Nitrogen and phosphorus additions negatively affect tree species diversity in tropical forest regrowth trajectories

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Abstract. Nutrient enrichment is increasingly affecting many tropical ecosystems, but there is no information on how this affects tree biodiversity. To examine dynamics in vegetation structure and tree species biomass and diversity, we annually remeasured tree species before and for six years after repeated additions of nitrogen (N) and phosphorus (P) in permanent plots of abandoned pasture in Amazonia. Nitrogen and, to a lesser extent, phosphorus addition shifted growth among woody species. Nitrogen stimulated growth of two common pioneer tree species and one common tree species adaptable to both high- and low-light environments, while P stimulated growth only of the dominant pioneer tree Rollinia exsucca (Annonaceae). Overall, N or P addition reduced tree assemblage evenness and delayed tree species accrual over time, likely due to competitive monopolization of other resources by the few tree species responding to nutrient enrichment with enhanced establishment and/or growth rates. Absolute tree growth rates were elevated for two years after nutrient addition. However, nutrient-induced shifts in relative tree species growth and reduced assemblage evenness persisted for more than three years after nutrient addition, favoring two nutrient-responsive pioneers and one early-secondary tree species. Surprisingly, N + P effects on tree biomass and species diversity were consistently weaker than N-only and P-only effects, because grass biomass increased dramatically in response to N + P addition. The resulting intensified competition probably prevented an expected positive N + P synergy in the tree assemblage. Thus, N or P enrichment may favor unknown tree functional response types, reduce the diversity of coexisting species, and delay species accrual during structurally and functionally complex tropical forest secondary succession.

Key words: biodiversity loss; clay Oxisol; ecosystem fertilization; mixed-effects models; N and P co-limitation; N + P synergism; niche dimensionality; spatial heterogeneity; tree–grass interactions; tree species richness; tropical moist forest recovery; vegetation development.

INTRODUCTION

Human food production and energy use have increased nitrogen and phosphorus fluxes in ecosystems by up to 10-fold in some regions, while other regions are experiencing nutrient impoverishment (Falkowski et al. 2000, Howarth et al. 2005). Both nutrient-enriched and impoverished regions occur in the tropics near or at hotspots of global biodiversity (Howarth et al. 2005). Enrichment with the nutrients limiting primary productivity plays an important role in biodiversity loss in temperate ecosystems (Wassen et al. 2005, Clark et al. 2007). In biodiversity-rich, functionally complex, and highly dynamic ecosystems such as tropical secondary forests the relationship between biodiversity and nutrient limitation remains unknown.

Low soil P availability limits primary productivity on the widespread, deeply weathered lowland tropical Oxisols (Vitousek and Sanford 1986, Elser et al. 2007). However, after land use change the resulting ecosystems may become limited by nitrogen (Davidson et al. 2007). Deforestation, subsequent land use, and repeated burning can cause large losses through volatilization of N and through ash transport, leaching, and erosion of both N and P (Kaufman et al. 1995, Williams and Melack 1997). Nutrient impoverishment may thus slow post-abandonment biomass accumulation during secondary succession (Gehring et al. 1999, Davidson et al. 2004) and, thereby, accrual of mature forest species, which are most at risk among tropical forest species biodiversity (Martinez-Garza and Howe 2003, Chazdon 2008).

By contrast, enrichment with limiting nutrients accelerates biomass accumulation and might thereby
accelerate secondary succession and forest recovery. However, enrichment with limiting nutrients in relatively simple ecosystems has widely been reported to cause species loss in herbaceous, shrub, epiphyte, and fungal assemblages, impairing biodiversity conservation and perhaps ecosystem function, although evidence is almost entirely restricted to temperate ecosystems (Bobbink et al. 1998, 2010, Fynn and O’Connor 2005, Wassen et al. 2005, Clark et al. 2007). Conversely, P fertilization in tropical montane forest dramatically increased abundance and diversity of epiphytes across functional groups without concomitant species loss (Benner and Vitousek 2007). In N-fertilized tropical montane forest, Ostertag and Verville (2002) found lower understory plant species richness than in controls, but pre-fertilization data were unavailable. We did not find reports of nutrient enrichment affecting trajectories of tree assemblage composition or diversity over time in any biome.

Mechanisms for N-induced species loss may include reduced heterogeneity of soil nutrient availability (Gilliam 2006) and soil acidification (Bobbink et al. 1998, Fynn and O’Connor 2005, Clark et al. 2007). Competition for other nutrients or light may also be intensified, potentially favoring species efficient at their acquisition and use (Herbert et al. 2004), thereby increasing local extinction of and/or reducing colonization by other species (Wilson and Tilman 2002), for instance those adapted to low nutrient availability (Bobbink et al. 1998). Similarly, P enrichment may cause competitive exclusion of species, although species responses to N and P addition may vary due to fundamental differences in plant strategies for N and P acquisition (Lambers et al. 2008).

Theory predicts that simultaneous enrichment with co-limiting nutrients reduces the number of simultaneously limiting resources, thereby reducing niche dimensions among which species may coexist (Harpole and Tilman 2007). Thus, it may be hypothesized that simultaneous enrichment with N and P in tropical secondary forests causes greater biomass response and greater loss in tree species diversity than single-nutrient enrichment (Tanner et al. 1998, Lawrence 2003).

In tropical secondary forests, species favored by nutrient enrichment may be fast-growing pioneers with high photosynthetic rates (Ellsworth and Reich 1996, Lawrence 2003). The resulting potential delays in species replacement during succession may hinder the recovery of large-seeded mature forest species by prolonging the persistence of early-successional tree species (Mesquita et al. 2001, Martinez-Garza and Howe 2003) in tropical landscapes increasingly dominated by secondary vegetation (Chazdon 2008).

The short-term nature of previous nutrient addition experiments in tropical secondary forests (Harcombe 1977, Gehring et al. 1999, Davidson et al. 2004) precludes interpretation of nutrient effects on the dynamics of composition or diversity. The high taxonomic, structural, and largely unknown functional diversity of tropical secondary forests (Chazdon 2008) may limit the applicability of mechanistic explanations demonstrated for nutrient enrichment affecting temperate plant communities. Furthermore, highly differentiated strategies of nutrient acquisition and use (Lambers et al. 2008) complicate the relationship between ecosystem nutrient limitation and nutrient-induced shifts in community composition and diversity (Tanner et al. 1998). We hypothesized that nutrient enrichment negatively affects tree species diversity by disproportionately promoting few, nutrient-responsive pioneer species.

To assess how N and P enrichment affects tree community dynamics, we related trajectories of vegetation structure, tree species diversity, and evenness with absolute shifts in species woody biomass estimates for six years, in response to factorial N and P addition in lowland Amazonia. We chose a well-studied pasture site, abandoned after prolonged use and frequent fires on naturally infertile, deeply weathered clay Oxisol.

METHODS

Site description

The study was conducted at Fazenda Vitória, Paragominas, Pará, Brazilian Amazonia (2°59’ S, 47°31’ W), where many studies have been completed on the ecology of abandoned pastures and forests (e.g., Uhl et al. 1988, Nepstad et al. 1994, 1996, Vieira et al. 1994, Verchot et al. 1999, Davidson et al. 2004, 2007, Markewitz et al. 2004). Annual rainfall of 1800 mm is highly seasonal, i.e., <100 mm monthly for six months and 35 mm monthly during the driest consecutive three months of July–September (Jipp et al. 1998). Mature and secondary forests in the region remain evergreen, despite the pronounced dry season, by extracting water from deep subsoil layers (Nepstad et al. 1994), although some early-successional species shed a substantial portion of their foliage during the dry season (Davidson et al. 2004).

The Kaolinitic Yellow Latosol soils (USDA: Haplustox) in the region have developed on Belterra clay and Tertiary Barreiras formations (Sombroek 1966). These deeply weathered sediments consist mainly of kaolinite, quartz, and hematite, with 60–80% clay and <9% sand content at the study site. Pasture soil pH in water decreases with depth from 5.7 at the surface. Most actively managed pastures in this region have been fertilized with P starting in the late 1980s and early 1990s (Dias-Filho et al. 2001). However, the degraded cattle pasture used in this study was never fertilized prior to the experiment and was abandoned before fertilization became common. Soil solution N fluxes and soil available P in abandoned pastures and in a 19-year-old secondary forest in the region were very low compared to mature forest. (For detailed nutrient budgets of these land use systems within a few kilometers of our plots, see Markewitz et al. [2004].)
The study was undertaken in 7.25 ha of abandoned pasture that was originally planted in 1971 with the grass *Panicum maximum* and abandoned in 1984. The pasture was grazed by cattle in a rotation regime, with 1 animal/ha, and was periodically burned and then sprayed with herbicide. Accidental fires have burned the area at least three times since abandonment, the last time being 1993. We started the experiment six years after the last fire, when forbs, shrubs, and trees dominated and grass cover had diminished.

**Nutrient addition: experimental design**

Six years after the last fire (i.e., 15 years after pasture was abandoned), four plots (20 × 20 m) were established in each of three replicate blocks in a randomized complete block design. The four plots within each block received the following treatments at the beginning of two consecutive rainy seasons (January 2000 and February 2001): “C” (control), “N” (100 kg N/ha as urea), “P” (50 kg P/ha as simple superphosphate), and “N + P” at these same rates. Fertilizers were broadcast by hand in each of four 10 × 10 m subplots per fertilized 20 × 20 m plot to homogenize distribution. The full factorial design is composed of two crossed binary factors of nutrient addition (“-N” vs. “+N” and “-P” vs. “+P”), with n = 6 plots, i.e., a combined area of 0.24 ha in each level for the main effects. At the end of the dry season before the first nutrient addition, the baseline census was carried out. Taxonomic identity at the species level, height, and diameter at breast height (dbh, at 1.3 m above ground level) of all trees ≥2 cm dbh were measured and permanently marked with individual ID numbers at breast height. Height and dbh of all individual trees, including recruits into the plots nested within blocks, using the “nlme” package, version 3.1-83 in R 2.6.0 (Pinheiro et al. 2007). We compared the simplest linear model with ecologically meaningful alternatives by including quadratic terms for time and covariates (i.e., density of individuals or initial condition of response variable prior to nutrient addition). Subsequently, we selected the simplest model of best fit that (1) satisfied assumptions of independence, normality, and homogeneity of variances and that (2) had the lowest Akaike’s information criterion (AIC). The detailed statistical procedure is reported in Appendix A.

We report effects of nutrient addition where we observed a significant effect on slopes of the response variable, i.e., where interactions were significant between factors of nutrient addition (N or P) and time, or squared time (time2) in second-order polynomial models (e.g., N × time, P × time, N × P × time, N × time2, etc.). This interpretation of interactions with time (e.g., N × time) is valid, because the central assumption was satisfied that none of the main effects of factors of nutrient addition (i.e., N, P, N × P) were significant. For reported significant model terms, we give t values of Wald tests (in parentheses), subscripted by the corresponding degrees of freedom (df), followed by the significance level (fitted equations may be found in Appendices A, B, and C). All graphs were produced using the “graphics” package in R 2.6.0 (R Development Core Team 2007). All diversity calculations were performed using the “vegan” package, version 1.8-8 in R 2.6.0 (Oksanen et al. 2007).

**Results**

Initial variation among blocks (and plots) exceeded (fixed) effects of nutrient addition. Therefore, initial variation among blocks and plots is modeled here as random effects, based on repeated measures of all plots. This approach enabled testing for nutrient effects on slopes (rather than means) of response variables as they varied with time, thereby allowing visualization of changes in tree density and woody biomass, grass biomass, tree species richness, and evenness, as well as absolute shifts in growth rates among tree species.

**Tree density and biomass and grass biomass**

Between 6 and 12 years after the last stand-clearing fire (i.e., 15–21 years since pasture abandonment), total densities of individual trees generally increased over time, but varied widely among plots (Fig. 1a). Net
recruitment of individuals leveled off at approximately year 11 in half of all plots, but this occurred at densities ranging from 1300 to 3100 trees/ha. Model selection resulted in the second-order polynomial fit without covariates. Consistently greater slopes in plots amended with N (N × time, $t_{10} = 2.65, P < 0.01$) revealed higher rates of accrual of individuals in response to N addition (Fig. 1a), but this effect diminished over time (N ×
time², \( t_{64} = -2.24, P < 0.05 \)), while accrual of individuals also diminished in plots amended with P (P × time², \( t_{64} = -2.18, P < 0.05 \)).

Between 6 and 12 years after the last fire, total tree aboveground woody biomass estimates showed, on average, a fivefold increase (Fig. 1b). To permit comparison with other studies on fertilization effects, we present the simplest linear model of total woody biomass with initial biomass as covariate. To this end we distinguish separate models for: (1) short-term linear responses within two wet seasons of last nutrient addition, on which other published studies report (Gehring et al. 1999), and (2) subsequent midterm trajectories from three wet seasons after last nutrient addition, for which we are unaware of any published comparison. The short-term fit indicated positive effects of N addition (N × time, \( t_{28} = 3.14, P < 0.01 \)), i.e., after accounting for the effects of initial biomass on subsequent trajectories, N addition resulted on average in faster total woody biomass growth within two wet seasons of last nutrient addition (Fig. 1b, short-term). However, an equivalent linear fit of the subsequent midterm trajectories revealed no difference due to N addition (P > 0.1; Fig. 1b, midterm). By contrast, P addition did not significantly increase the rates of short-term biomass accumulation (P × time, P > 0.1), possibly partly due to high variation among +P plots.

Total aboveground grass biomass was also modeled as separate short-term and midterm linear fits (Fig. 1c; cases with missing values omitted). Within 1.5 years of first nutrient addition, grass biomass increased fivefold in N + P plots (N × P × time, \( t_{30} = 3.57, P < 0.01 \)). Within the same period, grass biomass tended to double after P-only addition (P × time, P > 0.1) and was unaffected by N-only addition.

Three wet seasons after last nutrient addition, grass biomass in +P plots was several-fold higher than at baseline census, though highly variable (Fig. 1c), and subsequently collapsed to baseline levels. Model selection for the midterm period included covariates of grass biomass at year 9.7 (grass0.7), and a variance power covariate, which accounted for heteroscedasticity observed in the midterm (Fig. 1c). The rate of midterm grass biomass decline was primarily predicted by its initial value (grass0.7 × time, \( t_{18} = -4.41, P < 0.001 \)), without significant effects of nutrient addition. This sudden collapse of grass biomass coincided with a significant decline over time in the negative N × P interactions of woody biomass of both Zanthoxylum rhoifolium Lam. (Rutaceae) and Bania guianensis Aubl. (Flacourtiaceae) and of rarefied species richness (N × P × time², \( t_{52} = 2.62, P < 0.05 \); \( t_{52} = 2.57, P < 0.05 \); and \( t_{63} = -2.09, P < 0.05 \), in Figs. 3b, 3c, and 2a, respectively).

**Tree species richness and evenness**

The observed tree species richness increased from 36 to 54 in the total pooled 0.48 ha of experimental plots between 6 and 12 years after the last fire. We accounted for the observed high variation in numbers of individual trees per plot (cf. Fig. 1a) by individual-based rarefaction curves (not shown; rationale in Appendix B). The resulting rarefied species richness represents the number of species in a random subsample of 13 individuals (\( R_{13} \)). Model selection for \( R_{13} \) led to a second-order polynomial fit with initial \( R_{13} \) as covariate (Fig. 2a). Both N and P addition had a negative effect of similar size on increases in \( R_{13} \) over time (N × time, \( t_{63} = -2.89, P < 0.01 \); P × time, \( t_{63} = -3.14, P < 0.01 \)). However, effect sizes diminished significantly over time (N × time², \( t_{63} = 2.34, P < 0.05 \); P × time², \( t_{63} = 3.12, P < 0.01 \)). In N + P plots the negative effect of nutrient addition was weaker than the sum of main effects (N × P × time, \( t_{63} = 2.11, P < 0.05 \)).

Model selection for the other component of tree species diversity, measured as the Simpson evenness measure (\( E_{1/D} \)), resulted in a second-order polynomial fit with density of individuals as covariate (Fig. 2b). Nitrogen addition had a greater, negative effect on \( E_{1/D} \) trajectories (N × time, \( t_{52} = -3.27, P < 0.01 \)) than P addition (P × time, \( t_{52} = -2.31, P < 0.05 \)). Similar to \( R_{13} \), less-than-additive N × P interactions were significant (N × P × time, \( t_{52} = 2.33, P < 0.05 \)). However, there was no evidence of a diminishing effect size over time (both N × time² and P × time², P > 0.1).

The Simpson diversity index (1 − D) showed a delaying effect of P addition on tree species diversity increases over time (second-order polynomial fit; covariate initial 1 − D; P × time, \( t_{52} = -2.89, P < 0.01 \); P × time², \( t_{52} = 2.59, P < 0.05 \); Fig. 2c). This matched the response of \( R_{13} \) to P addition, but was nonsignificant for N addition.

**Species woody biomass**

Fitting mixed models to individual species’ aboveground woody biomass estimates revealed important interspecific differences in nutrient responses. The dominant tree species, Rollinia exsucca A.DC. (Annonaceae), accounted for 64% of initial total woody biomass and was the only species to respond to both N and P additions (N × time, \( t_{52} = 3.77, P < 0.001 \); P × time, \( t_{52} = 3.93, P < 0.001 \); Fig. 3a), but with less-than-additive effects of N + P (N × P × time, \( t_{52} = -2.42, P < 0.05 \)). Negative interactions of both N and P with the quadratic term of time (N × time², \( t_{52} = -2.46, P < 0.05 \); P × time², \( t_{52} = -2.14, P < 0.05 \)) indicated diminishing effect sizes for both nutrients on woody biomass of R. exsucca. Overall, R. exsucca reached a biomass plateau by 12 years after the last fire.

Codominant Zanthoxylum rhoifolium and initially rare Bania guianensis contributed 12% and 0.8% of total initial woody biomass, respectively. Both species responded positively to N addition (Z. rhoifolium, N × time, \( t_{52} = 3.04, P < 0.01 \); B. guianensis, N × time, \( t_{52} = 3.82, P < 0.001 \); in Figs. 3b and 3c, respectively), but
also showed diminishing N responses over time (Z. rhoifolium, $N \times time^2$, $t_{52} = -3.28, P < 0.05$; B. guianensis, $N \times time^2$, $t_{52} = -3.76, P < 0.001$).

Furthermore, both species also exhibited a less-than-additive effect of $N \times P$ (Z. rhoifolium, $N \times P \times time$, $t_{52} = 2.48, P < 0.05$; B. guianensis, $N \times P \times time$, $t_{52} = 3.82, P < 0.001$).

All other 55 observed tree species either showed no significant nutrient response or had abundances too low across treatments and censuses to permit univariate fits.
without violating model assumptions. The most abundant nonresponsive species was *Vismia guianensis* (Aubl.) Choisy (Clusiaceae). Pooled biomass increments of these 55 species generally increased over time, regardless of nutrient addition (N x time, P x time, and N x P x time, P > 0.1; Fig. 3d).

**DISCUSSION**

We observed high inter-block and inter-plot variation, presumably attributable to spatial variation in soil conditions and in seed dispersal, as well as to plot size. Based on the pre-fertilization baseline census, our mixed-model analysis was able to account for this spatial heterogeneity among blocks and plots, which was independent of treatment application. This enabled us to detect the effects of nutrient addition on the rate of change (i.e., slope) and the duration of these effects (based on changes in slope over time) of tree species biomass and diversity. Regular remeasurement of floristic attributes, over several years beyond the previously reported short-term nutrient enrichment responses, revealed short-lived effects of the original nutrient additions on woody biomass accumulation. However, nutrient enrichment effects on assemblage evenness persisted for more than three years.

The scope to generalize from these findings is limited, because this study intentionally focused on a single region to control extraneous variables known to affect successional trajectories, such as climate, soil type, land use history, and the landscape matrix (Chazdon 2003). This enabled sensitive direct detection of treatment response trajectories and consistent negative N x P synergies associated with tree-grass competition. Assessment of the generality of these findings across regions and ecosystem types will only be possible once other long-term field experiments become available.

**Patchy forest regrowth after lag phase**

Pasture use was abandoned 15 years prior to our baseline census. However, for the first nine years, repeated accidental fires kept reverting tree woody biomass accumulation back to near 0 Mg ha\(^{-1}\) yr\(^{-1}\), although fire intensity may have been spatially variable. During the first six years of postfire regrowth, total tree biomass accumulated slowly at a rate of <1 Mg ha\(^{-1}\) yr\(^{-1}\) (Davidson et al. 2004), which is comparable to other degraded sites after moderate to heavy pasture use in the region (Uhl et al. 1988). Similar post-abandonment lag phases with negligible biomass accumulation elsewhere in Amazonia have been attributed to intensity and/or duration of prior land use (Feldpausch et al. 2007).

Between 6 and 12 years postfire, total woody biomass accumulation averaged ~4 Mg ha\(^{-1}\) yr\(^{-1}\), with high inter-plot variation, regardless of nutrient enrichment (Fig. 1b). This rate is at the far low end of the range reported for secondary forest biomass accumulation on non-sandy soils in Amazonia (Houghton et al. 2000, Zarin et al. 2001).

**Short-lived growth responses to nutrient addition**

The N-induced increase in tree biomass was due to both increased net tree recruitment into the ≥2 cm dbh size class and to growth increments of larger trees of the two dominant species (*Rollinia exsucca* and *Z. rhoifolium*). Surprisingly, mean rates of total woody biomass accumulation (i.e., slopes in Fig. 1b) in control plots (no nutrients added) caught up with those in +N plots within five years following the last nutrient addition, despite the large amounts of N initially added. Davidson et al. (2004) reported that only 10% of the applied P and 20% of the applied N was incorporated into tree biomass during the first growing season after the first nutrient addition event. Apparently, this young secondary forest did not efficiently acquire and recycle enough of the added nutrients in bioavailable forms to maintain longer-term enhanced biomass accumulation relative to regrowth in the control plots.

Davidson et al. (2004) could not quantify the fate of most of the added N earlier during this experiment, because soil stocks of total N are large relative to the N amendments. Rapid immobilization of added nutrients in the soil was inferred from the strong net immobilization capacity and a highly conservative N cycle in these plots (Davidson et al. 2004) and in other secondary forests in the region (Verchot et al. 1999, Davidson et al. 2007). Immobilization of nutrients into soil pools unavailable to tree roots may be responsible for the short duration of the nutrient response. For instance, Nepstad et al. (1996) observed responses to added nutrients in planted trees at the same site only after soil tilling, which indicates that this compacted clay soil limits access by tree roots to added nutrients. Thus, in the present study, it appears that most of the added N had become immobilized into nonavailable pools of organic matter within a few years of addition, so that the initial nutrient response was not sustained.

These earlier studies at the same site support the notion that biomass accumulation reverted to be limited by N within a few years of our two pulse nutrient additions. Indeed, chronosequence studies suggest that N limitation in young secondary forests slowly declines relative to P limitation over several decades, as forests accumulate actively cycling N (Davidson et al. 2007). However, tree biomass accumulation of individual species may also be co-limited by other factors (Tanner et al. 1998). This was supported by increases in leaf P concentrations of *Z. rhoifolium* and leaf N of both dominant tree species induced by nutrient addition earlier in the present experiment (Davidson et al. 2004) and in other tree species (Gehring et al. 1999).

**Shifts in tree species biomass**

Responses of tree species to added nutrients contrast with findings by Gehring et al. (1999), 200 km north of
our site. For instance, at their sandy site *R. exsucca* did not respond to N or P addition, but *V. guianensis* was primarily N-limited, and *B. guianensis* responded vigorously not only to N, but also P. These differences relate to species that all occurred at sufficiently high abundances at our site to allow detection of treatment effects. Possible explanations for the contrasting results may relate to immediate nutrient addition after fire, high sand content (Zarin et al. 2001), and/or arable cropping history (Moran et al. 2000) at the site studied by Gehring et al. (1999). Given such variation among sites, the results cannot be generalized at the species’ response level. By contrast, at the community level, we speculate that further research could demonstrate that nutrient enrichment may commonly delay species accrual in secondary forests, as observed in the present study.

**Negative effects on tree species diversity**

Consistent negative effects of nutrient addition on all tree species diversity measures examined provide strong evidence that both N-only and P-only addition delayed increases in tree species richness and reduced evenness. The duration of these effects coincided with the positive biomass responses of three common tree species. This suggests that in tropical secondary forests, enrichment with a (co-)limiting nutrient may increase dominance of species that are capable of monopolizing other resources limiting to coexisting tree species. To our knowledge, such effects have so far only been demonstrated in relatively simple ecosystems and have been predicted from theory and from simulations (Fahey et al. 1998, Wilson and Tilman 2002, Herbert et al. 2004).
Nitrogen limitation of primary productivity is an important predictor of plant species loss following N enrichment, partly due to intensified competitive exclusion (Clark et al. 2007). By contrast, P enrichment did not significantly stimulate biomass growth of the pooled tree assemblage, yet had a disproportionately large negative impact on tree species diversity. This demonstrates that P enrichment might reduce species diversity without measurably increasing overall productivity, possibly by directly inhibiting some of the diverse range of species adapted to mobilizing P from unavailable pools in soils with low P availability (Lambers et al. 2008).

However, we found no obvious functional grouping among responsive species, as successional characteristics, leaf nutrient concentrations, and specific leaf areas varied within responsive species, as well as within nonresponsive species (data not shown). Further research is needed to identify ecophysiological characteristics and functional traits differentiating species favored from species disadvantaged by nutrient enrichment and underlying competitive mechanisms responsible for tropical species loss. Such understanding may help predict and ultimately prevent prolonged persistence of pioneers delaying colonization by mature forest species (Mesquita et al. 2001, Martínez-Garza and Howe 2003).

**Negative $N \times P$ interactions**

Surprisingly in this context, $N + P$ addition had a consistently weaker effect than $N$ only and $P$ only, both on tree species biomass and diversity measures. In contrast, a strong positive $N \times P$ interaction in enhancing biomass accumulation has been widely demonstrated in aquatic and terrestrial ecosystems (Elser et al. 2007).

The timing of our observed negative $N \times P$ interaction in tree species biomass and diversity responses coincided with a dramatic initial $N + P$ response by remnant grass biomass, as well as grass leaf P concentrations (Davidson et al. 2004). Simultaneous to $N$ and $P$ effects on tree biomass, grass biomass responses were also reversed within five years and were not significant for $P$-only or $N$-only additions (Fig. 1c). Herbs, shrubs, and vines have previously been reported to effectively compete with tropical secondary forest trees for added nutrients,

![Graph showing N-responsive tree species](image1.png)

**Fig. 3.** Continued.
especially P, potentially retarding tree biomass accumulation (Harcombe 1977, Gehring et al. 1999). In contrast to other studies, our N + P addition markedly increased the grass:tree biomass ratio for several years, even though trees had already successfully competed with grasses for the previous six years postfire. We did not determine herbaceous species composition and thus cannot rule out a positive N × P interaction in grass and/or forb species loss, as has been demonstrated in grasslands (Fynn and O’Connor 2005, Harpole and Tilman 2007). However, our data highlight that synergistic effects of combined N + P enrichment on grass biomass responses may prevent synergistic N + P responses in tree biomass and tree species diversity. Such interactions could have important implications for ecological dynamics and biodiversity, both in regenerating forests, as well as in savannas after nutrient enrichment, for instance by intensive agriculture.

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Literature Cited


APPENDIX A

Univariate statistical procedure and detailed responses of vegetation structure (Ecological Archives E091-147-A1).

APPENDIX B

Quantification and detailed responses of tree species diversity (Ecological Archives E091-147-A2).

APPENDIX C

Detailed responses of absolute tree species woody biomass (Ecological Archives E091-147-A3).