.01NS</td>
<td>0.00</td>
<td>0.01NS</td>
<td>0.01</td>
</tr>
<tr>
<td>Subtropical</td>
<td>Root</td>
<td>0.05**</td>
<td>0.61</td>
<td>0.01*</td>
<td>0.37</td>
</tr>
<tr>
<td></td>
<td>Shoot</td>
<td>0.04*</td>
<td>0.38</td>
<td>0.01*</td>
<td>0.31</td>
</tr>
</tbody>
</table>
Figure 2.1. Kernel density curve of the offset in days between the maximum in root and shoot production. A kernel density curve is analogous to a histogram, but rather than showing counts of binned data, it estimates a probability density function of offset using sample data. The curve is smoothed using a kernel bandwidth of 1. Offset is defined as DOY_maximum_root – DOY_maximum_shoot where DOY refers to day of year from 1 to 365. Data include 63 tree, shrub, and herbaceous species from 40 studies (Table 2.5), grouped by biome. The black vertical dotted line is the grand mean of all offset values.
Figure 2.2. (a) Offset in days between maximum shoot and root production for 87 observations averaged across four biomes: boreal, temperate, Mediterranean and subtropical (Table 2.S2). Letters indicate a statistically significant difference in means ($\alpha = 0.05$) calculated using Tukey’s HSD after one way ANOVA (offset ~ biome + root collection method, biome: $F_{3,80} = 5.0, P = 0.0032$, root collection method: $F_{3,80} = 2.67, P = 0.053$, model $F_{6,80} = 3.83, P = 0.0021$). (b) Deciduous ($n = 20$) trees had a significantly smaller offset than did evergreen ($n = 14$) trees (ANOVA, $F_{1,32} = 7.52, P = 0.009$).
Figure 2.3. The proportion of maximum monthly root and shoot growth for each month in (a) boreal, (b) temperate, (c) Mediterranean, and (d) subtropical biomes (Table 2.8). In panels a-d, dark brown corresponds to root growth and light green is shoot growth. The blue dotted line is mean monthly precipitation (mm, right side y-axis). The color bar across the top is a heat map showing seasonal temperatures ranging from -10°C (purple) to 25°C (red), with 0°C as bright blue. Panels e-h plot the proportion of maximum monthly root vs. shoot growth. In these panels, black lines join consecutive months and the direction of the arrowheads indicates time from January to December. This approach assumes that shoot growth is a suitable proxy for the initiation of photosynthesis. Calculating the proportion of peak root or shoot growth rather than absolute growth rates enables us to plot the different types of data on the same y-axis (i.e., minirhizotron vs. soil coring). Note that these proportions are not a probability distribution function (i.e., area under the curve ≠ 1) and that no point equals 1 because multiple studies with differently timed maximum growth were averaged.
Supplementary Information

Paper selection

Web of Science was searched using the following keyword search: (belowground phenology OR root phenology OR root allocation OR root growth) AND (shoot phenology OR aboveground OR stem growth OR leaf out OR budburst OR greenness), in order to capture a variety of above and belowground measurements. There were 13,934 returns on Feb 3, 2014. I evaluated all papers at the abstract level for mention of seasonal root and/or shoot data. All candidate papers were read in full. Data including measurements of root and either leaf or stem production were included. Studies measuring transpiration rates, flowering, or single phenological events with no time series were not included.

Gross and net root production

In this study, offset is calculated using both gross and net root production. The phenology of net root production may be affected by seasonal variations in root mortality, especially when growth and mortality are asynchronous [1]. However, studies in temperate and boreal regions show that rates of root mortality are roughly synchronous with root growth [2, 3], implying that in these cases measurements of net root production may underestimate the magnitude of growth but not the timing of growth peaks. I found no significant difference between offset calculated using either gross or net root production (ANOVA, $F_{1,86} = 0.788, P = 0.38$) in the observations used in this study. These and all other analyses were performed in R [4].
Offset

Each observation was classified into one of four biomes based on Whittaker’s biome classification system (1970). When median annual temperature and mean annual precipitation were close to the edge of a biome, classification was determined using USDA Natural Resources Conservation Service Major Biomes Map [5]. Day of year for peak root and shoot production was calculated as a mean of replicates with varying sample size (n = 2–48). For samples that produced multiple shoot and root flushes, the day of year of the largest peak was used to represent the above or belowground peak. Offset was calculated as the number of days between the peak in shoot and root production \((\text{offset} = \text{DOY}_{\text{peak root}} - \text{DOY}_{\text{peak shoot}})\). Negative offset values indicate that peak root production preceded peak shoot production. To explore possible biome-specific effects, I used analysis of variance to test for the effect of biome, root collection methods, and study on offset. Differences in root and shoot data collection methods (ANOVA: offset ~ root collection method + shoot collection method, \(F_{6,80} = 2.17, P = 0.055\)), and study author did not affect offset (ANOVA: offset ~ study author, \(F_{1,85} = 0.00004, P = 0.99\)). Stepwise model selection using explanatory variables 1) biome, 2) data collection method, and 3) study author indicated that the [biome + root collection method] model had the lowest AIC value (stepAIC[library:MASS] Initial model: offset ~ biome + shoot collection method + root collection method + study author, AIC = 749.6731; Final model: offset ~ biome + root collection method, AIC=737.1790). Offset between the peak in root and shoot production in the subtropical biome was significantly different from other biomes (Tukey’s HSD after ANOVA: offset ~ biome + root collection method, biome:
F_{3,80} = 5.0, P = 0.0032, root collection method: F_{3,80} = 2.67, P = 0.053, model F_{6,80} = 3.83, P = 0.0021, Figure 2.S1a). Offset was not affected by plant growth form (herbaceous, tree or shrub; ANOVA: offset~growth form, F_{2,84} = 0.827, P = 0.44).

Monthly root and shoot growth

Measurements from literature were used to estimate monthly root and shoot growth rates for each biome. Growth rate of roots and shoots were tabulated from each study (excluding those that did not include seasonal data [i.e., measured DOY 1 as start of experiment rather than the Julian day]) and converted to a % of the maximum growth rate. Growth rates were averaged for each month with > 5 observations. Most rates do not exceed 1 because multiple studies with differently timed maximum growth were averaged (Table 2.S3). For all biomes, root and shoot growth together have a positive relationship with MMT and MMP (Growth = 0.0148*MMT + 0.123, F_{1,85} = 41.05, P < 0.0001, R_{adj}^2 = 0.32; Growth = 0.00190*MMP + 0.240, F_{1,85} = 14.19, P < 0.001, R_{adj}^2 = 0.13). There was no significant interaction effect between MMT and MMP when both were included in a model (MAT:MAP P = 0.15, model F_{3,83} = 17.22, P < 0.0001, R_{adj}^2 = 0.36).

Temperature and precipitation data

To explore whether the seasonal variation in organ growth was the result of changes in temperature or precipitation, I obtained monthly daily maximum and minimum temperature as well as monthly mean total precipitation data for the station
located nearest each study site from the World Weather Information Service [6]. Means are based on 30 year averages from 1959-1997 (Greece), 1961–1990 (Chile, Czech Republic, Malaysia, Sweden), 1971–2000 (Canada, Finland, France, the Netherlands, Spain), or 1981-2010 (Japan, United Kingdom, United States of America). Daily maximum and minimum temperatures for each month were averaged to obtain a median monthly temperature (°C). Precipitation was reported as monthly mean total precipitation (mm). Precipitation refers to all types of precipitation in Sweden, Canada, France, Netherlands, and Spain. Only rainfall was measured in Greece, Chile, Czech Republic, Malaysia, Finland, Japan, United Kingdom, and United States of America.
Figure 2. S1. Proportion root (a) and shoot (b) growth for data in all biomes plotted against median monthly temperature (°C). Color and symbol shape indicate biome. Regression statistics are found in Table 2.2.
Figure 2.S2. (a) The relationship between root growth and MMT in boreal evergreen tree roots only. Arrows indicate direction of time from April to November. (b) The relationship between root growth and MMT in boreal deciduous tree, shrub and herbaceous plant roots. Arrows indicate the direction of time from March to September.
Table 2.5. Phenological relationships between aboveground and belowground processes as well as potential drivers of belowground phenology, their direction and strength of effect, if given. ST, temperature; SWC, soil water content; E, endogenous.

<table>
<thead>
<tr>
<th>Biome</th>
<th>Location</th>
<th>Species</th>
<th>Relationship of above- to belowground phenology</th>
<th>Possible Drivers</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boreal</td>
<td>Northern Savonia, Finland</td>
<td><em>Pinus sylvestris</em></td>
<td>Shoot elongation before peak in root production</td>
<td>T (+/-, hysteresis), E</td>
<td>[7]</td>
</tr>
<tr>
<td></td>
<td>Northern Savonia, Finland</td>
<td><em>Picea abies</em></td>
<td>Root growth before and after shoot growth</td>
<td>ST (+)</td>
<td>[8]</td>
</tr>
<tr>
<td></td>
<td>Uppland, Sweden</td>
<td><em>Salix viminalis</em></td>
<td>Max height growth before max change in fine root number</td>
<td>n/a</td>
<td>[9]</td>
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<td></td>
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<td><em>Stipa comata</em></td>
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<td></td>
<td></td>
<td>Carex spp.</td>
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<td></td>
<td></td>
<td><em>Bouteloua gracilis</em></td>
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<td></td>
<td></td>
<td>Agropyron spp.</td>
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<tr>
<td></td>
<td></td>
<td><em>Koeleria gracilis</em></td>
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<td></td>
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</tr>
<tr>
<td></td>
<td>Saskatchewan, Canada</td>
<td>Poa spp.</td>
<td>Shoot elongation before peak in root production</td>
<td>T (grassland: ( r = -0.16, P &lt; 0.05 ), forest: ( r = -0.20, P &lt; 0.05 )), SWC (grassland: ( r = +0.58, P &lt; 0.001 ), forest: ( r = +0.60, P &lt; 0.001 ))</td>
<td>[10]</td>
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<td></td>
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<td><em>Selaginella densa</em></td>
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<td><em>Populus tremuloides</em></td>
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<td></td>
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<td><em>Symphoricarpos occidentalis</em></td>
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<td>Rubus idaeus</td>
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<td>Festuca rubra</td>
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<td><em>Koeleria gracilis</em></td>
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<tr>
<td></td>
<td>Saskatchewan, Canada</td>
<td>Poa compressa</td>
<td>Shoot elongation before peak in root production</td>
<td>T (herbaceous: ( r = +0.42 ) to ( +0.63 ), ( P &lt; 0.01 ) to ( P &lt; 0.001 ), woody: ( r = +0.23 ) to ( +0.62 ), ( P &lt; 0.05 ) to ( P &lt; 0.001 )), SWC (herbaceous: ( r = -0.01 ) to ( -0.78 ), ns to ( P &lt; 0.001 ), woody: ( r = -0.07 ) to ( -0.40 ), ns to ( P &lt; 0.01 ))</td>
<td>[11]</td>
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<tr>
<td></td>
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<td><em>Bouteloua gracilis</em></td>
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<td><em>Schizachyrium scoparium</em></td>
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<td><em>Symphoricarpos occidentalis</em></td>
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<td>Biome</td>
<td>Location</td>
<td>Species</td>
<td>Relationship of above- to belowground phenology</td>
<td>Possible Drivers</td>
<td>Reference</td>
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<td>---------------------------</td>
<td>----------------------------------------------</td>
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<td>-----------</td>
</tr>
<tr>
<td>Boreal</td>
<td>Northern Savonia, Finland</td>
<td><em>Pinus sylvestris</em>&lt;br&gt;<em>Picea abies</em></td>
<td>Synchronous for pine seedlings, root growth initiated before shoot elongation in spruce seedlings</td>
<td>ST (+)</td>
<td>[12]</td>
</tr>
<tr>
<td>Mediterranean</td>
<td>California, USA</td>
<td><em>Pinus ponderosa</em></td>
<td>Synchronous</td>
<td>ST = (+), SWC = (+)</td>
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</tr>
<tr>
<td></td>
<td>Huesca, Spain</td>
<td><em>Echinochloa horridum</em>&lt;br&gt;<em>Salvia lavandulifolia</em>&lt;br&gt;<em>Lepidium subulatum</em>&lt;br&gt;<em>Linum suffruticosum</em></td>
<td>Shoot elongation before peak in root production, offset</td>
<td>T(+), SWC(+), E</td>
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<tr>
<td></td>
<td>California, USA</td>
<td><em>Pinus ponderosa</em></td>
<td>Variable</td>
<td>ST (+)</td>
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<td></td>
<td>Nevada, USA</td>
<td><em>Larrea tridentata</em>&lt;br&gt;<em>Ambrosia dumosa</em>&lt;br&gt;<em>Ephedra nevadensis</em>&lt;br&gt;<em>Lycium pallidum</em></td>
<td>n/a</td>
<td>SWC (r = +0.245 to +0.8, P = 0.217 to P&lt;0.001)</td>
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<td>Subtropical</td>
<td>Florida, USA</td>
<td><em>Citrus sinensis x Citrus jambhiri</em>&lt;br&gt;<em>Citrus sinensis</em> x <em>Poncirus trifoliata</em></td>
<td>Alternating shoot and root flushes</td>
<td>ST (r = +0.74 to +0.87, P &lt;0.05), SWC (+)</td>
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<td><em>Roystonea regia</em>&lt;br&gt;<em>Cocos nucifera</em>&lt;br&gt;<em>Syagrus romanzoffiana</em>&lt;br&gt;<em>Phoenix roebelenii</em></td>
<td>Root growth before shoot growth</td>
<td>ST (r = +0.354 to +0.697, P &lt; 0.0001)</td>
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<td>Northern Territory, Australia</td>
<td><em>Eucalyptus miniata</em>&lt;br&gt;<em>Eucalyptus tetrodonta</em>&lt;br&gt;<em>Sorghum spp.</em></td>
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<td>SWC (+)</td>
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<td>Species</td>
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<td>Possible Drivers</td>
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<td><strong>Subtropical</strong></td>
<td>Sabah, Malaysia</td>
<td>Dipterocarpaceae, Meliaceae, Tilliaceae, Lauraceae, Euphorbiaceae</td>
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<td>SWC ($r = +0.39$, $P &lt; 0.05$)</td>
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<td></td>
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<td>Persea americana</td>
<td>Root elongation before shoot elongation</td>
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<td>Barro Colorado Island, Panama</td>
<td>Tropical moist forest</td>
<td>n/a</td>
<td>SWC (+)</td>
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<td><strong>Temperate</strong></td>
<td>New York, USA</td>
<td>Acer saccharum, Fagus grandifolia, Betula alleghaniensis, Acer rubrum</td>
<td>Initiation of root growth synchronous with budswell</td>
<td>T ($r = 0.871$, $P &lt; 0.01$), E</td>
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<tr>
<td></td>
<td>Connecticut, USA</td>
<td>Quercus rubra</td>
<td>Alternating shoot and root flushes</td>
<td>E</td>
<td>[24]</td>
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<tr>
<td></td>
<td>Quebec, Canada</td>
<td>Acer saccharum, Fagus grandifolia, Betula alleghaniensis, Populus grandidentata, Acer rubrum</td>
<td>Offset, root growth before peak in diameter increment in one year, after in another</td>
<td>ST (-), SWC (+), diameter increment (-), N mineralization (-)</td>
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<td>Dumfries, United Kingdom</td>
<td>Picea sitchensis</td>
<td>n/a</td>
<td>ST, SWC (+, $P &lt; 0.05$), E</td>
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<tr>
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<td>Cumbria, United Kingdom</td>
<td>Festuca ovina, Juncus squarroso, Nardus stricta</td>
<td>n/a</td>
<td>model $r = +0.608$, radiation flux (regression coefficient: +5.75, $P &lt; 0.001$), surface temp (regression coefficient: +2.86, $P &lt; 0.01$), air temp (regression coefficient: -2.59, $P &lt; 0.01$)</td>
<td>[27]</td>
</tr>
<tr>
<td></td>
<td>Cumbria, United Kingdom</td>
<td>Grassland dominated by Festuca ovina and Agrostis capillaris</td>
<td>n/a</td>
<td>radiation flux ($r = +0.520$)</td>
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<td>Possible Drivers</td>
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<tr>
<td>Temperate</td>
<td>Michigan, USA</td>
<td><em>Quercus alba</em></td>
<td>Some root production before leaf expansion, 80% of annual production before August</td>
<td>n/a</td>
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<tr>
<td></td>
<td>Tennessee, USA</td>
<td><em>Quercus alba</em> <em>Quercus prinus</em></td>
<td>Offset, root production after leaf expansion, little root production after first week in August</td>
<td>SWC ($r = +0.25, P &lt; 0.01), E (correlation with known phenology patterns: $r = -0.4, P &lt; 0.0001$)</td>
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<td>Washington, USA</td>
<td><em>Pseudotsuga menziesii</em> <em>Abies amabilis</em> <em>Abies procera</em> <em>Pinus contorta</em> <em>Pinus ponderosa</em></td>
<td>n/a</td>
<td>ST (optimum)</td>
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<td>Berkshire, United Kingdom</td>
<td><em>Brassica napus</em> <em>Hordeum vulgare</em></td>
<td>n/a</td>
<td>ST (+ with high temp asymptote)</td>
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<td>Missouri, USA</td>
<td><em>Quercus alba</em> <em>Quercus velutina</em> <em>Quercus marilandica</em></td>
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<td>North Carolina, USA</td>
<td><em>Glycine max</em></td>
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<td>root temperature (optimum)</td>
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<td><em>Quercus alba</em></td>
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<td>ST (optimum), SWC (+)</td>
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Table 2.52. Measurements from literature used to quantify the offset between aboveground and belowground growth peaks in Figures 1 and 2. Methods used to measure root production varied by study (change in root biomass, \(^7\)minirhizotron, \(^8\)field rhizotron or root box, \(^9\)soil CO\(_2\) efflux). AP, day of year aboveground peak; #AP, number of aboveground production peaks; BP, day of year belowground peak; #BP, number of belowground production peaks; H, herbaceous; W, woody; MMT, median monthly temperature (°C); MMP, mean monthly precipitation (mm); REF, reference.

<table>
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<th>D or E</th>
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<th>#AP</th>
<th>Type of belowground measurement</th>
<th>BP</th>
<th>#BP</th>
<th>Offset</th>
<th>MMT</th>
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<tr>
<td>Boreal</td>
<td>Northern Savonia, Finland</td>
<td>Picea abies</td>
<td>change in seedling height (mm)</td>
<td>W</td>
<td>E</td>
<td>176.5</td>
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<td>change in root biomass (g*time(^{-1})) (^{1})</td>
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<td>Picea abies</td>
<td>change in height increment (cm*time(^{-1}))</td>
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<td>Salix viminalis</td>
<td>maximum shoot height (cm)</td>
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Table 2.S3. Monthly root and shoot growth rates for each biome estimated from literature. Growth rate of roots and shoots were tabulated from each study (excluding those that did not include seasonal data [i.e. measured DOY 1 as start of experiment rather than the Julian day]) and converted to a % of the maximum growth rate. Growth rates were averaged for each month with > 5 observations. These rates do not exceed 1 because many studies with differently timed maximum growth were averaged. MMT, median monthly temperature (°C); MMP, mean monthly precipitation (mm).

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<th>MMP</th>
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<td>root</td>
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<td>22.38</td>
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<td>- ± -</td>
<td>root</td>
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<td>16.85</td>
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<tr>
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<tr>
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<tr>
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<tr>
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**Mediterranean (n=20)**
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<th>MMP</th>
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<td>0.20 ± 0.05</td>
<td>shoot</td>
<td>6.2986</td>
<td>94.68</td>
</tr>
<tr>
<td></td>
<td>Dec</td>
<td>0.08 ± 0.05</td>
<td>shoot</td>
<td>1.273</td>
<td>89.37</td>
</tr>
</tbody>
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Supplementary Information Reference Guide


CHAPTER THREE: SEASONALITY AND PARTITIONING OF ROOT ALLOCATION TO RHIZOSPHERE SOILS IN A MID-LATITUDE FOREST

Abstract

Fine roots are a seasonally dynamic carbon (C) pool that accounts for a large proportion of net primary production. Roots affect the composition and function of the soil microbial community through turnover of root tissues, exudation of labile organic compounds and allocation of C to mycorrhizal fungi. As such root growth, mortality, and exudation are important components of biogeochemical cycles, yet data on the timing and partitioning of C to these processes are rare. The objective of this study is to estimate the seasonality, magnitude, and partitioning of C allocated belowground across the growing season in three mid-latitude hardwood and conifer stands at Harvard Forest in central Massachusetts using a minirhizotron camera, infrared gas analyzer, and cuvette collection to measure root production, respiration and exudation, respectively.

Fine root growth and respiration were both positively correlated with temperature, but differed in their phenology. Root phenology was characterized by multiple flushes of growth and mortality, whereas root respiration was unimodal across the growing season. Deciduous hardwood stands allocated C belowground earlier in the season compared to a conifer-dominated stand. Total belowground C flux (TBCF) was highest in the red oak stand, consistent with the observed increase in aboveground biomass of red oak at Harvard Forest over the past 20 years. TBCF was lowest in the eastern hemlock stand due to a decline in allocation to root production over the study period. This decline is coincident with the arrival and spread of the hemlock woolly adelgid in this stand. This
study elucidates the seasonal partitioning of belowground C within the context of long-term stand dynamics.
Introduction

Fine roots are a dynamic carbon (C) pool that accounts for 10-60% of net primary production (NPP, Keyes & Grier 1981, Helmisaari et al., 2002, Ostonen et al., 2005). Allocation of C to roots is the main conduit by which C is transported belowground and incorporated into soil organic matter (SOM). Roots affect the composition and function of the soil microbial community through tissue turnover, exudation of labile organic compounds and allocation of C to mycorrhizal fungi. Root exudation stimulates microbial biomass growth and extracellular enzyme activity, thereby increasing the decomposition rate of SOM, liberating plant available nitrogen (N), and releasing carbon dioxide (CO$_2$) to the atmosphere (Kuzyakov, 2010, Phillips et al., 2011). As such root growth, mortality and exudation are important components of biogeochemical cycles (Finzi et al., 2015).

Root respiration can account for up to 70% of total belowground C allocation and over half of soil CO$_2$ efflux (Burton et al., 2008, Drake et al., 2011, Fahey et al., 2005). Soil respiration is a large flux that integrates information about autotrophic and heterotrophic processes that together follow the seasonal cycle of air temperature and photosynthesis, and as a result it is relatively easy to detect the seasonal pattern of soil respiration. In contrast, measuring individual processes that contribute to belowground C allocation and soil respiration, such as autotrophic respiration, is far more challenging because of disturbance effects associated with sampling, small sample volume and small fluxes. A compromise approach measures the rate of root respiration indirectly by subtracting soil respiration rates in trenched and untrenched plots. While seasonally resolved, this method may over- or under-estimate autotrophic respiration as a result of
root death and decomposition (Drake et al., 2012, Savage et al., 2013) or the loss of priming effects (Finzi et al., 2015). Perhaps as a result of these difficulties, few studies assess seasonal patterns of belowground C allocation, making it difficult to understand how the belowground C flux is partitioned among roots, exudates, and mycorrhizal fungi (Drake et al., 2011). There are a handful of studies that have included seasonally resolved measurements of root respiration using chamber measurements on intact or excavated roots (Burton et al., 2002, Davidson & Holbrook, 2009, Davidson et al., 2006). Similarly, there are few seasonally-resolved measurements of fine root nonstructural carbohydrates (NSC) pools despite their potential importance to ecosystem-scale C cycling (Boldingh et al., 2000, Li et al., 1996). Even less is known about the seasonality of root exudation (Phillips et al., 2008, Pritchard et al., 2008).

Despite the paucity of process-level data, there are sufficient studies that document notable variations in the phenology of belowground C allocation. For example, Cardon et al. (2002) report multiple flushes of root production in oak species throughout the growing season. There are also observations of differences in the phenology of root growth between different tree species in the same geographical location, indicating that belowground phenology depends in part on internal cues (McCormack et al., 2014). At the biome scale, root production is commonly asynchronous with aboveground growth, particularly where coniferous tree species are present (Abramoff & Finzi, 2015).

In previous work I found that the phenology of belowground autotrophic activity is sensitive to variations in temperature, precipitation, and substrate supply (Abramoff & Finzi, 2015, Giasson et al., 2013). One observation common to both studies is a peak in
soil respiration or maximum root growth occurring after the peak in ecosystem respiration and maximum shoot growth, respectively. Neither of these studies, however, directly measured the components of belowground C allocation and their phenology using the same methods through time.

The objective of this study is to assess the phenology, magnitude, and partitioning of C allocated belowground across the growing season in mid-latitude, hardwood and conifer stands at the Harvard Forest in central Massachusetts. Based on the results of my previous work, I hypothesize that (1) deciduous stands allocate C belowground earlier in the growing season than the conifer stand; (2) root growth, mortality and respiration are positively correlated with soil temperature and precipitation; and the (3) total belowground C flux (TBCF) is highest overall in the red oak stand compared to the white ash and eastern hemlock stand, consistent with the observed increase in red oak aboveground biomass throughout the Harvard Forest over the past 20 years (Urbanski et al., 2007), and consistent with the arrival of the invasive pest the hemlock woolly adelgid to the hemlock stand (Ellison et al., 2015).

Materials and Methods

Study Site

This study was conducted in the Prospect Hill tract of the Harvard Forest Long Term Ecological Research site, a 120+-year-old secondary-growth forest located in Petersham, MA (42’N, 72’W, elevation 340 m; Wofsy et al., 1993, Goulden et al., 1996). The dominant species are northern red oak (Quercus rubra) and red maple (Acer rubrum), with smaller populations of eastern hemlock (Tsuga canadensis), white ash
(Fraxinus americana), white pine (Pinus strobus) and red pine (Pinus resinosa). The site is located on former agricultural land that was abandoned in the mid-1800s allowing forest regrowth beginning late in the 19th century (Foster et al., 2003). Forest uptake of C has increased since 1990 from ~ 2 Mg C ha\(^{-1}\) yr\(^{-1}\) to ~ 5 Mg C ha\(^{-1}\) yr\(^{-1}\) (Keenan et al., 2012, Urbanski et al., 2007). Soils are Typic Distrochrepts derived from glacial deposits of granite, schist and gneiss.

Plots were established in three mono-dominant stands: white ash, red oak, and eastern hemlock (Figure 3.1). Each stand occupies an area of 3.4, 8.3, and 10.9 hectares, respectively. I established 6 biometry plots per stand (N = 18) and 10 minirhizotron tube plots per stand (N = 30). The basal area in each 8-m-radius biometry plot is composed of 80% dominant tree species, with the inner 5-m basal area containing only the dominant species. The three stands differ in soil chemistry and biogeochemistry (Brzostek & Finzi, 2012). The white ash stand has a lower ratio of C-to-N content in the soil than the oak and hemlock stand (Table 3.1).

**Root Production and Biomass**

Root production and turnover were measured April–December 2012, March–November 2013, and April–November 2014 using a BTC-100x high magnification minirhizotron camera system (Bartz Technology Company, Carpenteria, CA). Measurements were made bi-weekly during the growing season in 2012 and monthly in 2013, 2014, and during the snow-free dormant season of each of the three years. There was no sampling from December 2012 to March 2013 and November 2013 to April 2014.
The camera system was inserted into cellulose acetate butyrate tubes installed at a 45° angle to a vertical soil depth of 40 cm. Thirteen tubes were installed in the center of each minirhizotron tube plot at Harvard Forest 10+ years ago (n = 4 in red oak, n = 9 in eastern hemlock). Seventeen tubes were installed in November 2012 (n = 6 in red oak, n = 1 in eastern hemlock, and n = 10 in white ash) for a sample size of n = 10 for each stand in 2013 and 2014. Minirhizotron tubes installed in November 2012 likely severed existing roots during placement and may have increased root growth rates in the following seasons. To test for this effect I conducted an analysis of variance using growth or mortality as the dependent variable and sample date and whether or not the sample came from a recently installed minirhizotron tube as fixed effects for 2013 and 2014.

The camera captures thirty-nine sequential images that are 13.5 x 17 mm in size along the upper axis of each tube at each sampling interval. The resulting images were processed using the open source imaging software Rootfly (Rootfly Development Team, Version 2.0.2, GNU General Public License). In each image, every root or root segment’s length and diameter was annotated. Length (mm) and diameter (mm) were scaled to mass (g) using a site-specific relationship based on n = 20 per species. For each root sample I recorded length and diameter, and dried it to constant mass at 60°C for 4 days. The polynomial fit to mass as a function of length and diameter was of the form:

\[
\text{mass} = \text{length} \times [a \times (\text{diameter})^2 - b \times (\text{diameter})]
\]  

[1]

where the coefficients [a, b] were [3·10^{-4}, 1·10^{-5}], [6·10^{-4}, 5·10^{-5}], and [4·10^{-4}, 9·10^{-5}], for white ash, red oak, and eastern hemlock respectively. The R^2 value for this relationship was greater than 0.94 for each species. Root biomass was estimated from the
images assuming that the viewing depth is 0.7848 mm and calculating the diameter of the imaged root using the method of Taylor et al. (2014). Assuming that roots are cylindrical, the relationship between the true diameter (D) and diameter perceived (p) at depth (f) is:

\[
D = \frac{\left(f^2 + \frac{1}{4}p^2\right)}{f}\]  [2]

Daily fine root growth and mortality (g root d\(^{-1}\)) in each minirhizotron image for each sampling interval were calculated as:

\[
Growth \ (g \ root \ d^{-1}) = \frac{m_{t2} - m_{t1}}{t_2 - t_1} \text{ when } m_{t2} - m_{t1} > 0
\]  \[3\]

\[
Mortality 
(g \ root \ d^{-1}) = \frac{m_{t2} - m_{t1}}{t_2 - t_1} \text{ when } m_{t2} - m_{t1} < 0
\]  \[4\]

where \(m\) is the total mass of roots traced on day of year (DOY) \(t_1\) and \(t_2\). Gross root production refers to total fine root growth summed over an interval of time. Net root production is the sum of growth and mortality for that interval. Root production measurements were scaled to g C m\(^{-2}\) d\(^{-1}\) using the assumption that each minirhizotron image is representative of a 0.173 cm\(^3\) (13 mm x 17 mm x 0.7848 mm) volume of soil (Taylor et al., 2014).

The standing biomass of roots was estimated from field samples. I collected three 10 x 10 cm samples of the organic horizon and three 5 cm diameter mineral soil samples to a depth of 15 cm monthly in each plot. Roots were removed and sorted into fine (< 2 mm), coarse (> 2 mm), live and dead pools. Roots were then dried and weighed to obtain standing biomass for live fine roots (g m\(^{-2}\)). Subsamples of roots from monthly soil coring were assayed for carbon content (%C) using an elemental analyzer (model NC2500; CE Instruments, Milan, Italy). Fine root standing biomass for the organic
horizon down to a depth of 15 cm in the mineral soil was scaled up to g C m\(^{-2}\) by adjusting for the horizontal area of the soil core, the carbon content of roots in each stand, and rock content.

**Root Respiration**

CO\(_2\) efflux was measured directly on recently severed roots using an infrared gas analyzer (LI6400, LiCor Biosciences, Lincoln, NE). Measurements were made monthly from March to October 2013 on three samples per stand. Respiration rates have been measured successfully using severed roots in previous studies (Burton *et al.*, 2012, Burton & Pregitzer, 2003). Furthermore, I compared respiration measurements of an attached and severed root system for each stand on three separate days and confirmed that respiration rates were similar between the two types of roots (correlation coefficient = 0.78) and stable up to approximately 7 hours.

Two measurements per sample were made on each sample date within 7 hours of collection, one in the field at ambient temperature and one in the lab at a constant temperature. Field measurements of CO\(_2\) efflux were fit to the Arrhenius equation,

\[
R_S = A * e^{-\frac{E_a}{RT}}
\]

where \(R_S\) is the respiration rate (\(\mu\)mol CO\(_2\) s\(^{-1}\) g\(^{-1}\)), \(E_a\) is the activation energy (kJ mol\(^{-1}\)), \(A\) is a pre-exponential factor (\(\mu\)mol CO\(_2\) s\(^{-1}\) g\(^{-1}\)), \(T\) is temperature (Kelvin) and \(R\) is the gas constant (kJ Kelvin\(^{-1}\) mol\(^{-1}\)). The parameters \(E_a\) and \(A\) were estimated using non-linear curve fitting in SigmaPlot (Version 10.0, Systat Software, San Jose, CA). I then estimated growing season rates of \(R_S\) using daily soil temperature measured at 10 cm depth from HOBO data loggers installed in March 2013 in each stand. Mass-specific \(R_S\)
(μmol CO₂ g root⁻¹ s⁻¹) was scaled to g C m⁻² s⁻¹ using the mass of root per square meter ground surface area and converting from μmol to μg.

**Nonstructural Carbohydrates**

The pool of NSC was estimated as the sum of the concentration of sugars (assumed to be glucose:fructose:galactose in 1:1:1 ratio) and starch using the method of Chow & Landhausser (2004). I collected three ~1-2 g root samples from each biometry plot monthly from May to November 2011 and four times from March to November 2012. Roots were excavated, washed, and frozen in liquid nitrogen until analysis. Sugars were extracted from dried and finely ground root tissue using a 12:5:3 methanol:chloroform:water solution before being developed with 2% phenol and concentrated sulfuric acid. Absorbance was measured at 490 nm using a digital spectrophotometer (Spectronic 20D+, Thermo Scientific). Starch was extracted using a 0.005 N sulfuric acid solution at 95ºC and developed as described above.

**Root Exudation**

Root exudates were collected from six root systems per stand in June and August 2012, and April, May, July and October 2013 following the method of Phillips *et al.* (2008, 2011). In brief, roots were excavated 48 hours prior to collection, washed, and incubated in a moist soil-sand mixture. Roots were placed into cuvettes with glass beads and a C-free nutrient solution 24 hours prior to collection. At the time of collection, exudate-containing nutrient solution was extracted with two additional flushes of C-free nutrient solution to ensure that exudates adhering to glass beads were flushed into
solution. Samples were transported back to the lab on ice and analyzed for non-purgeable organic carbon content using an elemental analyzer (Shimadzu TOC-VCSH analyzer, New Haven, CT). Exudation rate (g C g root$^{-1}$ d$^{-1}$) was scaled to g C m$^{-2}$ d$^{-1}$ using root biomass (g root m$^{-2}$) from soil cores.

**Total Belowground C Flux**

To estimate the quantity of C allocated belowground during the growing season (herein abbreviated as “gs”), I define a simplified belowground C flux budget (sensu Litton et al., 2007, Drake et al., 2011) using two different approaches. First, a top-down estimate of total belowground C flux (TBCF$_{\text{top}}$, g C m$^{-2}$ gs$^{-1}$) is defined as:

$$TBCF_{\text{top}} = F_{\text{efflux}} + F_{\text{leaching}} - F_{\text{litter}} + \Delta(C_{\text{roots}} + C_{\text{soil}})$$

[7]

where $F_{\text{efflux}}$ is the growing season rate of soil respiration, $F_{\text{leaching}}$ is the flux of dissolved organic C into streamwater, $F_{\text{litter}}$ is litterfall, and $\Delta(C_{\text{roots}} + C_{\text{soil}})$ is the growing season change in the C pool associated with fine roots and soil. $F_{\text{efflux}}$ is estimated from a 22-year synthesis of soil CO$_2$ efflux data using a range of methods across multiple stand types (Giasson et al., 2013). $F_{\text{litter}}$ is measured using litter baskets (Barker Plotkin, 2010, Brzostek, 2012, Frey & Ollinger, 1999, Hadley, 2009, Lemos, 2013, Munger & Wofsy, 1999). I estimated $\Delta C_{\text{roots}}$ using net root production from this paper. I set $\Delta C_{\text{soil}} = 0$, assuming that it is small relative to the timescale of this study (Gaudinski et al., 2000).

Second, a bottom-up estimate (TBCF$_{\text{bottom}}$, g C m$^{-2}$ gs$^{-1}$) is defined as:

$$TBCF_{\text{bottom}} = F_{\text{roots}} + F_{\text{resp}} + F_{\text{exudates}}$$

[8]

where $F_{\text{roots}}$ is gross fine root production, $F_{\text{resp}}$ is fine and coarse root respiration, and $F_{\text{exudates}}$ is root exudation. $F_{\text{roots}}$, $F_{\text{exudates}}$, and fine root respiration are estimated using
measurements from this study. Coarse root respiration is estimated using the observation that mass-specific rates of coarse root respiration are ca. 70% lower than fine root respiration rates in the same stand (Desrochers et al., 2002, Fahey et al., 2005, Pregitzer et al., 1998). Coarse root respiration (μmol CO$_2$ g root$^{-1}$ s$^{-1}$) was scaled to g C m$^{-2}$ s$^{-1}$ using coarse root biomass to 15 cm from soil pits in evergreen and hardwood stands at Harvard Forest and the Harvard Conservation Trust (Harvard, MA, 42°31’N 71°32’W), respectively (Lemos, 2013).

I did not include transpiration of dissolved inorganic C, fungal production or throughfall leaching in the calculation of C outputs or inputs as these terms are generally < 5% of the C budget in other mid-latitude forest stands (Fahey et al., 2005, Drake et al., 2011). Coarse root production may account for 5-10% of total C inputs (Fahey et al. 2005; Drake et al. 2011), but I had no data on coarse root production at Harvard Forest and therefore did not include it.

Belowground C pools were derived from the Harvard Forest Data Archive and this study. Soil C content data were from both published (Bowden et al., 2009, Brzostek, 2012, Frey et al., 2014, Lemos, 2013, Nadelhoffer et al., 1999, Orwig & Foster, 2009) and unpublished sources (Drake, unpublished data; Sorensen, unpublished data).

Standing biomass of roots was estimated using data from this study as well as a previous study conducted on the same plots (Lemos, 2013). Soil microbial biomass in the hemlock stand was measured in June, July, August, and September of 2012 (Averill, unpublished data). Soil microbial biomass in the ash stand was measured in July of 2011 (Averill,
unpublished data). Microbial biomass data in oak-dominated hardwoods stands were taken from Drake et al. (2013) and more recent studies (Sorensen, unpublished data).

Monthly values of gross primary production (GPP) were estimated using partitioned NPP measured in 2012 at the eddy-covariance towers located in a mixed hardwood stand and a hemlock stand at Harvard Forest (ORNL-DAAC, 2013). Data were not available for ash stand GPP or soil CO₂ efflux; for the purpose of this budget I assumed these values were equivalent to oak-dominated hardwoods, although I recognize this makes the ash belowground C budget less certain. I used PhenoCam data (http://phenocam.sr.unh.edu/) to determine maximum canopy greenness in red oak and eastern hemlock stands.

Data Analysis

All statistical analyses were performed in R Statistical Software (R Development Core Team, 2013). Stand-specific and seasonal-to-interannual variations in fine root growth, mortality, respiration, NSC concentration, and exudation were modeled using a mixed-effects model with stand, sample date, and year as fixed effects and plot as a random effect. A Tukey’s HSD post-hoc test was used to test for differences between stands. Analysis of variance (ANOVA) was used to test for the effect of newly installed tubes on root growth and mortality, to test for the effect of stand on root tissue [N], to test for the effect of stand and soil horizon on root biomass, and to test for a significant difference between TBCF_{bottom} and TBCF_{top}. I also used ANOVA to test for differences in C content, N content, and organic horizon mass between stands, averaging across
subsamples. All mixed effects and ANOVA models were constructed using the `aov` function in base R.

Linear regression was used to correlate root growth with root mortality, and to correlate root growth with temperature for each stand separately. Multiple linear regression was used to model mass-specific respiration as a function of soil temperature, precipitation, and stand. I used a linear-mixed-effects model to model root mortality as a function of temperature, precipitation, stand, year, and day of year with tube as a random effect. Precipitation and day of year were not significant effects and were dropped from the final model. I also used a linear-mixed-effects model to model root growth as a function of temperature, precipitation, stand, year, and day of year with tube as a random effect using the `lmer` function in the `lme4` package in R (Bates et al., 2014). In mixed-effects, ANOVA, and regression models, fine root growth and mortality were log-transformed to meet assumptions of normality.

**Results**

*Root Production and Biomass*

There was a measurable but transient effect of tube installation on root production in red oak. In 2013, mean red oak root growth was 0.34 g C m\(^{-2}\) d\(^{-1}\) higher in newly installed tubes compared to tubes established a decade earlier (\(F_{2,79} = 2.7, P = 0.07\)), but this difference diminished in 2014 to 0.09 g C m\(^{-2}\) d\(^{-1}\) (\(P = 0.12\)). There was no detectable installation effect on red oak root mortality or on the single hemlock tube installed. All white ash tubes were newly installed in November 2012 so it was not possible to establish an effect of installation on white ash root production.
Root growth was positively correlated with soil temperature in each stand ($P < 0.001$, Table 3.2), but was not correlated with precipitation (Table 3.3). As a result root growth was concentrated in mid-summer, but with distinct stand-level differences (Figure 3.2, a-f). Red oak root growth occurred in 1–3 flushes over the growing season, with highest mortality in mid to late-summer (Figure 3.2, a-c). Root mortality was not correlated either with soil temperature ($P = 0.13$) or precipitation ($P = 0.76$). Hemlock root growth was low throughout the growing season with smaller production peaks occurring in the fall in 2013 and 2014. White ash root growth peaked in mid-summer with high mortality in late-summer. In red oak and white ash stands the peak in maximum canopy greenness (*data not shown*) occurred ~20 days earlier than in the eastern hemlock stand, but the peak in root growth occurred ~50 days earlier. As a result, the deciduous stands had a smaller offset between maximum canopy greenness and peak root growth than did the hemlock stand. Similar to the phenology of gross production, net fine root production increased in early to mid-summer in the deciduous stands. In 2012 hemlock fine root NPP was positive but in 2013 and 2014 there was no net production of fine roots.

Root biomass was significantly higher in the red oak ($P < 0.001$) and eastern hemlock ($P < 0.1$) stand compared to ash, largely because of a surface organic horizon ($F_{1,111} = 47.9$, $P < 0.001$). White ash stands at Harvard Forest lack an organic horizon, possibly the result of bioturbation by exotic earthworms that are not present in the red oak or eastern hemlock stands. In this stand, surface litter is incorporated directly into the
mineral soil horizon in less than 1 year. As a result, there is relatively little variation in
the root depth profile in this stand (Table 3.4).

In all years, red oak and white ash stands allocated more C to fine roots compared
to eastern hemlock (Figure 3.2, g-i). Averaged across growing seasons 301 ± 76 g C m⁻²
gs⁻¹ and 133 ± 42 g C gs⁻¹ were allocated to fine roots in red oak and white ash stands,
respectively. C allocation to gross fine root production in eastern hemlock was 42 ± 13 g
C m⁻² gs⁻¹.

Root Respiration

The phenology of fine root respiration was similar between stands, with highest
mass-specific respiration rates in mid-summer (Figure 3.3a). This follows from a positive
correlation with soil temperature (Figure 3.3b, R²_adj = 0.68, P < 0.001). Mass-specific
rates of fine root respiration were significantly higher in the white ash stand than in oak
and hemlock (Figure 3.3b). Arrhenius fits to mass-specific data indicate that white ash
root respiration had a lower apparent activation energy (Eₐ = 20 kJ mol⁻¹) across the
growing season compared to that of red oak (Eₐ = 40 kJ mol⁻¹) and eastern hemlock (Eₐ =
29 kJ mol⁻¹). The pre-exponential constant A (umol CO₂ g⁻¹ s⁻¹) varied over three orders
of magnitude between stands and was highly correlated with Eₐ (R² > 0.99).

Root respiration measured in the lab increased across the growing season (Figure
3.4). The average rate of fine root respiration for the six month growing season was 191 ±
24, 167 ± 33, and 205 ± 27 g C m⁻² gs⁻¹ for red oak, eastern hemlock, and white ash,
respectively (Figure 3.3c).
Nonstructural Carbohydrates and Root Exudation

There was large inter-annual variability in the concentration of NSC in fine roots (Figure 3.5). In 2012, there was a decline in NSC concentration mid-summer relative to the spring and fall. In 2011, there was a slight but significant increase in NSC concentration across the growing season (P < 0.001). Red oak roots had significantly lower NSC concentration than white ash and eastern hemlock (P < 0.001, Figure 3.5).

Exudation rate was highly variable and there was no clear stand level difference or seasonal pattern, although there were significantly lower exudation rates in early spring (DOY = 106) compared to summer and fall (Table 3.5). Exudation rates for red oak, eastern hemlock, and white ash were 47 ± 24, 55 ± 25, and 46 ± 11 g C m$^{-2}$ gs$^{-1}$, respectively.

Total Belowground C Flux

In the red oak stand, total belowground C flux was 577 ± 85 g C m$^{-2}$ gs$^{-1}$ based on TBCF$_{\text{bottom}}$. Of this, 52% was allocated to root production, 40% to root respiration, and 8% to exudation (Figure 3.6, orange bars). TBCF$_{\text{top}}$ was 791 ± 94 g C m$^{-2}$ gs$^{-1}$. In the eastern hemlock stand, TBCF$_{\text{bottom}}$ was 340 ± 51 g C m$^{-2}$ gs$^{-1}$. Approximately 13% of this flux was allocated to root production, 71% to root respiration, and 16% to exudation (Figure 3.6, purple bars). TBCF$_{\text{top}}$ was 474 ± 29 g C m$^{-2}$ gs$^{-1}$. In the white ash stand, TBCF$_{\text{bottom}}$ was 449 ± 57 g C m$^{-2}$ gs$^{-1}$. Thirty percent of TBCF$_{\text{bottom}}$ was allocated to root production, 60% to root respiration, and 10% to exudation (Figure 3.6, blue-green bars). TBCF$_{\text{top}}$ was 633 ± 58 g C m$^{-2}$ gs$^{-1}$. 
TBCF\textsubscript{top} was larger than TBCF\textsubscript{bottom} in all of the stands, but this difference was not significant (Table 3.6). In the red oak stand, TBCF\textsubscript{top} was 214 ± 127 g C m\textsuperscript{-2} gs\textsuperscript{-1} larger than TBCF\textsubscript{bottom}.

The phenology of TBCF\textsubscript{bottom} differed between stands (Figure 3.7). Red oak stands allocated more C belowground earlier in the growing season compared to white ash and eastern hemlock. The peak in TBCF\textsubscript{bottom} in red oak was coincident with the spring ramp-up of GPP. In eastern hemlock stands, the phenology of TBCF\textsubscript{bottom} was not pronounced. In both oak and hemlock stands, soil respiration peaked later in the season than either GPP or TBCF\textsubscript{bottom}.

**Discussion**

Data on timing and partitioning of C to belowground processes are rare (Dohleman *et al.*, 2012, Fahey *et al.*, 2013). This paper provides a first look at the seasonal dynamics of multiple root processes over 2-3 years in a mid-latitude forest. I observed significant stand-level differences in belowground C flux and its partitioning to root growth, exudation and respiration. The phenology of TBCF also differed broadly between stands, especially in the evergreen hemlock stands compared to the deciduous red oak and ash stands where the peak in TBCF was coincident with the peak in GPP. I found broad support for my three hypotheses (discussed below) and a proximate explanation for the increase in the biomass of red oak throughout the Harvard Forest over the last two decades. The emergence of a forest pest, the hemlock woolly adelgid, during the course of this study negatively affected belowground C allocation and root production in hemlock from 2012 to 2014. The data reported here suggest that belowground C flux is
sensitive to variations in temperature and C supply. Follow up studies are needed to assess the impact of belowground phenology on soil biogeochemistry.

**Phenology of Root Growth and Mortality**

In a recent meta-analysis I found that deciduous trees have more synchronous above- and belowground phenology than evergreen trees (Abramoff & Finzi, 2015). Consistent with the meta-analysis, I found that the offset between maximum canopy greenness, a proxy for aboveground phenology, and root growth was about 30 days shorter in the deciduous stands. Consistent with my first hypothesis, fine root growth was initiated earlier in the growing season in the hardwood stands compared to the hemlock stand (Figure 3.2, a-c).

There was not a strong correspondence between root growth and mortality despite statistical significance (Figure 3.2, $R^2_{adj} = 0.01$, $P < 0.05$). At best, a visual inspection of the data suggests a lag in mortality relative to growth for most stands and years. A more striking observation was the multiple flushes of root growth observed in red oak, particularly in 2012 (Figure 3.2a). The direct observation of multiple root flushes confirm the results inferred by Cardon *et al.* (2002), who based their inference on a negative correlation between shoot elongation and soil respiration in a common garden experiment with red oak saplings. There were fewer flushes in the 2013 and 2014 data that may reflect sampling intensity. The minirhizotrons were sampled biweekly in 2012 and monthly in 2013 and 2014. It is possible that the lower sampling frequency missed flushing episodes. Alternatively, there may have been fewer flushes in these years.
Environmental Controls over Root Growth and Respiration

Consistent with hypothesis 2, root growth and respiration were sensitive to variations in soil temperature (Table 3.3). Contrary to my second hypothesis, there was no effect of precipitation on growth or respiration, and mortality was unrelated to variations in temperature or precipitation. The positive correlation between root growth and temperature observed here has been observed in a variety of ecosystems using both observational (Bevington & Castle, 1985, Teskey & Hinckley, 1981) and experimental approaches (Lahti et al., 2005, Tryon & Chapin III, 1983). The positive relationship between root growth and temperature reflects a number of processes including enhanced C supply due to photosynthesis, increases in the rate of cell division and lower resistance to water uptake favoring cell expansion (Lambers et al., 2008). Though very high temperatures (> 30°C) can inhibit root growth, I did not observe soil temperatures greater than 24°C (Barney, 1951, Graves et al., 1991).

There was a strongly seasonal cycle to the rate of fine root respiration that reflected the sensitivity of respiration to temperature (Figure 3.3a,b). The apparent temperature sensitivity (E_a) of root respiration was lowest in the ash stands and highest in the red oak stand (Table 3.7). The measurement of temperature sensitivity reported here convolves many potential sources of variability such as differences in root chemistry and substrate supply. Thus it is difficult to ascribe control(s) for the greater temperature sensitivity in red oak compared to the other species.

When incubated at a common temperature the rate of root respiration increased across the growing season in all three species (Figure 3.4). This suggests an increase in
photosynthate allocation to roots through time and that the decline of respiration rates in the fall is related to temperature rather than acclimation of root respiration or substrate limitation. This observation is qualitatively similar to that of Burton & Pregitzer (2003) who did not find acclimation of root respiration across the growing season in sugar maple stands in Michigan. Both studies contrast with the apparent acclimation of root respiration in response to experimental soil warming at the Harvard Forest (Burton et al., 2008).

Fine root respiration in ash was significantly higher than oak and hemlock and appears to be the result of high root [N]. The rate of N mineralization is high in ash compared to oak and hemlock stands (Brzostek & Finzi, 2012, Finzi et al., 1998). The presence of the invasive European earthworm *Lumbricus terrestris* in these plots further accelerates the rate of N cycling because they rapidly incorporate leaf litter into the mineral soil and decrease the turnover time of organic matter (Marhan & Scheu, 2005, Scheu, 1987). High N availability is most likely the reason root [N] is significantly higher in ash compared to oak and hemlock (F$_{2,68}$ = 90.3, P < 0.001). Given that root N concentration is correlated with respiration rate (Burton et al., 2002, Reich et al., 2002), the high mass-specific rates of root respiration in this species are most likely explained by their high [N]. Notably at the plot scale, total fine root respiration (g C m$^{-2}$ gs$^{-1}$) was highest in the ash stand despite it having ~50% lower biomass than that in red oak.

### Nonstructural Carbohydrates and Root Exudation

Root NSC concentrations varied significantly between stands, years, and within each growing season (Figure 3.5). In 2012, there was a strong apparent seasonal decline
in root NSC mid-summer, which may reflect the metabolism of NSC for root growth and maintenance respiration (Lynch et al., 2013). In 2011, there was no evidence for a seasonal decline. The difference between years may reflect inter-annual variation in allocation to root NSC pools. It is also possible that NSC concentrations vary over timescales finer than the monthly sampling interval used here, in contrast to stemwood NSC, for which monthly sampling appears sufficient to capture the seasonal trend in the NSC pool (Richardson et al., 2013).

In contrast to root NSC there was no significant difference in exudation rate among stands (Table 3.8). Brzostek et al. (2013) similarly found no difference in the rate of root exudation among four stand types, including the hemlock and ash plots studied here. Exudation did, however, vary significantly at the seasonal time-scale with the main distinction being significantly lower rates in the spring compared to summer and fall collection dates (Table 3.5). This pattern of exudation mirrors the seasonal increase in mass-specific rates of root respiration and supports the idea of a progressive increase in C allocation belowground across the growing season (Figure 3.4).

I caution that the detectability of variations in exudation may prove difficult given the methods presently available for research in the field. The collection of exudation requires the physical extraction of intact root systems from the soil that necessarily disrupts microbial interactions (Phillips et al., 2008). Similarly, sample collection requires submerging root tips in a liquid medium with a chemical composition different from the soil solution. This may affect the rate at which C is released from roots into the
solution. Indeed, other studies of exudation across the growing season did not detect a clear seasonal pattern (Phillips et al., 2008, Phillips et al., 2011).

**Total Belowground C Flux**

TBCF varied between 28 and 35% of GPP (Table 3.6). Consistent with my third hypothesis, red oak had the highest rate of TBCF and the greatest proportional allocation to fine root production (Figure 3.6). Eddy-covariance estimates of net ecosystem production at the Harvard Forest mixed hardwood tower site suggest a near doubling of C uptake from the atmosphere over the last 20 years (Keenan et al., 2012, Urbanski et al., 2007). The increase in C uptake is correlated with an increase in red oak productivity and biomass. Of the dominant species within the tower footprint, the concentration of N in red oak foliage is among the highest, and this species has the most rapid rate of light-saturated net photosynthesis (Bassow & Bazzaz, 1997). Foliar [N] concentration in red oak remains unchanged throughout, suggesting that the rate of annual N uptake from the soil has also increased. Given that belowground C allocation is required to acquire soil N, this analysis suggests that high TBCF in this red oak stand relative to hemlock, the second most dominant tree species in the tower footprint, has facilitated its emergence as the dominant species at this site (Figure 3.6, Table 3.6).

The modest C investment in root production in hemlock stands may be an attribute of this species in particular or evergreen trees in general, which allocate a substantial fraction of C to ectomycorrhizal symbionts (Clemmensen et al., 2013, Hobbie, 2006). Confounding this interpretation of the results, however, is the recent infestation of the invasive pest hemlock woolly adelgid (HWA), which became
widespread at the Harvard Forest in 2012 with visible signs of crown thinning and HWA-induced tree mortality recorded in 2014. Compared to 2012, the allocation of C to root production in hemlock stands dropped by 11% in 2013 and 72% in 2014, suggesting a negative effect of the HWA on root C allocation. Surveys of hemlock roots in infested stands in Connecticut found that ectomycorrhizal colonization, bacterial abundance in the adjacent rhizosphere, and root C:N all declined (Vendettuoli et al., 2015).

The phenology of TBCF\textsubscript{bottom} differs between stands and relative to that of GPP or soil respiration (Figure 3.7). In the deciduous red oak stand, the peak in TBCF\textsubscript{bottom} precedes the peak in soil respiration and coincides with spring ramp-up of GPP. This suggests that belowground C allocation in oak is strongly dependent upon the supply of photosynthate. This result is consistent with pulse-chase experiments demonstrating rapid transfer of C belowground (Hogberg et al., 2008) and girdling experiments showing steep declines in springtime soil respiration owing to the absence of active roots (Högberg et al., 2001).

TBCF\textsubscript{top} was consistently greater than TBCF\textsubscript{bottom}. This suggests an over-estimate of $\Delta C_{\text{root}}$ in TBCF\textsubscript{top}, an underestimate of root GPP, respiration, or exudation in TBCF\textsubscript{bottom}, or a combination of both. The largest difference between TBCF\textsubscript{top} and TBCF\textsubscript{bottom} ($\sim$240 gC m\textsuperscript{-2} gs\textsuperscript{-1}) was observed in the red oak stand. The differences were smaller in the ash and hemlock stands. Root NPP was estimated from minirhizotron images and was highest in red oak (Figure 3.2, g-i). High NPP contributes to a large estimate of $\Delta C_{\text{root}}$ and hence the estimate of TBCF\textsubscript{top} (equation [1]). In contrast to the high minirhizotron estimate of root NPP, there was no significant change in root biomass in
soil cores collected across the growing season in 2012 (*data not shown*). Because of high spatial variability (Taylor *et al.*, 2013), the absence of a change in root biomass in one year does not exclude the possibility of high root NPP. However, the three-growing-season average root NPP of 225 g C m\(^{-2}\) estimated from minirhizotron imaging is about one-half of the standing crop (Figure 3.3c), and should be measurable in soil cores. TBCF in red oak is therefore likely to reside between the two estimates.

**Summary**

Climatic and atmospheric perturbations alter the magnitude of C inputs belowground (e.g., Drake *et al.*, 2011) and thus at the inter-annual time-scale it is likely that changes in the quantity and phenology of belowground C inputs will influence soil biogeochemical cycling. Warming-induced increases in growing season length have significantly increased annual ecosystem C uptake at Harvard Forest, a large portion of which is hypothesized to be allocated belowground (Keenan *et al.*, 2014). Whether and how this enhanced allocation belowground will affect long-term soil-C cycling and C storage remains an open question.

**Acknowledgements**

I would like to thank Colin Averill, John Drake, William Munger, and Patrick Sorensen for use of data. I also thank Andrew Richardson for making data from the PhenoCam network publicly available. Development of the PhenoCam network has been supported through the National Science Foundation (award EF-1065029). This research was supported by the American Association of University Women (AAUW) American
Dissertation Fellowship, the Office of Science (BER), the US Department of Energy (grant no. 10-DOE-1053) and the National Science Foundation (DEB-0743564).
Table 3.1. Stand-level characteristics. Total C content refers to organic and mineral horizon soil C down to 15 cm. Letters indicate a statistically significant difference in means using analysis of variance at the P < 0.05 level.

<table>
<thead>
<tr>
<th>Stand</th>
<th>Total C content (g m$^{-2}$)</th>
<th>Total N content (g m$^{-2}$)</th>
<th>O horizon mass (g m$^{-2}$)</th>
<th>pH</th>
<th>Soil C:N Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>White ash</td>
<td>5989 ± 330$^{b}$</td>
<td>431 ± 24$^{a}$</td>
<td>-</td>
<td>5.1</td>
<td>13.9 ± 0.17$^{b}$</td>
</tr>
<tr>
<td>Red oak</td>
<td>7709 ± 438$^{a}$</td>
<td>354 ± 23$^{b}$</td>
<td>131 ± 8$^{a}$</td>
<td>5.2</td>
<td>21.8 ± 0.72$^{a}$</td>
</tr>
<tr>
<td>Eastern hemlock</td>
<td>6922 ± 246$^{a}$</td>
<td>290 ± 13$^{b}$</td>
<td>105 ± 7$^{b}$</td>
<td>3.7</td>
<td>23.9 ± 0.70$^{a}$</td>
</tr>
</tbody>
</table>
Table 3.2. Summary of multiple linear regression statistics that model log(fine root growth) for each stand as a function of soil temperature. Fine root growth was log-transformed to meet assumptions of normality. *** P < 0.001, ** P < 0.01, * P < 0.05, . P < 0.1

<table>
<thead>
<tr>
<th>Soil temperature</th>
<th>β</th>
<th>F</th>
<th>$R_{adj}^2$</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>White ash</td>
<td>0.08</td>
<td>48.9</td>
<td>0.26</td>
<td>***</td>
</tr>
<tr>
<td>Red oak</td>
<td>0.11</td>
<td>89.2</td>
<td>0.30</td>
<td>***</td>
</tr>
<tr>
<td>Eastern hemlock</td>
<td>0.08</td>
<td>43.8</td>
<td>0.15</td>
<td>***</td>
</tr>
</tbody>
</table>
Table 3.3. Summary of multiple linear regression statistics that model fine root growth, mortality, and mass-specific respiration as a function of soil temperature and precipitation. Growth and mortality were log-transformed to meet assumptions of normality. *** $P < 0.001$, ** $P < 0.01$, * $P < 0.05$, ns $P < 0.1$

<table>
<thead>
<tr>
<th></th>
<th>Soil temperature</th>
<th>Precipitation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$\beta$ F $R^2_{adj}$ P-value</td>
<td>P - value</td>
</tr>
<tr>
<td>Growth</td>
<td>1.5 202.9 0.47 ***</td>
<td>ns</td>
</tr>
<tr>
<td>Mortality</td>
<td>1.3 3.4 0.19 *</td>
<td>ns</td>
</tr>
<tr>
<td>Respiration</td>
<td>0.29 32.2 0.68 ***</td>
<td>ns</td>
</tr>
</tbody>
</table>
Table 3.4. Median root biomass (5th percentile, 95th percentile) in g C m⁻² estimated from minirhizotron tubes binned in 10 cm depth increments. Uppercase letters indicate significant differences between columns, and lowercase letters indicate significant differences between rows at the P < 0.01 level.

<table>
<thead>
<tr>
<th>Depth</th>
<th>White ash</th>
<th>Red oak</th>
<th>Eastern hemlock</th>
</tr>
</thead>
<tbody>
<tr>
<td>O horizon</td>
<td>-</td>
<td>3.46 (2.29, 6.65)³⁶⁶</td>
<td>2.82 (0.80, 7.33)³⁶⁶⁶</td>
</tr>
<tr>
<td>0 - 10 cm</td>
<td>0.98 (0.20, 1.84)³⁶⁶</td>
<td>3.03 (0.71, 6.17)⁶</td>
<td>1.17 (0.53, 1.46)⁶</td>
</tr>
<tr>
<td>10 - 20 cm</td>
<td>0.79 (0.45, 1.08)³⁶⁶</td>
<td>0.77 (0.38, 1.66)³⁶⁶</td>
<td>1.16 (0.42, 3.24)³⁶⁶</td>
</tr>
<tr>
<td>20 - 30 cm</td>
<td>0.79 (0.41, 2.62)³⁶⁶</td>
<td>0.49 (0.25, 4.19)³⁶⁶</td>
<td>0.50 (0.01, 1.29)³⁶⁶</td>
</tr>
<tr>
<td>30 - 40 cm</td>
<td>0.45 (0.16, 2.22)³⁶⁶</td>
<td>0.71 (0.08, 2.01)³⁶⁶</td>
<td>0.65 (0.06, 1.92)³⁶⁶</td>
</tr>
<tr>
<td>40 - 50 cm</td>
<td>0.19 (0.07, 0.66)³⁶⁶</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Table 3.5. Exudation rate (± SE) for samples collected in 2012 and 2013. There is a significant effect of sample date ($F_{1,106}=8.0, P < 0.01$), but no differences between years or stands. The exudate rate on the April sample date (DOY = 106) was significantly lower than the exudation rate measured on June, July, and October sample dates at the $P < 0.05$ level.

<table>
<thead>
<tr>
<th>Stand</th>
<th>Date</th>
<th>N</th>
<th>Exudation Rate (mg C g root$^{-1}$ day$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>White ash</td>
<td>6/19/2012</td>
<td>6</td>
<td>1.35 (0.43)</td>
</tr>
<tr>
<td></td>
<td>8/26/2012</td>
<td>6</td>
<td>1.14 (0.25)</td>
</tr>
<tr>
<td></td>
<td>4/16/2013</td>
<td>6</td>
<td>0.29 (0.23)</td>
</tr>
<tr>
<td></td>
<td>5/30/2013</td>
<td>7</td>
<td>1.03 (0.07)</td>
</tr>
<tr>
<td></td>
<td>7/27/2013</td>
<td>7</td>
<td>2.69 (0.68)</td>
</tr>
<tr>
<td></td>
<td>10/7/2013</td>
<td>6</td>
<td>1.39 (0.34)</td>
</tr>
<tr>
<td>Red oak</td>
<td>6/19/2012</td>
<td>7</td>
<td>2.65 (1.34)</td>
</tr>
<tr>
<td></td>
<td>8/26/2012</td>
<td>7</td>
<td>0.26 (0.10)</td>
</tr>
<tr>
<td></td>
<td>4/16/2013</td>
<td>7</td>
<td>-0.47 (0.33)</td>
</tr>
<tr>
<td></td>
<td>5/30/2013</td>
<td>6</td>
<td>0.52 (0.09)</td>
</tr>
<tr>
<td></td>
<td>7/27/2013</td>
<td>6</td>
<td>1.37 (0.54)</td>
</tr>
<tr>
<td></td>
<td>10/7/2013</td>
<td>6</td>
<td>0.70 (0.24)</td>
</tr>
<tr>
<td>Eastern hemlock</td>
<td>6/19/2012</td>
<td>5</td>
<td>0.78 (0.35)</td>
</tr>
<tr>
<td></td>
<td>8/26/2012</td>
<td>6</td>
<td>0.36 (0.07)</td>
</tr>
<tr>
<td></td>
<td>4/16/2013</td>
<td>6</td>
<td>0.02 (0.17)</td>
</tr>
<tr>
<td></td>
<td>5/30/2013</td>
<td>6</td>
<td>0.56 (0.25)</td>
</tr>
<tr>
<td></td>
<td>7/27/2013</td>
<td>6</td>
<td>2.23 (0.99)</td>
</tr>
<tr>
<td></td>
<td>10/7/2013</td>
<td>6</td>
<td>3.59 (1.77)</td>
</tr>
</tbody>
</table>
Table 3.6. Total belowground carbon flux (TBCF, g C m\(^{-2}\) gs\(^{-1}\)) for each stand. TBCF\(_{\text{top}}\) is defined as \(F_{\text{efflux}} + F_{\text{leaching}} - F_{\text{litter}} + \Delta(C_{\text{roots}} + C_{\text{soil}})\). TBCF\(_{\text{bottom}}\) is \(F_{\text{roots}} + F_{\text{resp}} + F_{\text{exudates}}\).

<table>
<thead>
<tr>
<th>Flux</th>
<th>White ash</th>
<th>Red oak</th>
<th>Eastern Hemlock</th>
</tr>
</thead>
<tbody>
<tr>
<td>TBCF(_{\text{top}})</td>
<td>633 (58)</td>
<td>791 (94)</td>
<td>474 (29)</td>
</tr>
<tr>
<td>TBCF(_{\text{bottom}})</td>
<td>449 (57)</td>
<td>577 (85)</td>
<td>340 (51)</td>
</tr>
</tbody>
</table>
Table 3.7. Activation energy ($E_a$, kJ mol$^{-1}$), pre-exponential constant (A, nmol CO$_2$ s$^{-1}$ g$^{-1}$), and $R^2$ of mass-specific rates of root respiration (nmol CO$_2$ g$^{-1}$ s$^{-1}$) for each stand over the growing season.

<table>
<thead>
<tr>
<th></th>
<th>$E_a$</th>
<th>A</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>White ash</td>
<td>20</td>
<td>20872</td>
<td>0.60</td>
</tr>
<tr>
<td>Red oak</td>
<td>40</td>
<td>36498539</td>
<td>0.81</td>
</tr>
<tr>
<td>Eastern hemlock</td>
<td>29</td>
<td>572865</td>
<td>0.56</td>
</tr>
</tbody>
</table>
Table 3.8. Statistical summary of mixed-effects models with fine root growth, mortality, respiration, NSC concentration, or exudation as the dependent variable. The fixed effects were stand, sample date, and year with plot as a random effect. Growth and mortality were log-transformed to meet assumptions of normality. *** $P < 0.001$, ** $P < 0.01$, * $P < 0.05$, . $P < 0.1$

<table>
<thead>
<tr>
<th></th>
<th>Growth</th>
<th>Mortality</th>
<th>Respiration</th>
<th>[NSC]</th>
<th>Exudation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>df</td>
<td>MS</td>
<td>F</td>
<td>df</td>
<td>MS</td>
</tr>
<tr>
<td>Stand</td>
<td>2</td>
<td>147.7</td>
<td>40.4 ***</td>
<td>2</td>
<td>106.5</td>
</tr>
<tr>
<td>Day of Year</td>
<td>1</td>
<td>13.8</td>
<td>3.8 *</td>
<td>1</td>
<td>0.01</td>
</tr>
<tr>
<td>Year</td>
<td>1</td>
<td>274.7</td>
<td>75.2 ***</td>
<td>1</td>
<td>25.3</td>
</tr>
<tr>
<td>Error</td>
<td>584</td>
<td>3.65</td>
<td>497</td>
<td>4.0</td>
<td>57</td>
</tr>
</tbody>
</table>
Figure 3.1. Map of plot locations at the Harvard Forest’s Prospect Hill Tract. There are 6 biometry plots (circles) and 10 minirhizotron plots (stars) per stand. Colored symbols indicate stand type. Blue-green symbols are white ash (*Fraxinus americana*), orange are red oak (*Quercus rubra*), and purple are eastern hemlock (*Tsuga canadensis*).
Figure 3.2. Growth (a-c), mortality (d-f), and net primary production (NPP, g-i) of fine roots. In 2012, n = 4 and n = 9 for red oak and eastern hemlock, respectively. In subsequent years, n = 10 for each stand. Error bars are standard error of the mean. There was a significant effect of year and stand on both growth and mortality (Table 3.8). The eastern hemlock stand had significantly less growth and mortality than red oak and white ash stands (P < 0.001), but red oak and white ash stands were not different from each other (growth was only marginally different, P < 0.1).
Figure 3.3. (a) Field measurements of monthly mean mass-specific fine root respiration (± SE) in 2013 (left y-axis). The shaded area represents soil temperature (right y-axis). (b) Relationship of fine root respiration to the temperature of the measurement chamber (n = 61). Lines are nonlinear fits of the Arrhenius function to data for each stand. (c) Fine root respiration (± SE) scaled to the growing season using fine root biomass from soil coring. In panels a and b, asterisk indicates that white ash had significantly higher respiration rates than red oak and eastern hemlock. In panel c, letters indicate significant differences between stands at the P < 0.001 level.
Figure 3.4. Fine root respiration measured in the lab at temperatures ranging between 16°C and 25°C, with a mean of 22.8 ± 0.5°C. There was a significant positive relationship between fine root respiration and sample date for white ash ($\beta = 0.016$, $F_{1,11} = 15.3$, $P < 0.01$, $R^2_{\text{adj}} = 0.54$), red oak ($\beta = 0.012$, $F_{1,9} = 17.6$, $P < 0.01$, $R^2_{\text{adj}} = 0.62$), and eastern hemlock ($\beta = 0.011$, $F_{1,9} = 32.1$, $P < 0.001$, $R^2_{\text{adj}} = 0.76$). Dash lines are 95% confidence intervals around the regression fit.
Figure 3.5. Total nonstructural carbohydrates (± SE) from n = 6 samples from each stand. There were significant differences between years ($F_{1,1089} = 23.7, P < 0.001$), sample dates ($F_{1,1089} = 66.9, P < 0.001$), and stands ($F_{2,1089} = 67.0, P < 0.001$). Each stand was significantly different from the other two stands in both years.
Figure 3.6. Gross primary production and belowground fluxes (litterfall, TBCF, fine root production, root respiration, exudation), heterotrophic respiration, soil CO$_2$ efflux) and pools (microbial biomass, root biomass, organic horizon soil C, mineral horizon soil C) for each stand. Heterotrophic respiration is estimated as the difference between soil CO$_2$ efflux and root respiration. Standard error is reported in parentheses next to each pool or flux value, and all units are g C m$^{-2}$ gs$^{-1}$. The growing season is defined as the six month period from May-October.
Figure 3.7. Time series of gross primary production estimated using data from the Harvard Forest Environmental Measurement Site flux tower and the hemlock tower in 2012 (circles). Soil respiration for the eastern hemlock and red oak stands (squares) and TBCF bottom (diamonds) for red oak, eastern hemlock and white ash stands calculated using root GPP, respiration and exudation from May to October. For Jan–Apr and Nov–Dec when data were not available (shaded areas), I used the median ratio of TBCF:GPP to extrapolate TBCF from GPP data. This ratio ($25^{th}$, $95^{th}$ percentiles) was 0.42 (0.23, 0.5) for hardwoods and 0.36 (0.28, 0.57) for hemlock. The orange and pink colored bars show the maximum canopy greenness in red oak and eastern hemlock stands, respectively, for this study period determined using PhenoCam data (http://phenocam.sr.unh.edu/).
CHAPTER FOUR: WHERE DOES THE RHIZOSPHERE END? SPATIALLY RESOLVED MEASUREMENTS OF IN SITU SOIL EXTRACELLULAR ENZYME ACTIVITY

Abstract

Rhizosphere soils are hotspots for soil organic matter decomposition, with soil decomposing enzymes stimulated up to 100%, but there are few estimates of rhizosphere spatial extent in the field. Zymography (i.e., 2-D visualization of enzyme activity) allows for spatially resolved in situ measurements of soil extracellular enzyme activity (EEA), but there has been no quantitative analysis of the relationship between EEA and distance from a root. The objective of this work is to develop a quantitative framework for zymogram image analysis as a means for estimating the spatial extent of the rhizosphere. My analysis utilizes a spatial error model and break-point regression analysis to estimate the scale over which four enzymes – beta-glucosidase, N-acetyl-beta-D-glucosaminidase, aminopeptidase, and acid phosphatase – vary as a function of distance from a root. This analysis compares two different types of enzyme substrates (i.e., colorimetric and fluorometric) at three different image resolutions. For each assay, it was possible to visualize and estimate the size of the rhizosphere. The resolution of image analysis did not affect the estimate of the rhizosphere extent for fluorometric assays, but did affect the estimate of rhizosphere extent for colorimetric assays, where the break-point regression was only significant at very-fine-image resolution. I also note methodological concerns specific to the type of enzyme substrate, such as uneven background staining in the colorimetric assay and diffusion of the substrate in the fluorometric assay, and provide
recommendations to improve the method. Zymography is a promising methodology with the potential to increase the spatial resolution of rhizosphere studies compared to methods that measure enzyme activity by physically removing soil.
Introduction

Soil is the largest terrestrial carbon (C) pool, storing more C than the atmosphere and terrestrial vegetation combined (Schlesinger & Bernhardt, 2013). A substantial portion of this pool is associated with plant roots. In the root-associated zone, or rhizosphere, soil-organic-matter-decomposing enzymes are stimulated up to 100% relative to surrounding bulk soil (Kourtev et al., 2002, Tscherko et al., 2004). Though the rhizosphere occupies a small soil volume relative to bulk soil, up to 40% of heterotrophic respiration can be attributed to the rhizosphere (Finzi et al., 2015). As a result, the rhizosphere has a disproportionate effect on biogeochemical cycling.

Roots exude low-molecular-weight C compounds such as organic acids and amino acids into the soil (Jones, 1998, Phillips et al., 2004). This labile C stimulates microbial growth and nutrient demand, resulting in the production of extracellular enzymes that decompose soil organic matter (SOM; Brzostek & Finzi, 2011). Previous research shows that root exudates can be found anywhere from 0.1 to 1.2 cm from the root surface (De Neergaard & Magid, 2001, Jones, 1998, Sauer et al., 2006, zu Schweinsberg-Mickan et al., 2010). Where root exudates are present, microbial biomass, respiration, enzyme activity, N mineralization, and soil organic matter decomposition are all increased relative to soil that is unaffected by roots (Finzi et al., 2015, Phillips et al., 2012).

At present, measurements of microbial and extracellular enzyme activity (EEA) use coarse approaches to separate rhizosphere and bulk soil. For example, some studies consider the rhizosphere to be the soil adhering to the root after gentle shaking or soil that
is $\leq 2$ mm from the root surface (Landi et al., 2006, Phillips & Fahey, 2006). Of the studies that try to estimate diffusion distance of exudates in order to determine the rhizosphere extent (Darrah, 1991a,b, Kuzyakov et al., 2003, Raynaud et al., 2003), most assay exudate concentration at up to 10 distances from the root surface by cutting the soil into sections of $\sim 1 – 2$ mm and analyzing the homogenized soil sample (Nuruzzaman et al., 2006, Sauer et al., 2006, zu Schweinsberg-Mickan et al., 2010).

Two methods represent a recent advance in the visualization of enzyme activity using zymography, and allow for spatially resolved in situ measurements of soil EEA at finer resolution than traditional studies (Dong et al., 2007, Spohn et al., 2013). These methods produce an image where the brightness, hue, or intensity of coloration or fluorescence corresponds to the amount of enzyme activity in a given area of the image, and provide the opportunity to quantify the relationship between EEA and distance from a root at a very fine spatial resolution, without compromising the integrity of root-microbe interactions. To date, where these methods have been applied, there has been no quantitative analysis of the relationship between EEA and distance from a root (Dong et al., 2007, Spohn et al., 2013).

The objective of this study is to develop a quantitative framework for analyzing zymograms, in order to determine the relationship between EEA and distance from a root. This is part of a longer-term objective of quantifying rhizosphere volume for different species and ecosystems. Here I propose two statistical methods to quantify this relationship. The first is linear regression with a spatially autocorrelated error term in order to account for the lack of independence between neighboring pixels in an image.
The second is “break-point” regression analysis, which determines the location of a change in the slope of the linear regression. Within this context, I also compare enzyme substrates that leave either a visual or fluorescent tag when degraded. Colorimetric substrates, such as that for acid phosphatase and aminopeptidase, leave behind a colored stain following reaction with an enzyme, whereas fluorometric substrates, such as that for beta-glucosidase and N-acetyl-beta-D-glucosaminidase leave behind a fluorescent stain. The previous study that used both colorimetric and fluorescent substrates found qualitative differences in the staining pattern of the assays (Dong et al., 2007). Both types of assays were applied to rooted soil surfaces in three mono-dominant stands at the Harvard Forest in central Massachusetts. To determine whether image resolution affects the scaling of EEA and distance from a root, I compared images analyzed at three spatial scales.

Methods

Site description

The field portion of this study was conducted at the Harvard Forest Long Term Ecological Research Site in Petersham, MA (42°N, 72°W, elevation 340 m). The site is located on former agricultural land that was abandoned in the mid-1800s allowing forest regrowth beginning late in the 19th century (Foster et al., 2003). Soils are Typic Distrochrepts derived from glacial deposits of granite, schist and gneiss. The dominant tree species in this tract are northern red oak (*Quercus rubra*) and red maple (*Acer rubrum*), with smaller populations of eastern hemlock (*Tsuga canadensis*), white ash (*Fraxinus americana*), white pine (*Pinus strobus*) and red pine (*Pinus resinosa*).
Twelve root boxes were constructed at Boston University and installed in the field in March 2012. Each root box was built from 0.9 cm thick clear polycarbonate sheeting. Each side is 40 × 40 cm, with two sides beveled out at 15° to form a rhomboid. Root boxes were set in soil pits and backfilled with excavated soil, maintaining the mineral and organic horizon delineation. Each box was equipped with two 20 × 20 cm removable panels that allow for access to the rooted soil surface (Figure 4.1). Four root boxes (n = 8 panels) were installed in each of three monodominant stands: white ash, red oak, and eastern hemlock. I assume that the > 2 years between root box installation and data collection allowed sufficient time for soils to recover from disturbance and form dense root networks.

Each root box panel (n = 24) was assessed for the activity and spatial extent of four extracellular enzymes: acid phosphatase (AP), aminopeptidase (PR), beta-glucosidase (BG), and N-acetyl-beta-D-glucosaminidase (NAG) in late June – early July 2014. EEA was measured using zymograms prepared following Dong et al. (2007). Briefly, a 20 × 20 cm sheet of chromatography paper was soaked in a buffer solution containing colorimetrically (AP, PR) or fluorescently (BG, NAG) labeled substrate for each enzyme, alpha naphthyl phosphate (AP), L-leucyl 2-naphthylamide (PR), 4-methylumbelliferyl-beta-glucopyranoside dehydrate (BG), or 4-methylumbelliferyl-N-acetyl-beta-glucosaminide (NAG). Standards were made by applying acid phosphatase from wheat germ (AP), aminopeptidase from Aspergillus oryzae (PR), or fluorescent 4-methylumbelliferone (BG, NAG) to prepared zymograms. The units for acid phosphatase and aminopeptidase activity are enzyme units per milliliter (EU ml⁻¹), defined as the
amount of enzyme in one milliliter that is needed to convert 1μmol of substrate per minute, and for fluorescent 4-methylumbelliferone, millimolar (mM). Each zymogram (N = 96; 8 root box panels × 3 stands × 4 enzymes) was incubated against the soil surface for 30 minutes (BG, NAG) to 1 hour (AP, PR) depending on the type of assay. Immediately following incubation, zymograms were transported back to Boston University where they were dried, and imaged using a Ricoh 907Ex 600dpi color scanner (AP, PR; Ricoh Electronics, Inc., Tustin, CA) or an AlphaImager HP UV gel scanner (BG, NAG; ProteinSimple, San Jose, CA).

All image processing was done in the open source software ImageJ (Rasband, 1997). Digital photographs of open root box panels taken at the time of incubation were cropped, corrected for image distortion, and converted to 8-bit black and white. The background of each image was detected using a rolling ball algorithm (Sternberg, 1983). This algorithm averages over a 50 pixel area around each pixel to compute a local background value which is then subtracted from that pixel. Roots were selected by pixel brightness and false positives (e.g., light colored soil aggregates, insects, rocks) were removed manually using the selection tool. The selection of rooted area was converted to a thresholded mask, where pixels containing roots were given a value of 0 (i.e., black) and non-root pixels were given a value of 255 (i.e., white). The zymogram images were also converted to 8-bit black and white. Digital photographs and zymogram images were compressed to 30 × 30, 100 × 100, or 1500 × 1500 pixels using bilinear interpolation, corresponding to an effective pixel length of 6.7 mm, 2 mm, or 0.13 mm, respectively. The x- and y-coordinate position of each pixel was used to overlay the root threshold
image onto zymogram images. Image pixel brightness values and their $x$- and $y$-coordinate position were output to R Statistical Software (R Development Core Team, 2013), where all subsequent analyses took place.

In order to determine the relationship between EEA (EU ml$^{-1}$ or mM) and distance from a root (mm), Euclidian distance from the nearest root was determined for each pixel in the zymogram image by creating a vector with the location of each root-containing pixel, and finding the element in this vector with the minimum distance to each pixel. Image pixel brightness values were converted to EU ml$^{-1}$ or mM using a linear relationship between the standards and their pixel brightness values for AP ($\beta = -4.9$, $F_{1,3} = 28.0$, $R^2_{\text{adj}} = 0.87$, $P < 0.05$), PR ($\beta = -1.3$, $F_{1,3} = 15.9$, $R^2_{\text{adj}} = 0.79$, $P < 0.05$), NAG ($\beta = -45$, $F_{1,3} = 60.5$, $R^2_{\text{adj}} = 0.93$, $P < 0.01$), and BG ($\beta = -47$, $F_{1,2} = 22.3$, $R^2_{\text{adj}} = 0.88$, $P < 0.05$; Figure 4.2). High brightness values correspond to low enzyme activity due to the convention of setting brightness values to 0 for black and 255 for white pixels. I considered the total area of all pixels containing roots to be the rooted area for each stand. I used ANOVA and Tukey’s HSD to test for differences in rooted area among the three stands.

I tested for the presence of spatial autocorrelation between neighboring pixels using the Moran’s I statistic. I used the `moran.test` function in the `spdep` package for R Statistical Software (Bivand et al., 2014). To establish a relationship between EEA and distance from a root while accounting for spatial autocorrelation between pixels and their neighbors, I used a spatial simultaneous autoregressive error model of the form:
\[ Y = \beta_1 x_1 + \beta_2 x_2 + \beta_3 + \varepsilon, \text{ and} \]
\[ \varepsilon = \rho W e + u \]

where \( Y \) is EEA (EU ml\(^{-1}\) or mM), \( x_1 \) is distance from root (mm), \( x_2 \) is depth (mm), and \( \beta_1, \beta_2, \) and \( \beta_3 \) are linear regression coefficients. The error term, \( \varepsilon \), contains a spatially structured error term, \( \rho W e \), where \( \rho \) is a fit coefficient, \( W \) is the spatial weight matrix, \( e \) is the spatial error term, and \( u \) is a random residual. This model assumes that neighboring pixels are not independent of one another, and as a result the predicted value for a given pixel depends in part on the value of its neighbors. The fit parameter \( \rho \) corresponds to the strength of autoregression between neighboring pixels on model residuals. I fit the spatial error model to data from each image using the `errorsarlm` function in the `spdep` package for R Statistical Software (Bivand et al., 2014).

To test for the location of a change in the slope of the relationship, or break point, between EEA and the distance from a root, I used a weighted “break-point” regression. I used the break point as a proxy for the rhizosphere extent. EEA was fit as a function of distance from a root using the equation:

\[ Y = \beta_1 x + \beta_2 (x - c) + \gamma I(x > c) \]  

where \( x \) is the Euclidian distance from the root (mm), \( \beta_1 \) and \( \beta_2 \) are slope and intercept parameters, \( c \) is the break point, and \( I(x > c) \) is an index variable that is equal to zero when \( x < c \). The model minimizes \( \gamma \), a measure of the distance between the two segments in order to constrain the segments to be close to continuous.

I observed that the sample size (i.e., number of pixels represented at each root distance) decreased as distance from a root increased (Figure 4.3). As a result, rather than
minimizing the sum of squared residuals (SSE) as in ordinary least squares, I minimized the weighted SSE:

\[
\text{Weighted SSE} = \sum_{i=1}^{n} w_i (y_i - \hat{y}_i)^2 \tag{4}
\]

where \(w_i\) is a vector of weights proportional to the sample size such that

\[
w_i = \frac{N_i}{\sum_{i=1}^{n} N_i} \tag{5}
\]

and \(N_i\) is the number of pixels at each root distance \(i\) to \(n\). This model was fit using the segmented function in the segmented package for R statistical software (Muggeo, 2008).

I used the spatial autoregressive error model (Eq. [1]) on images at coarse and fine resolution in order to determine if there was an effect of image resolution on the relationship between EEA and distance from a root. At coarse image resolution (30 × 30 pixels) each pixel was 6.7 mm in length, corresponding to an area of 0.44 cm\(^2\) per pixel. At fine image resolution (100 × 100 pixels) each pixel was 2 mm in length, corresponding to an area of 0.04 cm\(^2\) per pixel.

I used break-point regression (Eq. [3]) on images at coarse, fine, and very fine resolution in order to determine if there was a relationship between pixel size and break-point value. For very-fine-resolution image analysis, I examined a subsection of a 1500 × 1500 pixel-resolution image focusing on a rooted area in hemlock soil (Figure 4.4a, Box 4.1). In this image, the length of each pixel was 0.13 mm, corresponding to an area of 1.7×10\(^{-4}\) cm\(^2\).
Results

Visual inspection

White ash and red oak stands were more sparsely rooted than eastern hemlock with 14 ± 1.0 cm\(^2\) and 15 ± 1.1 cm\(^2\) of rooted area out of the 400 cm\(^2\) sample area, respectively (\(F_{2,93} = 34.5, P < 0.001\), Table 4.3). Eastern hemlock images contained 27 ± 1.7 cm\(^2\) of rooted area. The mean area covered by roots across the three stands is 18.7 cm\(^2\), approximately 5% of the sample area.

The staining pattern on scanned zymogram images was visibly different for colorimetric compared to fluorometric assays. The colorimetric acid phosphatase assay had the most localized staining (Figure 4.4a). Aminopeptidase zymograms developed the least amount of staining (60% of pixels had no detectable EEA) compared to AP (10%), BG (1.7%), or NAG (3.2%). However, when PR did stain, it was 20–40% brighter than the other zymograms (Figure 4.4b). Beta-glucosidase and N-acetyl-beta-D-glucosaminidase were fluorometrically assayed. I found this assay to be relatively diffuse, often with large areas of high EEA covering >50% of the zymogram (Figure 4.4c,d).

At coarse image resolution, clusters of roots and EEA were visible, but not fine root morphology (Figure 4.5a). At fine resolution, individual fine roots were visible (Figure 4.5b), but it was not possible to determine the diameter of fine roots < 2 mm. At both image resolutions, there appeared to be fewer pixels with high EEA at 0 – 10 mm from the top of the image in BG and NAG images (data not shown). At very fine resolution, it was possible to distinguish between small- and large-diameter roots (Figure
There was visible overlap in many images between locations of rooted pixels and locations with high enzyme activity at fine resolution. However, in some cases there was a 1–5 mm misalignment of images, where EEA associated with a root was offset from the thresholded mask of the root as a result of manual error in the image analysis steps that were not fully automated (i.e., root thresholding, image distortion correction).

**Coarse image resolution**

EEA ranged between 0–10 EU ml\(^{-1}\) for AP, 0–40 EU ml\(^{-1}\) for PR, and 0–4.9 mM for BG and NAG. EEA generally declined as distance from a root increased (Figure 4.6). For all images, there was significant spatial autocorrelation (Moran’s I, P < 0.05). Of the 96 images analyzed, 54 had a significant relationship between EEA and distance from a root, including 51 which had a significant negative relationship (Table 4.1). Red oak had the greatest proportion of significant negative relationships between EEA and root distance (78%) of the three stands. Acid phosphatase had the greatest proportion (88%) of significant negative relationships of the four enzymes, and aminopeptidase had the fewest (25%). Forty-three out of 48 significant spatial error models predicted that EEA declined with depth (Table 4.1). Eastern hemlock had the greatest proportion of significant and negative relationships with depth (60%) of the three stands.

In the break-point regression model, the relationship between EEA and distance from a root was not significant for the colorimetrically assayed enzymes (AP, PR). Fluorometrically assayed enzymes (BG, NAG) had break points of 7.5 cm and 1.8 cm, respectively (Figure 4.6). Averaging across the enzymes with a significant relationship between EEA and distance to a root (BG, NAG), I identified break points at 5.8 cm and
4.7 cm for white ash and red oak (Figure 4.7). The break-point regression model for
eastern hemlock was not significant.

*Fine image resolution*

At fine image resolution, computing time increased 72-fold compared to coarse
resolution. The increased resolution allowed us to observe a greater range of brightness
values compared to coarse resolution. EEA ranged between 0-30 EU ml$^{-1}$ for AP, 0-170
EU ml$^{-1}$ for PR, and 0-5.3 mM for BG and NAG. The break point was closer to the root
surface than at coarse resolution, reflecting the closer association between pixel size and
root size (Figure 4.8).

A spatially autoregressive error model fit to the data found 84 of 96 significant
relationships between EEA and distance from a root, of which 39 were negative (Table
4.2). BG and NAG images had the greatest number of negative regression coefficients,
while AP and PR had a majority of positive coefficients, indicating that EEA increased
with distance from a root. Sixty-eight of 96 images had a significant relationship with
depth. Of these, 57 were positive relationships.

In the break-point regression model, the relationship between EEA and root
distance was not significant for AP and PR. The break points for BG and NAG were 6.7
and 1.5 cm, respectively (Figure 4.8). The break-point regression between EEA and
distance from a root for eastern hemlock was not significant. Break points for white ash
and red oak were identified at 7.1 and 1.2 cm, respectively (Figure 4.9).
Very fine image resolution

In a very-fine-resolution subsection of an acid phosphatase assay in the hemlock stand (Figure 4.4, Box 4.1), the maximum observable EEA was 36 EU ml\(^{-1}\), compared to 32 EU ml\(^{-1}\) in the fine resolution and 9 EU ml\(^{-1}\) in the coarse resolution analysis of the same enzyme. The median EEA (1.6 EU ml\(^{-1}\)) was identical to that at coarse resolution. EEA was high within 10 mm of the root, but declined quickly thereafter. Break-point regression placed the break-point location at 2.6 mm (Figure 4.10).

Discussion

There were differences in the localization of EEA in colorimetric compared to fluorometric assays, with colorimetric assays resulting in more specific staining compared to fluorometric assays. This is likely the result of the fluorometric tag’s ability to diffuse when wet (Spohn & Kuzyakov, 2013). The colorimetric aminopeptidase assay resulted in an average of 60% of the image with no detectable EEA, indicating that there was little aminopeptidase enzyme activity in the soils that I sampled. In the colorimetric acid phosphatase assay, there was an increase in EEA with distance from a root at the fine resolution that I did not observe at coarse resolution. I did not expect this result given the visual correspondence between EEA and roots in this assay. However, the acid phosphatase assay developed an orange-colored background. Uneven background color and image misalignment are two methodological concerns that would be exacerbated at fine resolution compared to coarse resolution, given the wider range of EEA and smaller pixel size, respectively. Analysis of a subset of a very-fine-resolution image suggests that a negative relationship between EEA and root distance is observable in colorimetric
assays at high resolution if the background color is even and the zymograph and thresholded root mask are well-aligned.

Rhizosphere extent across stand types and enzyme classes

For the fluorometric assays, EEA declined with increasing distance from a root. Break point analysis predicted that the rhizosphere extent ranged from 0.26 to 7.5 cm, depending on the resolution of analysis. Direct sampling and isotopic labeling studies place the rhizosphere extent between 0.1 and 1.2 cm (De Neergaard & Magid, 2001, Dessureault-Rompré et al., 2007, Falchini et al., 2003, Sauer et al., 2006, zu Schweinsberg-Mickan et al., 2010), but another study using a similar method to the one used here recorded enhanced EEA up to 6 cm away from the root surface (Spohn & Kuzyakov, 2014). It is therefore possible that recent estimates of rhizosphere contributions to decomposition and nutrient mineralization based on an exudate diffusion distance of 2 mm are very conservative (Finzi et al., 2015).

Effects of image analysis resolution

The resolution of the image analysis affected the spatial distribution of EEA, the break-point location, and the influence of artifacts and image alignment. As resolution increased, the maximum observable EEA increased because there was a larger sample size and less averaging of high EEA pixels with pixels of intermediate and low EEA. In addition, the distance of the break point to the root surface declined due to greater visibility of root morphology relative to staining of EEA (Figure 4.5). Generally, this
suggests that matching the scale of resolution to the scale of fine root architecture will increase the accuracy of both EEA and break point estimates.

At fine image resolution, the spatial error model predicted a mixture of positive and negative relationships between EEA and distance from a root. There were a surprising number of instances where the relationship between EEA and distance from a root was positive. This was more common for the colorimetric assays, AP and PR (Figure 4.8). For AP in particular, background staining developed from the substrate solution, a mix of α-naphthyl phosphate and Fast Red TR. Though care was taken to apply the substrate evenly, there was variation in the background coloring of the zymograms. At high resolution, this variation in background color may have introduced artifacts into the assessment of EEA using pixel brightness. This was somewhat accounted for by subtracting the background using a rolling ball algorithm, but with very unevenly stained backgrounds such as that in Figure 4.4a, there may have been some pixels with under- or over-estimated EEA.

Image misalignment was another error that was exacerbated at fine resolution. Images were scaled to identical size and overlayed automatically, but root images were corrected for pixel distortion and cropped manually, and no image-aligning algorithm was used. Any misalignment of images would be on the scale of < 1 mm to 5 mm, so coarse image resolution that averages over large root clusters may alleviate errors that are a result of misalignment.

I wanted to explore whether or not there was value in investing the additional computing time to analyze very-fine-resolution images in order to more accurately
represent fine root morphology. I selected a subset of the zymogram pictured in Figure 4.4a (Box 4.1) that appeared to have even background staining and be well aligned with the image of the root. In this subset, EEA declined steeply as distance from the root increased up to 2.6 mm (Figure 4.10). From 2.6 mm to 7 cm the relationship was relatively flat. This analysis is not replicated, but suggests that high resolution images can be used to estimate relationships between EEA and roots if methodological concerns are alleviated.

Fluorescent clustering of EEA in the center of the image may result from the diffusion of the substrate when wet. In colorimetric zymograms, there was more EEA at the surface of the soil than at depth, corresponding with high root abundance. In the fluorescent zymograms, however, there was comparatively less EEA at the soil surface, suggesting that as the zymograms are incubated in a vertical position, the fluorescent tag diffused downward in response to gravity. Zymograms were moistened before incubation, and the 4-methylumbelliferone tag is known to diffuse readily when wet (Spohn & Kuzyakov, 2013). Because the zymogram staining is diffuse, coarse-resolution image processing is sufficient to represent the correspondence between EEA and root distribution. Thus, fine resolution is only necessary when the zymogram staining also has fine spatial resolution, as in the colorimetric acid phosphatase assay.

Methodological considerations

This methodology can be improved with image-analysis techniques. I counted the number of black pixels in the root image to estimate the proportion of sample area covered by roots and found that red oak stands were more sparsely rooted than hemlock.
From a visual inspection of the root boxes, it is clear that red oak root boxes are heavily rooted, but that these roots are very fine and similar in color to the soil. In contrast, hemlock roots are thick and light-colored compared to the soil (Figure 4.1). As a result, a greater number of roots in the hemlock stand were successfully thresholded relative to the red oak stand. A different annotation method or an edge-detection algorithm may improve root detection in stands with roots that are similar in color to their background (Canny, 1986, Lobet et al., 2011). Higher resolution images may also aid in detecting very fine roots that may be smaller than the pixel size.

Some patches of high EEA were unrelated to root activity. This method cannot yet annotate mycorrhizal hyphae or detect other possible microbial hotspots associated with invertebrate casts or litter that may not be near a root surface. Additionally, not all roots produce an active rhizosphere. Exudates are primarily released from growing root tips, and assuming an average root growth rate of 1 μm s⁻¹ and a 30-minute half-life for exudates, then at 30 mm behind the growing root tip there will be < 1% of the original exudate remaining (Hirsch et al., 2013). Finally, my thresholding function does not distinguish between suberized roots, dead roots, absorptive roots, or growing root tips, and therefore areas of high root proliferation with low EEA may reflect different demographic stages of the root. For this reason, coarse resolution averaging over thickly rooted areas may partially alleviate this issue. An alternative would be to manually annotate fine absorptive roots and root tips.
Conclusions

In order to scale up measurements of microbial activity (e.g., enzyme activity, microbial biomass) made in bulk and rhizosphere soils to an ecosystem scale estimate, some assumption of the volume of the rhizosphere is necessary. Finzi et al. (2015) used a numerical model to estimate the volume of the rhizosphere, and demonstrated that changing the assumption of the exudate diffusion distance can cause estimates of the proportion of soil that is in the rhizosphere to vary by >20%.

Soil zymography provides the opportunity to generate statistical relationships between EEA and distance from a root, in order to determine the rhizosphere extent of a given soil. Zymograms are easy to prepare and deploy in the field, and allow for spatial analysis of multiple enzymes across heterogeneous soil surfaces. Many techniques in automatic annotation, background correction, and image alignment are well-developed and can be applied to this method (Amat et al., 2008, Nobis & Hunziker, 2005, Ritchie et al., 2007). The main methodological concerns for this method are evenness of background staining and image alignment for the colorimetric assay and diffusion of the tag in the fluorometric assay. Methodological improvements such as using agarose gel to keep soil particles from quenching the fluorescent tag are also promising (Spohn et al., 2013). In this method, enzymes can diffuse through the gel to the zymogram, but soil particles do not contact the zymogram, resulting in high staining accuracy.

To determine the rhizosphere extent of enzymes that degrade a colorimetric substrate, I make the following recommendations:

- Use very fine image resolution (~0.13 mm)
- Use an image-aligning algorithm
- Annotate root tips and absorptive roots

To determine the rhizosphere extent of enzymes that degrade a fluorometric substrate, I make the following recommendations:

- Use coarse or fine image resolution (2 mm – 6 mm)
- Dry zymogram flat after substrate application and soil incubation to reduce vertical diffusion
- Prepare zymogram using agarose gel layer, following Spohn *et al.*, (2013)

I have provided a quantitative framework to analyze and estimate the rhizosphere extent from image data using a zymographic method. This analysis could be expanded to include uncertainty associated with the linear parameters estimated in both the spatial error regression and the break-point regression using Bayesian or Monte Carlo techniques. In addition, nesting images within stands, sites, and region may eventually allow for geographically specific estimates of rhizosphere extent interpolated between locations for which zymographic data has been collected. Given access to a rooted soil surface, preparation and incubation of zymograms is a simple and relatively inexpensive process. Once a quantitative framework for image analysis is adopted, determination of rhizosphere extent using zymograms will be much easier than cutting, extracting, and measuring enzyme activity from soil directly.
Table 4.1. Summary of spatial error model regression coefficients estimated at coarse resolution. Enzyme activity (EU ml\(^{-1}\) or mM) is modeled as a function of distance from a root (mm) and soil depth (mm). \(\rho\) is the spatial error coefficient. AIC is the Akaike information criterion, a model-selection score that balances goodness-of-fit against the number of parameters. Regression coefficients, \(\rho\), and AIC for each image were averaged to obtain one value for each stand and enzyme. AP = acid phosphatase, PR = aminopeptidase, BG = beta-glucosidase, NAG = N-acetyl-beta-D-glucosaminidase, F = white ash, Q = red oak, T = eastern hemlock.

<table>
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<th>Enzyme</th>
<th>Stand</th>
<th>Intercept</th>
<th>Distance</th>
<th>Depth</th>
<th>(\rho)</th>
<th>AIC</th>
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<tr>
<td>Q</td>
<td>1.60</td>
<td>-0.009</td>
<td>-0.08</td>
<td>0.98</td>
<td>1283</td>
<td></td>
</tr>
<tr>
<td>T</td>
<td>2.84</td>
<td>-0.005</td>
<td>-0.16</td>
<td>1.00</td>
<td>2131</td>
<td></td>
</tr>
</tbody>
</table>
Table 4.2. Summary of spatial error model regression coefficients estimated at fine resolution. Enzyme activity (EU ml⁻¹ or mM) is modeled as a function of distance from a root (mm) and soil depth (mm). ρ is the spatial error coefficient. AIC is the Akaike information criterion, a model-selection score that balances goodness-of-fit against the number of parameters. Regression coefficients, ρ, and AIC for each image were averaged to obtain one value for each stand and enzyme. AP = acid phosphatase, PR = aminopeptidase, BG = beta-glucosidase, NAG = N-acetyl-beta-D-glucosaminidase, F = white ash, Q = red oak, T = eastern hemlock.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Stand</th>
<th>Intercept</th>
<th>Distance</th>
<th>Depth</th>
<th>ρ</th>
<th>AIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>AP</td>
<td>F</td>
<td>0.58</td>
<td>0.058</td>
<td>0.87</td>
<td>0.96</td>
<td>36234</td>
</tr>
<tr>
<td></td>
<td>Q</td>
<td>0.30</td>
<td>0.060</td>
<td>0.85</td>
<td>0.96</td>
<td>39796</td>
</tr>
<tr>
<td></td>
<td>T</td>
<td>1.69</td>
<td>0.029</td>
<td>0.01</td>
<td>0.96</td>
<td>33604</td>
</tr>
<tr>
<td>PR</td>
<td>F</td>
<td>-2.23</td>
<td>0.185</td>
<td>0.06</td>
<td>0.96</td>
<td>59046</td>
</tr>
<tr>
<td></td>
<td>Q</td>
<td>-4.19</td>
<td>0.190</td>
<td>0.10</td>
<td>0.96</td>
<td>58653</td>
</tr>
<tr>
<td></td>
<td>T</td>
<td>-2.19</td>
<td>0.209</td>
<td>0.08</td>
<td>0.95</td>
<td>61021</td>
</tr>
<tr>
<td>BG</td>
<td>F</td>
<td>-4.46</td>
<td>-0.010</td>
<td>0.05</td>
<td>1.00</td>
<td>24530</td>
</tr>
<tr>
<td></td>
<td>Q</td>
<td>-5.63</td>
<td>-0.019</td>
<td>0.05</td>
<td>1.00</td>
<td>27673</td>
</tr>
<tr>
<td></td>
<td>T</td>
<td>-5.46</td>
<td>-0.005</td>
<td>0.03</td>
<td>1.00</td>
<td>23748</td>
</tr>
<tr>
<td>NAG</td>
<td>F</td>
<td>-2.30</td>
<td>-0.004</td>
<td>-0.01</td>
<td>1.00</td>
<td>19855</td>
</tr>
<tr>
<td></td>
<td>Q</td>
<td>-8.07</td>
<td>-0.006</td>
<td>0.04</td>
<td>1.00</td>
<td>16399</td>
</tr>
<tr>
<td></td>
<td>T</td>
<td>-0.44</td>
<td>-0.005</td>
<td>-0.01</td>
<td>1.00</td>
<td>24662</td>
</tr>
</tbody>
</table>
Table 4.3. Tukey’s HSD test on an ANOVA model of rooted area (cm$^2$) per 20 $\times$ 20 cm panel as a function of stand. Different superscript letters denote significant differences at P < 0.05.

<table>
<thead>
<tr>
<th>Stand</th>
<th>Rooted area (cm$^2$)</th>
<th>6mm</th>
<th>2mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>White ash</td>
<td></td>
<td>13.9 ± 0.84$^b$</td>
<td>13.6 ± 1.0$^b$</td>
</tr>
<tr>
<td>Red oak</td>
<td></td>
<td>17.1 ± 1.1$^b$</td>
<td>14.9 ± 1.1$^b$</td>
</tr>
<tr>
<td>Eastern hemlock</td>
<td></td>
<td>26.6 ± 1.9$^a$</td>
<td>27.4 ± 1.7$^a$</td>
</tr>
</tbody>
</table>
Figure 4.1. Plexiglass root box designed and constructed for this study at Boston University. Roots can be accessed by lifting a hinged panel. The second hinged panel is partially visible in the lower right hand corner.
Figure 4.2. Linear relationship between enzyme activity and brightness values for the zymogram area, determined by staining the zymogram with a range of acid phosphatase, protease, or 4-MUB concentrations. AP = acid phosphatase, PR = aminopeptidase, BG = beta-glucosidase, NAG = N-acetyl-beta-D-glucosaminidase.
Figure 4.3. Sample size (number of pixels) in each 1 mm distance from root bin at coarse (6.7 mm) and fine (2 mm) image resolution.
Figure 4.4. Zymograms measuring the activity of (a) acid phosphatase, (b) aminopeptidase, (c) beta-glucosidase, and (d) N-acetyl-beta-D-glucosaminidase on the same hemlock stand root box panel. Thresholded mask of roots is overlaid on each zymogram at 70% transparency. Box 4.1 is the image subset used in very-fine-resolution analysis.
Figure 4.5. Level of detail visible across a range of spatial resolution. Length of the side of one pixel is, (a) 6.7 mm, (b) 2 mm, or (c) 0.13 mm (resolution of original 8-bit image).
Figure 4.6. Enzyme activity (EU ml⁻¹ or mM) versus distance from root (mm) for four extracellular enzymes: acid phosphatase (AP), aminopeptidase (PR), beta-glucosidase (BG), and N-acetyl-beta-D-glucosaminidase (NAG) at coarse resolution. The broken solid line is the best-fit break-point regression line. The dotted vertical line indicates the location of the break point. The shaded gray area is the interquartile range of the data.
Figure 4.7. Enzyme activity (EU ml\(^{-1}\) or mM) versus distance from root (mm) for white ash, red oak, and eastern hemlock stands at coarse resolution. The broken solid line is the best-fit break-point regression line. The dotted vertical line indicates the location of the break point. The shaded gray area is the interquartile range of the data.
Figure 4.8. Enzyme activity (EU ml$^{-1}$ or mM) versus distance from root (mm) for four extracellular enzymes, acid phosphatase (AP), aminopeptidase (PR), beta-glucosidase (BG), and N-acetyl-beta-D-glucosaminidase (NAG) at fine resolution. The broken solid line is the best-fit break-point regression line. The dotted vertical line indicates the location of the break point. The shaded gray area is the interquartile range of the data.
Figure 4.9. Enzyme activity (EU ml⁻¹ or mM) versus distance from root (mm) for white ash, red oak, and eastern hemlock stands at fine resolution. The broken solid line is the best-fit break-point regression line. The dotted vertical line indicates the location of the break point. The shaded gray area is the interquartile range of the data.
Figure 4.10. Enzyme activity (EU ml\(^{-1}\)) in a 400 × 400 pixel subsection of a high-resolution photo (1500 × 1500 pixels). Break point for this subsection is 2.6 mm.
CHAPTER FIVE: A PARSIMONIOUS MODULAR APPROACH TO BUILDING
A MECHANISTIC BELOWGROUND C AND N MODEL

Abstract

More carbon (C) is stored in soil than in the atmosphere and terrestrial vegetation combined, and microbial decomposition of soil organic matter (SOM) makes up the largest proportion of the flux of carbon dioxide from the soil to the atmosphere. As a result, microbial activity can have a large effect on the global C cycle. Microbial decomposition and uptake of dissolved substrate is sensitive to temperature and substrate supply, the latter of which can be limited by diffusion of substrate to microbial extracellular enzymes or by oxygen limitation. Many decomposition models use temperature and soil moisture in a linear decomposition coefficient, and few explicitly model the processes that limit substrate supply on the micro-site level. In order to build process-level representation of the effect of temperature and substrate supply on microbial physiology, I merged a model that uses temperature and substrate supply to predict depolymerization rate with a C and nitrogen (N) microbial physiology model.

I tested the performance of the combined model relative to each model alone using measurements of heterotrophic respiration in a mid-latitude forest located in central Massachusetts, USA. I then applied the combined model to theoretical concepts such as the response of microbial activity to varying ratios of C-to-N (C:N), and global change-inspired perturbations to mean annual temperature and soil moisture. The combined model predicted realistic C efflux during transient wet-up events, but over-predicted C efflux in constant high soil moisture conditions. Model predictions of SOM
decomposition in response to variable C:N inputs are consistent with the observation that wide C:N stands store more C than narrow C:N stands. The combined model reproduced realistic responses to temperature perturbation, and thus may be useful for investigating the interaction between substrate and temperature controls on soil C storage at regional and global scales.
Introduction

Soil is the largest terrestrial carbon (C) pool, and the flux of CO₂ from the soil to the atmosphere is dominated by microbial decomposition (Schlesinger & Bernhardt, 2013). Hence, the rate of C mineralized by soil micro-organisms affects the global C cycle. A number of biotic and abiotic factors control the amount of C that microbes release via decomposition. Soil microbes produce extracellular enzymes in order to depolymerize soil organic matter (SOM), but they can also utilize C and N substrates released by plant roots (Brzostek & Finzi, 2011, Frey et al., 2013). Labile organic C and N exuded from plant roots can induce microbial population growth and nutrient limitation, which results in enhanced production of microbial extracellular enzymes that decompose SOM (Brzostek et al., 2013, Kuzyakov, 2010).

The carbon-to-nitrogen (C:N) ratio of root exudates has been shown to affect depolymerization rates, with root inputs of C and N together stimulating decomposition more than additions of C alone (Drake et al., 2013). Since litter and root inputs affect microbial activity, differences in the C:N of inputs owing to differences in species traits may affect the function of soil microbial communities. For example, a stand with acidic soils that supports ectomycorrhizal fungi such as eastern hemlock (Tsuga canadensis) has a wider litter and root C:N ratio as well as a higher fungal-to-bacterial (F:B) ratio relative to a stand such as white ash (Fraxinus americana), that has narrow litter C:N and a low F:B ratio (Finzi et al., 1998, Strickland & Rousk, 2010).

Microbial activity is also temperature sensitive, with depolymerization of SOM and uptake of dissolved organic carbon (DOC) conforming to Arrhenius kinetics (Lloyd
& Taylor, 1994). However, soil temperature is only the dominant driver of microbial activity when substrate supply is not limiting. When substrate supply is limited by diffusion or oxygen ($O_2$) limitation, then decomposition conforms to Michaelis-Menten kinetics (Davidson & Janssens, 2006). Substrate and $O_2$ limitation have been demonstrated using field experiments measuring carbon dioxide ($CO_2$) efflux under different soil moisture conditions (Davidson et al., 1998, McNicol & Silver, 2015). Physical separation between substrate and enzyme is cited as a mechanism by which SOM persists through time (Schmidt et al., 2011).

Despite well-known effects of temperature and substrate supply on the activity of soil microbes, microbial processes have not been modeled explicitly in terrestrial biosphere models. In these models, decomposition rate is determined using a linear rate constant that may vary as a function of temperature or soil moisture (Bolker et al., 1998, Jenkinson et al., 1990). Indeed, none of the models in the Fifth Coupled Model Intercomparison Project (CMIP5), used by the Fifth Assessment Report of the Intergovernmental Panel on Climate Change (2013), have process-level representation of microbial physiology (Todd-Brown et al., 2013). Where they have been included, the models suggest that process-level representation of microbial physiology influences soil C storage at the global scale (Hararuk et al., 2014, Tang & Riley, 2015, Wieder et al., 2013).

Linear rate constants do have the advantage of being computationally and mathematically tractable, and do not result in the oscillatory behavior that is commonly reported in models that incorporate non-linear processes such as Arrhenius and
Michaelis-Menten relationships (Hararuk et al., 2014, Wieder et al., 2014). Given the importance of microbial physiology to global C cycling, I believe that process-level representation is necessary, but the resulting model should be as parsimonious as possible and modular enough to easily exclude features that are not necessary to a particular user or for which data is not available at a particular site.

In order to build process-level representation of the effect of temperature and substrate supply on microbial physiology, I merged a model that uses temperature and substrate supply to predict depolymerization rate (Davidson et al., 2012) with a C and N microbial physiology model (Finzi et al., 2015). Merging two models involves a number of challenges including combining model equations, parameterization, and model assessment. To simplify the presentation of these processes I describe two overarching objectives here. My first objective is to test the performance of the combined model relative to each model alone using measurements of heterotrophic respiration in a mid-latitude forest located in central Massachusetts, USA. My second objective is to apply the combined model to theoretical concepts such as the response of microbial activity to varying ratios of C and N substrate additions, and global-change-inspired perturbations to mean annual temperature and soil moisture. To link these modeling activities to my prior research on belowground C allocation and rhizosphere processes, I hypothesize that forest stands characterized by a high C-to-N ratio [litter, root inputs, microbial biomass] will have lower C efflux rates, and store more soil C than narrow C:N stands (Averill et al., 2014). My second hypothesis is that warming will increase C efflux, but that added
soil moisture will not increase microbial activity as a result of the cancellation of a positive effect on C supply and a negative effect on O₂ supply.

**Methods**

*Model description*

The DAMM-MCNiP model is a parsimonious belowground C and N model with core processes-level representation of microbial and exoenzymatic activity. This model was developed from the merger of the Dual-Arrhenius Michaelis Menten (DAMM) model of Davidson *et al.* (2012) with the Microbial Carbon and Nitrogen Physiology (MCNiP) model of Finzi *et al.* (2015). DAMM explicitly simulates the effects of temperature, soil moisture and substrate supply on the kinetics of soil organic matter (SOM) depolymerization. Outputs of the MCNiP model include microbial maintenance respiration, biomass production, exoenzyme production, and C and N uptake and mineralization.

Soil temperature and moisture inputs constrain the rate of unprotected SOM depolymerization and uptake of dissolved organic matter, including carbon and nitrogen (DOM) to the microbial biomass pool. Microbes depolymerize SOM and take up DOM according to Arrhenius and Michaelis-Menten kinetics. Further, SOM availability is limited by soil moisture according to DAMM principles of substrate diffusion, and DOM uptake is limited by soil moisture via O₂ availability. Microbes allocate C to maintenance respiration, enzyme production, growth, and overflow C and N mineralization. Enzyme production and growth are dependent on the C:N ratio of the DOM taken up, as well as the carbon use efficiency of microbial metabolism. Microbial biomass and enzymes turn
over at a constant rate, and this carbon is partitioned to SOM and DOM pools (Figure 5.1).

In DAMM-MCNIp, a series of differential equations determines the change in each pool size at each time step. The change in the soluble C and N pool is modeled as:

\[
\begin{align*}
\frac{dDOC}{dt} &= \text{inputDOC} + \text{DEPOLY}_C + \text{DEATH} \times (1 - \text{MIC}_{C \text{ to } SOM}_C) + (\text{CNe}/(1+\text{CNe})) + \text{ELOSS}_C - \text{UPT}_C \\
\frac{dDON}{dt} &= \text{inputDON} + \text{DEPOLY}_N + \text{DEATH} \times (1 - \text{MIC}_{N \text{ to } SOM}_N) + (1/\text{CNe}) + \text{ELOSS}_N - \text{UPT}_N
\end{align*}
\]

where \( \frac{dDOM}{dt} \) is the change in the DOM pool size, \( \text{inputDOM} \) is the root input [exudation, turnover], \( \text{DEPOLY} \) is the depolymerization rate of SOM, \( \text{DEATH} \) is microbial turnover, \( \text{MIC}_{C \text{ to } SOM} \) is the fraction of dead microbial biomass C or N returned to the SOM pool, \( \text{CNe} \) is the C:N ratio of exoenzymes, \( \text{ELOSS} \) is the enzyme turnover rate, and \( \text{UPT} \) is the rate of DOM uptake by the microbial biomass pool.

Depolymerization rate is modeled as:

\[
\begin{align*}
\text{DEPOLY}_C &= V_{maxC} \times a \times \text{Enz} \times \text{avail}_\text{SOC}/(K_{mC} + \text{avail}_\text{SOC}) \\
\text{DEPOLY}_N &= V_{maxN} \times (1-a) \times \text{Enz} \times \text{avail}_\text{SON}/(K_{mN} + \text{avail}_\text{SON})
\end{align*}
\]

where \( V_{max} \) is the maximum reaction rate when the enzyme is saturated with substrate, \( a \) is the proportion of the enzyme pool acting on the SOC pool (i.e., \( 1-a \) is the proportion acting on the SON pool), \( \text{Enz} \) is the enzyme pool size, \( \text{avail}_\text{SOC} \) is unprotected SOM, and \( K_{m} \) is the half-saturation constant for depolymerization.

Unprotected SOM and \( V_{max} \) are modeled as:

\[
\begin{align*}
\text{avail}_\text{SOC} &= \text{SOM} \times \text{frac} \times \text{Dliq} \times \text{soilM}^3 \\
V_{maxC,N} &= A_{C,N} \times \exp(-Ea_{C,N} / RT)
\end{align*}
\]
where \( \text{frac} \) is the fraction of unprotected SOM, using soluble substrate estimated from Magill et al. (2000), \( D_{\text{liq}} \) is the diffusion coefficient for unprotected SOM in liquid, \( \text{soilM} \) is the volumetric water content, \( A \) is the pre-exponential constant for SOM depolymerization, \( E_a \) is the activation energy for SOM depolymerization, \( R \) is the universal gas constant, and \( T \) is temperature.

Microbial biomass is modeled as:

\[
d\text{MIC}_C/ dt = C_{Nm} \times \text{GROWTH} - \text{DEATH}_C \tag{7}
\]

\[
d\text{MIC}_N/ dt = \text{GROWTH} - \text{DEATH}_N \tag{8}
\]

where \( \text{MIC} \) is the microbial biomass, \( C_{Nm} \) is the C:N ratio of microbial biomass, and \( \text{GROWTH} \) is microbial biomass growth, and:

\[
\text{GROWTH} = \frac{(1-p) \times \text{UPT}_C \times \text{CUE} + \text{Enz}_C - \text{CNe} \times \text{EPROD}}{C_{Nm}} \quad \text{[C limited]} \tag{9}
\]

\[
\text{GROWTH} = (1-q) \times \text{UPT}_N \times \text{Enz}_N - \text{EPROD} \quad \text{[N limited]} \tag{10}
\]

\[
\text{DEATH} = \text{r}_{\text{death}} \times \text{MIC}_{C,N} \tag{11}
\]

where \( p \) and \( q \) are the proportion of assimilated C or N, respectively, allocated to enzyme production, \( \text{CUE} \) is carbon use efficiency, \( \text{EPROD} \) is enzyme production, and \( \text{r}_{\text{death}} \) is the microbial turnover rate.

Microbial uptake is defined as:

\[
\text{UPT}_{C,N} = \text{MIC}_{C,N} \times \frac{\text{Vmax}_{\text{uptC,N}} \times \text{DOC}_N}{\text{Km}_{\text{uptC,N}} + \text{DOC}_N} \times \frac{\text{O}_2}{\text{Km}_{\text{O}_2} + \text{O}_2} \tag{12}
\]

where \( \text{Vmax}_{\text{upt}} \) and \( \text{Km}_{\text{upt}} \) are the maximum reaction rate and half-saturation constants for uptake, respectively. \( \text{O}_2 \) is the oxygen concentration and \( \text{Km}_{\text{O}_2} \) is the half-saturation constant for \( \text{O}_2 \) as a substrate. \( \text{Vmax}_{\text{upt}} \) and \( \text{O}_2 \) are determined by:

\[
\text{Vmax}_{\text{uptC,N}} = \text{A}_{\text{uptC,N}} \times \exp \left(-\frac{E_{\text{uptC,N}}}{RT}\right) \tag{13}
\]
\[ O_2 = D_{gas} \cdot O_{2airfrac} \cdot [(\text{porosity} - \text{soilM})^{4/3}] \]  
\[ \text{porosity} = 1 - \frac{BD}{PD} \]

where \( A_{upt} \) is the pre-exponential constant for DOC uptake, \( E_{aup} \) is the activation energy for DOC uptake, \( D_{gas} \) is the diffusion coefficient for \( O_2 \) in air, \( O_{2airfrac} \) is the volume fraction of \( O_2 \) air, \( BD \) is bulk density and \( PD \) is particle density of the soil.

Enzyme production can be C or N limited depending on the stoichiometry of the DOM taken up:

\[ E_{PROD} = p \cdot (\text{CUE} \cdot \text{UPT}_C) / \text{CNe} \quad [\text{C limited}] \]
\[ E_{PROD} = q \cdot \text{UPT}_N \quad [\text{N limited}] \]

C mineralization, overflow C, and N mineralization are modeled as:

\[ \text{CMIN} = \text{UPT}_C \cdot (1 - \text{CUE}) \]  
\[ \text{NMIN} = \text{GROWTH}_N - \text{GROWTH} \]  
\[ \text{OverflowC} = \text{GROWTH}_C - \text{CNm} \cdot \text{GROWTH} \]

The enzyme pool is a balance between production and turnover, defined as a first order process:

\[ \frac{d\text{Enz}}{dt} = E_{PROD} - E_{LOSS} \]
\[ E_{LOSS} = r_{enzloss} \cdot \text{Enz} \]

where \( r_{enzloss} \) is the enzyme turnover rate.

Lastly, the change in the SOM pool is modeled as:

\[ \frac{d\text{SOM}_{C,N}}{dt} = \text{Litter}_{C,N} + \text{DEATH} \cdot \text{MIC}_{C,N} \cdot \text{SOM}_{C,N} - \text{DEPOLY}_{C,N} \]

where \( \text{Litter}_{C,N} \) is the litter input to the SOC pool.
Inputs to the model include litter and root exudate C and N, temperature and soil moisture. Outputs used in this study are the rate of C mineralization (i.e., C efflux) and N mineralization. Litter (leaf, root) is partitioned to SOM and DOM pools at each timestep (h⁻¹). Root exudates enter the DOM pool only.

Model parameters were identical to original DAMM and MCNiP parameters, excepting the following: \(A_{\text{uptC}_N}, \ E_{\text{uptC}_N}, \ A_{\text{C}_N}, \ E_{\text{C}_N}, \ K_m_{\text{C}_N}\). I estimated \(A_{\text{uptC}_N}, \ E_{\text{uptC}_N}, \ A_{\text{C}_N}, \ E_{\text{C}_N}\) from independent measurements of \(\beta\)-glucosidase activity from organic and mineral soil (Davidson et al., 2012, Finzi et al., 2015). Because I cannot experimentally distinguish between depolymerization kinetics and uptake kinetics, each pair of \(A\) and \(E_a\) values are identical. \(K_m_{\text{C}_N}\) was estimated such that at standard temperature, 293 K (20ºC), and the mean soil moisture value for this site, 0.229 cm³ H₂O cm⁻³ soil, \(K_m_{\text{C}_N}\) was equal to the initial available substrate SOM concentration (sensu Davidson et al., 2012). In contrast to Allison et al. (2010) and MCNiP, I did not choose a \(K_m\) larger than the available SOM pool. Microbes in MCNiP have access to the entire SOM pool, so without a \(K_m\) value that is high relative to the SOM pool size, substrate will always be saturating. My parameterization allows for substrate to saturate a reaction site, for example, if substrate is temporarily mobilized during a wet-up event (Birch, 1958, Davidson et al., 2014). Model parameters are found in Table 5.1.

Model parameters in MCNiP are based on Allison et al. (2010). One exception is the C:N of soil, taken from Schimel and Weintraub (2003). Default initial pool sizes were determined after model spin up for 2000 years using spin up parameters from Allison et al. (2010) for the C pool. The N pool was parameterized using the following principles,
SON = SOC/27.6 [C:N ratio of soil], DON = DOC/15 [mid-range of DOC:DON from Hopkinson et al. (1997) and Neff & Hooper (2002)], microbial biomass N = microbial biomass C/10 [C:N ratio of microbes]. For the default model, I assume that litter and root inputs have C:N of 27.6 (Table 5.2).

**Model comparison**

I compared model performance of the DAMM-MCNiP model against three other models, DAMM alone, MCNiP alone, and DAMM-MCP (without any mechanistic linkage between C and N cycling). DAMM parameters are from Davidson et al. (2012). MCNiP’s enzyme kinetic parameters, $A_{uptC_N}$, $Ea_{uptC_N}$, $A_{C_N}$, $Ea_{C_N}$, $Km_{C_N}$ and initial values were parameterized as in Finzi et al. (2015). DAMM-MCNiP and DAMM-MCP have identical parameter values (Table 5.1). In DAMM-MCP, N limitation is not possible because I removed the conditional statements that determine whether or not the system is N-limited. As a result, C dynamics control the model and N pools follow C pools according to stoichiometry. I ran DAMM and MCNiP for one year. I ran DAMM-MCNiP and DAMM-MCP for 200 years to achieve stable efflux values.

To test model performance, I used measurements of soil temperature, moisture, and C efflux from a trenching experiment at Harvard Forest, MA. A 5 x 5 m trench was dug to 1 m depth in November 2008 in a mixed hardwood stand on the Prospect Hill tract of Harvard Forest. Automated measurements of C efflux were collected from April through October 2009. I used C efflux from trenched plots as an estimate of heterotrophic respiration. This dataset is described in greater detail in Davidson et al. (2012).
Linear regression was used to fit the output predicted by each model with C efflux measurements from trenching, and to fit model-predicted N mineralization with soil moisture measurements. I computed the correlation coefficient (Pearson’s ρ) between model-predicted C efflux and temperature or soil moisture. All analyses were conducted in R Statistical Software (R Development Core Team, 2013).

**Root input measurements**

Root C inputs were measured in three stands at Harvard Forest, white ash (Fraxinus americana), red oak (Quercus rubra), and eastern hemlock (Tsuga canadensis). Root exudates were collected from six root systems per stand in June and August of 2012, and April, May, July and October of 2013 following the method of Phillips et al. (2008, 2011). Root turnover was measured April–December 2012, March–November 2013, and April–November 2014 in ten minirhizotron tubes per stand using a BTC-100x high magnification minirhizotron camera system (Bartz Technology Company, Carpenteria, CA). I averaged across the three stands and across replicate samples to determine the base rate of root input, 120 g C m⁻² yr⁻¹, which was comprised of an exudation (102 g C m⁻² yr⁻¹) and root turnover (18 g C m⁻² yr⁻¹) flux component. I assume that exudation occurs primarily during the growing season and follows a unimodal Gaussian distribution with a standard deviation of 42 days. I used this root input rate and phenology for the following simulations that manipulate C:N, temperature, and soil moisture.
Model simulation of microbial physiology response to C:N ratio

I conducted model simulations of DAMM-MCNiP that test the importance of stand-level differences in C:N on microbial physiology. C and N mineralization was simulated in two theoretical forests differing in their litter, root input, and microbial biomass C:N, based on ranges reported in literature for an arbuscular-mycorrhizal hardwood stand where decomposition is dominated by bacteria (herein Narrow C:N), and an ectomycorrhizal conifer stand where decomposition is dominated by fungi (herein Wide C:N; Table 5.3). In order to compare the effects of varying input and microbial C:N, I conducted three tests. First, I ran the model using either high or low values of litter and root input C:N. Second, I ran the model using high or low values of microbial biomass C:N. Finally, I varied both input and biomass C:N (Table 5.3).

To determine the effect of root input C:N on depolymerization and microbial efficiency, I varied root input C:N between 1 and 100 and recorded the SOC pool size at different quantities of root input and root input C:N. I used the ratio of depolymerization rate to microbial biomass as a measure of microbial efficiency.

Model simulations of global change

In order to estimate the effect of global change on annual C and N efflux, I perturbed the temperature and soil moisture inputs for the last year of each model run of DAMM-MCNiP. First, I increased and decreased the temperature at each time point by 5°C. Second, using ambient temperature forcing, I increased and decreased soil moisture at each time point by 50%. I calculated the absolute difference between each temperature
and soil moisture treatment relative to ambient, and cumulatively summed the C or N efflux over the year.

**Results**

*Model comparison*

From a visual inspection of the model output, DAMM captures the seasonality of measured C efflux well, though it slightly over-predicts C efflux during mid-summer (DOY 220-250), and under-predicts efflux during large precipitation events (e.g., DOY 170, 240; Figure 5.2a). The RMSE of the correlation between DAMM and measured C efflux is 31.7 ($\beta = 0.85$, $F_{1,2903} = 3102$, $P < 0.001$, $R^2_{adj} = 0.52$; Figure 5.3a). MCNiP has an RMSE of 7.6 with little change in amplitude between summer and winter efflux (Figure 5.2b). The slope ($\beta$) of the relationship between MCNiP and measured efflux is lower than 1 ($\beta = 0.20$, $F_{1,2965} = 2998$, $P < 0.001$, $R^2_{adj} = 0.50$; Figure 5.3b). The RMSE of DAMM-MCNiP is 26.7, and is lower than that of DAMM. The slope of the relationship between predicted and measured efflux for DAMM-MCNiP is lower than 1, but higher than MCNiP ($\beta = 0.59$, $F_{1,2965} = 2156$, $P < 0.001$, $R^2_{adj} = 0.42$; Figure 5.3c). DAMM-MCP lacks any linkage to the N cycle, and has an RMSE of 25.9 ($\beta = 0.57$, $F_{1,2965} = 2096$, $P < 0.001$, $R^2_{adj} = 0.41$; Figure 5.3d).

DAMM-MCNiP model residuals indicate that DAMM-MCNiP slightly over-predicts C efflux across the range of soil moisture values and under-predicts some C efflux values between 0.3 and 0.5 cm$^3$ H$2$O cm$^{-3}$ soil (Figure 5.4a). DAMM alone over-predicts C efflux at low and high soil moisture values and under-predicts C efflux between 0.3 and 0.5 cm$^3$ H$2$O cm$^{-3}$ soil (Figure 5.4b). At soil moisture values > 0.68 cm$^3$
H₂O cm⁻³ soil, DAMM cannot solve for a C efflux value, because soil pores are completely filled with water. In DAMM-MCNiP, however, I implemented a drainage subroutine that allows water to exit soil pores, alleviating the mathematical impossibility of having more water in the soil than available pore space.

DAMM is more strongly correlated with temperature (ρ = 0.90) than soil moisture (ρ = 0.15). DAMM-MCNiP is correlated with temperature (ρ = 0.69) and soil moisture (ρ = 0.33). In DAMM-MCNiP, the seasonal pattern of temperature and soil moisture is similar to the pattern of depolymerization rate, C uptake rate, C mineralization rate and the size of DOC pool (Figure 5.5a,b,e,f,g,j). The phenology of the SOC pool is opposite that of the microbial biomass and enzyme C pools, such that SOC is most depleted (i.e., DOY 250) when the microbial biomass pool reaches its annual peak (Figure 5.5i,k,l). N mineralization is not confined to the growing season and has a significant negative relationship with soil moisture (F₁,4559 = 925, P < 0.001, R²_adj = 0.17; Figure 5.5b,d).

Overflow C is the release of excess C from microbes after allocation to maintenance respiration, enzyme production, and biomass growth. Overflow C is an intermittent flux that only occurs when all available N is immobilized (i.e., when N mineralization is low or absent; Figure 5.5h; Figure 5.6).

Model simulation of microbial physiology response to C:N

The theoretical Narrow C:N stand had higher C and N mineralization rates compared to the Wide C:N stand, by 12 g C m⁻² yr⁻¹ and 0.9 g N m⁻² yr⁻¹, respectively. When input C:N was changed from default settings but microbial biomass C:N was held constant, the Narrow C:N stand mineralized 27 g C m⁻² yr⁻¹ and 1.4 g N m⁻² yr⁻¹ more than
the Wide C:N stand (Figure 5.7a,d). In contrast, the Wide C:N stand had higher C and N mineralization by 12 g C m$^{-2}$ yr$^{-1}$ and 0.9 g N m$^{-2}$ yr$^{-1}$, respectively, when only microbial biomass C:N was changed from default settings (Figure 5.7b,e). As a result, the difference in C and N mineralization between the Wide and Narrow C:N stands was small when both input and microbial biomass C:N were varied (Figure 5.7c,f; Table 5.3).

SOC depolymerization increased with the magnitude of root input to the DOC pool, depleting the SOC pool. This effect appeared to exhibit asymptotic behavior at high values of exudate inputs. Low root input C:N resulted in more depolymerization than high input C:N (Figure 5.8a). At low C:N, there was an increase in microbial efficiency with root inputs (Figure 5.8b, red line). At high C:N, SOC was depleted as a result of an increase in microbial biomass (Figure 5.8c).

**Model simulations of global change**

A 5°C warming of seasonal temperature inputs increased C efflux by 119 g C m$^{-2}$ yr$^{-1}$. An equivalent decrease in ambient temperature caused C efflux to fall by 71 g C m$^{-2}$ yr$^{-1}$ (Table 5.4; Figure 5.9a). N mineralization increased by 0.37 g N m$^{-2}$ yr$^{-1}$ with warming and decreased 0.32 g N m$^{-2}$ yr$^{-1}$ with cooling (Figure 5.9b).

A 50% increase in seasonal soil moisture caused a large increase in C (220 g C m$^{-2}$ yr$^{-1}$) efflux and a smaller increase in N mineralization (0.9 g C m$^{-2}$ yr$^{-1}$), while a 50% decrease in soil moisture decreased C efflux by 156 g C m$^{-2}$ yr$^{-1}$ and N mineralization by 0.21 g N m$^{-2}$ yr$^{-1}$ (Figure 5.9c,d). There were also large changes in the seasonal pattern of N mineralization, with more variation in the wet treatment compared to the dry treatment (Figure 5.9d). The absolute difference in C efflux in the warming and added soil moisture
treatments were greater than the reduced temperature and soil moisture treatments, but the effect of each treatment was similar for N mineralization (Figure 5.10). Cumulative sums of mineralization measured over the growing season demonstrate that temperature and soil moisture treatments had a larger effect on annual C efflux than on N mineralization (Figure 5.11). In addition, N mineralization in the dry treatment exceeded both ambient and wet treatments for large portions of the year (Figure 5.10d, 5.11d).

**Discussion**

As currently parameterized, DAMM-MCNiP predicts realistic C efflux during transient wet-up events (Figure 5.4a), an area where many present-day models struggle to represent the increase in C efflux with soil moisture (Ise & Moorcroft, 2006, Rodrigo et al., 1997). However, DAMM-MCNiP over-predicts C efflux in constant high soil moisture conditions. DAMM-MCNiP produces realistic responses to temperature perturbation, and thus may be useful for investigating the interaction between substrate and temperature controls on microbial activity. Consistent with hypothesis 1, model predictions of SOM decomposition in response to variable C:N inputs show that N-rich root exudates increase the decomposition rate of SOM and that wide C:N stands store more C than narrow C:N stands (Averill et al., 2014).

**Model comparison**

DAMM alone captures the seasonality of C efflux. The slope of the C efflux predicted by DAMM plotted against measured C efflux is the closest to 1 of the four models. However, DAMM parameters were fit to these data using parameter optimization, and therefore is it not surprising that model bias is low. The RMSE of
DAMM, however, is highest of the four models. In contrast, MCNiP alone has the lowest RMSE of the four models, but does not capture the observed seasonal pattern of C efflux, with the slope between observed and predicted efflux equal to 0.2. This indicates that while MCNiP has high precision, alone it has low predictive accuracy. In contrast to DAMM, MCNiP is currently parameterized from literature values as a theoretical model, and has not been previously fit to data (Allison et al., 2010, Schimel & Weintraub, 2003). That it performs as well as it does is notable.

Together, DAMM and MCNiP capture the seasonal cycle of soil C efflux. Though the RMSE of DAMM-MCNiP is much higher than that of MCNiP, DAMM-MCNiP has the ability to predict realistic seasonal patterns in response to fluctuations in temperature and soil moisture. That is, the slope of the predicted compared to observed efflux is closer to 1 for DAMM-MCNiP compared to MCNiP.

The correlation coefficients between C efflux predicted by DAMM and temperature ($\rho = 0.90$) or soil moisture ($\rho = 0.15$) indicate that DAMM alone is more sensitive to temperature than to soil moisture. DAMM-MCNiP is relatively more sensitive to soil moisture ($\rho = 0.33$) than is DAMM. As a result, DAMM-MCNiP successfully predicts the large increase in C efflux after wet-up events, which DAMM alone cannot recreate.

The major benefit of DAMM-MCNiP in comparison to DAMM and MCNiP alone is the process-level representation of microbial physiology coupled with the ability to represent substrate availability at the micro-site scale. Another benefit is the coupling of the C and N cycle, which allows for investigation into nutrient limitation and priming
effects. Although, DAMM-MCP is decoupled from the N cycle, model performance is similar to DAMM-MCNiP [slightly lower RMSE and slope], though the lack of a N cycle limits its utility. The fact that it does perform as well as DAMM-MCNiP suggests the need to assess both models against other datasets. The stoichiometric coupling in DAMM-MCNiP may also benefit from data-model assimilation using Harvard Forest datasets, modifications to the model’s core structure, or both.

From a visual inspection of the model inputs, pools and fluxes, it is clear that temperature and soil moisture together determine the rate of depolymerization, which then constrains the DOC pool, C uptake, and C efflux (Figure 5.5e,f,g). N mineralization occurs throughout the growing season, and is negatively correlated with soil moisture. Overflow C occurs when the system is N-limited, and as a result overflow C and N mineralization are nearly mutually exclusive (Figure 5.6). This is an important model diagnostic which demonstrates that the coupling of C and N in the model is working properly.

*Model simulation of microbial physiology response to C:N*

When I narrowed the C:N ratio of root inputs and microbial biomass, I observed a small increase in the amounts of C and N mineralized, supporting my first hypothesis that stands with narrow C:N will have higher C efflux rates (Figure 5.7). This response to low C:N is consistent with my understanding of priming based on MCNiP (Finzi et al., 2015) and related models (Drake et al., 2013), where adding N increased C efflux and SOM decomposition (Figure 5.8a).
The mechanism by which more SOM is decomposed with added exudates at root C:N > 3 is an increase in microbial biomass rather than an increase in microbial efficiency (Figure 5.8b,c). This understanding of priming is not necessarily counter to findings that suggest that low N inputs stimulate foraging for N by inducing N limitation (Craine et al., 2007). Adding more N to the system supports a larger microbial biomass, but these microbes can still be N limited, since the C:N of litter and root inputs is still higher than the C:N of microbial biomass (Table 5.3).

When only microbial biomass C:N was varied, the wide C:N stand predicted a slightly higher C efflux than the narrow C:N stand, because microbial biomass with high C:N is more closely matched to the C:N of litter and root inputs. Varying input and microbial biomass C:N together does not change the overall C efflux by a large amount, consistent with idea that microbial activity is limited by the stoichiometry of its substrate (Sinsabaugh et al., 2009).

*Model simulations of global change*

Warming seasonal temperatures by 5°C increased C efflux by about 55%, supporting the first part of the second hypothesis that warming increases heterotrophic respiration. C efflux increased ~50% from cold to ambient treatments, suggesting that the temperature response to 5°C warming is consistent regardless of the starting temperature. A 50 – 55% increase in heterotrophic respiration with warming was greater than the observed increase in total soil CO₂ efflux in response to 5°C warming in two studies at the Harvard Forest by 11 and 27% (Contosta, 2011, Melillo et al., 2002). Soil warming in the field usually dries the soil along with heating, which may account for the smaller
increase in C efflux observed in the field. In fact, when I simultaneously warmed the soil by 5°C and decreased soil moisture by 10%, which corresponds to the amount of drying observed by Contosta et al. (2011) in the soil organic horizon, I predicted an increase in C efflux increase of only 17% over ambient conditions (Figure 5.12).

Warming and cooling treatments had similar effects on N mineralization (Figure 5.10b), with a slightly positive effect of warming, but many instances over the growing season where N mineralization decreased with additional warming. At the seasonal time-scale, N mineralization was not very sensitive to temperature with perhaps marginally more N mineralization during the growing season, so it is not surprising that at the annual time-scale, an increase in temperature also had a small effect on N mineralization.

In contrast to the second part of hypothesis 2, soil moisture manipulations had a larger effect on C efflux than did either warming or cooling. I hypothesized that a 50% increase in soil moisture would have a relatively small positive effect, because the increase in diffusive transport of substrates would be moderated by O₂ limitation. However, I found that O₂ limitation had a small effect relative to diffusion because C efflux was strongly stimulated in the 1.5x soil moisture treatment and strongly limited in the 0.5x soil moisture treatment. Suseela et al. (2012) found that heterotrophic respiration was reduced in a mesic suburban old-field site when volumetric soil moisture was outside of a 15-26% window. Davidson et al. (1998) found high soil respiration at similar volumetric soil moisture values. The mean volumetric soil moisture for the dry and wet treatments in the simulation was outside of this window (15% and 41%, respectively), suggesting that DAMM-MC NiP may overestimate C efflux at high soil moisture values.
Soil moisture manipulations had a large impact on N mineralization. At low soil moisture, N mineralization had a lower coefficient of variation (CV = 0.41) over the growing season than it had at ambient (CV = 0.96) or added soil moisture (CV = 1.1; Figure 5.9d). During dry conditions there is little substrate available for depolymerization and microbial uptake. Because substrate supply is low throughout the growing season, this limits the rate N mineralization. Conversely, when substrate is plentiful during wet conditions, N mineralization is widely variable depending upon whether microbial growth is C or N limited. Since the wet treatment resulted in a large pool of available SOC, I expected that N mineralization would decrease because microbial growth becomes N limited. This was not the case, however, because there were many time periods throughout the simulation when microbial growth was C limited (Figure 5.10d). This suggests that the model may be operating under greater C limitation than expected.

Summary

The DAMM-MCNiP model is a parsimonious model that simulates decomposition and C and N mineralization using process-level representation of microbial physiology and stoichiometry. It is one of the first microbial physiology models that can represent the C and N cycle together with plant substrate supply, making DAMM-MCNiP a candidate for linkage with terrestrial biosphere models that allocate a fraction of fixed C belowground at each time-step. The model inputs are similar to CENTURY (i.e., soil temperature, moisture, C and N), but rather than combining temperature and soil moisture effects into a linear decomposition factor, DAMM-MCNiP
explicitly models the processes by which temperature and moisture affect substrate availability and enzyme kinetics (Bolker et al., 1998).

DAMM-MCNiP requires parameter estimation and sensitivity analyses beyond those performed here (i.e., by varying C:N, soil temperature and soil moisture) in order to reproduce measured C efflux without bias. Future work will focus on parameterizing DAMM-MCNiP using Bayesian data assimilation, first at the Harvard Forest in Petersham, MA and then at other sites. I will nest parameter value estimates and their associated uncertainty at each site using a hierarchical framework that can then estimate broader regional-scale parameters. A broadly parameterized microbial physiology model such as DAMM-MCNiP may improve terrestrial biosphere model predictions of SOC pools under global change.
Table 5.1. DAMM-MCNiP default model parameter values.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>Default Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>rootDOC</td>
<td>mg cm$^{-3}$</td>
<td>input</td>
<td>Root exudates</td>
</tr>
<tr>
<td>T</td>
<td>K</td>
<td>input</td>
<td>temperature in Kelvin</td>
</tr>
<tr>
<td>$\theta$</td>
<td>cm$^3$ H$_2$O cm$^{-3}$</td>
<td>soil</td>
<td>input volumetric water content</td>
</tr>
<tr>
<td>BD</td>
<td>g cm$^{-3}$</td>
<td>0.8</td>
<td>bulk density</td>
</tr>
<tr>
<td>PD</td>
<td>g cm$^{-3}$</td>
<td>2.52</td>
<td>particle density</td>
</tr>
<tr>
<td>O$_2$airfrac</td>
<td>L O$_2$/L air</td>
<td>0.209</td>
<td>volume fraction of O$_2$ air</td>
</tr>
<tr>
<td>frac</td>
<td>g C cm$^{-3}$/g C cm$^{-3}$</td>
<td>0.000414</td>
<td>fraction of unprotected SOM, using soluble substrate estimated from Magill et al., 2000</td>
</tr>
<tr>
<td>$D_{liq}$</td>
<td>-</td>
<td>3.17</td>
<td>diffusion coefficient for unprotected SOM and DOM in liquid</td>
</tr>
<tr>
<td>$D_{gas}$</td>
<td>-</td>
<td>1.67</td>
<td>diffusion coefficient for O$_2$ in air</td>
</tr>
<tr>
<td>$K_{mO2}$</td>
<td>cm$^3$ O$_2$ cm$^{-3}$</td>
<td>air</td>
<td>Michaelis constant for O$_2$</td>
</tr>
<tr>
<td>R</td>
<td>kJ K$^{-1}$ mol$^{-1}$</td>
<td>0.0083145</td>
<td>universal gas constant</td>
</tr>
<tr>
<td>endTime</td>
<td>h</td>
<td>2000000</td>
<td>number of hours simulated in model run</td>
</tr>
<tr>
<td>p</td>
<td>-</td>
<td>0.5</td>
<td>proportion of assimilated C allocated to enzyme production</td>
</tr>
<tr>
<td>q</td>
<td>-</td>
<td>0.5</td>
<td>proportion of assimilated N allocated to enzyme production</td>
</tr>
<tr>
<td>a</td>
<td>-</td>
<td>0.5</td>
<td>proportion of enzyme pool acting on SOC pool (1-a = proportion acting on SON pool)</td>
</tr>
<tr>
<td>initSOC</td>
<td>mg cm$^{-3}$</td>
<td>144.5986</td>
<td>initial SOC pool</td>
</tr>
<tr>
<td>initSON</td>
<td>mg cm$^{-3}$</td>
<td>5.4413</td>
<td>initial SON pool</td>
</tr>
<tr>
<td>initDOC</td>
<td>mg cm$^{-3}$</td>
<td>0.00091631</td>
<td>initial DOC pool</td>
</tr>
<tr>
<td>initDON</td>
<td>mg cm$^{-3}$</td>
<td>0.00049421</td>
<td>initial DON pool</td>
</tr>
<tr>
<td>initBiomassC</td>
<td>mg cm$^{-3}$</td>
<td>1.1957</td>
<td>initial microbial biomass C</td>
</tr>
<tr>
<td>initBiomassN</td>
<td>mg cm$^{-3}$</td>
<td>0.1196</td>
<td>initial microbial biomass N</td>
</tr>
<tr>
<td>Litter$_{C,N}$</td>
<td>mg cm$^{-1}$ hr$^{-1}$</td>
<td></td>
<td>litter input to SOC pool</td>
</tr>
<tr>
<td>LitterCN</td>
<td>mg cm$^{-3}$ hr$^{-1}$</td>
<td>27.6</td>
<td>C:N of litter input to SOC pool</td>
</tr>
<tr>
<td>initEnz</td>
<td>mg cm$^{-3}$</td>
<td>0.0381</td>
<td>initial enzyme pool</td>
</tr>
<tr>
<td>inputDOC</td>
<td>mg cm$^{-1}$ hr$^{-1}$</td>
<td>0.0005</td>
<td>root input to DOC pool</td>
</tr>
<tr>
<td>inputDOCN</td>
<td>mg cm$^{-3}$ hr$^{-1}$</td>
<td>27.6</td>
<td>C:N of litter input to DOC pool</td>
</tr>
<tr>
<td>r_death</td>
<td>hr$^{-1}$</td>
<td>0.00015</td>
<td>microbial turnover rate</td>
</tr>
<tr>
<td>r_EnzLoss</td>
<td>hr$^{-1}$</td>
<td>0.001</td>
<td>enzyme turnover rate</td>
</tr>
<tr>
<td>MIC$<em>{C,N}$toSOC$</em>{C,N}$</td>
<td>mg mg$^{-1}$</td>
<td>0.5</td>
<td>fraction of dead microbial biomass</td>
</tr>
<tr>
<td>Parameter</td>
<td>Units</td>
<td>Default Value</td>
<td>Description</td>
</tr>
<tr>
<td>-------------</td>
<td>--------------------------------------------</td>
<td>---------------</td>
<td>--------------------------------------------</td>
</tr>
<tr>
<td>$A_{C,N}$</td>
<td>mg SOM cm$^{-3}$ (mg Enz cm$^{-3})^{-1}$ h$^{-1}$</td>
<td>$1.0815 \times 10^{11}$</td>
<td>Pre-exponential constant for SOM depolymerization</td>
</tr>
<tr>
<td>$A_{uptC,N}$</td>
<td>mg DOC cm$^{-3}$ (mg biomass cm$^{-3})^{-1}$ h$^{-1}$</td>
<td>$1.0815 \times 10^{11}$</td>
<td>Pre-exponential constant for DOC uptake</td>
</tr>
<tr>
<td>$K_{mC,N}$</td>
<td>mg cm$^{-3}$</td>
<td>0.0025</td>
<td>Half-saturation constant for SOM depolymerization</td>
</tr>
<tr>
<td>$K_{muptC,N}$</td>
<td>mg cm$^{-3}$</td>
<td>0.3</td>
<td>Half-saturation constant for DOC uptake</td>
</tr>
<tr>
<td>CUE</td>
<td>mg mg$^{-1}$</td>
<td>0.31</td>
<td>Carbon use efficiency</td>
</tr>
<tr>
<td>$E_{aC,N}$</td>
<td>kJ mol$^{-1}$</td>
<td>61.77</td>
<td>Activation energy for SOM depolymerization</td>
</tr>
<tr>
<td>$E_{auptC,N}$</td>
<td>kJ mol$^{-1}$</td>
<td>61.77</td>
<td>Activation energy for DOC uptake</td>
</tr>
<tr>
<td>CN$_s$</td>
<td>-</td>
<td>27.6</td>
<td>C:N of soil</td>
</tr>
<tr>
<td>CN$_l$</td>
<td>-</td>
<td>27.6</td>
<td>C:N of litter</td>
</tr>
<tr>
<td>CN$_m$</td>
<td>-</td>
<td>10</td>
<td>C:N of microbial biomass</td>
</tr>
<tr>
<td>CN$_e$</td>
<td>-</td>
<td>3</td>
<td>C:N of enzymes</td>
</tr>
<tr>
<td>CN$_ex$</td>
<td>-</td>
<td>27.6</td>
<td>C:N of root inputs</td>
</tr>
</tbody>
</table>
Table 5.2. Initial pool sizes (Spin Up) and the resulting default initial pool sizes (Default) after 2000 years of spin-up.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Abbreviation</th>
<th>Units</th>
<th>Spin Up</th>
<th>Default</th>
</tr>
</thead>
<tbody>
<tr>
<td>SOC pool</td>
<td>initSOC</td>
<td>mg cm⁻³</td>
<td>100</td>
<td>144.5986</td>
</tr>
<tr>
<td>SON pool</td>
<td>initSON</td>
<td>mg cm⁻³</td>
<td>3.6232</td>
<td>5.4413</td>
</tr>
<tr>
<td>DOC pool</td>
<td>initDOC</td>
<td>mg cm⁻³</td>
<td>0.5</td>
<td>0.00091631</td>
</tr>
<tr>
<td>DON pool</td>
<td>initDON</td>
<td>mg cm⁻³</td>
<td>0.0333</td>
<td>0.00049421</td>
</tr>
<tr>
<td>microbial biomass C</td>
<td>initBiomassC</td>
<td>mg cm⁻³</td>
<td>0.5</td>
<td>1.1957</td>
</tr>
<tr>
<td>microbial biomass N</td>
<td>initBiomassN</td>
<td>mg cm⁻³</td>
<td>0.05</td>
<td>0.1196</td>
</tr>
<tr>
<td>enzyme pool</td>
<td>initEnz</td>
<td>mg cm⁻³</td>
<td>0.01</td>
<td>0.0325</td>
</tr>
</tbody>
</table>
Table 5.3. Parameter values for the C:N of litter, root inputs, and microbial biomass for two theoretical stands.

<table>
<thead>
<tr>
<th></th>
<th>Narrow C:N</th>
<th>Wide C:N</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Litter input</td>
<td>50</td>
<td>100</td>
<td>Finzi et al. 2001</td>
</tr>
<tr>
<td>Root input</td>
<td>10</td>
<td>100</td>
<td>Drake et al. 2013, Finzi et al. 2015</td>
</tr>
<tr>
<td>Microbial</td>
<td>6</td>
<td>12</td>
<td>Wallenstein et al. 2006, Bengtson et al. 2012</td>
</tr>
</tbody>
</table>
Table 5.4. Annual C and N mineralization predicted by DAMM-MCNP in four simulations that manipulate either temperature or soil moisture. Here θ refers to volumetric soil moisture.

<table>
<thead>
<tr>
<th></th>
<th>Ambient</th>
<th>+5°C</th>
<th>-5°C</th>
<th>1.5x θ</th>
<th>0.5x θ</th>
</tr>
</thead>
<tbody>
<tr>
<td>C mineralization</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(g C m$^{-2}$ yr$^{-1}$)</td>
<td>217</td>
<td>337</td>
<td>146</td>
<td>437</td>
<td>61</td>
</tr>
<tr>
<td>N mineralization</td>
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<tr>
<td>(g N m$^{-2}$ yr$^{-1}$)</td>
<td>2.9</td>
<td>3.2</td>
<td>2.5</td>
<td>3.7</td>
<td>2.6</td>
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</tbody>
</table>
Figure 5.1. Conceptual figure of the merger of DAMM (blue and orange boxes) and MCNiP (box and arrow diagram). SOM depolymerization occurs using Arrhenius and Michaelis-Menten kinetics, substrate diffusion, and a temperature-dependent $V_{\text{max}}$ (blue box). I held $K_m$ constant for the model runs in this study, but a linear temperature-sensitivity relationship can be applied to it if desired. DOC uptake is controlled by DAMM kinetics as well as $O_2$ concentration at the reaction site (orange box; see Table 5.1 for DAMM equation parameter definitions). MCNiP has four pools: SOM, DOM, Microbial biomass, and Enzymes. Litter and root inputs enter the SOM and DOM pools, respectively, and outputs are $CO_2$ as a product of respiration and inorganic $N$ as a product of $N$ mineralization.
Figure 5.2. Model inter-comparison of (a) DAMM (pink), (b) MCNiP (blue), (c) DAMM-MCNiP (red), and (d) DAMM-MCP (green) overlayed on soil C efflux measurements from trenched plots in Harvard Forest, MA.
Figure 5.3. Relationship between predicted and measured C efflux for (a) DAMM, (b) MCNiP, (c) DAMM-MCNiP, and (d) DAMM-MCP. The solid line is the regression fit, and the dotted line is the 1:1 line.
Figure 5.4. (a) DAMM-MCNIp and (b) DAMM model residuals plotted as a function of soil moisture ($\theta$).
Figure 5.5. DAMM-MCniP model inputs (black), fluxes (blue) and pools (pink).
Figure 5.6. Plot of overflow C as a function of N mineralization.
Figure 5.7. Annual C and N mineralization in stands with narrow (pink) or wide (blue) C:N using parameters defined in Table 5.3. (a) C and (d) N mineralization when only input C:N was varied between stands and microbial biomass was held at the default value. (b) C and (e) N mineralization when only microbial biomass C:N was varied and inputs were held at their default value. (c) C and (f) N mineralization when both input C:N and microbial biomass C:N were varied between stands.
Figure 5.8. (a) SOC as a function of root inputs. (b) Microbial efficiency (depolymerization per unit microbial biomass) as a function of root inputs. (c) Microbial biomass as a function of the SOC pool. In panels a-c, root input C:N varies between 1 and 100 (colored lines).
Figure 5.9. Effect of 5°C warming (pink) and 5°C cooling (blue) on (a) C and (b) N mineralization. Effect of 1.5x soil moisture (pink) and 0.5x soil moisture (blue) on (c) C and (d) N mineralization. Black lines indicate C or N mineralization at ambient temperature and soil moisture conditions.
Figure 5.10. Absolute difference in (a) C and (b) N mineralization between ambient and 5°C of warming (pink) or cooling (blue). Absolute difference in (c) C and (d) N mineralization between ambient and 1.5x (pink) or 0.5x soil moisture (0; blue).
Figure 5.11. Cumulative sums of (a) C and (b) N mineralization rates under 5°C warming (pink) and cooling (blue). Cumulative sums of (c) C and (d) N mineralization rates with 1.5x soil moisture (pink) and 0.5x soil moisture (blue). Black lines indicate cumulative sums of C or N mineralization at ambient temperature and soil moisture conditions.
Figure 5.12. C mineralization under ambient temperature and soil moisture (θ; black line), 5°C warming with no change in soil moisture (pink line), and 5°C warming with a 10% decrease in soil moisture (0.90θ; green line).
CHAPTER SIX: CONCLUSION

The research in this dissertation followed C from aboveground allocation through roots to soil microbes at seasonal and inter-annual timescales. I used data compilation, field studies, and computer simulation to study global, plot-level, and micro-site spatial scales, respectively. One overarching objective of this work was to scale up measurements of local or micro-site processes to make ecosystem scale inferences. This was accomplished using plot averaging in the field study in Chapter 3, and using computer simulation to scale micro-site microbial activity to recreate plot-level phenomena (i.e., heterotrophic respiration) in Chapter 5. I explored a new method for quantifying rhizosphere extent in Chapter 4, which could then be used to scale up microbial activity in models that represent the distribution of roots in soil and root-derived substrate supply (Finzi et al., 2015).

A second overarching theme was the effect of substrate supply on plant and microbial activity, such as the temporal coupling between photosynthesis and root growth or respiration, or the exudation of labile C to support microbial growth and soil organic matter (SOM) decomposition. My primary focus on C cycling was motivated by the fundamental role of C in organizing biological systems, as well as its role as a potent greenhouse gas that is altering the Earth’s climate, with largely negative impacts on ecosystem services (Field & Van Aalst, 2014).

**Phenology and C allocation**

In both a meta-analysis across multiple biomes and field studies in a mid-latitude temperate forest, I found a difference in the phenology of evergreen compared to
deciduous trees. Deciduous root growth occurred earlier in the growing season relative to evergreen trees. Moreover, the timing of maximum root production in deciduous trees was more synchronous with measurements of aboveground phenology than that of evergreen trees. If this difference in phenology is consistent among the many species that are represented in each growth form, then there are implications for the ecosystem models that currently assume all phenology is synchronous (Medvigy et al., 2009, Oleson et al., 2010, Woodward & Lomas, 2004). Based on the findings of Chapters 2 and 3, this assumption of synchronicity is accurate for the temperate broadleaf deciduous plant functional type (PFT). However, a more accurate representation of the temperate needleleaf evergreen PFT is a lagged allocation of C belowground to roots.

It is difficult to determine one set of parameters for the all of species represented in a PFT because of the diversity of plant trait values among different species. As a result of this limitation, recent proposals are focusing on replacing PFTs altogether with species-specific or phylogenetically hierarchical plant traits that may vary as a function of temperature, soil moisture, clay content, or other widely available correlates of plant function (Wullschleger et al., 2014). Though existing plant trait databases are very sparse (Kattge et al., 2011), advances in matrix imputation may be able to use phylogenetic relationships to fill in missing plant traits with traits of closely-related taxa (Swenson, 2014). My data are immediately applicable as part of a plant trait database. Chapter 2 meta-analysis data have already been submitted to TRY, a global plant trait database initiated in 2007 (Kattge et al., 2011).
Temperate deciduous trees have high growing season photosynthetic rates relative to temperate conifers (Hadley et al., 2008), resulting in similarly high rates of belowground C allocation. I found that red oak (*Quercus rubra*) stands allocated ~250 g m\(^{-2}\) more C belowground than eastern hemlock (*Tsuga canadensis*) during the growing season. This discrepancy is driven by the difference in root production between stands. This observation suggests that long-term increases in C uptake and biomass noted in red oak trees at the Harvard Forest are facilitated by root production to meet nutrient demand (Keenan et al., 2012, Urbanski et al., 2007).

In red oak, increasing root production was a confirmation of high stand productivity, but in eastern hemlock, decreasing root production suggested a stand-level decline. The year-over-year decline in root production is coincident with the arrival of the hemlock woolly adelgid in 2012, its spread throughout Harvard Forest in subsequent years, and visible signs of crown thinning and HWA-induced tree mortality recorded by 2014. From these data, I conclude that disease-related declines in productivity are manifested belowground. Notably, live roots sampled from the hemlock stand had similar nonstructural carbohydrate, exudation, and respiration measurements to the other stands. Both the tight coupling of above and belowground phenology in red oak and the rapidity of disease-related decline in hemlock suggest that newly fixed C is important to root growth and longevity.

There is still little known about the phenology of roots. The National Ecological Observatory Network (NEON), founded in 2006 to collect standardized observational data across the United States, is building soil arrays that include minirhizotron cameras
permanently mounted in tubes installed into their terrestrial sites (Pennisi, 2010). These cameras can rotate and move vertically on a track in order to take photos and wirelessly transmit data back to a field station (Roberti et al., 2014). This will be the first large-scale implementation of this technology, and will also create the need to improve automated root detection, as the number of root images proposed cannot be annotated manually (Vamerali et al., 1999, Zeng et al., 2008). I am optimistic that NEON’s national implementation of minirhizotron cameras as well as ancillary soil and environmental measurements will produce data that greatly improve estimates of root production and its environmental drivers.

**Modeling the micro-site at global scales**

I developed a model that represents the effect of temperature and substrate supply on microbial activity by merging the Dual Arrhenius Michaelis-Menten model of Davidson et al. (2012) with the Microbial Carbon and Nitrogen Physiology Model of Finzi et al. (2015). The combined model, DAMM-MCNiP, reproduced measured rates of heterotrophic respiration over the growing season, and performed particularly well compared to each model alone when confronted with wet-up events. I show that the stoichiometry of root exudates influences both the amount and the mechanism by which priming occurs.

In order to scale up DAMM-MCNiP’s rhizosphere and bulk soil model outputs to an ecosystem flux, I had to make an assumption about the rhizosphere volume. Currently, DAMM-MCNiP is parameterized for 1 cm$^3$ of soil that is assumed to be exactly 20% rhizosphere, determined using a modeling exercise based on the assumption that exudates
diffuse 2 mm away from the root surface (Finzi et al., 2015). I developed a quantitative framework for estimating the spatial extent of the rhizosphere using image analysis of 2-D zymographs that may be able to improve estimates of the exudate diffusion distance.

An alternative to using a single value for the rhizosphere volume in models is to explicitly represent fluid transport of root exudates. I am preparing a depth-resolved version of DAMM-MCNiP that can include vertical or lateral transport of root exudates that are dissolved into the DOC pool, as well as declining root inputs with depth based on the depth distribution of root biomass (Finzi et al., 2015). Given adequate data on the enzyme activity of bulk versus rhizosphere soil, it may also be possible to use a future version of DAMM-MCNiP to estimate an effective rhizosphere distance by fitting it as a parameter to measurable outputs such as C and N mineralization.

The main contribution of DAMM-MCNiP is that it explicitly represents the processes that control the availability of SOM. However, SOM availability is also determined by the chemical composition of the soil, which influences aggregate formation, adsorption, and desorption of organic matter to minerals (Conant et al., 2011, Grandy & Neff, 2008). DAMM-MCNiP does not currently represent physical or chemical protection of SOM dynamically. Rather, DAMM-MCNiP considers a fixed fraction of SOM at each time step to be “unprotected”. This unprotected SOM is available for microbial depolymerization if it can diffuse to an extracellular enzyme. While DAMM-MCNiP can reproduce C efflux after a wet-up event by accurately representing the large amount of available C released by wet-up (Parton et al., 2012), it
over-predicts C efflux in constant high soil moisture conditions. DAMM-MCNiP’s representation of soil moisture dynamics may improve with parameter optimization, but it is also possible that there is a temperature control on SOM protection that is not currently represented.

Some microbial physiology models have also started to incorporate some representation of sorption to soil minerals (Tang & Riley, 2015, White et al., 2014). One microbial physiology model that has an intriguing simplification of SOM protection is MIMICS, which omits soil moisture but calculates the half-saturation constant of depolymerization as a function of soil clay content. This approximation of SOM “protection” simulates changes in SOM pools better than DAYCENT, a model lacking microbial physiology (Wieder et al., 2014).

There are a large number of decomposition models, many of which have some representation of microbial physiology (Allison et al., 2010, Drake et al., 2013, Moorhead et al., 2012, Schimel & Weintraub, 2003, Tang & Riley, 2015). Though DAMM-MCNiP is the only microbial physiology model to include representation of temperature, substrate supply, C and N cycling, it may not necessarily have the most predictive power. In order to determine which models perform well when confronted with new data, there needs to be a framework for model testing and comparison. In CMIP5, the predicted pools or fluxes of different process-models are plotted together and a moving average of these models is considered the most likely scenario (Ahlström et al., 2012). If parameter uncertainty can be propagated through to outputs, then it is possible
to compare model outputs to each other and also identify which parameters require more data constraints. This type of error propagation could be done in a Bayesian framework (Tang & Zhuang, 2009) or in a computationally simpler Monte Carlo framework (Keenan et al., 2013).

**Summary**

The research presented in this dissertation focuses on the seasonality and partitioning of belowground C allocation to roots and soil microbes. I found that above and belowground phenology are often asynchronous, and that root growth drives the phenology of belowground C allocation. Root growth and respiration are positively correlated with temperature, and C allocation to root production in a deciduous mid-latitude forest is tightly coupled with photosynthesis. Continued burning of fossil fuels will increase the temperature and CO$_2$ concentration in the atmosphere, resulting in an increase in belowground C allocation (Drake et al., 2011). Root allocation and elevated temperature both stimulate soil microbial activity, and ultimately, the flux of CO$_2$ into the atmosphere via heterotrophic respiration. This cycle constitutes a positive feedback to climate change, one that may be mitigated by C sequestration in plant biomass, acclimation of microbial metabolism, or depletion of substrate supply (Frey et al., 2013, Norby et al., 2005).

DAMM-MCNiP is one of the first microbial physiology models that can represent the C and N cycle together with plant substrate supply, making it a candidate for linkage with terrestrial biosphere models that are used to estimate feedbacks between plants, microbes, soil and the atmosphere. Though I have only tested DAMM-MCNiP against
one data set thus far, others have demonstrated that including microbial processes in C cycle models substantially improves predictions of the spatial distribution of SOC (Hararuk et al., 2014). Inclusion of these processes can change the prediction of future SOC pool size by > 300 Pg C (Tang & Riley, 2015, Wieder et al., 2013), indicating that plant–microbe–soil feedbacks are critical to the global C cycle.
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CURRICULUM VITAE

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EDUCATION

Boston University  May 2015
Ph.D. Biology
- Certificate in Terrestrial Biogeoscience
- Advisor: Professor Adrien Finzi

Amherst College  May 2009
B.A. Biology, Theater and Dance
- Advisors: David Ratner, Wendy Woodson

RESEARCH EXPERIENCE

Finzi Lab, Boston University  March 2011-May 2015
Dissertation Research
- Literature review: Are root and shoot phenology in sync?
- The seasonality of C allocation to roots in a mid-latitude forest: Measurements of root growth, mortality and nonstructural carbohydrate content, respiration, and exudation on seasonal timescales; study system is three common northeastern temperate tree species, red oak (Quercus rubra), white ash (Fraxinus americana), and eastern hemlock (Tsuga canadensis)
- Where does the rhizosphere end? Measurements of spatially resolved in situ enzyme activity using zymograms incubated in root boxes: Image analysis is used to determine the spatial extent of enzyme activity and their proximity to roots
- Effects of soil warming and N addition on belowground phenology: Temperature and nitrogen manipulation experiment to determine the effect of warming and N fertilization on root mass, growth, mortality, mycorrhizal colonization, and emergence of soil fauna
- DAMM-MCNiP-RRM model fusion: Modular integration of three models that estimate 1) temperature, soil moisture and substrate supply effects on SOM depolymerization, 2) microbial physiology with C and N cycling, 3) rhizosphere and bulk soil spatial distribution and depth scaling
Elsevier  
Editorial Assistant for Biological Conservation  
Jan 2010- June 2011  
- Invited appropriate candidates to review submitted manuscripts  
- Addressed reviewer and author questions via e-mail on behalf of the journal

Amherst College Biology Department  
Summer 2007  
Independent Undergraduate Research  
- Detected GFP (green fluorescent protein) fluorescence in *Dictyostelium discoideum* to determine localization of DG17 gene in the cell  
- Transformed a synthetic gene into *Dictyostelium discoideum*  
- Trained in use of fluorescent microscopy, confocal microscopy, Western blot, and flow cytometry

Amherst College Chemistry Department  
Laboratory Assistant  
- Synthesized organic chemicals, specifically a novel PTP (protein tyrosine phosphatase) inhibitor  
- Trained in use of $^{13}$C-NMR (nuclear magnetic resonance) spectroscopy

National Center for Microscopy and Imaging Research  
Summer 2003, 2006  
Laboratory Assistant  
- Traced regions of interest in the hippocampus of the mouse brain, using xvoxtrace and imod tracing programs  
- Created 3-d medium spiny dendrite model  
- Performed in situ hybridization on hippocampal, cerebellar, and forebrain sections of mouse brain  
- Trained in use of in situ hybridization, immunocytochemistry, cryostat operation

TEACHING EXPERIENCE

Pomona College  
May 2013- May 2014  
Undergraduate thesis advisor  
- Served as thesis advisor and first reader for Johanna Recalde’s senior honors thesis in root-mycorrhizal interactions  
- Helped Johanna develop and execute an independent research project to fulfill honors thesis requirements

Harvard Forest REU Program  
May 2013-August 2013, June 2012- August 2012  
Mentor  
- Mentored a total of three students for 11 weeks
- Helped students develop and execute an independent research project

**Boston University**  
**Sep 2012-Present, Sep 2010- May 2011**

- Teaching Fellow
  - Led two laboratory classes per week supervising a total of 42 undergraduate students
  - Organized and presented lab curriculum
  - Generated learning activities and discussion

**Finzi Lab**  
**Undergraduate Research Intern Mentor**  
**Jan 2011- August 2012**

- Coordinated up to 7 undergraduate research interns assisting with lab work
- Managed the independent research project of Amanda Alon, senior Biology major

**NSF GK-12 GLACIER**  
**Teaching Fellow**  
**June 2011- June 2012**

- Created and taught science and math curriculum for 5th graders for 10 hours per week

**OUTREACH**

**Advocates for Literacy in Environmental Sciences**  
**April 2012-present**

- Founding member, Secretary
- Graduate Student Organization Award for Excellence in Student Activities (2013)

**Pierce School Climate Change Summit**  
**January 2013**

- Moderator

**Curley K-8 School Science Fair**  
**January 2012**

- Judge

**NSF GK-12 GLACIER Garage Sale Fundraiser**  
**September 2011**

- Lead Organizer

**Summer Pathways Program: Tech Savvy**  
**July 2011**

- Program Coordinator

**Biology Inquiry & Outreach with Boston University Graduate Students**  
**May 2011**

- Student teacher

**POSITIONS HELD**
- LTER Higher Education Working Group Member (2013-present)
- Harvard Forest Graduate Student Representative (2013-2015)
- BU Biogeosciences Program Digital Media Committee Member (2014-2015)
- BU Biogeosciences Program Outreach Committee Member (2014-2015)
- Ecological Society of America Member (2012-present)
- American Geophysical Union Member (2012-present)

AWARDS
- Howard Hughes Medical Institute Fellowship (2007)
- Boston University Dean’s Fellowship (2009)
- NSF EAPSI Fellowship - Japan (2010)
- Amherst College Fellowship for Graduate Study: Forris Jewett Moore Fellowship (2009-2011)
- NSF GK-12 GLACIER Fellowship (2011)
- Graduate Writing Fellowship (waiting list, 2011)
- Boston University Biogeosciences Fellowship Research Award (2013)
- American Geophysical Union Outstanding Student Paper Award (2013)
- Department of Energy Student Travel Fellowship (2014)
- George R. Bernard, Jr. Travel Award (2012, 2013, 2014)
- American Association of University Women Dissertation Fellowship (2014)
- American Geophysical Union Student Travel Grant Award (2012, 2014)
- Boston University Biogeoscience Symposium Outstanding Oral Presentation Award (2015)

SKILLS
- Computer: Git/GitHub, R, Matlab, SAS, SPSS, JMP, STELLA, ArcGIS, RootFly, ImageJ/Fiji, SigmaPlot, Linux
- General: root viewing box and minirhizotron tube design and installation

PUBLICATIONS


Abramoff RZ, Finzi AF. Seasonality and partitioning of root allocation to rhizosphere soils in a mid-latitude forest (*in prep*)

Abramoff RZ, Finzi AF. Where does the rhizosphere end? Spatially resolved measurements of in situ soil extracellular enzyme activity (*in prep*)

Abramoff RZ, Davidson EA, Finzi AF. A parsimonious modular approach to building a mechanistic belowground carbon and nitrogen model (*in prep*)

**SELECTED PRESENTATIONS**

Long Term Ecological Research All Scientists’ Meeting (2012), September – **Poster: Phenology of belowground carbon allocation in a mid-latitude forest**

American Geophysical Union Fall Meeting (2013), December – **Presentation: Root phenology at Harvard Forest and beyond**

Northeast Natural History Conference (2014), April – **Invited Presentation: Linking above and belowground phenology at Harvard Forest and beyond**

Department of Energy Terrestrial Ecosystem Science PI Meeting (2014), May – **Poster: Seasonality of carbon allocation to roots in a mid-latitude forest**

Boson University (2014), May – **Guest lecture: Measuring C in terrestrial ecosystems**

Ecological Society of America Annual Meeting (2014), August – **Presentation (Organized Session): What is the relationship between aboveground and belowground phenology? A meta-analysis and case study**

American Geophysical Union Fall Meeting (2014), December – **Invited Presentation: A Parsimonious Modular Approach to Building a Mechanistic Belowground C and N Model**