



Changing litter composition following the dual invasion of Amur honeysuckle and the emerald ash borer alters fungal driven decomposition in Midwestern forests

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Abstract Midwestern forests are currently impacted by two prominent invaders, the emerald ash borer (EAB; *Agrilus planipennis*) and Amur honeysuckle (AHS; *Lonicera maackii*). The loss of ash (*Fraxinus* spp.) trees due to EAB invasion can further facilitate AHS invasion, driving changes in the composition of forest leaf litter to reflect a greater portion of labile, more easily decomposed litter. To evaluate the extent to which these changes alter ecosystem function, we conducted litter bag and culture-based decomposition experiments using leaf litter from sugar maple (*Acer saccharum*), oak (*Quercus* spp.), black ash (*Fraxinus nigra*), green ash (*Fraxinus pennsylvanica*), spicebush (*Lindera benzoin*) and AHS. To further understand the mechanism driving differences in decay rates, we inoculated six species of decomposing fungi separately onto both single species and multispecies (half AHS and half native species) leaf litter and measured decomposition rate, fungal growth and enzymatic activity in laboratory-based cultures. AHS leaf litter decomposed faster, had increased fungal growth, and had higher activity for carbon degrading enzymes

compared to native species leaf litter. Furthermore, multispecies mixtures followed the same patterns as AHS, suggesting that the addition of AHS to leaf litter to native litter will accelerate ecosystem functions related to carbon breakdown. Consequently, forests that experience the invasion of AHS and EAB induced loss of ash are likely to have faster rates of decomposition, potentially resulting in an influx of available nutrients.

Keywords *Lonicera maackii* · *Agrilus planipennis* · Fungi · Leaf litter decomposition · Enzyme activities

Introduction

Forests throughout the world have undergone major changes that affect entire ecosystems and will have lasting impacts on their overall composition and function. These changes include dominant species loss, habitat destruction and exotic species introductions and invasions, each of which greatly alter ecosystem functioning (Tilman et al. 1994; Pyšek and Richardson 2010). In eastern deciduous forest of the United States, forests are greatly impacted by the introduction of the emerald ash borer (EAB; *Agrilus planipennis*), which has significantly decreased the abundance of ash (*Fraxinus* spp.) trees as well as the introduction of the invasive shrub Amur honeysuckle (AHS; *Lonicera maackii*). While the independent roles these two events play in structuring plant communities are

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widely studied (i.e., Gould and Gorchov 2000; Collier et al. 2002; Hartman and McCarthy 2004; Schradin and Cipollini 2012; Klooster et al. 2014; Spei and Kashian 2017), the interactive effects of these events on ecosystem functions such as nutrient cycling and microbial processes are largely unexplored. Here we investigate the potential for forests experiencing a large degree of EAB induced ash mortality with a concurrent expansion of the invasion of AHS to experience changes nutrient cycling as a result of changes in fungal driven decomposition.

Originally from Asia, EAB is an invasive pest with limited natural predators in North America (see Duan et al. 2012; Jennings et al. 2013; Hooie et al. 2015). Since its discovery in 2002 in southeast Michigan, it has spread to 35 states within the United States as well as five neighboring Canadian provinces (<http://www.emeraldashborer.info>; Nisbet et al. 2015). Ash mortality rates are near 100% in the most infected areas (Klooster et al. 2014). Loss of dominant tree species such as ash trees from forests are likely to impact key functions in forest ecosystem functions like decomposition. In turn, these losses are likely to lead to significant changes in nutrient availability due to interspecific differences in the quality of leaf litter which drive differences in the decay rates of leaves since litter from individual plant species has unique chemical properties. Labile leaves with higher N content decompose quicker compared to recalcitrant leaves, which have higher C content (Cotrufo et al. 2013). Leaves are more labile when they have a lower ratio of lignin, a polymer for cellular wall structure and cellulose, to N, which leads to a faster decomposition rate (Melillo et al. 1993; Bhatnagar et al. 2018). In general, leaf litter from invasive plant species has a quicker decay rate compared to native species due to a combination of having both higher N content and a lower amount of lignin (Supplementary Table 1; Arthur et al. 2012; Ehrenfeld 2003; Ashton et al. 2005). For example, European ash (*Fraxinus excelsior*) leaf litter has higher amounts of nitrogen (N), magnesium and calcium leading to faster decomposition and a higher litter turnover rate than both European beech (*Fagus sylvatica*) and lime (*Tilia cordata* Mill. or *Tilia platyphyllos*; Langenbruch et al. 2012). In North American forest ecosystems, decomposition of ash leaf litter leads to a lower soil carbon to N ratio than decomposition of other native species including red oak (*Quercus rubra*), sugar maple

(*Acer saccharum*), red maple (*Acer rubrum*), American beech (*Fagus grandifolia*) and eastern hemlock (*Tsuga canadensis*; Finzi et al. 1998). This change in soil nutrients reflects interspecific differences in the quality of leaf litter which drive differences in the decay rates of leaves. Therefore, losing ash from forest ecosystems can potentially lead to a large change in soil nutrient availability in forest ecosystems.

The loss of ash from forest ecosystems is also likely to cause increased turnover in plant species composition. Regeneration of white, black and green ash following EAB induced mortality has been limited (Klooster et al. 2014), potentially as a result of depleted seed banks for these species (Kashian and Witter 2011; Knight et al. 2012); however, blue ash has lower rates of EAB infestation (Anulewicz et al. 2007) and mortality rates (Tanis and McCullough 2012), and appears to be regenerating at rates consistent with canopy self-replacement, suggesting it may still be a component of forests in which EAB is introduced (Spei and Kashian 2017). Forests where EAB has completely depleted white, green and black ash species are expected to experience canopy gaps, leading to higher light availability in the understory, which could lead to an increase in shrubs and saplings with high light tolerance (Dolan and Kilgore 2018). Replacement of ash by other native species can also potentially happen (Smith et al. 2015; Margulies et al. 2017) and maples (particular sugar maple) appear to be a prime candidate as they exhibit increased growth in areas with greater ash loss (Hoven et al. 2020). Replacement by an invasive species such as AHS, autumn olive (*Elaeagnus umbellata*) or common buckthorn (*Rhamnus cathartica*) is also possible since such species are better adapted to higher light availability than native species (McNeish and McEwan 2016; Hoven et al. 2017; Baron and Rubin 2021). Indeed, in an EAB infected forest with high rates of ash mortality, 18% of the remaining cover was from invasive plant species (Hoven et al. 2017). Similarly, when dead ash was removed from a forest, invasive plant species made up 18.7% of the total herbaceous cover (Hausman et al. 2010). When ash saplings are able to recover in a forest, there are declines in invasive and weedy native saplings (Margulies et al. 2017), suggesting forests where blue ash remain or regenerate may not experience the same degree of change to their species composition.

AHS is an invasive shrub that was introduced into the United States in 1896 but is now present throughout the understory of Midwestern forests and at least one Canadian province and 33 US states (Hutchinson and Vankat 1997; PLANTS 2023). As a light specialist, it is one of the prime candidates for replacing ash species in forests experiencing significant mortality due to EAB (Hoven et al. 2017). When a forest is invaded by AHS, it disrupts forest ecosystems in a variety of ways. When AHS is present, overall plant species richness and diversity declines (Gould and Gorochov 2000; Collier et al. 2002; Hartman and McCarthy 2004; Schradin and Cipollini 2012; McNeish and McEwan 2016). It can also reduce the growth of native plant species by limiting space and resources, such as light, water and nutrient availability (Arthur et al. 2012; Lieurance and Cipollini 2013). Through these effects AHS can both directly and indirectly alter nutrient cycling. The leaves of AHS have a higher N concentration and decompose faster than native species, potentially driving an increase of N availability in invaded forests (Arthur et al. 2012). There is also evidence that AHS increases decay rates for the litter of native species (Blair and Stowasser 2009). This may reflect differences in leaf chemical concentrations or differences in the microorganism community associated with its leaves. AHS has a unique community of microbes associated with its leaves that are not present on native ash or hickory (*Carya* spp.) leaves in Midwestern forests (Arthur et al. 2012). Since leaf microbial community can be an important driver for decomposition, the introduction of a new microbial community to a leaf litter community may drive changes overall decomposition rates and consequently drive changes in nutrient cycling. In whole, the combination of increased decomposition rates and reduction in native vegetative growth from AHS invasion could have a major impact on nutrient cycling in affected forests.

Despite differences in their time of introduction, the potential for EAB induced ash mortality to change forest dynamics in such a way as to facilitate expansion of the invasion by AHS is strong. For example, one study examining the potential for EAB induced ash mortality to facilitate AHS invasion found a positive relationship between AHS basal area growth and ash decline (Hoven et al. 2017) but the extent to which these invasions interact to alter ecosystem functions like decomposition are largely unexplored.

To do so, experiments which simultaneously compare leaf litter decay for native and introduced species are needed. AHS litter should decompose faster than litter from many native species due to having a low C:N compared to native trees, and within native tree species, oaks should have a slower decomposition rate than maple or ash species (Table S1; Howard and Howard 1974; Madritch and Cardinale 2007). When native species are combined with invasive species, decomposition rates of native species are usually accelerated (Ashton et al. 2005; Arthur et al. 2012), but litter decomposition can be slower when mixed with native species litter (Grossman et al. 2020). When AHS litter is combined with other litter from other species, we expect three potential outcomes. First, AHS could increase the overall rate of decomposition for the mixtures via additive properties, such as through “priming effects” which can occur when labile C additions accelerate microorganism decomposition of recalcitrant C sources (Rousk et al. 2015). Second, the mixture could decompose at a rate reflective of the average values for the mixed species when decomposing independently. Finally, the overall rate of decomposition could be slowed. The mechanisms driving non-additive effects of combining litter species on decomposition rates are not fully understood but generally revolve around mixtures possessing a greater diversity of litter chemistry which may result in the transfer of toxic compounds and/or phenolics between litter components, richer microorganism diversity or improved water retention (Wardle et al. 2003; Freschet et al. 2012; Otsing et al. 2018; Porre et al. 2020). Since decomposition dynamics for mixed species litter are often not easily predicted based on the decomposition patterns of single species (Gartner and Cardon 2004; Porre et al. 2020), to fully understand changes in decomposition dynamics of forests experiencing both a loss of ash due to EAB and an expansion of AHS invasion, decomposition of different leaf species combinations should be directly evaluated.

Fungi in particular are important drivers of decomposition and do so through the secretion of a specialized set of extracellular enzymes that allow for the breakdown of different components of organic matter (Hankin and Anagnostakis 1975). Plant species invasion can lead to increases in enzyme activity, most noticeably for N and P decomposing enzymes (Zhou and Staver 2019), but can also contribute to

an increase in C degrading enzymes such as peroxidase and polyphenol oxidase (Woods et al. 2019), potentially as a result of shifts in fungal communities driven by invasive species (Liu et al. 2019; Yang et al. 2019). Consequently, changes in plant composition will drive changes in the composition of leaf litter, potentially altering fungal community composition and function, and thereby changing nutrient availability in invaded ecosystems.

Here we use both field and laboratory-based decomposition experiments to investigate how the addition of AHS to leaf litter in forests that have also experienced an EAB induced loss of ash changes fungal facilitated decomposition. We hypothesized that (1) AHS litter will decompose faster than all native species, including ash; (2) combinations of AHS litter and native litter will decompose faster than individual species litter; and (3) decomposition rates will be driven by changing fungal traits, such as increased fungal biomass, hyphal growth rate, and enzyme activities. Outcomes from this work will inform predictions related to nutrient cycling in forests experiencing EAB induced loss of ash with concurrent expansions of AHS.

Materials and methods

Study site

This study utilized material from a relatively undisturbed section of the Runkle Woods at Wright State University in Dayton, Ohio (39.785253°N, 84.05424°W). This section represents primary woods which are approximately 127 years old and has never been clear-cut (DeMars and Runkle 1992). The over-story of the woods consists primarily of oaks (*Quercus* spp.) and maples (*Acer* spp.), but also contains other species such as hickory (*Carya* spp.) and American elm (*Ulmus americana*; DeMars and Runkle 1992). EAB is believed to have been introduced into the woods in 2011 (Rigsby et al. 2014), and has subsequently reduced populations of white ash (*F. americana*), green ash (*F. pennsylvanica*) and blue ash (*F. quadrangulata*) such that only blue ash trees remain (Cipollini and Runkle, *personal communication*). Like other Midwestern forests, the understory has been invaded by and is now primarily composed of AHS (Dorning and

Cipollini 2006), but spicebush (*Lindera benzoin*) is also present in the shrub layer.

Leaf litter collection

This study focused on six woody species which are representative of a typical Midwestern forest: sugar maple (*A. saccharum*), northern red oak (*Q. rubra*), white oak (*Q. alba*), black ash (*F. nigra*), green ash (*F. pennsylvanica*), spicebush (*L. benzoin*) and AHS (*L. maackii*). Because there was not enough leaf litter of each individual *Quercus* spp. to include in the experiment independently, we combined litter from *Q. rubra* and *Q. alba* in a 1:1 mix to create a single ‘oak’ mixture. Additionally, green ash was only used to assess litter decomposition in culture as we could not collect enough to use for both experiments. These species vary in decomposition rate and leaf litter chemistry (Table S1; Petersen and Cummins 1974; Kominoski et al. 2007; Blair and Stowasser 2009; Swan et al. 2009; Arthur et al. 2012; Poulette and Arthur 2012; Nisbet et al. 2015; Jo et al. 2016; Stoler et al. 2017).

Leaves from sugar maple, both oak species, spicebush and AHS were hand collected in the WSU woods following natural senescence but leaves from black ash, originating from Baileys Nursery in St. Paul, Minnesota, and green ash, originating from the WSU woods, were collected from trees maintained in the WSU greenhouse. These trees were kept in pots with commercial soil outside the WSU greenhouse to prevent EAB infection (Friedman et al. 2020; Peterson et al. 2020). Following collection, leaf litter was dried in a drying oven at 80 °C for 2 days. To kill any microbes present, leaf litter was autoclaved for 20 min at 121 °C twice within 24 h. Litter was stored at room temperature until the start of the experiments, approximately seven days. All leaves were collected after abscission from September–November 2017 and September–November 2018 from multiple locations and pooled.

Litter bag decomposition

Experimental set-up

To evaluate field rates of litter decomposition, we performed a standard litter bag decomposition experiment. We placed 10 g of leaf litter from each of five

focal species (sugar maple, oak, black ash, spicebush and AHS) in individual litter bags constructed by folding 300 µm nylon mesh into 20.32 × 15.24 cm squares and stapling all four edges (Harmon et al. 1999). Decomposition bags with mixed species were created by combining 5 g of native plant leaf litter from sugar maple, oak or spicebush and 5 g of AHS leaf litter. A total of 56 litter bags (5 single species bags + 3 mixed species bags × 7 replicates) were placed in primary forested areas of the Runkle woods for 100 days (17 November 2018–25 February 2019). Litter bags were grouped into seven blocks so each species combination was represented in each block and arranged adjacent to one another. They were collected after 100 days since previous work indicated AHS leaf litter decomposed completely by this time (McEwan et al. 2012). Leaf litter was removed from litter bags and dried at 80 °C for 2 days to obtain dry weight to determine final leaf mass. The decomposition rate (k) was calculated in g/year by taking the natural log of the final weight of the leaf litter (W_f) divided by the initial weight of the leaf litter (W_o) (Equation 1; Olson 1963).

$$k = -\frac{\ln\left(\frac{W_f}{W_o}\right)}{t}$$

Equation 1: Exponential decay model (Olson 1963).

Litter decomposition in culture

To better understand the role fungi play in driving changes in decomposition rates, we created a culture-based experiment with six species of fungi representing three fungal guilds (white rot, brown rot, ectomycorrhizal fungi) originally collected from the WSU Runkle Woods: *Mycena galericulata* (brown rot), *Amanita parcivolvata* (ectomycorrhizal fungi), *Schizophyllum commune* (white rot), *Laetiporus sulphureus* (brown rot), *Pseudosperma rimosum* (aka, *Inocybe rimososa*; ectomycorrhizal fungi), and *Marasmius rotula* (white rot; Table S2). These fungi represent dominant members of the fungal community of the Runkle Woods (Rúa, unpublished data) and are generalist fungi expected to be present across temperate forests. Fungi were isolated and maintained on Modified Melkin-Norkans (MMN) agar for approximately 21 days prior to use. Streptomycin sulfate was added to the plates to prevent bacterial contamination.

To obtain enough fungal biomass for the experiment, we first placed 1 cm³ plugs from pure fungal cultures of each species onto 12 plates and incubated them for four weeks at 22 °C. We then inoculated 1 cm³ plugs from these plates onto leaf litter for the experiment.

Experimental set-up

To isolate the effect of fungi on leaf litter decomposition, we performed a culture-based decomposition experiment using different species of leaf litter and fungi. Cultures consisted of deep plate Petri-dishes (100 mm × 20 mm) filled with 30 ml MMN agar. Each plate received 1 g of a single leaf species (sugar maple, oak, black ash, green ash, spicebush or AHS) or a 1 g combination of 0.5 g of AHS and 0.5 g of the native leaf species (sugar maple, oak or spicebush), which were crushed by hand while wearing gloves and homogenized before being placed on each plate. Each fungal species by leaf litter type combination was replicated 7 times for a total of 378 plates [6 fungal species x (6 single species litter cultures + 3 mixed species litter cultures) × 7 replicates)]; however, some plates were discarded due to contamination resulting in a total of 367 plates (Table S3).

Each plate was inoculated with a single species of fungi. Four 1 cm³ plugs of fungi were removed from four-week-old cultures with a sterilized scalpel blade and placed on top of leaves. After a 100-day incubation period, fungal material was scraped from the leaves and placed into the agar to measure fungal biomass as described below. The remaining leaf litter was collected and weighed to determine rate of decomposition. Approximately 0.25 g of remaining litter was stored at –20 °C until ready for enzyme assays. For logistical purposes, the culture-based experiment was performed in two rounds with three different species of fungi in each round.

Fungal hyphae

To determine the growth rate of fungi in culture, fungal hyphal length was measured from the edge of the initial fungal plug to the end of the hyphae with a caliper twice a week until the hyphae reached the edge of the plate. Hyphal growth rate (mm/day) was calculated by taking the natural log of the difference

between initial hyphal length and final hyphal length over the time maximum hyphal length was reached.

Fungal biomass

To assess total fungal biomass, both the agar and the fungal material that was removed from the leaf litter were melted in a beaker in an autoclave following a procedure outlined by Maynard et al. (2017). Cultures were autoclaved for 20 min at 121 °C to separate the fungal material from the agar, then poured through a 45 µm sieve to isolate the fungal material from the agar. Fungal material was further separated from agar by rinsing with ~100 mL of 90 °C DI water. The remaining fungi was placed in a drying oven at 65 °C for 12–24 h until dry and weighed to determine fungal biomass in mg.

Enzyme activities

To determine differences in fungal function between leaf species, we tested five enzymatic activities commonly assessed in decomposition. We measured the activities of β -glucosidase (BG), cellobiohydrolase (CBH), leucine aminopeptidase (LAP), peroxidase (PER) and polyphenol oxidase (PPO) following the procedures outlined in Woods et al. (2019). Each enzyme assay was conducted using homogenous leaf slurries made with 13 mg of leaf and 4 ml of 50 mM sodium acetate buffer at a pH of 5.6 and incubated in the dark at 4 °C. For the fluorometric enzymes BG, CBH, and LAP, we measured assay absorbance and fluorescence values using a BioTex Synergy HT microplate reader (BioTek, Winooski, VT, USA). For the colorimetric enzymes, PPO and PER, we measured assay absorbance using a Molecular Devices Corporation SpectraMax 190 microplate reader (Molecular Devices Corporation, Sunnyvale, CA, USA). Incubation times were selected to maximize the potential of each enzymatic activity for leaf litter. Due to a lack of leaf litter, not all of the assays from CBH, PER and LAP could be completed, resulting in sample sizes for CBH, PER, and LAP that are lower than for the other fungal traits (Table S3). For univariate analyses described below, analyses were conducted without these missing values but for multivariate analyses, values were imputed.

Statistical analysis

All statistical analyses were performed in the statistical programming environment R version 4.2.0 (R Core Team 2022). All results were visualized using *ggplot2* (Wickham 2016) unless otherwise noted. Significant interactions for all models were tested with ANOVA and post hoc analysis using the *emmeans* package with adjustment for Tukey HSD (Lenth 2022).

To understand how litter decay rates differed among plant species in the field, decay rate was tested as a function of plant species using a linear model created with the *lm* function from the *stats* package (R Core Team 2022) and one-way ANOVA. The effect of block was tested as an interaction with plant species but was found to be non-significant; therefore, analyses did not include this effect moving forward. Separate models were created with data from the mixed species bag and the two single species bags that made up those mixes to assess differences in decay rates for natives when AHS is present.

To evaluate the extent to which fungi drive differences in decay rate, we tested decay rate as a function of either hyphal growth rate, fungal biomass or enzyme activity with an interaction for plant species using a linear mixed effects model from the package *nlme* (Pinheiro et al. 2023) and a random effect for experimental round. We used principal component analysis (PCA) to condense enzyme activities into two linear principal components using *prcomp* from the *stats* package. PERMANOVA using the *adonis2* function in the *vegan* package with 1000 permutations was then used to determine significant differences in enzyme activities based on decay rates (Oksanen et al. 2022). Prior to multivariate analyses, missing data for CBH ($n=13$), PER ($n=20$), and LAP ($n=1$) was interpolated using the *na.approx* function from the *zoo* package (Zeileis and Grothendieck 2005).

Structural equation modeling was used to compare the relative importance of the different fungal traits on decomposition rates for each litter species. Partial least squares path modeling (PLS-PM) was conducted using the package *plspm* to assess the direct and indirect effects of fungal hyphal growth rate, fungal biomass and enzyme activities (CBH, PER, PPO, LAP, BG) on decomposition rates (Sanchez et al. 2015). As with the PCA, interpolated values were used for missing CBH, PER, and LAP data. An *a priori* model

was constructed to include all possible paths among these factors (Fig. S1). Estimates of path coefficients, coefficients of determination (R^2) and goodness-of-fit (GoF) values were validated using 999 bootstraps on the path model. The GoF statistic was used to assess the reliability of the models. Independent models were created for each plant species and plant species combinations to compare differences in decomposition drivers.

Results

Data not shown for non-statistically significant results.

Plant species identity and decay

Litter bag decay rates

For single species litter bags, litter decay rate significantly varied by plant species ($F_{4,30}=83.51$, $P<0.0001$, Table 1). AHS leaf litter decayed ~100% faster than spicebush and black ash, ~200% faster than sugar maple and ~250% faster than oak (Table 1). In

multispecies mixtures, decay rates were intermediate between AHS and its associated native species (Table 1). AHS + spicebush mixtures decayed ~43% faster than spicebush alone ($F_{2,18}=19.82$, $P<0.0001$, Table 1), AHS + sugar maple mixtures decayed ~90% faster than sugar maple alone ($F_{2,18}=62.57$, $P<0.0001$, Table 1) and AHS + oak mixtures decayed ~90% faster than oak alone ($F_{2,18}=63.41$, $P<0.0001$, Table 1).

Culture based decay rates

For cultures with a single plant species, litter decay rate significantly varied by plant species ($F_{5,238}=27.2$, $P<0.0001$, Table 1). AHS leaf litter decayed ~65% faster than spicebush and black ash, 75% faster than green ash and ~80% faster than sugar maple and oak (Table 1). In multispecies mixtures, decay rates were intermediate between AHS and its associated native species (Table 1). AHS + spicebush mixtures decayed ~37% faster than spicebush alone ($F_{2,118}=54.52$, $P<0.0001$, Table 1), AHS + sugar maple mixtures decayed 49% faster than sugar maple alone ($F_{2,120}=62.08$, $P<0.0001$, Table 1) and

Table 1 Mean decay rates (g/year) and standard error by litter treatment and experiment

Litter treatment	Common name	Field experiment	Culture based experiment
<i>Single species</i>			
<i>Lonicera maackii</i>	Amur honeysuckle	3.04 ± 0.2^a	1.86 ± 0.1^a
<i>Lindera benzoin</i>	Spicebush	1.72 ± 0.09^b	0.94 ± 0.05^b
<i>Acer saccharum</i>	Sugar maple	1.10 ± 0.05^c	0.78 ± 0.04^b
<i>Quercus</i> spp.	Oak	0.84 ± 0.03^c	0.77 ± 0.05^b
<i>Fraxinus nigra</i>	Black ash	1.79 ± 0.1^b	0.94 ± 0.06^b
<i>Fraxinus pennsylvanica</i>	Green ash	—	0.82 ± 0.06^b
<i>Mixed Species</i>			
<i>Lonicera maackii</i>	Amur honeysuckle	3.04 ± 0.2^a	1.86 ± 0.1^a
<i>Acer saccharum</i>	Sugar maple	1.10 ± 0.05^c	0.78 ± 0.04^c
<i>Lonicera maackii</i> + <i>Acer saccharum</i>	Amur Honeysuckle + Sugar maple	2.22 ± 0.06^b	1.3 ± 0.07^b
<i>Lonicera maackii</i>	Amur honeysuckle	3.04 ± 0.2^a	1.86 ± 0.1^a
<i>Lindera benzoin</i>	Spicebush	1.72 ± 0.09^c	1.38 ± 0.06^c
<i>Lonicera maackii</i> + <i>Lindera benzoin</i>	Amur Honeysuckle + Spicebush	2.14 ± 0.13^b	0.94 ± 0.05^b
<i>Lonicera maackii</i>	Amur honeysuckle	3.04 ± 0.2^a	1.86 ± 0.1^a
<i>Quercus</i> spp.	Oak	0.84 ± 0.03^c	1.31 ± 0.07^c
<i>Lonicera maackii</i> + <i>Quercus</i> spp.	Amur Honeysuckle + Oaks	2.39 ± 0.14^b	0.77 ± 0.05^b

a–c Different letters indicate significant difference from Tukey's honest significant difference (HSD) test

AHS + oak mixtures decayed 52% faster than oak alone ($F_{2,130}=95.1, P<0.0001$, Table 1).

Culture based litter decay rates

Role of hyphal growth rate and plant species identity on decay rates

The interaction of hyphal growth rate and plant species identity significantly explained leaf litter decay rates in single species litter cultures ($F_{5,232}=8.467, P<0.0001$). AHS, green ash, oak and black ash litter decayed slower with increased hyphal growth rate, but decay rates increased with an increase in hyphal growth rate for spicebush and sugar maple litter (Fig. 1A).

Patterns for decay rate as a function of fungal hyphal growth rate and plant species identity for multispecies mixtures consistently fell between patterns for AHS and the relevant single species culture (Fig. 1B–D). Each multispecies culture followed similar trends to AHS litter such that decay rates decreased with increasing hyphal growth rate: AHS + spicebush ($F_{2,115}=14.03, P<0.0001$, Fig. 1B), AHS + maple ($F_{2,117}=11.89, P<0.0001$, Fig. 1C) and AHS + oak ($F_{2,117}=4.068, P=0.0196$, Fig. 1D).

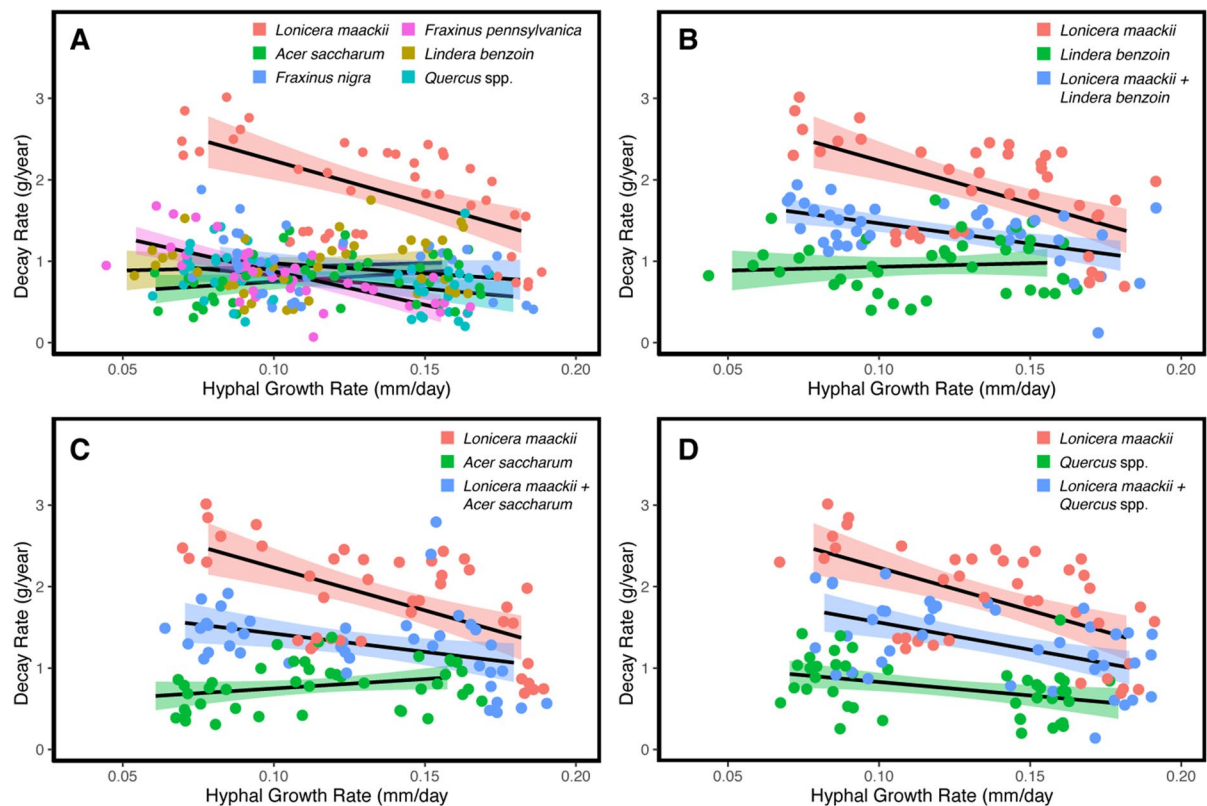


Fig. 1 Hyphal growth rate, plant species identity and fungal guild drive litter decay rate for **A** single species cultures ($P<0.0001$), **B** spicebush ($P<0.0001$), **C** maple ($P<0.0001$), and **D** oak ($P=0.0196$). All plant species had decreased decay with increasing hyphal growth rate except for spicebush and sugar maple. Decay rates for multispecies cultures followed similar patterns to AHS with decreasing decay rates with

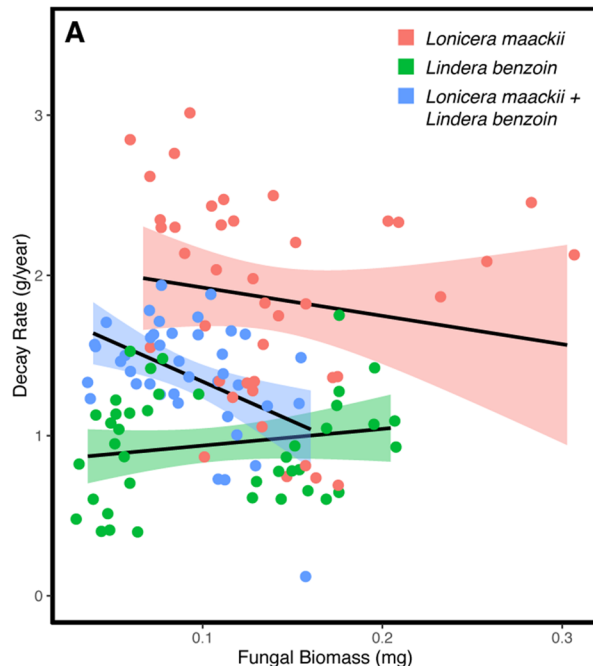
increased hyphal growth rate. Shaded regions represent 95% confidence intervals for AHS (pink), spicebush (brown), sugar maple (green), oak (light blue), black ash (dark blue), and green ash (purple) in single species cultures and AHS (pink), native species (green) and mixtures (blue) in multispecies cultures

Role of fungal biomass and plant species identity on decay rates

In single species models, litter decay rate increased with increasing fungal biomass ($F_{1,232}=25.15$, $P<0.0001$) but this was not affected by plant species identity ($F_{5,232}=1.375$, $P=0.2343$). In multispecies cultures, litter decay rates decreased with increasing fungal biomass for AHS + spicebush ($F_{2,115}=4.065$, $P=0.0197$, Fig. 2A) and AHS + maple ($F_{2,117}=4.004$, $P=0.0208$, Fig. 2B) models. However, fungal biomass did not significantly predict decay rate as an interaction with plant species in AHS + oak mixed models ($F_{2,117}=1.769$, $P=0.1750$).

Role of enzyme activities and plant species identity on decay rates

Individual enzyme activities and plant species identity predicted litter decay rate for single species cultures for CBH ($F_{5,221}=2.211$, $P=0.0542$), PPO ($F_{5,232}=6.151$, $P<0.0001$), LAP ($F_{5,231}=4.753$, $P=0.0004$) and PER ($F_{5,218}=2.242$, $P=0.0512$).



Decay rates increased with increasing CBH activity for all plant species except for spicebush litter (Fig. 3A). Decay rates increased with increasing PPO activity for AHS, spicebush, oak, and black ash but decreased with increasing PPO activity for maple and green ash (Fig. 3B). Decay rates increased with increasing LAP activity for AHS and oaks, but decreased for spicebush, maple, black ash and green ash (Fig. 3C). Finally, decay rates decreased with increasing PER activity for AHS, oak, and green ash and increased for black ash but had no relationship for spicebush and maple (Fig. 3D).

Decay rate increased with increasing BG activity ($F_{1,232}=42.10$, $P<0.0001$) but this did not interact with plant species identity ($F_{1,232}=0.8883$, $P=0.4896$).

Multispecies cultures

Spicebush

Decay rates increased with increasing enzyme activity for AHS, spicebush, and their mixed litter for

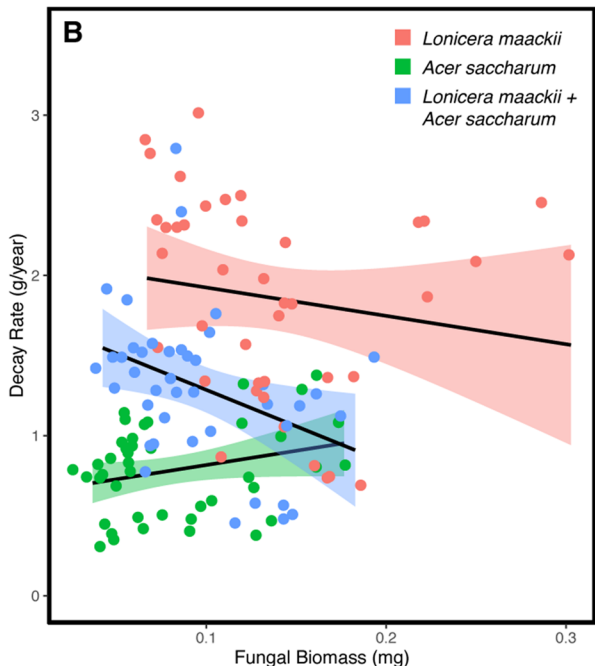


Fig. 2 Fungal biomass and plant species identity drive litter decay rate in multispecies cultures for **A** spicebush ($P=0.0197$) and **B** maple ($P=0.0208$). Similar to AHS, in multispecies litter cultures, decay rates decreased with increasing fungal biomass compared to native single species litter where decay rates increased with increasing fungal biomass. Shaded regions represent 95% confidence intervals for AHS (pink), native species (green) and mixtures (blue)

ing fungal biomass compared to native single species litter where decay rates increased with increasing fungal biomass. Shaded regions represent 95% confidence intervals for AHS (pink), native species (green) and mixtures (blue)

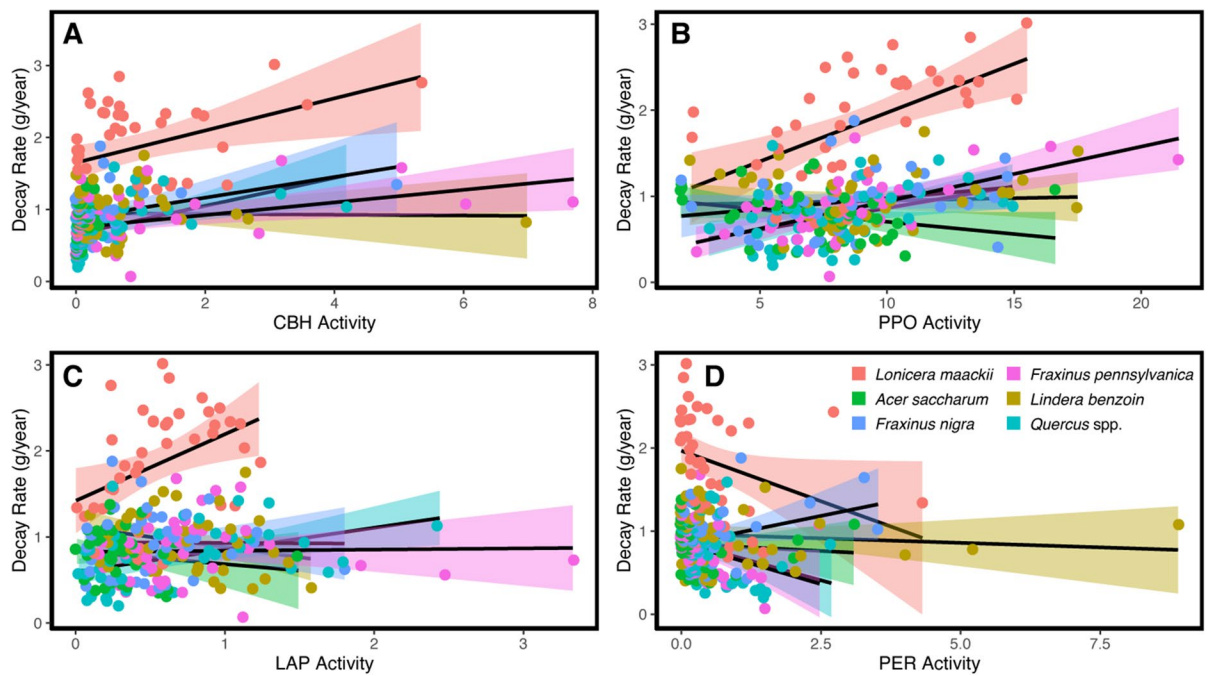


Fig. 3 Enzyme activity and plant species identity interacted to drive decay rate in single species cultures for **A** cellobiohydrolase (CBH) activity ($P<0.0001$), **B** leucine aminopeptidase (LAP) activity ($P=0.0004$), **C** polyphenol oxidase

(PPO) activity ($P<0.0001$), and **D** peroxidase (PER) activity ($P=0.0512$). Shaded regions represent 95% confidence intervals for AHS (pink), spicebush (brown), maple (green), oak (light blue), black ash (dark blue), and green ash (purple)

CBH activity ($F_{2,114}=4.358$, $P=0.0150$; Fig. 4A) and PPO activity ($F_{2,115}=5.582$, $P=0.0049$; Fig. 4B). The relationship between decay rates and LAP activity varied such that increasing LAP activity increased decay rates of AHS litter, decreased decay rates of spicebush litter and increased decay rates for mixed litter ($F_{2,115}=8.584$, $P<0.0001$; Fig. 4C). Enzyme activity failed to predict decay rates as a function of plant species identity in AHS + spicebush cultures for BG activity ($F_{2,115}=0.7934$, $P=0.4548$) and PER activity ($F_{2,111}=1.557$, $P=0.2154$).

Maple

Decay rates increased with increasing PPO activity for both AHS litter and maple + AHS mixed litter but decreased for maple litter alone ($F_{2,117}=7.528$, $P<0.0001$; Fig. 5A). In contrast, decay rates increased with increasing LAP activity for AHS litter, decreased with increasing LAP activity for maple + AHS mixed litter and no effect for maple litter alone ($F_{2,116}=10.41$, $P=0.0001$; Fig. 5B). Enzyme activity failed to predict decay rates as a

function of plant species identity in AHS + maple cultures for BG activity ($F_{2,117}=0.6995$, $P=0.4989$), CBH activity ($F_{2,110}=0.9211$, $P=0.4011$), PER activity ($F_{2,107}=0.8731$, $P=0.4206$).

Oak

Decay rates increased with increasing PPO activity for AHS litter, oak litter, and mixed AHS-oak litter ($F_{2,117}=3.355$, $P=0.0383$, Fig. S3). Enzyme activity failed to predict decay rates as a function of plant species identity in AHS + maple cultures for BG activity ($F_{2,117}=1.244$, $P=0.2920$), CBH activity ($F_{2,116}=1.425$, $P=0.2448$), LAP activity ($F_{2,117}=2.618$, $P=0.0772$), PER activity ($F_{2,112}=0.3474$, $P=0.7073$).

Role of hyphal growth rate, fungal biomass and enzyme activities on decay rates

Independent of litter species identity, hyphal growth rate did not significantly predict litter decay rates ($F_{1,364}=0.0611$, $P=0.8049$) but fungal biomass did

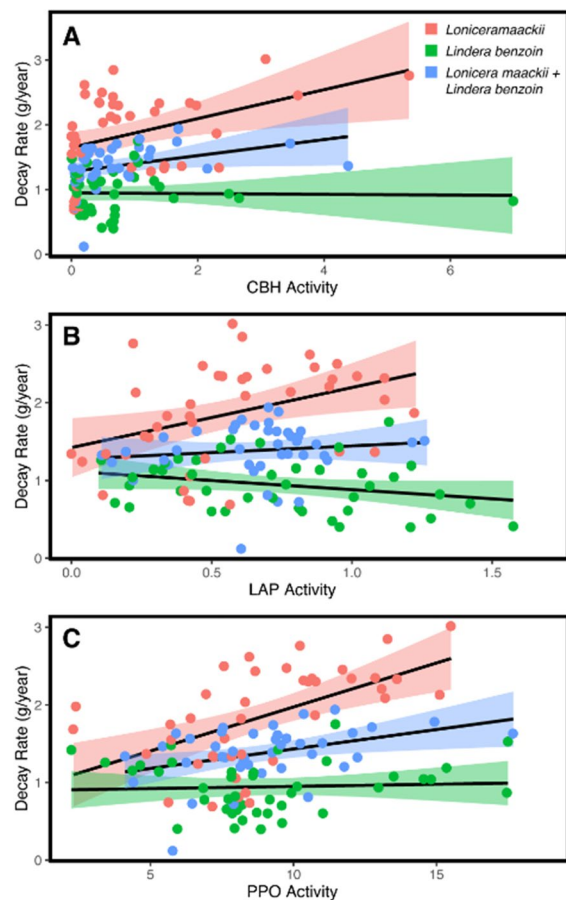


Fig. 4 The relationship between decay rates and enzyme activity with plant species identity varied for **A** cellobiohydro-lase (CBH) activity ($P=0.0150$), **B** polyphenol oxidase (PPO) activity ($P=0.0049$) and **C** leucine aminopeptidase (LAP) activity ($P<0.0001$). Shaded regions represent 95% confidence intervals for AHS (pink), spicebush (green) and AHS + spicebush mixtures (blue)

such that decay rates increased with increasing fungal biomass ($F_{1,364}=6.701, P=0.01$).

The relationship between decay rate and enzyme activities across litter species identity varied by enzyme. Decay rate increased with increasing BG activity ($F_{1,364}=28.87, P<0.0001$, Fig. S3A), CBH activity ($F_{1,351}=23.26, P<0.0001$, Fig. S3B), PPO activity ($F_{1,364}=18.02, P<0.0001$, Fig. S3C), and LAP activity ($F_{1,363}=5.94, P=0.0153$, Fig. S3D) but decay rate decreased with increasing PER activity ($F_{1,344}=4.138, P=0.0427$, Fig. S3E). PERMANOVA results suggest decay rate significantly affects enzymes activities ($R^2=0.08$,

$F_{1,364}=29.735, P=0.001$) but separation appeared weak (Fig. S3F).

Fungal traits as drivers of decay rates

PLS-PM analysis explained 22% of total variance in decay rates. In models with only the single species cultures, goodness of fit (GOF) values ranged from 0.35 for spicebush and sugar maple to 0.46 for oak, AHS, and green ash (Fig. 6). Oak, AHS, black ash, and green ash PLS-PM models all had similar outputs such that enzyme activities and fungal biomass jointly regulated decay rates, of which enzyme activities showed stronger direct and total impacts (Fig. 6A,C-E). The PLS-PM model for sugar maple also showed that enzyme activities have strong direct and total impacts on decay rates (Fig. 6B). In contrast, in spicebush PLS-PM models, the moderator variable fungal traits, of which fungal biomass and enzyme activities loaded most strongly, had the strongest effect on decay rates while fungal hyphal growth rate also had a significant effect on decay rate despite not mapping onto the fungal traits variable (Fig. 6F).

Models with mixed species cultures revealed interesting patterns relative to single species patterns. In AHS single species models, decay rate was regulated by both fungal biomass and most strongly, enzyme activities, but in sugar maple single species models, only enzyme activities regulated decay rates (Fig. 6A, C). For sugar maple+AHS PLS-PM models, decay rate was jointly and strongly regulated by enzyme activities and fungal biomass (Fig. 7A).

Oak and AHS single species PLS-PM models all had similar outputs such that decay rates were regulated by fungal biomass, and most strongly, enzyme activities (Fig. 6A, B). However, in mixed species PLS-PM models with oak+AHS, decay rates were regulated by fungal traits as represented by fungal biomass and enzyme activities and fungal growth rate (Fig. 7B).

Finally, decay rates were regulated differently for AHS and spicebush single species PLS-PM models such that enzyme activities and fungal biomass regulated decay rates for AHS while the moderator variable fungal traits as represented by fungal biomass and enzyme activities, had the strongest effect on decay rates, while fungal hyphal growth rate also had a significant effect on decay rate despite not mapping onto the fungal traits variable for spicebush (Fig. 6A,F).

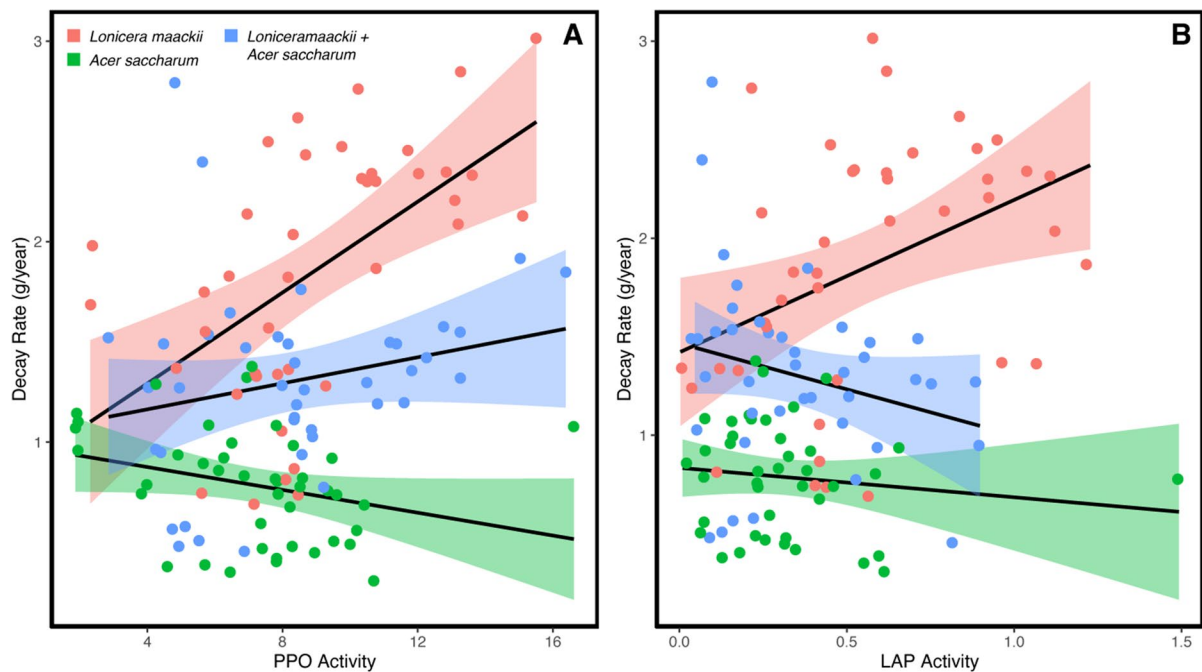


Fig. 5 **A** Polyphenol oxidase (PPO) activity and **B** leucine aminopeptidase (LAP) activity drives litter decay rates as a function of plant species identity. Litter decay rates in mixed AHS-maple cultures increased with increased PPO activity but

decreased with increased LAP activity for the mixes compared to the single species litter. Shaded regions represent 95% confidence intervals

In mixed species PLS-PM models with AHS + spicebush, enzyme activities strongly and directly regulated decay rates, but no other variables had an impact on decay rate (Fig. 7C).

Discussion

In the work presented here, we found that the addition of AHS litter not only increased decomposition rates when combined with native species litter, but also altered fungal biomass and enzyme activities, all of which contributed to increased decomposition rates. The addition of AHS further increased enzyme activity for several key enzymes important for nutrient cycling, particularly C associated enzymes. Across all native species, litter from both species of ash decayed the fastest and had higher CBH, LAP and PPO activities compared to the other native species. Taken together, these results suggest that forests incurring both a loss of ash and an expansion of the invasion of AHS may further experience increased rates of nutrient release from decaying litter,

potentially influencing the rate of C sequestration in these invaded forests. They also suggest decomposition rates in forests which lose ash from EAB invasion but do not experience AHS invasion will generally decline if sugar maple and/or oaks fill the void.

In general, invasive species leaf litter decomposes faster than native species leaf litter in both field and lab studies (Arthur et al. 2012; Nisbet et al. 2015; Jo et al. 2016). In this study, AHS litter decomposed the fastest among the six leaf litter species for both in situ litter bag and laboratory culture experiments. The accelerated decomposition of AHS litter compared to that of the native species supports the general finding that invasive species litter is more labile with faster decomposition rates than native species (Ehrenfeld 2003; Ashton et al. 2005; Arthur et al. 2012). Furthermore, mixing AHS and individual native species leaf litter increased decay rates compared to native species decay alone in both litter bags and lab cultures, although perhaps not to a degree different than you would expect given the mean of the decomposition rates of the species independently. This may reflect “priming effects” (Rousk et al. 2015)

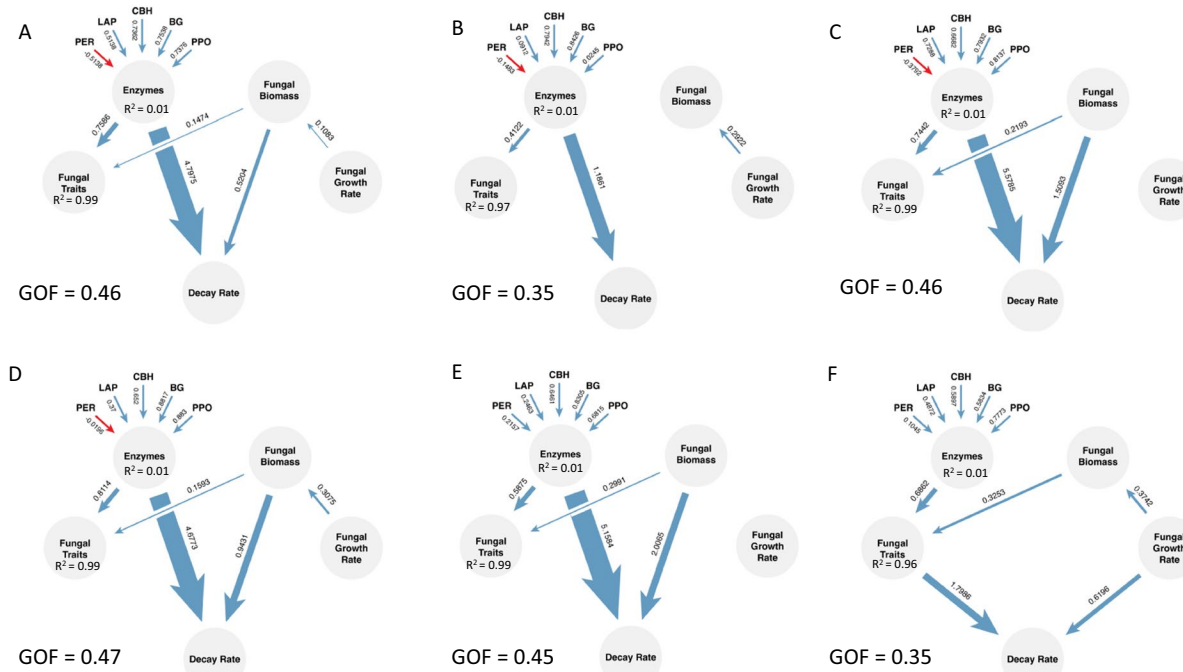


Fig. 6 Directed graph of the partial least squares path model (PLS-PM) analysis for decay rates for **A** AHS, **B** sugar maple, **C** oak, **D**, green ash, **E** black ash, and **F** spicebush. Each circle represents the observed (Fungal Biomass, Fungal Growth Rate) or latent variables (Enzymes, Fungal Traits). The latent variable Enzymes is created by the enzyme activities for β -glucosidase (BG), cellobiohydrolase (CBH), leucine aminopeptidase (LAP), peroxidase (PER) and polyphenol oxidase (PPO). The latent variable Fungal Traits is created from the latent variable Enzymes, Fungal Biomass and Fungal Growth Rate. Path coefficients and explained variability (R^2) are reflected in the width of the arrow were calculated after 1000 bootstraps. Blue and red arrows represent positive and negative effects. Goodness of fit (GOF) values represent model fit. Only significant paths ($P < 0.05$) are represented

(PPO). The latent variable Fungal Traits is created from the latent variable Enzymes, Fungal Biomass and Fungal Growth Rate. Path coefficients and explained variability (R^2) are reflected in the width of the arrow were calculated after 1000 bootstraps. Blue and red arrows represent positive and negative effects. Goodness of fit (GOF) values represent model fit. Only significant paths ($P < 0.05$) are represented

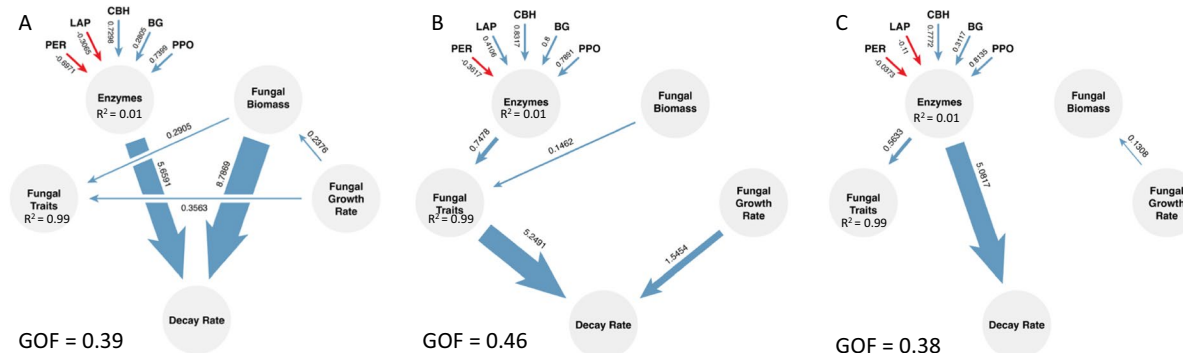


Fig. 7 Directed graph of the partial least squares path model (PLS-PM) analysis for decay rates for **A** AHS + sugar maple, **B** AHS + oak and **C** AHS + spicebush. Each circle represents the observed (Fungal Biomass, Fungal Growth Rate) or latent variables (Enzymes, Fungal Traits). The latent variable Enzymes is created by the enzyme activities for β -glucosidase (BG), cellobiohydrolase (CBH), leucine aminopeptidase (LAP), peroxidase (PER) and polyphenol oxidase (PPO). The

latent variable Fungal Traits is created from the latent variable Enzymes, Fungal Biomass and Fungal Growth Rate. Path coefficients and explained variability (R^2) are reflected in the width of the arrow were calculated after 1000 bootstraps. Blue and red arrows represent positive and negative effects. Goodness of fit (GOF) values represent model fit. Only significant paths ($P < 0.05$) are represented

of adding highly labile litter to more recalcitrant litter. In previous studies, the addition of a non-native labile leaf litter to recalcitrant native litter has led to mixed results for the direction of change in decomposition rates for the native leaf litter. Some studies have reported an overall increase in decomposition rate, which could be the result of increased N (Ashton et al. 2005; Arthur et al. 2012). Other studies have reported a decrease in decomposition rate, which may be driven by an increase in litter with diverse chemical traits (Zhang et al. 2014; Grossman et al. 2020). Our results may also reflect changes in decomposer community composition due to the addition of AHS (McEwan et al. 2012), however, we did not explicitly test phyllosphere microorganisms of any of the litter species and instead manipulated members of the decomposer community known to be important. Future research which explicitly identifies members of the phyllosphere is needed to rule out this mechanism. Overall, our results support an overall increase in litter decomposition due to the addition of AHS, which potentially reduces fungal N limitation during decay since its litter is so labile.

Invasive species can alter fungal growth and performance and thus act as primary drivers of changes in decomposition (Vitousek et al. 1997). In this study, fungi inoculated on AHS litter grew faster and had higher biomass than on native species litter; however, increased hyphal growth rates and fungal biomass were both associated with slower decay rates for AHS, green ash, black ash and oaks. This pattern suggests that fungi did not invest nutrients acquired from decomposition into new growth but instead invested in other avenues. One possible avenue is spore production, which we observed for both brown rot fungi but did not measure, as it can be important in the breakdown of labile litter components (van der Wal et al. 2013). Another possible avenue for fungi to invest nutrients during the decomposition process is investment in degradative enzyme production (Sinsabaugh 1994; Hättenschwiler et al. 2005). This seems likely in our experiment as evidenced by the strong relationship between decay rates and individual enzymes and enzymes as part of PLS-PM models.

The primary way fungi facilitate decomposition is through the release of extracellular enzymes (Hankin and Anagnostakis 1975). Here, enzymatic activity levels were key drivers of decay rates both as a member of a conglomerate variable for fungal traits and

independently. Additionally, their effects on decay rate differed by litter species such that enzymatic activities related to C and N breakdown increased decay rates more for species associated with labile litter compared to species associated with recalcitrant litter. Specifically, LAP, CBH and PPO had higher activities on AHS leaf litter compared to native species leaf litter. Higher enzyme activity for enzymes associated with N on AHS supports previous studies with invasive plants which suggests an increase in activity levels due to a larger pool of available N in invasive plants (Liao et al. 2008; Vilà et al. 2011; Zhou and Staver 2019). However, the results presented here with C associated enzymes do reflect previous studies that demonstrated increased C enzyme activity for C associated enzymes such as PPO when AHS is present (Liao et al. 2008; Woods et al. 2019).

Ash litter decomposed slower than AHS litter but decomposed faster than other native species. This finding supports ash litter being more labile than other native species, but less so than AHS (Nisbet et al. 2015). Just as with AHS, decomposition rates are driven by fungal traits. Specifically, decay rates decreased with increasing hyphal growth rates and tracked with enzyme activities. Ash litter had higher LAP, CBH, and PPO activities compared to litter from the other native species. This trend was particularly prominent for green ash, which had the overall highest enzyme activities compared to the other plant species. In total, these results support previous research that ash have an outsized effect on soil nutrient availability in forests where they are present (Langenbruch et al. 2012).

With greater decomposition, there is increased nutrient availability in forest soils (Sinsabaugh and Moorhead 1994). Increased nutrient availability can lead microorganisms to allocate greater enzymatic activity towards C degradation instead of N and P degradation, leading to increased C cycling (Allison and Vitousek 2004). In this system, multispecies litter with AHS decomposed faster than the native litter alone for all tested species, suggesting that forest systems where AHS is invading would cycle C faster than uninvaded ecosystems. This effect may be counterbalanced in systems also experiencing EAB induced loss of ash since ash represents a large percentage of biometric net primary production in forests where they are present (Flower et al. 2013). Therefore, the dual invasion of EAB and AHS potentially

leads to no net change in C despite the loss of C from the loss of ash. In forest systems which have experienced EAB induced loss of ash but where AHS is not invading, we expect to see lower rates of C cycling since natives like sugar maple and oaks which are expected to replace ash in these systems have lower decomposition rates and consequently release less C (Arthur et al. 2012; Marshall 2020).

In summary, the addition of AHS litter to systems experiencing EAB induced loss of ash is altering litter decomposition through several fungal traits: increased hyphal growth rates, increased fungal biomass, increases in activities of enzymes associated with C and decreases in activities of enzymes associated with both N and P. Forests that previously had abundant ash populations are increasingly becoming overtaken by invasive shrubs (Hoven et al. 2017). The consequent changes to the leaf litter layer are likely to have lasting impacts on overall soil nutrient cycling. This study represents an important first step in understanding how fungal driven responses to this changing litter layer will change in response to the alteration in leaf litter from the transition of ash to AHS in Midwestern US forests.

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Author contributions All authors contributed to the study conception and design. Material preparation, data collection and initial data analyses were performed by AMR with input from MAR. Subsequent data analyses were performed by MAR. CR supported AMR in data collection. The first draft of the manuscript was written by AMR with subsequent drafts written by MAR. All authors commented on previous versions of the manuscript and read and approved the final manuscript.

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Availability of data and materials The datasets generated in this study and R code used to analyze that data have been uploaded to the Environmental Data Initiative Data Repository (<https://doi.org/10.6073/pasta/e87910d2313e269c3e2124b85dc03011>).

Declarations

Conflict of interest The authors have no relevant financial or non-financial interests to disclose.

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