Negative density dependence is stronger in resource-rich environments and diversifies communities when stronger for common but not rare species

Abstract
Conspecific negative density dependence is thought to maintain diversity by limiting abundances of common species. Yet the extent to which this mechanism can explain patterns of species diversity across environmental gradients is largely unknown. We examined density-dependent recruitment of seedlings and saplings and changes in local species diversity across a soil-resource gradient for 38 woody-plant species in a temperate forest. At both life stages, the strength of negative density dependence increased with resource availability, becoming relatively stronger for rare species during seedling recruitment, but stronger for common species during sapling recruitment. Moreover, negative density dependence appeared to reduce diversity when stronger for rare than common species, but increase diversity when stronger for common species. Our results suggest that negative density dependence is stronger in resource-rich environments and can either decrease or maintain diversity depending on its relative strength among common and rare species.

Keywords
Density dependence, diversity maintenance, diversity–environment relationship, Janzen–Connell hypothesis, natural enemies, resource availability, seedling and sapling recruitment, species coexistence, species relative abundance, temperate forest.

INTRODUCTION
Conspecific negative density dependence (CNDD) is a widespread population process thought to maintain high species diversity by imposing intrinsic limits on population densities of individual species (Harms et al. 2000; HilleRisLambers et al. 2002; Comita et al. 2014; Bever et al. 2015). A classic model invoking CNDD is the Janzen–Connell hypothesis, which predicts that density- or distance-dependent specialised enemies reduce recruitment near conspecific adults, making space available for other species and enhancing local diversity (Janzen 1970; Connell 1971; Hubbell 1979; Carson et al. 2008). Many studies have found support for density- or distance-dependent growth, survival, or recruitment near conspecific adults (reviewed in Carson et al. 2008; Comita et al. 2014). Yet the ways by which CNDD maintains diversity remain unclear, likely because few studies have explicitly examined the relationship between CNDD and diversity (Harms et al. 2000; Johnson et al. 2012; Bagchi et al. 2014).

CNDD can influence diversity through two non-mutually exclusive mechanisms operating at different levels (Kobe & Vriesendorp 2011; Lin et al. 2012). First, if common species encounter higher local conspecific densities than rare species, CNDD will more strongly limit populations of common species and maintain diversity by allowing abundances of rare species to increase via a ‘community-compensatory trend’ (sensu Connell et al. 1984). Second, the strength of CNDD on a per-neighbour basis may differ among common and rare species due to life-history or other trait differences (Comita et al. 2010; Mangan et al. 2010; Kobe & Vriesendorp 2011). If so, the influence of CNDD on local diversity will depend on how both the strength of per-neighbour CNDD and local conspecific densities vary among species. For example, rare species can often be more spatially aggregated than common species (Hubbell 1979; Condit et al. 2000), yielding similar or higher local conspecific densities for rare species. In this case, stronger per-neighbour CNDD for common than for rare species should still limit common species and maintain local diversity, but stronger per-neighbour CNDD for rare than for common species might limit populations of rare species and decrease local diversity. Yet studies examining the relative strength of per-neighbour CNDD among common and rare species have found mixed results (Comita et al. 2010; Mangan et al. 2010; Johnson et al. 2012; Bagchi et al. 2014; Zhu et al. 2015a). This complexity highlights the need to examine factors that might change the relative strength of CNDD across species and how these changes contribute to community assembly (Paine et al. 2012), species coexistence (Yenni et al. 2012) and gradients of species diversity (HilleRisLambers et al. 2002; Johnson et al. 2012).

Changes in the relative strength of per-neighbour CNDD (hereafter CNDD) among species might depend on underlying environmental conditions. Recent studies suggest that the strength of CNDD in woody-plant species increases with precipitation and productivity at continental to global scales (Johnson et al. 2012; Comita et al. 2014). This pattern might reflect stronger intraspecific competition, increased virulence and/or abundance of host-specific pathogens, or increased pressure from species-specific herbivores in resource-rich environments (Mangan et al. 2010; Bever et al. 2012; Terborgh 2012). These same processes might alter the relative importance of CNDD as a mechanism underlying patterns of species composition and diversity across local resource gradients. However, the extent to which CNDD increases with local
resource availability and contributes to variation in species diversity across local resource gradients remains untested. Moreover, relative increases in the strength of CNDD across local resource gradients may differ among common and rare species, with important implications for diversity. For example, greater increases in the strength of CNDD for rare than for common species across local resource gradients may lead to local extinction of rare species and decrease diversity in resource-rich relative to resource-poor environments. Alternatively, greater increases in the strength of CNDD for common than for rare species across local resource gradients may increase diversity in resource-rich environments by reducing the strength of interspecific competition (Huston 2014). To the extent that the strength of CNDD changes in different ways for common and rare species across local resource gradients, this mechanism could help explain why patterns of local species diversity often show variable responses to changes in productivity and resource availability within and among landscapes (Chase & Leibold 2002; Chalcraft et al. 2008; Adler et al. 2011).

The relative strength of CNDD can also differ among life stages, potentially altering the relative strength of CNDD among common and rare species and influencing species diversity. CNDD can be stronger at earlier than at later life stages (Comita et al. 2014; Zhu et al. 2015b), and processes that structure communities at earlier life stages are generally thought to have a disproportionately strong influence on the maintenance of local diversity (Harms et al. 2000; HilleRisLambers et al. 2002; Comita et al. 2010; Green et al. 2014; Zhu et al. 2015b). However, CNDD at later life stages may contribute more to local patterns of species diversity if CNDD becomes relatively weaker for rare or common species at later life stages (Wright 2002). Thus, changes across life stages in the relative strength of CNDD among common and rare species might impact patterns of diversity, but empirical tests of this idea are lacking.

We examined the strength of CNDD at two life stages (seedling and sapling recruitment) across a soil-resource gradient for 38 woody-plant species in a large (20 ha) stem-mapped temperate forest. We also examined changes in local species diversity across the soil-resource gradient at four life stages (seed, seedling, sapling and adult). To assess whether stronger CNDD could be responsible for observed changes in species diversity across the soil-resource gradient, we predicted seedling and sapling diversity in resource-poor and resource-rich environments using only our estimates of CNDD at both ends of the soil-resource gradient. At both life stages, the strength of CNDD increased with local resource availability. Our results suggest that CNDD is stronger in resource-rich environments and can either decrease or maintain diversity depending on its relative strength among common and rare species.

**MATERIALS AND METHODS**

**Study site and data collection**

Our study was conducted at the Tyson Research Center Forest Dynamics Plot located on the northeastern edge of the Ozark Plateau, 40 km southwest of St. Louis, MO, USA (38° 31’ N, 90° 33’ W). This late-successional oak-hickory-dominated forest has been relatively undisturbed for ~80 years, and tree cores collected in the early 1980s from large individuals of dominant species indicated tree ages of 120–160 years (Zimmerman & Wagner 1979; Hampe 1984). In 2013, we identified, tagged, measured and mapped all free-standing stems of woody species greater than 1 cm diameter at breast height (dbh) in a 20-ha (480 × 420 m) section of a 25-ha plot following CTFS-ForestGEO protocols (Condit 1998).

In 2013, we measured 13 soil variables and four topographic variables across the 20 × 20 m quadrat and the 10 × 10 m subquadrat grid across the 25 ha plot (Fig. 1; Spasojevic et al. 2014). We used principle component analysis to determine the major axis of environmental variation at the 20 × 20 m scale for adults and saplings and at the 10 × 10 m scale for seeds and seedlings. The first principle component (PC1) at both scales described 52.8% and 59.8% of total environmental variation respectively (Fig. S1). As expected, loadings for PC1 were highly correlated between the two scales (r = 0.99), indicating that PC1 described the same environmental axis at both scales. Per cent soil moisture by mass was also collected from 254 sampling locations across the plot in 2013 and was positively correlated with PC1 (r = 0.66). Thus, PC1 described a change from drier acidic soil with high concentrations of iron and aluminium to moist pH-neutral soil with high concentrations of nitrogen, manganese, phosphorus and base cations. Drier acidic soil was associated with southwest-facing slopes and ridges, and moist pH-neutral soil was associated with northeast-facing slopes and valleys (Fig. 1). This topographic gradient is widespread across the Ozark region and linked to changes in slope, aspect and geological transitions from more acidic chert to cherty-limestone to more pH-neutral limestone (Hampe 1984; Nelson 2010).

Seed densities (m⁻² year⁻¹) were calculated from collections at 200 0.5 m² traps stratified across the soil-resource gradient (Fig. 1). Seeds were collected from 140 traps during 2012 and from all 200 traps between 2013 and 2015. We collected and identified potentially viable seeds from March to December each year (six to seven seed collections/year). Seedling densities (m⁻² year⁻¹) were measured during 2014–2015 from 600 1 m² plots. Three seedling plots were paired with each seed trap. We defined seedling as any individual shorter than 50 cm, which includes younger individuals that have retained cotyledons as well as older individuals. Seed traps and paired seedling plots were arranged on the landscape to minimise the correlation between geographic distance and environmental dissimilarity among plots (r = 0.12; Fig. 1).

**CNDD and resource availability**

We examined CNDD at the 10 × 10 m scale for seedling recruitment and at the 20 × 20 m scale for sapling recruitment because CNDD effects are known to decay strongly beyond distances of 10–20 m from a given adult tree in both tropical and temperate forests (Hubbell et al. 2001; Johnson et al. 2014). Saplings were generally defined as trees smaller than 10 cm dbh, but 5 cm dbh or 2 cm dbh were used for small-stature understory species (e.g. Cornus florida or Lindera
the effect of conspecific seed density to vary among species and incorporated random interactions that allowed the interaction between conspecific seed density and the resource gradient to vary among species. These random effects therefore estimated the strength of CNDD for seedling recruitment (random slopes) and changes in the strength of CNDD across the soil-resource gradient (random interactions) for each species. Seedling plots were excluded from analyses when paired seed traps lacked conspecific seed, as including these can bias estimates of density dependence (Harms et al. 2000). We evaluated spatial auto-correlation in our model with variograms that test for patterns in residual semi-variance with increasing distance.

We used a similar approach to examine CNDD at the sapling stage. Because adults are hypothesised to have strong negative density-dependent effects on saplings (Janzen 1970; Connell 1971), we used conspecific adult density to estimate CNDD at the sapling stage across the entire 20 ha plot (Johnson et al. 2012). While adult density in the 10 × 10 m sub-quadrats that contained seed traps was positively correlated with conspecific seed density (r = 0.21, P < 0.001), we note that this analysis encompasses all processes acting on sapling recruitment between the adult and sapling stages (e.g. seed production, pre-dispersal seed predation, seedling survival). CNDD was measured as the slope between log-transformed sapling density and log-transformed conspecific adult density (Fig. S2), and we also included log-transformed heterospecific sapling density and heterospecific adult basal area in our model to test for effects of HNDD on sapling recruitment. Interactions with the soil-resource gradient, random effects and interactions for species, and assessment of spatial auto-correlation were also examined as detailed for seedling recruitment above (Table S1). We also tested for a positive association between the strength of CNDD at seedling and at sapling recruitment.

**CNDD in common and rare species**

We assessed if CNDD was stronger for rare or common species, and if changes in resource availability altered the relative strength of CNDD among species. To assess if CNDD was stronger in rare or common species, we examined regressions between estimates of CNDD and adult size-weighted abundance (basal area summed across the 20 ha plot) across species. These across-species tests were performed for both life stages (seedling and sapling recruitment) at average resource availability (mean of PC1). To assess if CNDD became relatively stronger with resource availability for common or rare species, we examined regressions between the estimates of the change in CNDD with resource availability and adult size-weighted abundance. Size-weighted abundance was used to measure commonness/rarity instead of numerical abundance due to differences in size-age distributions across species and because it better represents the influence that a given tree will have on its surrounding environment (Comita et al. 2010). Nonetheless, numerical abundance and basal area were correlated across species (r = 0.69, P < 0.001). These regressions were weighted by the error around CNDD estimates for each species at each life stage.
CNDD and species diversity

We evaluated if changes in the strength of CNDD during seedling and sapling recruitment were associated with changes in woody-plant species diversity along the soil-resource gradient. We first assessed changes in three measures of species diversity (Shannon’s diversity index, or \(S\); species richness; and rarefied species richness) across the soil-resource gradient at each of four life stages (seeds, seedlings, saplings and adults). We also measured the change in diversity across the resource gradient from the seed to the seedling stage (\(S_{\text{seedling}} - S_{\text{seed}}\)) as well as from the seedling to the sapling stage (\(S_{\text{sapling}} - S_{\text{seedling}}\)). To measure the change in diversity from seedlings to saplings, we only included saplings in the 10 × 10 m subplots that contained seedling plots. For all diversity analyses along the resource gradient, spatial autocorrelation was explicitly controlled using generalised least squares models.

Predicted effects of CNDD on species diversity

To assess whether CNDD could be responsible for observed changes in species diversity across the soil-resource gradient, we predicted seedling and sapling diversity in resource-poor and resource-rich environments using only our estimates of CNDD at both ends of the gradient. For seedling recruitment, we simply multiplied observed seed abundance of each species in each seed trap by the estimate of CNDD for that species in either resource-poor (minimum observed value of PC1) or resource-rich (maximum observed value of PC1) environments. This approach generated two predicted seedling communities under contrasting conditions: (1) if the entire forest plot was resource-poor; and (2) if the entire forest plot was resource-rich. We then compared community-wide species diversity (Shannon’s diversity index) of both communities and determined if CNDD increased or decreased diversity in resource-rich relative to resource-poor environments. We did the same for sapling recruitment, but used observed seedling abundances and estimates of the strength of CNDD during sapling recruitment for each species in either resource-poor or resource-rich environments. Estimates of CNDD controlled for heterospecific effects on seedling and sapling recruitment, and so differences in diversity among these predicted communities should only reflect the influence of CNDD on diversity and not generalised effects of heterospecifics (see supplemental methods for more detailed methods).

RESULTS

In total, 19,776 seeds and 5,715 seedlings from 29 species were used to estimate changes in CNDD during seedling recruitment and seed and seedling diversity across the soil-resource gradient. The top-ranked model for seedling recruitment (\(R^2 = 0.64\)) included effects of conspecific seed density, heterospecific seedling density, the soil-resource gradient, plus interactions of conspecific seed density and heterospecific seedling density with the resource gradient (Table S1, S2). In addition, 15,369 saplings and 14,863 adults from 35 species were used to estimate changes in CNDD during sapling recruitment and sapling and adult diversity across the soil-resource gradient. The top-ranked model for sapling recruitment (\(R^2 = 0.71\)) included effects of conspecific adult density, heterospecific sapling density, heterospecific adult basal area, the soil-resource gradient, plus interactions of conspecific adult density and heterospecific sapling density with the resource gradient (Table S1, S2). We found no strong evidence of spatial autocorrelation in the residuals for either model (Fig. S3), and variograms showed little to no spatial autocorrelation (Fig. S4).

CNDD and resource availability

CNDD during both seedling and sapling recruitment was stronger in resource-rich than in resource-poor environments across species (Fig. 2, 3a, c). CNDD during seedling recruitment was strong for nearly all species (Fig. 2a), whereas CNDD during sapling recruitment was more variable across species (Fig. 2b). Nonetheless, the strength of CNDD during sapling recruitment increased with the strength of CNDD during seedling recruitment among species (\(r = 0.48; P = 0.012\)). The strength of CNDD increased significantly along the soil-resource gradient at both stages after controlling for additive effects of resource availability and potential effects of heterospecific densities (Fig. 2, right panels). HNDD during seedling and sapling recruitment also became stronger along the soil-resource gradient at both life stages (Fig. 3b, d). However, HNDD was non-existent in resource-poor environments and relatively weak compared to CNDD in resource-rich environments (Fig. 3). Moreover, the strength of CNDD increased twice as much along the resource gradient as HNDD (Fig. 3b, d). Thus, the strength of CNDD increased with resource availability and was generally stronger than heterospecific effects. We therefore focus our remaining results on trends in CNDD.

Changes in CNDD across the resource gradient did not appear to be caused by changes in relative densities across the resource gradient. Seed densities increased with resources for seven of 29 species and decreased for one species (mean \(r\) among species = 0.18). Seedling densities increased with resources for five of 29 species and decreased for three species (mean \(r = -0.02\)). At the 20 ha scale, adult densities increased with resources for five of 35 species and decreased for four species (mean \(r = 0.03\)). Sapling densities increased with resources for three of 35 species and decreased for eight species (mean \(r = -0.06\)). Only three of 29 species had higher seed but not higher seedling densities in resource-rich environments, and only three of 35 species had higher adult but not higher sapling densities. This pattern indicates that increases in CNDD were generally not linked to higher initial densities in resource-rich environments.

CNDD in common and rare species

The relative strength of CNDD among common and rare species differed across life stages and changed along the soil-resource gradient at the seedling stage. CNDD during seedling recruitment was equally strong for common and rare species (Fig. 4a), but became relatively stronger for rare species as resources increased (Fig. 4b). In contrast, CNDD during
sapling recruitment was much stronger for common than for rare species across the entire soil-resource gradient (Fig. 4c). The strength of CNDD also increased equally for all species with resource availability at the sapling stage (Fig. 4d), meaning that common species were most suppressed near conspecifics in resource-rich environments. Thus, common species increasingly had a recruitment advantage over rare species as resources increased at the seedling stage, but recruitment of common species was increasingly suppressed as resources increased at the sapling stage.

**CNDD and species diversity**

Different effects of resource availability on the relative strength of CNDD for common and rare species (Fig. 4) corresponded to different effects of resource availability on species diversity across life stages (Fig. 5). At both seedling and sapling life stages, CNDD was generally stronger in resource-rich environments (Fig. 2). However, stronger CNDD corresponded to decreased seedling diversity but increased sapling diversity in resource-rich environments. As resources increased,
stronger seedling CNDD for rare species was associated with declines in seedling diversity relative to seed diversity (Figs. 5, 6a). This trend was also evident at a larger spatial scale when calculating changes in seedling and seed diversity across 12 clustered regions that were stratified across the soil-resource gradient (Fig. 1, S5). In contrast, stronger sapling CNDD for common than for rare species as resources increased was associated with increased sapling diversity relative to seedling diversity (Figs. 5, 6b). Similar patterns were observed for species richness and rarefied species richness (Fig. S6). These patterns indicate that CNDD reduces diversity when it becomes relatively stronger for rare than for common species (as observed for seedling recruitment in resource-rich environments) but maintains diversity when it becomes stronger for common species (as observed for sapling recruitment in resource-rich environments) in this temperate forest.

**Predicted effects of CNDD on species diversity**

Consistent with observed changes in diversity across the soil-resource gradient (Figs. 5 and 6), communities assembled using only observed differences in CNDD across the soil-resource gradient revealed that strong CNDD can both decrease and maintain diversity depending on its relative strength among species (Fig. S7). Predicted seedling diversity was lower in resource-rich environments (Shannon’s diversity index \( \pm SE = 1.49 \pm 0.05 \)) where CNDD became relatively stronger for rare than for common species, than in resource-poor environments (1.77 \( \pm 0.05 \)). In contrast, predicted sapling diversity was higher in resource-rich environments (2.67 \( \pm 0.05 \)) where CNDD became stronger for all species but was relatively stronger for common than for rare species, than in resource-poor environments (2.48 \( \pm 0.06 \)).

**DISCUSSION**

Our results support the hypothesis that the strength of per-neighbour CNDD increases across local resource gradients with implications for species diversity. A previous study of temperate forests at the continental scale (Johnson et al. 2012) and meta-analysis of temperate and tropical forests at the global scale (Comita et al. 2014) found that CNDD becomes stronger in more productive and wetter regions respectively. To our knowledge, our study is the first to demonstrate that...
this interaction between CNDD and resource availability scales down to influence landscape patterns of species diversity within local ecological communities. We found that the strength of CNDD generally increased across a local soil-resource gradient for woody-plant species during two important life-stage transitions (Fig. 2). Our approach contrasts with previous studies of CNDD that examined landscape-level averages without explicitly considering changes in the strength of CNDD across local environmental gradients (Comita et al. 2010; Mangan et al. 2010; Metz et al. 2010; Chen et al. 2010; Johnson et al. 2012; Zhu et al. 2015a). Moreover, increases in the strength of CNDD with soil resources appeared to reduce diversity when CNDD was relatively stronger for rare than common species, but increase diversity when stronger for common species (Figs. 4–6, S7). Overall, these results suggest that CNDD is stronger in resource-rich environments and can either decrease or maintain diversity depending on its relative strength among common and rare species.

Two non-mutually exclusive mechanisms might cause stronger CNDD in more resource-rich environments: (1) stronger host-specific antagonistic interactions or (2) stronger intraspecific competition. Experiments in tropical and temperate forests have shown that negative host–antagonist interactions, specifically species-specific interactions with soil pathogens and plant herbivores, are largely responsible for CNDD effects (Packer & Clay 2000; Mangan et al. 2010; Terborgh 2012; Liu et al. 2012; Bagchi et al. 2014). For example, survival of juvenile black cherry (Prunus serotina), the species in our study with the second strongest CNDD during sapling recruitment (Fig. 2b), decreases near conspecific adults as a result of soil pathogens (Packer & Clay 2000). Microbiologists have predicted that increased moisture or temperature should increase pathogen virulence (Bever et al. 2012), and faster plant growth in more fertile soil may trade-off with weaker pathogen or herbivore defences (Kardol et al. 2006; Fine et al. 2006). Soil properties such as pH have also been linked to changes in bacterial and fungal species diversity and composition (Fierer & Jackson 2006; Barberan et al. 2015), which may affect the strength of plant–soil feedbacks and CNDD. Our results largely support these ideas at a local scale, with stronger CNDD in sites associated with higher soil moisture and pH (Figs. 1 and 2). However, we cannot eliminate the possibility that changes in unmeasured microclimatic variables like temperature or stronger intraspecific competition led to our observation of stronger CNDD in resource-rich environments. Thus, increasingly negative host–antagonist interac-

![Figure 4](https://example.com/figure4.png)

Figure 4 The relative strength of conspecific negative density dependence (CNDD) and changes in the strength of CNDD with resource availability among common and rare species. (a) Strength of CNDD during seedling recruitment as a function of size-weighted abundance, measured as adult basal area (m²). (b) The change in seedling CNDD with a 1 SD change in the soil-resource gradient (PC1) as a function of adult basal area. (c) Strength of CNDD during sapling recruitment as a function of adult basal area. (d) The change in sapling CNDD with a 1 SD change in the soil-resource gradient (PC1) as a function of adult basal area. Note that adult size-weighted abundance is plotted on a log scale.
tions may combine with more intense intraspecific competition to strengthen CNDD in resource-rich environments, but experimental tests are needed to determine the relative importance of these two mechanisms.

We found that stronger per-neighbour CNDD in resource-rich environments can both maintain and decrease diversity depending on its relative strength across species (Figs. 4–6). Specifically, diversity is maintained when CNDD is stronger for common than for rare species (Figs. 4c, 6b, S7), but diversity declines when CNDD becomes stronger for rare species (Figs. 4b, 6a, S7). These findings support the hypothesis that stronger CNDD for common than for rare species maintains diversity, but not via a community-compensatory trend. The traditional compensatory paradigm holds that common species have higher local conspecific densities regardless of their per-neighbour CNDD, limiting populations of common species at the community level (Connell et al. 1984; Wills et al. 2006). Yet rare species can be more spatially aggregated than common species (Hubbell 1979; Condit et al. 2000), yielding similar or higher local conspecific densities for rare species. In our ecosystem, neither median nor maximal local conspecific densities increased systematically with species relative abundance at the seedling (median: \( r = 0.09, P = 0.658 \); maximal: \( r = -0.05, P = 0.809 \)) or sapling stage (median: \( r = 0.18, P = 0.305 \); maximal: \( r = 0.05, P = 0.788 \)). Thus, innate differences in CNDD among common and rare species due to physiological, morphological, immunocompetence or other life-history trait variation may have greater influence on diversity than compensatory mechanisms (Kobe & Vriesendorp 2011).

For example, stronger per-neighbour CNDD for rare species in resource-rich environments during the seedling stage was associated with decreased diversity. Since local conspecific
densities did not differ strongly among common and rare species, this pattern likely reflects increased recruitment limitation and local extinction of rare species as their per-neighbour CNDD increased along the soil-resource gradient. Thus, innate differences among species in per-neighbour CNDD can potentially have greater influence on diversity than community-compensatory mechanisms, maintaining diversity when stronger for common than for rare species but decreasing diversity when stronger for rare species.

Despite declines in species diversity linked to stronger CNDD for rare species during seedling recruitment, stronger CNDD for common species during sapling recruitment increased diversity in resource-rich environments. Moreover, this pattern of higher diversity in resource-rich environments relative to resource-poor environments was maintained into the adult life stage and likely carried over to influence the diversity of seed rain (Fig. 5). These results suggest a limited role for CNDD as a diversifying mechanism at the seed-to-seedling transition and potentially contrast with the idea that processes structuring communities at earlier life stages (e.g. seedling recruitment) have a disproportionately strong influence on species diversity and composition at later life stages (Harms et al. 2000; HilleRisLambers et al. 2002; Comita et al. 2010; Green et al. 2014; Zhu et al. 2015b). However, stronger CNDD for common species during sapling recruitment and associated increases in species diversity may have resulted from other early-life-stage processes (e.g. seed production, pre-dispersal seed predation, seedling survival). Our observation of stronger CNDD for rare species in resource-rich environments may also depend on the years in which we surveyed seedlings. We analysed 2 years of seedling data, but the strength of CNDD (Lin et al. 2012) and the relationship between CNDD and species abundance (Bachelot et al. 2015) can vary temporally. Overall, our results highlight the need for theoretical studies that generate predictions for when changes in the relative strength of CNDD among common and rare species and among different life stages should most strongly influence patterns of diversity.

Like CNDD, HNDD during seedling and sapling recruitment also became stronger in resource-rich environments (Fig. 3). However, heterospecifics generally had weaker influences on seedling and sapling recruitment relative to conspecific (Fig. 3, Table S2). These results are supported by previous studies that show weaker heterospecific relative to conspecific effects on recruitment, survival and growth in temperate and tropical forests (Comita et al. 2010; Johnson et al. 2012, 2014). Stronger HNDD in resource-rich environments may reflect increased interspecific competition and/or increased mortality from generalist herbivores or seed predators (Terborgh 2012). However, recent work supports the idea that generalist enemies or interspecific competition from older age classes may contribute more to HNDD than interspecific competition within a cohort (Wright 2002; Paine et al. 2008; Terborgh 2012; Bever et al. 2015). Thus, stronger HNDD during seedling and sapling recruitment in resource-rich environments may be the result of stronger negative interactions with generalist antagonists or older heterospecifics, although experimental tests are needed to confirm these hypotheses.

Our results have broad implications for understanding how the population-level consequences of negative density dependence scale up to influence community assembly and patterns of species diversity across ecological gradients. Specifically, the relative importance of biotic interactions as a mechanism structuring species diversity and composition likely increases with the availability of local resources. We found that both CNDD and HNDD were weaker in resource-poor than in resource-rich environments, and diversity was lowest in these environments for saplings, adults and seed rain (Figs. 3 and 5). These results suggest that only species with physiological tolerances for low-resource availability can potentially occupy resource-poor environments (Grime 2001; Pianka 2011). On the other hand, biotic interactions appear to be a more important factor determining species diversity, composition and relative abundance in resource-rich environments. Theory already suggests that one type of biotic interaction, interspecific competition, should increase with resource availability, yielding a stronger influence of competition on community assembly in resource-rich environments (Grace 1991; Grime 2001). Yet we found CNDD was much stronger than HNDD in resource-rich environments (Fig. 3), suggesting that negative interactions with host-specific antagonists (pathogens, herbivores or predators), rather than increased competition for resources, may contribute most strongly to changes in the relative importance of community assembly mechanisms across resource-poor and resource-rich environments.

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AUTHORSHIP

JAL and JAM conceived the study; JAM obtained the funding; MLW, JAM and JAL collected the data and BLT
analysed the soil samples. JAL executed the statistical analyses and wrote the first draft of the manuscript, and all authors contributed to revisions.

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Negative density dependence is stronger in resource-rich environments and diversifies communities when stronger for common but not rare species

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SUPPLEMENTAL METHODS

Study site and data collection

Our study was conducted at the Tyson Research Center Forest Dynamics Plot located on the northeastern edge of the Ozark Plateau, 40 km southwest of St. Louis, Missouri, USA (38° 31’ N, 90° 33’ W; mean annual temperature 13.5° C; mean annual precipitation 957 mm). This late-successional oak-hickory-dominated forest has been relatively undisturbed for ~80 years, and tree cores collected in the early 1980s from large individuals of dominant species indicated tree ages of 120-160 years (Zimmerman & Wagner 1979; Hampe 1984; Spasojevic et al. 2014). In 2013, we identified, tagged, measured, and mapped all free-standing stems of woody species greater than 1 cm diameter at breast height (dbh) in a 20-ha (480×420-m) section of a 25-ha plot following CTFS-ForestGEO protocols (Condit 1998).

In 2013, we measured 13 soil variables (exchangeable aluminum, calcium, iron, magnesium, manganese, potassium, and sodium, base saturation, effective cation exchange capacity, KCI-extractable nitrate, nitrate mineralization, pH, Bray-1 extractable phosphorus) and four topographic variables (convexity, mean elevation, northeastern aspect, and slope) across the 20×20-m quadrat and the 10×10-m subquadrat grid across the entire 25-ha plot (Fig. 1; detailed description of soil and topographic calculations provided in Spasojevic et al. 2014). We used principle-components analysis (PCA) to determine the major axis of environmental variation within the plot at the 20×20-m scale for adults and saplings and at the 10×10-m scale for seeds and seedlings. The first principle component (PC1) at both the 20×20-m and the 10×10-m scales
described 52.8% and 59.8% of total environmental variation respectively (Fig. S1). As expected, loadings for PC1 were highly correlated between the two scales ($r = 0.99$; see Fig. S1), indicating that PC1 described the same environmental axis at both scales. Percent soil moisture by mass was also collected from 254 sampling locations across the plot in 2013 and was positively correlated with PC1 ($r = 0.66$). Thus, PC1 described a change from drier acidic soil with high concentrations of iron and aluminum to moister pH-neutral soil with high concentrations of nitrogen, manganese, phosphorus, and base cations. Drier acidic soil was associated with southwest-facing slopes and ridges, and moister pH-neutral soil was associated with northeast-facing slopes and valleys (Fig. 1). This topoedaphic gradient is wide-spread across the Ozark region and linked to changes in slope, aspect, and geological transitions from more acidic chert to cherty-limestone to more pH-neutral limestone (Hampe 1984; Nelson 2010).

Seed densities ($m^2 yr^{-1}$) were calculated from seed collections at 200 0.5-m$^2$ seed traps stratified along the soil-resource gradient within the 20-ha section of the plot (Fig. 1). Seed traps were positioned at the center of 10×10-m sub-quadrats within the 20×20-m quadrats. Seeds were collected from 140 traps during 2012 and from all 200 traps between 2013 and 2015. Seed traps were constructed with fine-mesh (0.3-mm) fiberglass screen attached to square PVC frames elevated one meter off the ground. We collected seeds every month or every-other month from March to December each year (six to seven total seed collections/year). After each seed collection, we counted the total number of potentially-viable seeds of each species in each seed trap. We recorded a seed as viable if it was filled with endosperm and did not have obvious signs of predation. Seedling densities ($m^2 yr^{-1}$) were measured during 2014-2015 from 600 1-m$^2$ plots. Three seedling plots were paired with each seed trap and placed two meters from the seed trap in a random cardinal direction. We defined seedling as any individual shorter than 50 cm,
which includes younger individuals that have retained cotyledons as well as older individuals. Seed traps and paired seedling plots were arranged on the landscape to minimize the correlation between geographic distance and environmental dissimilarity among plots ($r = 0.12$; Fig. 1).

**CNDD and resource availability**

We examined CNDD at the 10×10-m scale for seedling recruitment and at the 20×20 m scale for sapling recruitment because CNDD effects are known to decay strongly beyond distances of 10–20 m from a given adult tree in both tropical and temperate forests (Comita & Hubbell 2009; Johnson et al. 2014). Saplings were generally defined as trees smaller than 10-cm dbh, but 5-cm dbh or 2-cm dbh were used for small-stature understory species (e.g., *Cornus florida* or *Lindera benzoin*) that never or rarely reach 10-cm dbh or 5-cm dbh respectively. Adults were defined as individuals larger than the sapling size class for each species. We tested for increases in the strength of CNDD along the soil-resource gradient using hierarchical mixed models (R package *lme4*; Bates et al. 2014) for each life stage (i.e., seedling and sapling recruitment; see Table S1 for complete model list). Following Harms et al. (2000), we estimated CNDD at the seedling stage as the slope of the line between log-transformed seed density and log-transformed seedling density, with slopes progressively lower than one representing stronger CNDD (Fig. S2). We added one to seed and seedling densities before log-transformation as done in Harms et al. (2000). To examine whether CNDD became stronger with increasing resource availability, we also tested for a significant negative interaction between log-transformed seed density and the soil-resource gradient (PC1). We also tested for additive effects of log-transformed heterospecific seed and log-transformed heterospecific seedling density as well as heterospecific adult basal area in our models to control for potential heterospecific density effects on seedling
density (heterospecific negative density dependence, or HNDD). To control for potentially stronger HNDD with increasing resource availability, we also tested for interactions between the soil-resource gradient (PC1) and all heterospecific variables. Models were compared using sample-size corrected AIC ($AIC_c$; Burnham & Anderson 2002). Data from all woody-plant species shared between seed traps and seedling plots were pooled in our analysis. We incorporated random effects which allowed the effect of conspecific seed density to vary among species and incorporated random interactions which allowed the interaction between conspecific seed density and the resource gradient to vary among species. These random effects therefore estimated the strength of CNDD for seedling recruitment (random slopes) and changes in the strength of CNDD along the soil-resource gradient (random interactions) for each species. We began with a full random effects structure and excluded random intercepts or random slopes for the additive effect of the resource gradient when their estimated variation among species was estimated to be at or near zero and they were correlated ($r > 0.7$) with other random terms that were of theoretical interest to our study (i.e., variation among species in CNDD and changes in the strength of CNDD along the resource gradient). This approach avoids over-parameterization of models while retaining the random terms of interest (Schielzeth & Forstmeier 2009; Barr et al. 2013). Seedling plots were excluded from analyses when paired seed traps lacked conspecific seed, as including these can bias estimates of density dependence (Harms et al. 2000). However, retaining seedling plots that lacked conspecific seed produced species-specific estimates of CNDD that were highly correlated with estimates of CNDD obtained after excluding seedling plots lacked conspecific seed ($r = 0.94, p < 0.0001$). We evaluated spatial auto-correlation in our model with variograms (function `variogram`) that test for patterns in residual semi-variance with increasing distance (R package gstat; Pebesma 2004).
We used a similar approach to examine CNDD at the sapling stage along the soil-resource gradient. Because adults are also hypothesized to have strong negative density-dependent effects on saplings (Janzen 1970; Connell 1971), we used conspecific adult density to estimate CNDD at the sapling stage across the entire 20-ha plot (Johnson et al. 2012). In addition, adult density in the 10×10-m sub-quadrats that contained seed traps was positively correlated with conspecific seed density in those seed traps ($r = 0.21$, $p < 0.001$). CNDD was measured as the slope between log-transformed sapling density and log-transformed conspecific adult density (Fig. S2), and we also included log-transformed heterospecific sapling density and heterospecific adult basal area in our model to test for effects of HNDD on sapling recruitment. Interactions with the soil-resource gradient, random effects and interactions for species, and assessment of spatial auto-correlation were also examined as detailed for seedling recruitment above (Table S1).

In both seedling and sapling recruitment analyses, densities were log-transformed to satisfy assumptions of normal distributions for explanatory variables in linear analyses. In addition, ranges of log-transformed heterospecific densities were similar to ranges of log-transformed conspecific densities in both seedling and sapling recruitment analyses, meaning that differences in range should not influence any observed differences in the relative strength of conspecific and heterospecific density effects in our analyses. We also tested for a positive association between the strength of CNDD at seedling and at sapling recruitment. This regression was weighted by the error around CNDD estimates for each species at each life stage.
CNDD in common and rare species

We assessed if CNDD was stronger for rare or common species, and if changes in resource availability altered the relative strength of CNDD among species. To assess if CNDD was stronger in rare or common species, we examined regressions between estimates of CNDD (calculated from models above; see CNDD and resource availability) and adult size-weighted abundance (basal area summed across the 20-ha plot) across species. These across-species tests were performed for both life stages (seedling and sapling recruitment) at average resource availability (mean of PC1). To assess if CNDD became relatively stronger with resource availability for common or rare species, we examined regressions between the estimates of the change in CNDD with resource availability (CNDD×PC1 interactions from models above; see CNDD and resource availability) and adult size-weighted abundance. Size-weighted abundance was used to measure commonness/rarity instead of numerical abundance due to differences in size-age distributions across species and because it better represents the influence that a given tree will have on its surrounding environment (Comita et al. 2010). Nonetheless, numerical abundance and basal area were correlated across species ($r = 0.69, p < 0.001$). We also considered that relative species abundance may differ along the resource gradient, but basal area measured at the full 20-ha scale was highly correlated with measurements of basal area from only the lower ($r = 0.935, p < 0.0001$) and upper ($r = 0.918, p < 0.0001$) quartiles of the resource gradient. Thus, we proceeded using measurements of basal area at the full 20-ha plot scale. For these analyses, we only included species with precise estimates of CNDD based on two criteria: (1) species sampled from at least a minimal proportion, or 4%, of sample sites (species sampled from at least eight seed trap-seedling plot pairs or from at least twenty 20×20-m quadrats); or (2) species whose estimated error for density dependence were in the upper 75th percentile of
species. However, results were not different if we included all species. These regressions were weighted by the error around CNDD estimates for each species at each life stage.

**CNDD and species diversity**

We evaluated if changes in the strength of CNDD during seedling and sapling recruitment were associated with changes in woody-plant species diversity along the soil-resource gradient. We first assessed changes in three measures of species diversity (Shannon’s diversity index, or $S$; species richness; and rarefied species richness) along the soil-resource gradient at each of four life stages (seeds, seedlings, saplings, and adults). We also measured the change in diversity along the resource gradient from the seed to the seedling stage ($S_{\text{seedling}} - S_{\text{seed}}$) as well as from the seedling to the sapling stage ($S_{\text{sapling}} - S_{\text{seedling}}$). To measure the change in diversity from seedlings to saplings, we only included saplings in the 10×10-m sub-quadrats that contained seedling plots. For all diversity analyses along the resource gradient, spatial auto-correlation was explicitly controlled using generalized least squares models (Gaussian spatial correlation structures using gls in R package nlme; Pinheiro et al. 2015).

**Predicted effects of CNDD on species diversity**

To assess whether CNDD could be responsible for observed changes in species diversity along the soil-resource gradient, we predicted seedling and sapling diversity in resource-poor and resource-rich environments using only our estimates of CNDD at both ends of the soil-resource gradient. For seedling recruitment, we simply multiplied the observed seed abundance of each species in each seed trap by the estimate of CNDD for that species in either resource-poor (minimum observed value of PC1) or resource-rich (maximum observed value of PC1)
environments. This approach generated two predicted seedling communities under contrasting conditions: (1) if the entire forest plot was resource-poor; and (2) if the entire forest plot was resource-rich. We then compared community-wide species diversity (Shannon’s diversity index) of both communities and determined if CNDD increased or decreased diversity in resource-rich relative to resource-poor environments. We did the same for sapling recruitment, but used observed seedling abundances and estimates of the strength of CNDD during sapling recruitment for each species in either resource-poor or resource-rich environments. Estimates of CNDD controlled for effects of heterospecific density on seedling and sapling recruitment, and so differences in diversity among these predicted communities should only reflect the influence of CNDD on diversity and not generalized effects of heterospecifics. All analyses were performed with R 3.2.0 (R Core Team 2015).

LITERATURE CITED


Table S1. Tables showing complete model lists for analyses of conspecific and heterospecific negative density dependence during seedling and sapling recruitment from the Tyson Research Center Forest Plot, St. Louis, Missouri, USA. Models are ranked by AIC<sub>c</sub>, and number of parameters in the model (k) as well as model weight (w) are shown for each. (a) Models for seedling recruitment include conspecific seed (C<sub>seed</sub>), heterospecific seedling (H<sub>seedling</sub>) and seed (H<sub>seed</sub>) density, heterospecific adult basal area (H<sub>adult</sub>), the soil-resource gradient (Env), and interactions with the resource gradient. (b) Models for sapling recruitment include conspecific adult (C<sub>adult</sub>), heterospecific sapling (H<sub>sapling</sub>) density, heterospecific adult basal area (H<sub>adult</sub>), the soil-resource gradient (Env), and interactions with the resource gradient.

(a) Models describing seedling recruitment

<table>
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<th>w</th>
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(b) Models describing sapling recruitment

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Table S2. Parameter estimates, SE, and $p$-values for fixed effects on (a) seedling density (m$^{-1}$ yr$^{-2}$) across 200 paired seedling plots-seed rain traps and on (b) sapling density across 504 20 × 20 m quadrats at the Tyson Research Center Forest Plot, St. Louis, Missouri, USA. Random slopes and interactions that allowed the effect of conspecific seed density (in panel a) and conspecific adult density (in panel b) as well as their interactions with the soil-resource gradient to vary among species were also included (see Fig. 2). All parameter estimates are beta coefficients, and effects must be interpreted in the context of their interactions with other effects. Overall model fit ($R^2$) is given for each model.

(a) Seedling Recruitment (29 species, 200 paired seed traps and seedling plots, $R^2 = 0.64$)

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<td>0.013</td>
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(b) Sapling Recruitment (35 species, 504 20m × 20m quadrats, $R^2 = 0.71$)

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<td>Log(Heterospecific sapling density)</td>
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<td>Soil-resource gradient</td>
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<td>0.005</td>
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Figure S1. Plots showing scores and vectors for first principle component describing correlated environmental variation at the $10 \times 10$ m and $20 \times 20$ m scales from the Tyson Research Center Forest Plot, St. Louis, Missouri, USA. (a) The first principle component at the $10 \times 10$ m scale was used for seedling analyses, and (b) the first principle component at the $20 \times 20$ m scale was used for sapling analyses. Vectors indicate the loadings for each environmental variable.
Figure S2. Plots showing seedling and sapling densities against conspecific seed and adult densities for red maple (*Acer rubrum*) and white oak (*Quercus alba*), respectively, in the Tyson Research Center Forest Plot, St. Louis, Missouri, USA. The strength of conspecific negative density dependence was measured as the slopes of these relationships after controlling for heterospecific density effects, with more negative slopes indicating conspecific negative density dependence (see Table S1, S2). The dashed line indicates a 1:1 relationship, which would be expected in the absence of conspecific negative density dependence. These methods are identical to those in Harms (2000), but we account for potential heterospecific density effects and test for interactions between woody-plant densities and the soil-resource gradient.
Figure S3. Maps showing spatial distribution of residuals from mixed models measuring negative density dependence during (a) seedling and (b) sapling recruitment at the Tyson Research Center Forest Plot, St. Louis, Missouri, USA.
Figure S4. Variograms showing auto-correlation among residuals at increasing distances along four different directional axes (i.e., 90° or east-west, 135° or southeast-northwest, 0° or north-south, and 45° or northeast-southwest) from mixed models measuring negative density dependence during (a) seedling and (b) sapling recruitment at the Tyson Research Center Forest Plot, St. Louis, Missouri, USA.
Figure S5. The change in large-scale seedling diversity relative to large-scale seed diversity at the Tyson Research Center Forest Dynamics Plot. We pooled seedlings and seeds across all traps within 12 clustered regions that were stratified along the soil-resource gradient across the forest plot (see Fig. 1). Like diversity at the smaller scale of the seed traps and seedling plots, diversity at this much larger scale declined in resource-rich environments relative to resource-poor environments during the seed-to-seedling transition. This decline in diversity was associated with stronger CNDD for rare than for common species as resource availability increased. Correlation coefficients, regression lines, and significance tests account for spatial auto-correlation among samples.

![Seed-to-seedling graph](image)

- $r = -0.63$
- $p = 0.036$
Figure S6. Changes in species richness for (a) seeds, (b) seedlings, (c) saplings, (d) and adults along the soil-resource gradient. Changes in rarefied richness along the soil-resource gradient were similar for the seed ($r = 0.22; p = 0.007$), seedling ($r = 0.13; p = 0.20$), sapling ($r = 0.16; p = 0.012$), and adult ($r = 0.26; p < 0.001$) life stages.
Figure S7. (a) Seedling and (b) sapling communities (species-abundance distributions) predicted from observed seed and seedling communities, respectively, using only estimated species-specific changes in negative density-dependent recruitment along the soil-resource gradient (see Fig. 2) at the Tyson Research Center Forest Plot, St. Louis, Missouri, USA. Shannon’s diversity index (± SE) (a) increased in resource-poor environments relative to resource-rich environments during the seed-to-seedling transition but (b) increased in resource-rich environments relative to resource-poor environments during the seedling-to-sapling transition.