30 Years of Post-disturbance Recruitment in a Neotropical Forest

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Abstract

The role of tree diversity for tropical forests functioning and services makes it crucial tree diversity and composition fate in the global changing context. Community long-term response to disturbance rely on tree recruitment, long seen as following deterministic successional pathways. These pathways however might be altered in the hyper-diverse tropical forests and of slight but recurrent disturbances induced by global changes. Post-disturbance recruitment trajectories would (i) disentangle the determinants of tree recruitment between stochastic and deterministic processes that enhance a restricted pool of species, and (ii) elucidate tropical forests taxonomic and functional resilience. We examined the trajectories over 30 years of recruited trees taxonomic and functional diversity in 12 plots of 6.25 ha of forest following a disturbance gradient. We analyzed taxonomic richness, evenness, and turnover, and functional diversity and composition (regarding 7 leaf, stem and life-history functional traits). We revealed a three-phased successional pathway defined by the interplay of stochastic and deterministic recruitment processes. The succession translated into (i) saplings growth maintaining pre-disturbance community diversity and composition, (ii) light-demanding species enhanced recruitment entailing, above a disturbance intensity threshold, the dominance of pioneers and (iii) the recovery of pre-disturbance taxonomic and functional characteristics and of stochastic recruitment processes. Although tangible, community taxonomic and functional resilience was decades-long. Post-disturbance recruitment relied on deterministic competition processes for light balancing the stochastic processes ruling undisturbed communities. Although resilient, recruitment taxonomic and functional characteristics remained altered in the long-term, calling caution for forest management.

Keywords

Disturbance Dynamics, Neotropical Forests, Recruitment, Resilience, Taxonomic and Functional Diversity, Tree Community

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1. Introduction

Determining the response of tropical forests to disturbance is key to predict their fate in the global changing context. In the last decades, tropical forests experienced a wide range of disturbance, from radical land-use changes for agriculture or mining (Dezécache et al., 2017a,b) to more insidious changes following climatic changes or anthropogenic activities like selective logging (Baraloto et al., 2012; Aubry-Kientz et al., 2015). In that respect a vast literature successfully modeled community response to disturbance in terms of tree growth, tree height and fluxes of carbon, water, and nutrients (Gourlet-Fleury & Houllier, 2000; Putz et al., 2012; Piponiot et al., 2016; Rutishauser et al., 2016). In terms of tree community diversity and composition, the ecological theory of succession assumes post-disturbance pathways driven by a succession of deterministic recruitment processes. These pathways would track the post-disturbance variations in resource availability, that imply a succession of recruitment processes depending on species resource use and competitive abilities (Clements, 1916; Meiners et al., 2015). The theory assumes that the deterministic succession shaped predictable trajectories gradually restoring communities pre-disturbance states (Chesson, 2000; Rees et al., 2001; Adler et al., 2007). Specifically, in forest ecosystem,

tree species community succession first comprises the recruitment of saplings benefiting from higher resources availability and lower competition. Then, stand maturation progressively enforces the exclusion of low-competitive species until the senescence of early-successional species, and the emergence of late-successional ones leading to the recovery of pre-disturbance composition and diversity (Denslow & Guzman, 2000). Empirical evidence, however, often show post-disturbance trajectories deviating from the predictable succession and resulting from an interplay of deterministic and stochastic recruitment processes. Post-disturbance trajectories would then also rely on stochastic processes, like dispersal limitations, and thus lead to new equilibrium states instead of recovering towards the pre-disturbance state (Hubbell, 2001; Chave, 2004; Norden et al., 2015). Hence, the question is to determine the balance between deterministic and stochastic recruitment processes. In the hyper-diverse tropical forests, this balance is specifically debated. While several studies revealed predictable and homogeneous successional patterns restoring the pre-disturbance community (Norden et al., 2009; Letcher et al., 2015), others showed diverging trajectories following disturbance and different equilibrium state (Longworth et al., 2014; Norden et al., 2015).

Post-disturbance trajectories may correspond either to recruited trees taxonomic characteristics, that refer to neutral species assemblages, or to functional characteristics, that account for species ecology and functioning (MacArthur & Levins, 1967; Violle et al., 2007; Kunstler et al., 2016). Deterministic processes correspond to a recruitment depending on species competitive ability determined by their functional characteristics (Rees et al., 2001; Perronne et al., 2017). Resulting recruitment trajectories would be similar among communities. Specifically, as light is the limiting resource in tropical forests, deterministic processes would drive a functional succession from fast-growing species with "acquisitive" resource use, able of important and fast light acquisition, to slow-growing, long-lived species with "conservative" resource use (Denslow, 1980; Molino & Sabatier, 2001; Bongers et al., 2009).

Key leaf, wood and life-history functional traits assessing species resources acquisition strategy and ecology would thus allow grasping the deterministic processes involved in post-disturbance trajectories (Wright et al., 2004; Chave et al., 2009; Hérault et al., 2011). Stochastic processes, in turn, correspond to a random species recruitment independant of their taxonomic identity and functional characteristics, and imply trajectories diverging among communities. Because of the high diversity and functional redundancy in tropical forests, tree community taxonomic trajectories may not track functional ones. The combination of taxonomic and functional approaches is then necessary to identify the different recruitment processes involved (Fukami et al., 2005; Chalmandrier et al., 2015; Cequinel et al., 2018) and to conclude about the recovery of pre-disturbance community (Clements, 1916; Diamond, 1975).

Here, we followed recruitment trajectories over 30 years of 75 ha cumulated area of Neotropical forest plots set up on a gradient of disturbance intensity, from 10 to 60% of above-

ground standing biomass removed. We examined recruited trees taxonomic composition, richness, and evenness, taxonomic turnover compared to pre-disturbance community, and functional composition and diversity based on seven major leaf, stem, and life-history traits. We compared the recruitment trajectories to neutral models corresponding to a stochastic recruitment and a randomization of species functional traits. Specifically, we (i) elucidated the successional pathway shaping community response to disturbance and the underlying ecological processes, and (ii) clarified the extent of community taxonomic and functional resilience, in the sense of pre-disturbance characteristics recovery, and its consequences for tropical forest management.

2. Material and Methods

2.1 Study Site

The Paracou station is located in a lowland tropical rainforest in French Guiana (518°N and 5253°W). Climate is tropical wet with mean annual precipitation averaging 2980 mm.yr⁻¹ (30-yr period) and a 3-months dry season (< 100 mm.mo⁻¹) from mid-August to mid-November, and a one-month dry season in March (Wagner *et al.*, 2011). The mean annual temperature is 26°C. Elevation ranges from 5 to 50 m.

Across all plots, the topography mainly corresponds to hilltops or hillsides, and bottomlands cover less than 1 % of the area. Plots are shallow ferralitic acrisols over a layer of transformed saprolite with low permeability and lateral drainage. Soil conditions are homogeneous, to the exception of the highest hilltops where the thick surface allows a free vertical drainage (Gourlet-Fleury *et al.*, 2004).

The experiment is a network of twelve 6.25 ha plots (Table 1) that underwent three disturbance treatments in 1987 according to a randomized plot design (Hérault & Piponiot, 2018).

The experiment comprised three replicates of three sylvicultural treatments (hereafter plots T1, T2 and T3) and three control plots (T0). All treatments T1, T2 and T3 comprised the logging of 10 trees/ha with 50 cm minimum DBH that belonged to a set of 58 commercially exploited species (Gourlet-Fleury et al., 2004). Treatment T2 additionally comprised a thinning treatment by poison-girdling of noncommercially exploited species, randomly selected species with an average 30 trees/ha with 40 cm minimum DBH. Treatment T3 additionally comprised the logging of 15 trees /ha with 40 cm minimum DBH and the poison-girdling of 20 trees/ha with a 50 cm minimum DBH, all belonging to non-commercial species. Considering the silvicultural treatments and the following damage, disturbance intensity was measured as the percentage of aboveground standing biomass of trees above 10 cm DBH (%AGB) lost between the first inventory in 1984 and five years after disturbance (Piponiot et al., 2016), estimated with the BIOMASS R package (Réjou-Méchain et al., 2018), and without accounting for lianas. The three treatments constituted a disturbance intensity gradient with increasing of above-ground biomass (AGB) lost and surface disturbed.

Table 1. Intervention table, summary of the disturbance intensity for the 4 plot treatments in Paracou. DBH refers to						
Diameter at Breast Height and AGB refers to aboveground standing biomass of trees above 10 cm DBH.						

Treatment	Timber	Thinning	Fuelwood	%AGB lost
Control	-	-	-	0
T1	DBH \geq 50 cm, commercial species, $\approx 10 \text{ trees.ha}^{-1}$	-	-	[12 - 33]
T2	DBH \geq 50 cm, commercial species, $\approx 10 \text{ trees.ha}^{-1}$	DBH \geq 40 cm, non-valuable species, $\approx 30 trees.ha^{-1}$	-	[33 – 56]
Т3	DBH \geq 50 cm, commercial species, $\approx 10 \text{ trees.ha}^{-1}$	DBH \geq 50 cm, non-valuable species, $\approx 15 \ trees.ha^{-1}$	$40 \text{ cm} \le \text{DBH} \le 50$ cm, non-valuable species, ≈ 15 $trees.ha^{-1}$	[35 – 56]

2.2 Inventories Protocol and Dataset Collection

Dominant families in the study site are *Fabaceae*, *Chrysobalanaceae*, *Lecythidaceae* and *Sapotaceae*. All trees above 10 cm DBH were mapped and measured annually since 1984. Trees are first identified with a vernacular name assigned by the forest worker team, and afterward with a scientific name assigned by botanists during regular botanical campaigns. Botanical campaigns have been carried out every five to six years from 2003 onwards, but identification levels varied between campaigns.

The variability of protocols in time raised methodological issues as vernacular names usually correspond to different botanical species. This variability resulted in significant taxonomic uncertainties that had to be propagated to composition and diversity metrics. The uncertainty propagation was done through a Bayesian framework reconstituting complete inventories at genus level from real incomplete ones on the basis of vernacular/botanical names association. Vernacular names were replaced through multinomial trials based on the association probability $[\alpha_1, \alpha_2, ..., \alpha_N]$ observed across all inventories between each vernacular name v and the species $[s_1, s_2, ..., s_N]$:

$$M_{\nu}\Big([s_1, s_2, ..., s_N], [\alpha_1, \alpha_2, ..., \alpha_N]\Big)$$

See appendix 1 and Aubry-Kientz *et al.* (2013) for the detailed methodology.

To minimize the remaining identification uncertainties, the simulated botanical inventories were reported at genus level

We considered six functional traits representing leaf economics, the leaf thickness, toughness, total chlorophyll content and specific leaf area, and stem economics, the wood specific gravity and bark thickness. These traits were obtained from the BRIDGE project ¹, which proved to reliably assess species functional traits and community functional diversity (Paine *et al.*, 2015). Trait values were assessed from a selection of individuals located in nine permanent plots

in French Guiana, including two in Paracou, and comprised 294 species belonging to 157 genera.

For the trait value at species level we took the median value of the several trees sampled in the database, which allowed accounting for intraspecific variability although we considered species-level trait values.

Whenever a species was in the dataset was missing some trait values (10% of the species), missing values were filled using multivariate imputation by chained equation (van Buuren & Groothuis-Oudshoorn, 2011). To account for the phylogenetic signal in the filling process, imputations based on samples from the same genus or from the same family. Whenever a species was not in the dataset, it was attributed a set of trait values randomly sampled among species of the next higher taxonomic level (same genus or family). Two life-history traits (maximum specific height and seed mass) came from the Mariwenn database ². The database compiles information from a vast literature on the flora of French Guiana (Ollivier et al., 2007) and comprises 362 species belonging to 188 genera. As seed mass information was classified into classes, no data filling process was applied and analyses were restricted to the botanical species recorded.

All composition and diversity metrics were obtained after 50 iterations of the uncertainty propagation framework.

2.3 Recruitment trajectories

Communities were split into surviving trees of pre-disturbance communities and trees recruited afterward. Trajectories of recruitment diversity (hereafter H^q_{obs}) and turnover started from the first year after disturbance and then considered the trees recruited by 2-years intervals (*i.e.* for the date Y, the trees that were under 10 cm DBH at Y-1 but above 10 cm DBH at Y).

Trajectories of taxonomic diversity were assessed through species richness and evenness (the Hill number translation of the Simpson index) (Chao & Jost, 2015; Marcon & Hérault, 2015). The two diversities belong to the set of

¹http://www.ecofog.gf/Bridge/

²https://www.ecofog.gf/mariwenn/

Tsallis or generalized entropy, respectively corresponding to the zero and two order q of diversity that emphasizes common species regarding their stem density. These two metrics assess community richness and evenness, that reveals the structure of dominance in the community and allow grasping the changes in community abundance distribution. Both metric are complementary to detect the changes in community structure (Magurran, 2004).

Functional diversity trajectories were assessed through the Rao index of quadratic entropy, which combines species abundance distribution and average pairwise dissimilarity based on all functional traits. Functional composition trajectories were assessed through the functional traits community weighted means (CWM), representing the average trait value in a community weighted by species relative abundance based on the number of stem (Díaz *et al.*, 2007). Seed mass trajectories were reported by the stems proportion of each class recorded in the inventories.

The taxonomic similarity between recruited trees and pre-disturbance forest was measured with the relativized abundance replacement measuring species turnover between communities (Podani *et al.*, 2013):

$$T_{ab} = \frac{\sum_{i=1}^{n} |x_i^a - x_i^b| - \left| \sum_{i=1}^{n} x_i^a - \sum_{i=1}^{n} x_i^b \right|}{\sum_{i=1}^{n} \max\left(x_i^a; x_i^b\right)}$$
(1)

With Tab the turnover between communities a and b, n the total number of species in the two communities and x_{ij} the abundance of species I in community j.

The correlation between the initial disturbance intensity and recruited community taxonomic richness and evenness, functional richness and taxonomic turnover were tested through Spearman rank correlation tests.

The observed recruitment trajectories were compared to null trajectories obtained from taxonomic and functional null models. A taxonomic null model was independently built for each plot and for each year. Taxonomic null trajectories were simulated by randomly sampling the recruited trees in the current living community, according to the observed number of recruits. Functional null trajectories were simulated by randomly reassigning the combinations of functional trait values among species. The functional null models maintained the community abundance distribution while randomizing the relative abundance of trait values (Mason $et\ al.$, 2013). Trajectories of null diversity (hereafter H^q_{null}) were similarly obtained after 50 iterations of the random sampling models.

Identification of indicative species

We identified characteristic species of post-disturbance recruitment by computing species indicator values for post-disturbance and undisturbed recruited communities. We used the methodology developed in Dufrêne & Legendre (1997). Indicator species are defined as the most characteristic species of a group: it would be species only found in that group and recorded in the majority of the plots belonging to this group. All disturbed plots were pooled in the disturbed group and opposed to the undisturbed plots. $IndVal_{ij}$ is the indicator value of species i relative to group

j. It is the product of A_{ij} , a measure of species specificity, by B_{ij} , a measure of species fidelity. A_{ij} is the mean abundance of species i in each plot j of the group compared to all groups in the study. A_{ij} is maximum when species i is only present in group j. B_{ij} is the relative frequency of occurence if species i in the plots of group j, it is maximum when species i is only present in all plots of group j.

$$A_{ij} = Nindividuals_{ij}/Nindividuals_{i.}$$
 (2)

$$B_{ij} = Nsites_{ij}/Nsites_{.j}$$
 (3)

$$IndVal_{ij} = A_{ij}xB_{ij}x100 (4)$$

In the formula for A_{ij} , $Nindividuals_{ij}$ is the number of individuals of species i in plots of group j and $Nindividuals_{i}$ is the sum of the mean number of individuals of species i over all groups. In the formula for B_{ij} , $Nsites_{ij}$ is the number of plots in group j where species i is recorded and $Nsites_j$ is the total number of sites. The Indicator Value of a species is the maximum value of $IndVal_{ij}$, expressed in percentage, observed across all groups. The indicator value is maximal when all individuals of species i are observed in all plots of one group.

The significance of species indicator values was tested through the comparison with a random permutaiton procedure of sites among groups. Observed values were compared to the mean values obtained from 50 random permutations with a z statistic.

3. Results

Indicative species and species repartition

Overall, 602 species were recorded along the 30 years of inventory and across the 12 plots. Among all species, 157 were only inventoried in one plot and 43 were inventoried in all plots. Species repartition is detailed in table 2.

The dominant recruited species were *Lecythis persistens*, *Licania alba*, *Oenocarpus bataua*, *Oxandra asbeckii*, and *Eperua grandiflora*. The dominant species recruited in disturbed plots were *Miconia tschudyoides*, *Inga sp.*, *Oenocarpus bataua*, *Licania alba*, and *Xylopia sp.*.

We defined the indicative species of a group when their indicative value was above 90%. Indicative species of undisturbed communities were *Couepia bracteosa* and species of the genus *Neea*. Indicative species of post-disturbance communities were species of the genuses *Cecropia*, *Cupania*, *Apeiba*, *Pourouma*, *Sterculia*, *Tapirira*, *Vismia*, and the species *Miconia tschudyoides*, *Schefflera decaphylla*, *Virola surinamensis*, and *Xylopia nitida*. Among disturbance-indicator species, genus *Cecropia*, *Apeiba*, *Vismia* and *Pourouma* were representative of the first years following disturbance while *Cupania*, *Xylopia*, *Miconia* and *Sterculia* were characteristic of later time after disturbance.

Table 2. Recruited species repartition across treatments: number of species inventoried and species only recruited in the treatment considered

	Control	T1	T2	Т3
Number of species by treatment	373	313	390	432
Number of species recruited only in one treatment	42	30	43	81

3.1 Taxonomic richness and evenness and functional diversity

The average number of recruited trees per plot and per year along the whole time survey increased with the increasing disturbance intensity. For a 2-year laps, the number of recruits per plot was around 74 in plots T0, 142 in plots T1, 211 in plots T2 and 273 in plots T3. In total, during the 30 years, 602 species were recruited across the 12 plots. Among thoses, 26% occured in one plot only and 7% occured in all plots.

In undisturbed communities the taxonomic richness and evenness of recruited trees remained stable over the 30 years, around values equivalent to those of the taxonomic null model (Figure (1)).

In disturbed communities the taxonomic richness followed hump-shaped trajectories first increasing until a maximum reached after around 15 years and positively correlated to the disturbance intensity ($\rho_{spearman}^{Richness} = 0.93$). The taxonomic richness then decreased and recovered the predisturbance values after 30 years. Compared to the null models, the observed taxonomic richness was lower than the richness of a random species recruitment. The difference between the observed and null model richness followed a hump-shaped trajectory with a maximum difference reached after 15 years. In all disturbed communities, the taxonomic evenness decreased whenever the disturbance intensity over the 30 years ($\rho_{spearman}^{simpson} = -0.35$). Compared to the null model, the observed taxonomic eveness was lower than the evenness of a random species recruitment. The difference between the observed and null model evenness followed a hump-shaped trajectory with a maximum difference reached around 25 years after disturbance.

The functional diversity in the undisturbed plots remained stable and equivalent to this of the functional null model over the 30 years. In the lowest disturbance plots the functional diversity remained stable or slightly increasing, and was higher than this of the null model for two of the T1 plots. In the disturbed plots of higher disturbance intensity (T2 and T3) the functional diversity decreased until 15 years after disturbance, when it started to recover towards initial values. The observed functional diversity remained lower than this of the null model over the 30 years.

3.2 Functional composition

In undisturbed plots functional traits values remained stable over the 30 years while it followed hump-shaped trajectories in all disturbed plots, to the exception of the leaf chlorophyll content. Trajectories of SLA and bark thickness first increased before decreasing towards initial values. Conversely, trajectories of leaf thickness, leaf toughness, wood specific gravity, and maximum height first decreased and then started returning towards initial values but their recovery remained unachieved after 30 years (Figure 2).

3.3 Recruitment Turnover

Over the 30 years in control plots the turnover of recruited species compared to initial community remained low (Figure 3). In disturbed plots the recruited species turnover followed a marked hump-shaped trajectory, with a maximum reached around 15 years after disturbance. The maximum turnover was positively correlated to the disturbance intensity ($\rho_{spearman} = 0.93$). Thirty years after disturbance the turnover had returned to low values.

4. Discussion

Our analysis of the taxonomic and functional response of tree recruitment over 30 years following a disturbance gradient revealed a three-phased successional pathway shaped by an interplay between stochastic recruitment and deterministic processes. Although the recruitment trajectories suggested community taxonomic and functional recovery, the return to the initial stage remained unachieved after 30 years.

4.1 A three-phased deterministic successional pathway

Post-disturbance recruitment trajectories followed a threephased successional pathway defined by the emergence of deterministic competition processes for light, that balanced the stochastic processes involved in undisturbed communities before disturbance. Indicative species of disturbed plots corresponded to pionners and light demanding species while indicative species of undistrubed plots were shade-tolerant, late successional species.

A first phase (*i.e.* 0-8 years) was defined by a very low taxonomic turnover between recruited trees and initial communities (Figure 3), and by similar taxonomic and functional trajectories matching the null model trajectories (Figure (1)). The first phase corresponded to the time lapse before recruitment processes markedly changed. During this phase, recruits comprised a few pioneer species, like *Cecropia obtusa*, and rather corresponded to the growth of predisturbance surviving saplings (with DBH < 10 cm), that

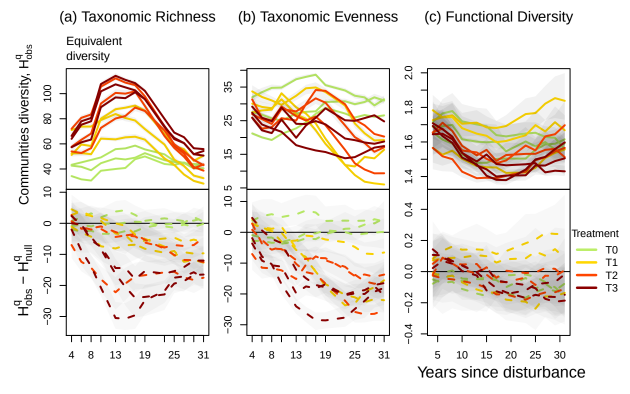


Figure 1. Upper panels, trajectories over 30 years of taxonomic richness (a), taxonomic evenness (b) and functional diversity (c) of observed 2-years laps recruitment H_{obs}^q . Lower panels, diversity differences to null models $H_{obs}^q - H_{null}^q$. Shaded areas are the credibility intervals.

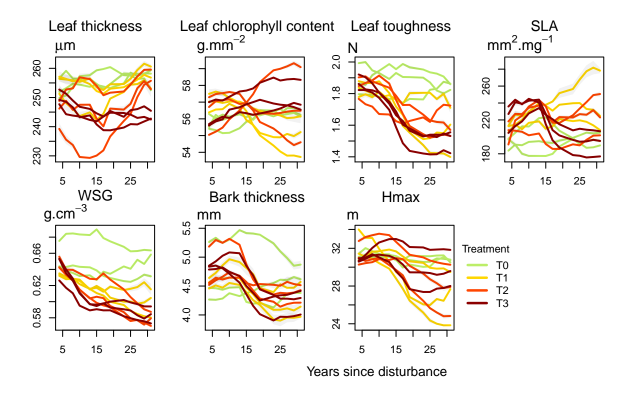


Figure 2. Community weighted means (CWM) of the 7 functional traits considered: leaf thickness, leaf chlorophyll content, leaf tougness, specific leaf area (SLA), wood specific gravity (WSG), bark thickness and maximum height at adult stage (Hmax). Shaded areas are the credibility intervals.

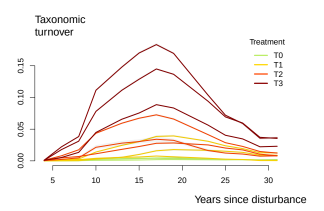


Figure 3. Trajectories over 30 years of the abundance-based turnover between 2-years laps recruited trees pre-disturbance communities.

were late-successional species like *Lecythis persistens* or *Licania alba*. These species displayed "resource-conservative" functional strategies, *i.e.* dense leaf and wood tissues and low exchange surface, favored in undisturbed communities where light levels are low (Peet, 1992; Denslow & Guzman, 2000). These saplings were the first to benefit from the alleviated competition and the higher light availability that is known to follow disturbance and foster plant productivity and growth (Monteith, 1972; Chazdon & Fetcher, 1984).

A second phase (i.e. 8-15 years) was marked by an increase of the taxonomic turnover between recruited trees and initial communities, and a decrease of recruits richness and evenness compared to the null model. The second phase also corresponded to a shift in recruits functional composition towards more "resource-acquisitive" strategies. Indicative species switched from pioneers of genus Cecropia or Vismia, to non-pioneer light demanding species of genuses Cupania, Xylopia or Miconia. The shift would mark the recruitment of trees germinated from seeds, eventually representing the main part of the recruitment (Lawton & Putz, 1988). This first phase was the time lapse before pioneers, like Miconia tschudyoides, Inga spp., Tapirira spp., and Cecropia spp., became dominant among recruits. The dominance of pioneers decreased the taxonomic richness and evenness of recruited trees and made the recruitment trajectories depart from those of the null model. Recruitment processes then changed after disturbance, with the emergence of deterministic processes favoring a restricted pool of species. During the second phase, the recruitment functional composition switched from a random trait assembly to more "resource-acquisitive" functional strategies (Figure (2)). Species displaying high SLA and low maximum height, leaf toughness and wood specific gravity became dominant (Wright et al., 2004; Chave et al., 2009; Hérault et al., 2011). During the second succession phase, a net advantage then emerged for species displaying efficient light acquisition, with high SLA and leaf chlorophyll content, and displaying inexpensive, short-lived tissues, with low leaf thickness, leaf toughness, small Hmax, low wood specific gravity, and low bark thickness. The advantage gained

by "light acquisitive" species confirmed the role of light availability in the post-disturbance successional pathways, as already demonstrated in temperate and tropical forests (Peña-Claros et al., 2008; Carreño-Rocabado et al., 2012; Kunstler et al., 2016; Both et al., 2019). The emergence of deterministic processes favoring pioneers balanced the neutral species assemblage where late-successional species were dominant, like in undisturbed communities. Deterministic processes made post-disturbance trajectories depart from the null trajectories, all the more so that the disturbance was intense (Figure (1)). Following low intensity disturbance (T1 plots), pioneers and late-successional species were balanced (Bongers et al., 2009). Following intense disturbance (T2 and T3 plots), deterministic processes prevailed and the composition of recruited communities rapidly differed from this of the pre-disturbance community.

A third recruitment ongoing phase (starting around 15 years) was delineated by a return towards pre-disturbance values of functional diversity and taxonomic richness, while taxonomic evenness and functional composition remained altered (Figure (1) & (2)). Although recruited species were mainly light-demanding and pioneers, like species from *Inga, Miconia*, or *Licania* genera, late-successional species progressively settled, like *Licania alba* (Fortunel *et al.*, 2014). The recruitment of late-successionals would translate the progressive closing of forest canopy and the increase competition for light and space [Peet (1992);Denslow & Guzman (2000);] Deterministic recruitment processes then gradually left room again to stochastic recruitment as observed in undisturbed forest (Lawton & Putz, 1988; Chave, 2004).

4.2 Community recovery

Following disturbance, community taxonomic and functional composition recovered towards pre-disturbance values. Such recovery confirmed previous results from the Paracou experiment, conducted 10 years (Molino & Sabatier, 2001) and 20 years (Baraloto *et al.*, 2012) after disturbance, where the early signs of the resilience of pre-disturbance taxonomic and functional composition recovery had been detected.

Thirty years after disturbance, the taxonomic and functional diversity and composition of the recruited community returned towards pre-disturbance states (Fukami *et al.*, 2005; Fortunel *et al.*, 2014). The recovery of pre-disturbance states meant the maintenance of community diversity and composition, and a low recruitment of species absent in the pre-disturbance community. It then confirmed the importance of dispersal limitations among tropical tree species (Svenning & Wright, 2005).

Although community taxonomic and functional recovery was ongoing, both remained altered 30 years after disturbance. Besides, the recovery of community taxonomic richness was faster, while this of the functional richness tracked the trajectory of the taxonomic evenness (Grime, 1998). The higher the disturbance intensity, the more persistent the dominance of light-demanding species. Such long-term impact raises questions for tropical forests management as most valuable species are late-successional species, so their exploitation would require cutting cycles longer than

30 years (Putz *et al.*, 2012). Besides, persistent changes in community composition likely alters community functioning (Díaz *et al.*, 2005), and increases the risk of losing keystone species, with unexpected ecological consequences (Jones *et al.*, 1994; Chazdon, 2003). Infrequent species might indeed have unique functional characteristics in the ecosystem, apart from the ones considered here, or be a key resource for some of the fauna (Schleuning *et al.*, 2016).

5. Conclusion

Post-disturbance recruitment trajectories revealed a threephased deterministic successional pathway driven by the emergence of trait-based deterministic processes balancing pre-disturbance stochastic processes, and enhancing lightacquisitive species. A first phase corresponded to the recruitment of pre-disturbance surviving saplings maintaining the taxonomic and functional diversity and composition of predisturbance communities. A second phase corresponded to the recruitment of true recruits from germinating seeds, belonging to a restricted pool of pioneers, that were favored by the emergence of competitive exclusion for light. Above a disturbance intensity threshold, the second recruitment phase saw the dominance of short-lived, fast growing pioneers that drastically changed community composition, diversity, and likely functioning. A third phase eventually corresponded to the recovery of stochastic recruitment processes and to the return towards pre-disturbance taxonomic and functional characteristics. Although resilient, recruits diversity and composition remained altered 30 years after disturbance, all the more so that disturbance was intense. Regarding forest management, our results supported cutting cycles longer than 30 years and demonstrated long-term impacts emphasising the need to evaluate forest management sustainability.

6. Acknowledgement

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7. Data availability

This article is based upon the dataset of the Paracou station, which is part of the Guyafor permanent plots network in French Guiana (Cirad-CNRS-ONF). The dataset is available upon request to the scientific director (https://paracou.cirad.fr).

References

Adler, P.B., HilleRislambers, J. & Levine, J.M. (2007) A niche for neutrality. *Ecology Letters*, **10**, 95–104.

- Aubry-Kientz, M., Rossi, V., Wagner, F. & Hérault, B. (2015) Identifying climatic drivers of tropical forest dynamics. *Biogeosciences*, **12**, 5583–5596.
- Aubry-Kientz, M., Hérault, B., Ayotte-Trépanier, C., Baraloto, C. & Rossi, V. (2013) Toward Trait-Based Mortality Models for Tropical Forests. *PLoS ONE*, 8.
- Baraloto, C., Hérault, B., Paine, C.E.T., Massot, H., Blanc, L., Bonal, D., Molino, J.F.F., Nicolini, E.a. & Sabatier, D. (2012) Contrasting taxonomic and functional responses of a tropical tree community to selective logging. *Journal of Applied Ecology*, 49, 861–870.
- Bongers, F., Poorter, L., Hawthorne, W.D. & Sheil, D. (2009) The intermediate disturbance hypothesis applies to tropical forests, but disturbance contributes little to tree diversity. *Ecology Letters*, **12**, 798–805.
- Both, S., Riutta, T., Paine, C.T., Elias, D.M., Cruz, R., Jain, A., Johnson, D., Kritzler, U.H., Kuntz, M., Majalap-Lee, N. *et al.* (2019) Logging and soil nutrients independently explain plant trait expression in tropical forests. *New Phytologist*, **221**, 1853–1865.
- Carreño-Rocabado, G., Peña-Claros, M., Bongers, F., Alarcón, A., Licona, J.C. & Poorter, L. (2012) Effects of disturbance intensity on species and functional diversity in a tropical forest. *Journal of Ecology*, **100**, 1453–1463.
- Cequinel, A., Capellesso, E.S., Marcilio-Silva, V., Cardoso, F.C. & Marques, M.C. (2018) Determinism in tree turnover during the succession of a tropical forest. *Perspectives in Plant Ecology, Evolution and Systematics*, **34**, 120–128.
- Chalmandrier, L., Münkemüller, T., Lavergne, S. & Thuiller, W. (2015) Effects of species' similarity and dominance on the functional and phylogenetic structure of a plant meta-community. *Ecology*, **96**, 143–153.
- Chao, A. & Jost, L. (2015) Estimating diversity and entropy profiles via discovery rates of new species. *Methods in Ecology and Evolution*, **6**, 873–882.
- Chave, J. (2004) Neutral theory and community ecology. *Ecology Letters*, **7**, 241–253.
- Chave, J., Coomes, D., Jansen, S., Lewis, S.L., Swenson, N.G. & Zanne, A.E. (2009) Towards a worldwide wood economics spectrum. *Ecology Letters*, **12**, 351–366.
- Chazdon, R.L. (2003) Tropical forest recovery: legacies of human impact and natural disturbances. *Perspectives in Plant Ecology, Evolution and Systematics*, **6**, 51–71.
- Chazdon, R.L. & Fetcher, N. (1984) Photosynthetic light environments in a lowland tropical rain forest in costa rica. *The Journal of Ecology*, pp. 553–564.
- Chesson, P. (2000) Mechanisms of maintenance of species diversity. *Annual review of Ecology and Systematics*, **31**, 343–366.

- Clements, F.E. (1916) *Plant succession: an analysis of the development of vegetation*. 242. Carnegie Institution of Washington.
- Denslow, J.S. (1980) Gap Partitioning among Tropical Rainforest Trees. *Biotropica*, **12**, 47–55.
- Denslow, J.S. & Guzman, G.S. (2000) Variation in stand structure, light and seedling abundance across a tropical moist forest chronosequence, Panama. *Journal of Vegetation Science*, **11**, 201–212.
- Dezécache, C., Faure, E., Gond, V., Salles, J.M., Vieilledent, G. & Hérault, B. (2017a) Gold-rush in a forested El Dorado: Deforestation leakages and the need for regional cooperation. *Environmental Research Letters*, **12**.
- Dezécache, C., Salles, J.M., Vieilledent, G. & Hérault, B. (2017b) Moving forward socio-economically focused models of deforestation. *Global Change Biology*, 23, 3484–3500.
- Diamond, J.M. (1975) Assembly of species communities. *Ecology and evolution of communities*, pp. 342–444.
- Díaz, S., Lavorel, S., de Bello, F., Quétier, F., Grigulis, K. & Robson, T.M. (2007) Incorporating plant functional diversity effects in ecosystem service assessments. *Proceed*ings of the National Academy of Sciences, 104, 20684– 20689.
- Díaz, S., Tilman, D., Fargione, J., Chapin III, F.S., Dirzo, R., Kitzberger, T., Gemmill, B., Zobel, M., Vilà, M., Mitchell, C., Wilby, A., Daily, G.C., Galetti, M., Laurance, W.F., Pretty, J., Naylor, R., Power, A., Harvell, A., Potts, S., Kremen, C., Griswold, T. & Eardley, C. (2005) Biodiversity Regulation of Ecosystem Services. *Trends and conditions*, pp. 297–329.
- Dufrêne, M. & Legendre, P. (1997) Species assemblages and indicator species: the need for a flexible asymmetrical approach. *Ecological monographs*, **67**, 345–366.
- Fortunel, C., Paine, C., Fine, P.V., Kraft, N.J. & Baraloto, C. (2014) Environmental factors predict community functional composition in amazonian forests. *Journal of Ecology*, **102**, 145–155.
- Fukami, T., Bezemer, T.M., Mortimer, S.R. & Van Der Putten, W.H. (2005) Species divergence and trait convergence in experimental plant community assembly. *Ecology Letters*, **8**, 1283–1290.
- Gourlet-Fleury, S., Guehl, J.M. & Laroussinie, O. (2004) Ecology & management of a neotropical rainforest. Lessons drawn from Paracou, a long-term experimental research site in French Guiana.
- Gourlet-Fleury, S. & Houllier, F. (2000) Modelling diameter increment in a lowland evergreen rain forest in French Guiana. Forest Ecology and Management, 131, 269–289.
- Grime, J.P. (1998) Benefits of Plant Diversity to Ecosystems: Immediate, Filter and Founder Effects. *Journal of Ecology*, pp. 902–910.

- Hérault, B., Bachelot, B., Poorter, L., Rossi, V., Bongers, F., Chave, J., Paine, C.E.T., Wagner, F. & Baraloto, C. (2011) Functional traits shape ontogenetic growth trajectories of rain forest tree species. *Journal of Ecology*, 99, 1431– 1440.
- Hérault, B. & Piponiot, C. (2018) Key drivers of ecosystem recovery after disturbance in a neotropical forest. *Forest Ecosystems*, **5**, 2.
- Hubbell, S.P. (2001) *The Unified Neutral Theory of Biodiversity and Biogeography*. Princeton University Press.
- Jones, C.G., Lawton, J.H. & Shachak, M. (1994) Organisms as ecosystem engineers. *Ecosystem management*, pp. 130–147. Springer.
- Kunstler, G., Falster, D., Coomes, D.A., Hui, F., Kooyman, R.M., Laughlin, D.C., Poorter, L., Vanderwel, M., Vieilledent, G., Wright, S.J., Aiba, M., Baraloto, C., Caspersen, J., Cornelissen, J.H.C., Gourlet-Fleury, S., Hanewinkel, M., Herault, B., Kattge, J., Kurokawa, H., Onoda, Y., Peñuelas, J., Poorter, H., Uriarte, M., Richardson, S., Ruiz-Benito, P., Sun, I.F., Ståhl, G., Swenson, N.G., Thompson, J., Westerlund, B., Wirth, C., Zavala, M.A., Zeng, H., Zimmerman, J.K., Zimmermann, N.E. & Westoby, M. (2016) Plant functional traits have globally consistent effects on competition. *Nature*, **529**, 204–207.
- Lawton, R.O. & Putz, F.E. (1988) Natural disturbance and gap-phase regeneration in a wind-exposed tropical cloud forest. *Ecology*, **69**, 764–777.
- Letcher, S.G., Lasky, J.R., Chazdon, R.L., Norden, N., Wright, S.J., Meave, J.A., Pérez-García, E.A., Muñoz, R., Romero-Pérez, E., Andrade, A. *et al.* (2015) Environmental gradients and the evolution of successional habitat specialization: a test case with 14 neotropical forest sites. *Journal of Ecology*, **103**, 1276–1290.
- Longworth, J.B., Mesquita, R.C., Bentos, T.V., Moreira, M.P., Massoca, P.E. & Williamson, G.B. (2014) Shifts in dominance and species assemblages over two decades in alternative successions in central amazonia. *Biotropica*, 46, 529–537.
- MacArthur, R. & Levins, R. (1967) The limiting similarity, convergence, and divergence of coexisting species. *The American Naturalist*, **101**, 377–385.
- Magurran, A.E. (2004) *Measuring Biological Diversity*. Blackwell Science Ltd.
- Marcon, E. & Hérault, B. (2015) entropart: An R Package to Measure and Partition Diversity. *Journal of Statistical Software*, **67**, 1–26.
- Mason, N.W.H., De Bello, F., Mouillot, D., Pavoine, S. & Dray, S. (2013) A guide for using functional diversity indices to reveal changes in assembly processes along ecological gradients. *Journal of Vegetation Science*, **24**, 794–806.

- Meiners, S.J., Cadotte, M.W., Fridley, J.D., Pickett, S.T. & Walker, L.R. (2015) Is successional research nearing its climax? New approaches for understanding dynamic communities. *Functional Ecology*, **29**, 154–164.
- Molino, J.F. & Sabatier, D. (2001) Tree diversity in tropical rain forests: a validation of the intermediate disturbance hypothesis. *Science*, **294**, 1702–1704.
- Monteith, J. (1972) Solar radiation and productivity in tropical ecosystems. *Journal of applied ecology*, **9**, 747–766.
- Norden, N., Angarita, H.A., Bongers, F., Martínez-Ramos, M., Granzow-de la Cerda, I., van Breugel, M., Lebrija-Trejos, E., Meave, J.A., Vandermeer, J., Williamson, G.B., Finegan, B., Mesquita, R. & Chazdon, R.L. (2015)
 Successional dynamics in Neotropical forests are as uncertain as they are predictable. *Proceedings of the National Academy of Sciences*, 112, 8013–8018.
- Norden, N., Chazdon, R.L., Chao, A., Jiang, Y.H. & Vílchez-Alvarado, B. (2009) Resilience of tropical rain forests: Tree community reassembly in secondary forests. *Ecology Letters*, **12**, 385–394.
- Ollivier, M., Baraloto, C. & Marcon, E. (2007) A trait database for guianan rain forest trees permits intra-and inter-specific contrasts. *Annals of forest science*, **64**, 781–786.
- Paine, C.T., Baraloto, C. & Díaz, S. (2015) Optimal strategies for sampling functional traits in species-rich forests. *Functional Ecology*, **29**, 1325–1331.
- Peet, R.K. (1992) Community structure and ecosystem function. *Plant succession: theory and prediction*, **11**, 103–citation_lastpage.
- Peña-Claros, M., Peters, E., Justiniano, M., Bongers, F., Blate, G.M., Fredericksen, T. & Putz, F. (2008) Regeneration of commercial tree species following silvicultural treatments in a moist tropical forest. *Forest Ecology and Management*, **255**, 1283–1293.
- Perronne, R., Munoz, F., Borgy, B., Reboud, X. & Gaba, S. (2017) How to design trait-based analyses of community assembly mechanisms: Insights and guidelines from a literature review. *Perspectives in Plant Ecology, Evolution and Systematics*, **25**, 29–44.
- Piponiot, C., Sist, P., Mazzei, L., Peña-Claros, M., Putz, F.E., Rutishauser, E., Shenkin, A., Ascarrunz, N., de Azevedo, C.P., Baraloto, C., França, M., Guedes, M.C., Coronado, E.N., D'Oliveira, M.V., Ruschel, A.R., da Silva, K.E., Sotta, E.D., de Souza, C.R., Vidal, E., West, T.A. & Hérault, B. (2016) Carbon recovery dynamics following disturbance by selective logging in amazonian forests. *ELife*, 5, e21394.
- Podani, J., Ricotta, C. & Schmera, D. (2013) A general framework for analyzing beta diversity, nestedness and related community-level phenomena based on abundance data. *Ecological Complexity*, **15**, 52–61.

- Putz, F.E., Zuidema, P.A., Synnott, T., Peña-Claros, M., Pinard, M.A., Sheil, D., Vanclay, J.K., Sist, P., Gourlet-Fleury, S., Griscom, B., Palmer, J. & Zagt, R. (2012) Sustaining conservation values in selectively logged tropical forests: the attained and the attainable. *Conservation Letters*, 5, 296–303.
- Rees, M., Condit, R., Crawley, M., Pacala, S. & Tilman, D. (2001) Long-term studies of vegetation dynamics. *Science*, **293**, 650–655.
- Réjou-Méchain, M., Tanguy, A., Piponiot, C., Chave, J. & Hérault, B. (2018) BIOMASS: Estimating Aboveground Biomass and Its Uncertainty in Tropical Forests. R package version 1.2.
- Rutishauser, E., Hérault, B., Petronelli, P. & Sist, P. (2016) Tree Height Reduction After Selective Logging in a Tropical Forest. *Biotropica*, **48**, 285–289.
- Schleuning, M., Fruend, J., Schweiger, O., Welk, E., Albrecht, J., Albrecht, M., Beil, M., Benadi, G., Bluethgen, N., Bruelheide, H. et al. (2016) Ecological networks are more sensitive to plant than to animal extinction under climate change. Nature communications, 7, 13965.
- Svenning, J.C. & Wright, S.J. (2005) Seed limitation in a panamanian forest. *Journal of ecology*, **93**, 853–862.
- van Buuren, S. & Groothuis-Oudshoorn, K. (2011) mice: Multivariate imputation by chained equations in r. *Journal of Statistical Software*, **45**, 1–67.
- Violle, C., Navas, M.L.L., Vile, D., Kazakou, E., Fortunel, C., Hummel, I. & Garnier, É. (2007) Let the concept of trait be functional! *Oikos*, 116, 882–892.
- Wagner, F., Hérault, B., Stahl, C., Bonal, D. & Rossi, V. (2011) Modeling water availability for trees in tropical forests. Agricultural and Forest Meteorology, 151, 1202– 1213.
- Wright, I.J., Reich, P.B., Westoby, M., Ackerly, D.D., Baruch, Z., Bongers, F., Cavender-Bares, J., Chapin, T., Cornelissen, J.H.C., Diemer, M., Flexas, J., Garnier, E., Groom, P.K., Gulias, J., Hikosaka, K., Lamont, B.B., Lee, T., Lee, W., Lusk, C., Midgley, J.J., Navas, M.L., Niinemets, &., Oleksyn, J., Osada, N., Poorter, H., Poot, P., Prior, L., Pyankov, V.I., Roumet, C., Thomas, S.C., Tjoelker, M.G., Veneklaas, E.J. & Villar, R. (2004) The worldwide leaf economics spectrum. *Nature*, **428**, 821–827.