

Accepted Article

# Second-growth and small forest clearings have little effect on the temporal activity patterns of Amazonian phyllostomid bats

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## Abstract

Secondary forests and human-made forest gaps are conspicuous features of tropical landscapes. Yet, behavioural responses to these aspects of anthropogenically-modified forests remain poorly investigated. Here, we analyse the effects of small human-made clearings and secondary forests on tropical bats by examining the guild- and species-level activity patterns of phyllostomids sampled in the Central Amazon, Brazil. Specifically, we contrast the temporal activity patterns and degree of temporal overlap of six frugivorous and four gleaning animalivorous species in old-growth forest and second-growth forest and of four frugivores in old-growth forest and forest clearings. The activity patterns of frugivores and gleaning animalivores did not change between old-growth forest and second-growth, nor did the activity patterns of frugivores between old-growth forest and clearings. However, at the species level we detected significant differences for *Artibeus obscurus* (old-growth forest vs. second-growth) and *Artibeus concolor* (old-growth forest vs. clearings). The degree of temporal overlap was greater than random in all sampled habitats. However, whereas for frugivorous species the degree of temporal overlap was similar between old-growth forest and second-growth, for gleaning animalivores it was lower in second-growth than in old-growth forest. On the other hand, forest clearings were characterized by increased temporal overlap between frugivores. Changes in activity patterns and temporal overlap may result from differential foraging opportunities and dissimilar predation risks. Yet, our analyses suggest that activity patterns of bats in second-growth and small forest clearings, two of the most prominent habitats in humanized tropical landscapes, varies little from the activity patterns in old-growth forest.

**Keywords:** Amazon, behaviour, forest gaps, niche partitioning, secondary forest, temporal overlap

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Across the tropics, human activities are moulding landscapes in which second-growth and forest gaps are conspicuous features (Chazdon 2014; Melo et al. 2013). The long-term persistence of tropical biodiversity, although primarily dependent on the conservation of vast tracts of old-growth forest habitat (Barlow et al. 2007; Gibson et al. 2011), will depend on the capacity of these novel ecosystems to accommodate old-growth forest-dwelling species. Secondary forests are now the largest forest cover type across the humid tropics (Chazdon 2014) and although the potential of second-growth to conserve tropical biodiversity remains hotly debated (Chazdon 2014; Gibson et al. 2011), mounting evidence suggests that areas of second-growth provide numerous ecosystem services and natural products to human populations, as well as crucial habitat for numerous taxa (Chazdon 2014; Melo et al. 2013; Rocha et al. 2018; Rozendaal et al. 2019). Alongside second-growth, man-made forest clearings, small to moderate in size, are increasingly widespread in many tropical landscapes (Tyukavina et al. 2018). This is reflected in the proportional decrease in large (>1,000 ha) deforested areas relatively to small clearings (6.25–50.00 ha) in the Brazilian Amazon, during the first decade of this millennium (Rosa et al. 2012). Indeed, in 2002–2009, small clearings accounted for over 80% of the annual deforestation in the region (Rosa et al. 2012). Since many old-growth specialist species avoid even narrow clearings (< 30 m wide) (Laurance et al. 2009), understanding the effects of these small-scale disturbances on primary forest species is paramount to frame effective conservation strategies.

Tropical bats are a species-rich and ecologically diverse group that plays key ecological roles in the maintenance of forest ecosystems (Farneda et al. 2018a; Kunz et al. 2011). Frugivorous bats act as effective seed dispersers, especially of pioneer plant species (de la Peña-Domene et al. 2014; Medellín and Gaona 1999), while insectivores play essential roles in the reduction of herbivory levels by controlling arthropod populations (Kemp et al. 2019; Maas et al. 2015). They are also highly responsive to human-induced landscape changes (Meyer et al. 2016), which, combined with their ecological diversity, abundance and relative ease of sampling make bats a potentially useful indicator group to investigate the effects of habitat disturbance on tropical ecosystems (Ávila-Gómez et al. 2015; Jones et al. 2009; Oliveira et al. 2017).

In the Neotropics, the responses of phyllostomid bats to second-growth and small-scale disturbances have received increasing attention (e.g. Avila-Cabadilla et al. 2012; Castro-Luna et al. 2007; Farneda et al. 2018b; Rocha et al. 2017b; Willig et al. 2007). The available evidence suggests that in second-growth and moderately disturbed habitats the abundance of frugivorous and nectarivorous phyllostomids tends to increase due to additional food resources (Farneda et al. 2015; Willig et al. 2007). In contrast, the abundance of gleaning animalivores tends to decrease in response to insufficient roosting and prey resources (Rocha et al. 2017a; Willig et al. 2019).

The magnitude of structural contrast between the original habitat and the habitat by which it is replaced has repeatedly been identified as a key factor driving changes in species abundance and richness, two commonly used metrics of species sensitivity to habitat disturbance (e.g. Mendenhall et al. 2014; Rocha et al. 2017a). However, analyses seldom address behavioural changes such as alterations in activity patterns and partitioning along the temporal niche axis, which may represent a more sensitive indicator of species' responses to habitat conversion.

How species distribute their activity over time is an important dimension of animal behaviour and provides valuable insights into their ecological niche and into how specific traits augment or constrain the accessibility to resources (Frey et al. 2019). Species' activity patterns are adapted to local biotic and abiotic conditions and can be adjusted in response to external stimuli such as predation risk, changes in resource availability and agonistic encounters with competitors (Halle 2000). Understanding how human-driven habitat changes impact temporal dynamics is key to uncover the mechanisms affecting niche partitioning and community structure. Furthermore, the discernment of the behavioural decisions that alter a species' activity can help unveil the degree of plasticity of animal assemblages to anthropogenic stressors. In tropical bats, changes in within-night activity patterns due to changes in habitat structure have been found

to be more pronounced in habitats which are structurally very different from old-growth forest, such as open agricultural areas (Presley et al. 2009a) and urban areas (Montaño-Centellas et al. 2015), than in more similar habitats, such as second-growth (Presley et al. 2009a) and forest subjected to reduced-impact logging (Castro-Arellano et al. 2009; Presley et al. 2009b).

Here, we used a periodical re-isolation of forest fragments at the Biological Dynamics of Forest Fragment Project in the Brazilian Amazon, to evaluate the effects of second-growth and small human-made forest clearings (i.e. areas created by the clearing of a 100 m wide strip of regrowth around old-growth forest fragments) on the within-night activity patterns of phyllostomid bats. Specifically, we hypothesized that: i) due to changes in resource availability and predation risk, temporal activity patterns should be habitat-specific; ii) due to different energetic requirements and foraging behaviours, changes in the activity budget should be guild- and species-specific; and, iii) within-guild patterns of temporal overlap should differ between habitat categories. Accordingly, we predicted that: i) decreasing levels of forest cover, from old-growth forest to second-growth and to forest clearings should lead bats to reduce their early evening activity to avoid periods of higher predation risk by visually oriented predators; ii) due to the lower energetic content of fruits compared with animal prey, frugivores should exhibit longer periods of foraging activity than gleaning animalivores; and iii) due to higher structural similarity, the level of temporal overlap should be more similar between old-growth forest and second-growth, than between old-growth forest and forest clearings.

## Materials and Methods

### Study area

This study was conducted at the Biological Dynamics of Forests Fragments Project (BDFFP), a whole-ecosystem fragmentation experiment located ca. 80 km north of Manaus (2°30'S, 60°W), Amazonas, Brazil (Supplementary Figure S1). The area is characterized by a mosaic of continuous old-growth rainforest (*terra firme* forest), old-growth forest fragments and secondary regrowth. Annual rainfall varies from 1,900 to 3,500 mm, with a dry season between July and November (Ferreira et al. 2017). In the early 1980s, 11 old-growth fragments (categorized into size classes of 1, 10 and 100 ha) were isolated from continuous old-growth forest by distances of 80-650 m. The fragments were initially surrounded by cattle pasture, but their rapid abandonment allowed pasture areas to develop into secondary forest. This second-growth matrix is dominated by *Vismia* spp. in areas that were cleared and burned, and by *Cecropia* spp. in areas that were cleared without fire (Mesquita et al. 2015). For a more detailed description of the study area and experimental manipulation see Laurance et al. (2011). Approximately every 10 years, a 100 m-wide strip of second-growth surrounding each of the 11 old-growth forest fragments is felled, forming clearings of areas ranging from 6.4 to 42 ha around each of the experimental forest fragments. Prior to this study, the most recent re-isolation of the old-growth forest fragments had occurred between 1999 and 2001 (Laurance et al. 2011). During this study, fragment re-isolation took place between November 2013 and March 2014, by clearing but not burning of a 100 m-wide strip of second-growth vegetation around each of the 11 old-growth forest fragments (Figure 1).

### Bat sampling

Before the creation of the experimental forest clearings, bats were sampled between August 2011 and June 2013 in 14 old-growth forest sites – six sites in continuous forest (located in Cabo Frio and Km 41 camps) and eight forest

fragments categorized according to their area as “1 ha” ( $n = 3$ ), “10 ha” ( $n = 3$ ) and “100 ha” ( $n = 2$ ) (located in Colosso, Porto Alegre and Dimona camps) (Supplementary Figure S1) – and eight second-growth sites, located 10 to 90 m from the forest fragments (in areas to be felled during the creation of the experimental forest clearings). The mean distance between old-growth forest sites was  $16,243 \pm 11,077$  (SD) m and between second-growth sites was  $12,780 \pm 9,336$  m. We estimated canopy cover (CC) as the average of four spherical densiometer readings within three  $100 \text{ m}^2$  ( $5 \times 20$  m) plots established 5 m from each side of the mist net transects (see Supplementary Table S1 for additional vegetation variables). Since CC was similar across old-growth sites in continuous forest ( $CC = 83.6 \pm 5.3$ , mean  $\pm 1SD$ ), 100 ha ( $CC = 83.6 \pm 0.9$ ), 10 ha ( $CC = 87.4 \pm 1.0$ ) and 1 ha fragments ( $CC = 84.6 \pm 2.0$ ) we clustered the six sites in continuous forest and eight forest fragments into the category “old-growth forest sites”.

After the creation of the experimental forest clearings, bats were sampled between April and November 2014 in the same 14 old-growth forest sites previously sampled in 2011-2013 and within the eight experimental forest clearings formed during the re-isolation of the old-growth fragments (these were the exact same sites covered by second-growth in 2011-2013) (Figure 1). In total 22 sites were sampled in both 2011-2013 (before the creation of the experimental clearings) and in 2014 (after the creation of the experimental clearings).

Each site was surveyed eight times before the creation of the experimental clearings and four times after the creation of the experimental clearings (respectively four and two times per season before and after the creation of the experimental clearings). Each site was surveyed using 14 ground-level mist nets ( $12 \times 2.5$  m, 16-mm mesh, ECOTONE, Poland), set next to each other along two transects of seven mist nets each. Transects were ca. 30-80 m apart. Mist nets were opened at dusk and remained open for six hours, being revised at intervals of 15 to 20 minutes. Species identification and taxonomy is in accordance with López-Baucells et al. (2016). Bat capture and handling was conducted following guidelines approved by the American Society of Mammalogists (Sikes and Gannon 2011) and in accordance with Brazilian conservation and animal welfare laws.

## Statistical analysis

We restricted species-level analyses to the ten most abundant phyllostomid species (for both 2011-2013 and 2014 only species with  $>30$  captures were evaluated – see results for species list). A measure of capture rate [bats per mist net hour (mnh); 1 mnh denoting one  $2.5 \times 12$  m mist net open for 1 hour] was obtained by standardizing the capture frequency of each site by the sampling effort (mnh). Captures were pooled into 12 sampling intervals of 30 min – e.g. bats captured between 18:00 h and 18:29 h were grouped into the same time interval. Due to unequal sampling sizes within habitat categories, we based the analyses on the per interval relative number of captures such that values for all 12 sampling intervals for each activity pattern amount to 1. Although we use capture frequency as a measure of capture rate, we note that captures might better reflect the distance travelled by bats than their activity *per se*.

Intraspecific and intraguild temporal activity patterns between habitat types were compared using Kolmogorov-Smirnov 2-sample tests. We used bat capture data from 2011-2013 to contrast activity patterns in old-growth forest and second-growth and bat capture data from 2014 to contrast activity patterns in old-growth forest and clearings. Additionally, to investigate any potential temporal changes in the activity patterns in old-growth forest (our baseline habitat), we contrasted the activity patterns in old-growth before the creation of the experimental clearings (2011-2013) and after the creation of the experimental clearings (2014). Furthermore, since the same sites were covered by

secondary forest and after the creation of the experimental clearings contrasted massively in habitat structure, we also compared the activity patterns in second-growth (2011–2013) to the activity patterns in clearings (2014). Kolmogorov-Smirnov 2-sample tests were also used to evaluate within-habitat pairwise differences in activity patterns, both between each pair of species within the same guild, and between guilds. The restriction of pairwise comparisons to members of the same trophic guild produces results that are ecologically more meaningful, as activity patterns are likely to reflect interspecific interactions and shared ecological constraints (Halle and Stenseth 2000). For analytical reasons only pairwise comparisons with an  $n > 30$  were evaluated. Kolmogorov-Smirnov 2-sample tests were conducted in R v3.0.2 (R Development Core Team 2013).

For each habitat category, the temporal overlap for frugivorous and gleaning animalivorous species was evaluated with Pianka (Pianka 1973) and Czechanowski (Feinsinger et al. 1981) indices. Assessment of all possible within-guild interspecific pairwise comparisons was conducted using the null model of assemblage-wide temporal overlap algorithm ROSARIO (Castro-Arellano et al. 2010). This algorithm was developed to analyse data of sequential and cyclical nature and works by shifting the continuous distribution of activity patterns over time, while retaining the empirical temporal autocorrelation in the activity of each species (Castro-Arellano et al. 2010). ROSARIO was implemented using the software TimeOverlap (Castro-Arellano et al. 2010) by means of 10,000 iterations. Significance ( $\alpha = 0.05$ ) was determined by relating each empirical index to the proportion of randomizations that obtained an overlap  $\geq$  to that of the empirical overlap value (observed). Significance therefore reflects higher temporal overlap than expected by chance.

## Results

During 22,070 mnh, we captured 3,971 phyllostomid bats (2,870 before and 1,101 after the creation of the experimental clearings) belonging to 46 species. Before the creation of the experimental clearings, six frugivorous (Brown Fruit-eating Bat *Artibeus concolor*, Great Fruit-eating Bat *A. lituratus*, Dark Fruit-eating Bat *A. obscurus*, Silky Short-tailed Bat *Carollia brevicauda*, Seba's Short-tailed Bat *C. perspicillata* and Dwarf Little Fruit Bat *Rhinophylla pumilio*) and four gleaning animalivorous species (White-throated round-eared bat *Lophostoma silvicolum*, Striped hairy-nosed bat *Mimon crenulatum*, Stripe-headed round-eared bat *Tonatia saurophila* and Fringe-lipped bat *Trachops cirrhosus*) met the criterion of  $>30$  captures established for species-level analyses. However, in the period after the creation of the experimental clearings, only four frugivorous species had over 30 captures (*Artibeus concolor*, *Carollia brevicauda*, *C. perspicillata* and *Rhinophylla pumilio*). For most of the aforementioned species, and for both periods (before and after the creation of the experimental clearings), there was a peak of activity within the first two hours after sunset, followed by a clear reduction of activity after that (Supplementary Figure S2 and S3).

### Temporal activity responses to habitat type

Differences in activity patterns between old-growth forest and second-growth (2011–2013) and between old-growth forest and clearings (2014) were negligible for most species and guilds (Table 1). At the species-level, the only two exceptions were the frugivorous *Artibeus obscurus* (Figure 2A), which exhibited significant differences between old-growth forest and second-growth (with activity being higher in the first two hours after sunset in old-growth forest compared to secondary forest), and *A. concolor* (Figure 2B), whose activity patterns differed between old-growth forest and clearings (with activity being higher in the first two hours after sunset in clearings compared to old-growth). Species-level activity patterns did not change significantly between old-growth forest sites sampled before (2011–2013)

and after (2014) the creation of the experimental clearings. Likewise, no differences were detected between sites that were second-growth in 2011–2013 and clearings in 2014 (Supplementary Table S2).

### Differences in temporal activity patterns between species and guilds

Significant within-guild interspecific differences in temporal activity patterns were restricted to the comparison between *A. obscurus* and *Rhinophylla pumilio* in second-growth (*R. pumilio*'s activity was relatively balanced through the six hours considered in our analyses, whereas *A. obscurus* exhibited a distinct peak of activity in the 2<sup>nd</sup> and 3<sup>rd</sup> hours after sunset; Figure 3A). At the guild-level, by contrast, differences were only observed for the comparison between frugivores and gleaning animalivores in old-growth and before the creation of the experimental clearings (activity levels of gleaning animalivores were more than double the activity levels of frugivores in the first 30 min after sunset; Figure 3B). Activity patterns for all species and guilds before (2011–2013) and after (2014) the creation of the experimental clearings are presented in Supplementary Figures S2 and S3.

### Temporal overlap

Frugivorous and gleaning animalivorous species presented greater temporal overlap than expected for all habitats both before (2011–2013) and after (2014) the creation of the experimental clearings (Table 2). For frugivores, the degree of temporal overlap (as given by the Pianka and Czechanowski indices) was always lower in old-growth forest than in second-growth and clearings. The contrast was particularly notable for the comparison between old-growth forest and clearings (Table 2). Compared to frugivores, the pattern of temporal overlap for gleaning animalivores between old-growth forest and second-growth was reversed (Table 2).

## Discussion

Numerous studies have reported detrimental effects of habitat modification on tropical forest bats (reviewed in Meyer et al. 2016). At the BDFFP, and despite the “soft” matrix composed of advanced secondary vegetation, bat species richness, abundance, evenness and assemblage composition varies across old-growth forest and the second-growth matrix (Farneda et al. 2015; Ferreira et al. 2017; Rocha et al. 2017a; Nuñez et al. in press). However, despite population-, guild-, and assemblage-level responses in occupancy and abundance, here we show that evidence for behavioural changes in the activity pattern of phyllostomid bats in response to variations in habitat structure between old-growth forest and both second-growth and forest clearings is limited.

### Temporal activity responses to habitat type

Intraspecific differences in temporal activity patterns between old-growth forest and modified habitats were restricted to *A. obscurus* in second-growth and *A. concolor* in forest clearings. In lowland Amazonian rainforest, Presley et al. (2009a) found the activity patterns of five out of eight frugivorous bat species differed between open agricultural areas and forested habitats in Iquitos (Peru), while Castro-Arellano et al. (2009) found that the temporal behaviour of three out of seven analysed bat species differed between reduced-impact logging sites and unlogged forest controls in Pará

(Brazil). In both studies, the greatest differences were observed soon after sunset and the authors proposed the alterations to reflect twilight avoidance of habitats with open canopy cover due to increased illumination and consequently higher susceptibility to bird predation. Reduced bat activity has been repeatedly linked with increased predation risk in habitats with reduced vegetation cover (Weinbeer et al. 2006), during twilight (Lima and O'Keefe 2013) and periods of brighter moonlight (Meyer et al. 2004; Ribeiro Mello 2009, but see Musila et al. 2019). Indeed, during the study period, the only three events of bird predation upon entangled bats took place in the first minutes after sunset (Rocha and López-Baucells 2014; Serra-Gonçalves et al. 2017). However, despite a noticeable reduction in activity of *A. obscurus* in the first 2 hours after sunset in second-growth relative to old-growth forest, we did not detect significant differences between the activity patterns in old-growth forest vs. clearings for any species other than *A. concolor*, for which the activity during the first two hours after sunset was higher in clearings than in old-growth forest. Our results thus do not lend strong support to the hypothesis of twilight avoidance in more open habitats due to increased predation risk.

Several, non-exclusive explanations can explain the apparent non-alteration of the activity patterns between old-growth forest and second-growth and particularly between old-growth forest and forest clearings. First, despite ample evidence of bat predation by nocturnal and diurnal birds and other visually oriented predators (de Moraes Costa et al. 2016; Mikula 2015; Mikula et al. 2016), levels of predation pressure might vary little between old-growth forest and second-growth, since the secondary forest matrix in our study system is <30 years old (Carreiras et al. 2014). Indeed, the maturation of the second-growth at the BDFFP is eroding the sharp differences in species composition previously found between old-growth and early second-growth habitats (Farneda et al. 2018b; Powell 2013; Quintero and Roslin 2005; Rocha et al. 2018; Stouffer et al. 2011) and, for at least two of the most common owl species (*Lophotrix cristata* and *Glaucidium hardyi*), occupancy levels are now indistinguishable between old-growth and second-growth forest (Sberze et al. 2010). Secondly, bats might adjust their flight patterns in order to avoid predation and fly closer to the understory vegetation, particularly in areas of reduced canopy cover. This “vegetation hugging” behaviour has been reported for several European vespertilionids (e.g. Schaub and Schnitzler 2007) and, in our system, could explain the increased capture rate of canopy species in the experimental clearings (Rocha et al. 2017c). This change in flight behaviour can be one of the possible explanations for the perceived higher activity of *A. concolor* in clearings than in old-growth forest during the first two hours after sunset. Lastly, given that surveys in 2014 began nearly immediately after the creation of the experimental clearings, local avian predators of bats might not have had enough time to adjust their predatory behaviour to the new landscape features.

### Differences in temporal activity patterns between species and guilds

Similarly to Presley et al. (2009a), no differences in activity patterns were detected between species pairwise comparisons in old-growth forest. However, at the guild-level, frugivores and gleaning animalivores were found to exhibit different activity dynamics, with the latter presenting a more marked activity peak during the first half-hour following sunset. In fact, more than 60% of the captured gleaning animalivores in old-growth forest were mist-netted within the 1<sup>st</sup> hour after dark whereas less than 40% of the sampled frugivores were captured over the same period. The poor nutritional and caloric content of fruits requires frugivores to consume the equivalent of twice their body weight in fruit each night (Charles-Dominique 1986) and, as a result, much of their activity budget is devoted to foraging (Henry and Kalko, 2007). On the other hand, gleaning animalivores feed on more energy-rich insects and small vertebrates (e.g.

Bernard 2002; Munin et al. 2012; Rocha et al. 2016) through a mix of hunting in continuous flight and sallying from perches (“hang-and-wait” strategy) (Kalko et al. 1999). Their activity peak seems to be related to the time of departure from the roost, which telemetry studies in Panama have revealed to be ~30 and ~60 min after sunset for respectively *Trachops cirrhosus* and *Lophostoma silvicolium*, two of our focal species (Kalko et al. 1999).

At the species level, the sole significant pairwise comparison was found for *A. obscurus* and *R. pumilio* in second-growth. The activity pattern of *R. pumilio* had already been recorded to differ from the temporal activity of other frugivorous bats in both open agricultural areas and second-growth in Peru (Presley et al. 2009a) and in closed canopy reduced-impact logging sites in the Brazilian Amazon (Presley et al. 2009b). Large frugivorous bats of the genus *Artibeus* are specialised on mass-producing fruiting trees that are often far apart, demanding long (>2-10 km) commuting distances. However, once a fruiting tree has been located and the bat initiates feeding, it seldom moves during the rest of the night (De Foresta et al. 1984; Henry and Kalko 2007). In contrast, the foraging strategy of the small *R. pumilio* (~9 g) relies on short (40 to 120 m) search flights for scattered epiphytic infructescences within a small foraging area (3.5-14.1 ha) (Henry and Kalko 2007). This, alongside smaller flight energetic efficiency in small-sized bats compared with larger ones (Speakman and Thomas 2003), explains the little variation in activity observed for *R. pumilio* during the six hours of nightly sampling.

### Temporal overlap

Temporal niche overlap was greater than expected by chance in all sampled habitats for both frugivorous and animalivorous bats. However, we found that the similarity between the temporal overlap in frugivores in old-growth forest and second-growth was greater than the similarity between old-growth forest and clearings. Additionally, there was a reduction in temporal overlap of gleaning animalivores in second-growth, comparatively to old-growth forest.

Temporal partitioning is one of the three primary mechanisms allowing the coexistence of closely related species (the other two being partitioning of space and food resources) (Schoener 1974) and, it is likely moulded by ecological determinants linked with species’ survival and reproduction (Presley et al. 2009a). The observed similar values for both Pianka and Czechanowski indexes in old-growth forest and second-growth suggests that for frugivorous bats, the second-growth matrix at the BDFFP may currently offer similar food availability to old-growth habitats. Consequently, frugivores may not need to adjust their activity budgets to compensate for scarcer or more scattered food resources. Indeed, the second-growth matrix at the BDFFP has a ~15 m tall canopy that although forming a dense layer that partly inhibits sunlight to reach the understory, still allows the understory to receive more sunlight than that of the old-growth (Bentos et al. 2013), therefore possibly allowing for higher fruit productivity. As such, potential higher predatory risks associated with reduced canopy cover might be counterbalanced by higher food availability, levelling the values for temporal niche overlap between old-growth forest and second-growth. The same justification can be given for the observed higher temporal overlap of this guild at clearings relative to old-growth forest. Forest gaps increase the abundance of fruits of early-successional plants targeted by frugivorous bats (Presley et al. 2009). The experimental clearings targeted by this study were surveyed too soon after tree felling to support a high abundance of fruits. However, increased light availability can augment growth and reproduction in tropical trees (Graham et al. 2003) and thus standing fruiting trees at the edge of the clearings might have increased fruit productivity due to additional exposition to sunlight. This food surplus might have lessened competition and consequently reduced temporal displacements. These results mimic the findings from bat assemblages inhabiting a disturbance gradient in the tropical Bolivian Andes, in

which temporal niche overlap among generalist frugivores was found to increase with disturbance (Montaño-Centellas et al. 2015).

For gleaning animalivores, the values of temporal overlap between old-growth forest and second-growth were considerably less similar than for the frugivore guild. This aligns with the findings of lower abundance of gleaning animalivorous bats in the BDFFP second-growth habitats (Rocha et al. 2017a) and indicates that while the *Vismia*- and *Cecropia*-dominated secondary vegetation allows for additional food resources for frugivorous bats, the regrowth habitats are still less suitable for gleaning animalivores than old-growth forest.

### Conservation implications

The analysis of behavioural responses in terms of changes in activity patterns in relation to forest disturbance complements studies focusing on the effects of habitat modification on species occupancy and abundance (Meyer et al. 2016). Consequently, this study contributes to a more comprehensive understanding of the impacts of the ongoing wave of forest loss and deterioration plaguing the humid tropics. Here, we show that although small differences can be detected for the disturbance-sensitive gleaning animalivorous bats, the activity patterns of most species in second-growth and small forest clearings, two of the most prominent habitat types in humanized tropical landscapes, vary little from the activity patterns in old-growth forest. This suggests that, at least for the most common species in our study landscape, regenerating second-growth seems to offer enough resources, allowing species to retain similar activity budgets between old-growth and regrowth habitats. Our results thus reinforce the conservation potential of secondary rainforests by showing that old (> 30 years) second-growth has little effect on the activity budgets of 10 of the most abundant Amazonian phyllostomids.

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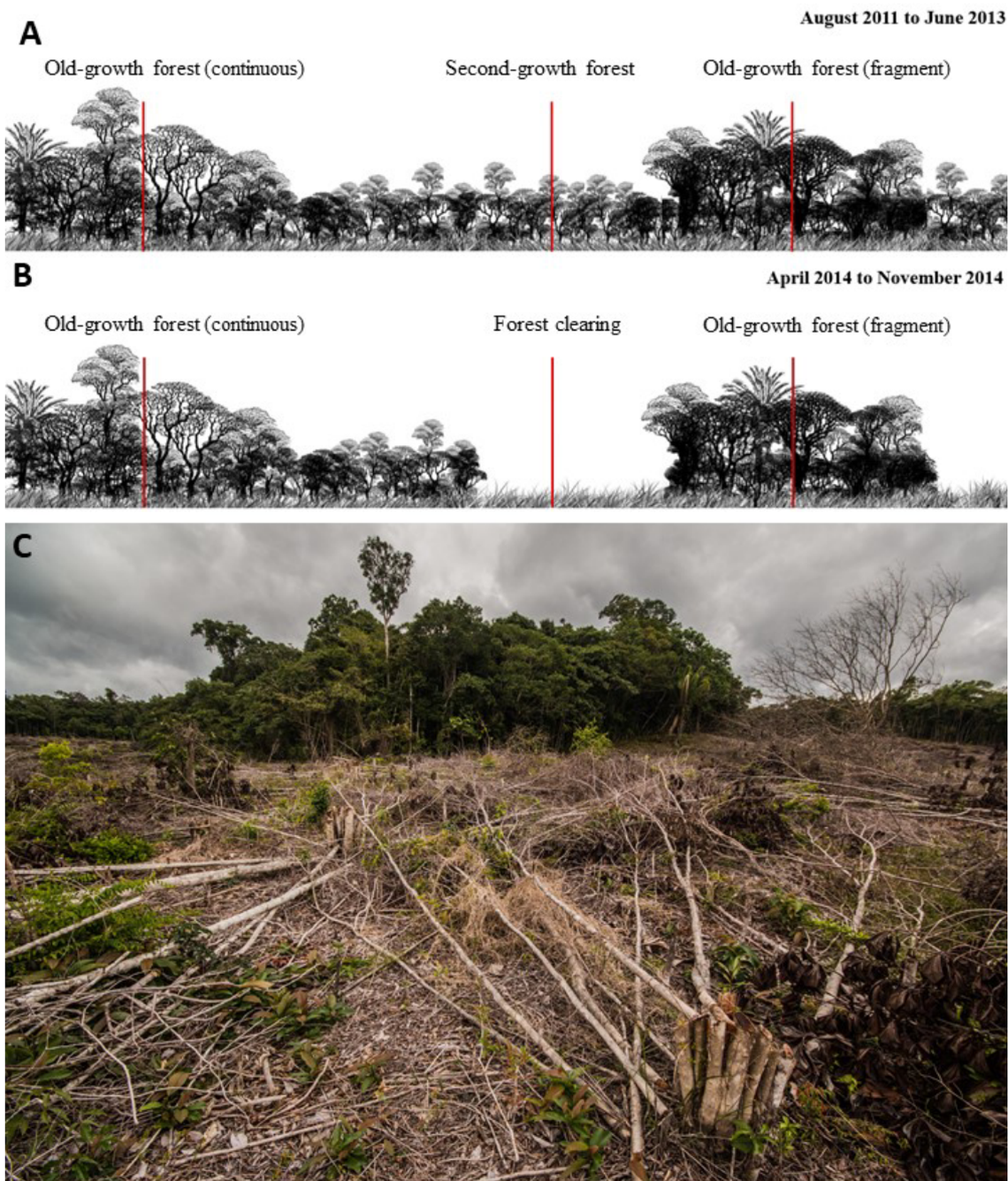
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**Table 1.** Comparison of bat activity patterns between old-growth forest and second-growth and between old-growth forest and clearings. Significant ( $P < 0.05$ ) results based on Kolmogoroff-Smirnov two-sample tests are highlighted in bold.

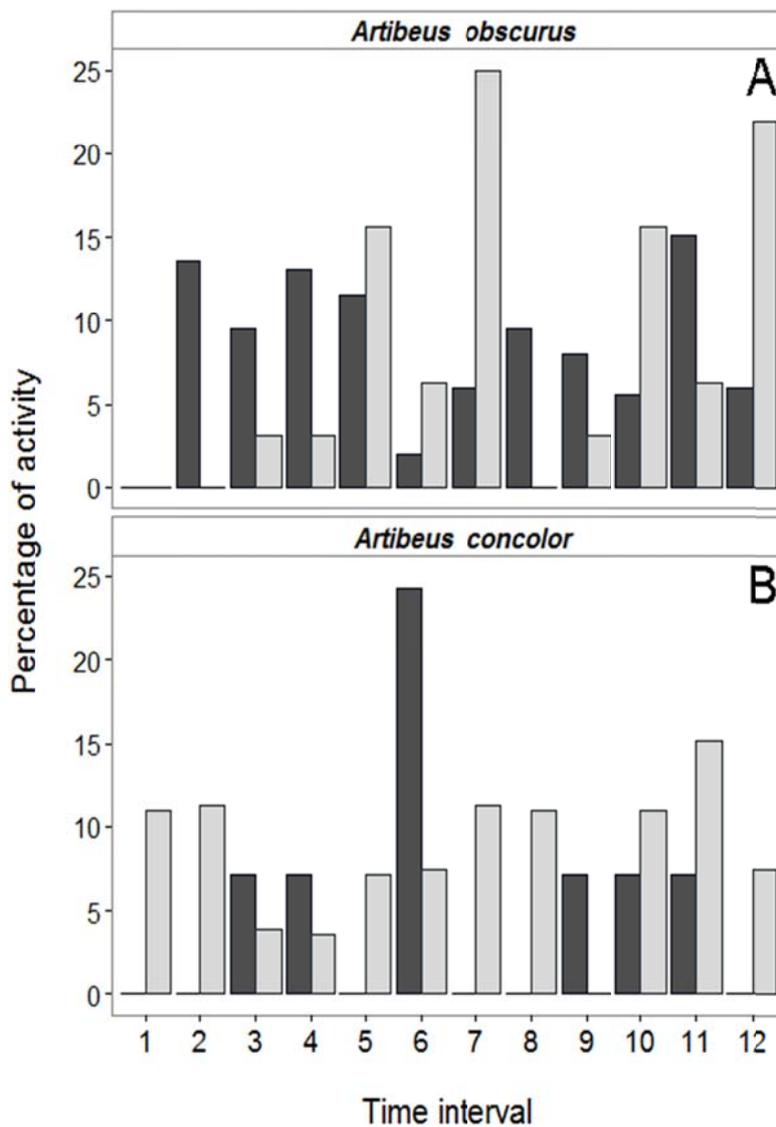
	Old-growth forest vs second-growth			Old-growth forest vs clearings		
	<i>n</i>	<i>D</i>	<i>P-value</i>	<i>n</i>	<i>D</i>	<i>P-value</i>
<b>Frugivores</b>						
<i>Artibeus concolor</i>	48	0.250	0.848	36	0.5833	<b>0.034</b>
<i>Artibeus lituratus</i>	31	0.417	0.249	-	-	-
<i>Artibeus obscurus</i>	111	0.583	<b>0.034</b>	-	-	-
<i>Carollia brevicauda</i>	84	0.333	0.518	35	0.250	0.848
<i>Carollia perspicillata</i>	1714	0.333	0.536	699	0.250	0.848
<i>Rhinophylla pumilio</i>	429	0.250	0.869	134	0.3333	0.518
<b>Gleaning animalivores</b>						
<i>Lophostoma silvicolum</i>	60	0.333	0.518	-	-	-
<i>Mimon crenulatum</i>	50	0.167	0.996	-	-	-
<i>Tonatia saurophila</i>	50	0.417	0.249	-	-	-
<i>Trachops cirrhosus</i>	81	0.333	0.517	-	-	-
TOTAL						
<i>Guild</i>						
Frugivores		0.250	0.869		0.250	0.869
Gleaning animalivores		0.1667	0.996		0.250	0.848

**Table 2.** Pianka and Czechanowski indices of temporal overlap for frugivores and gleaning animalivores in old-growth forest, second growth and forest clearings. Significant ( $P < 0.05$ ) results indicating greater temporal overlap than expected by chance are highlighted in bold.

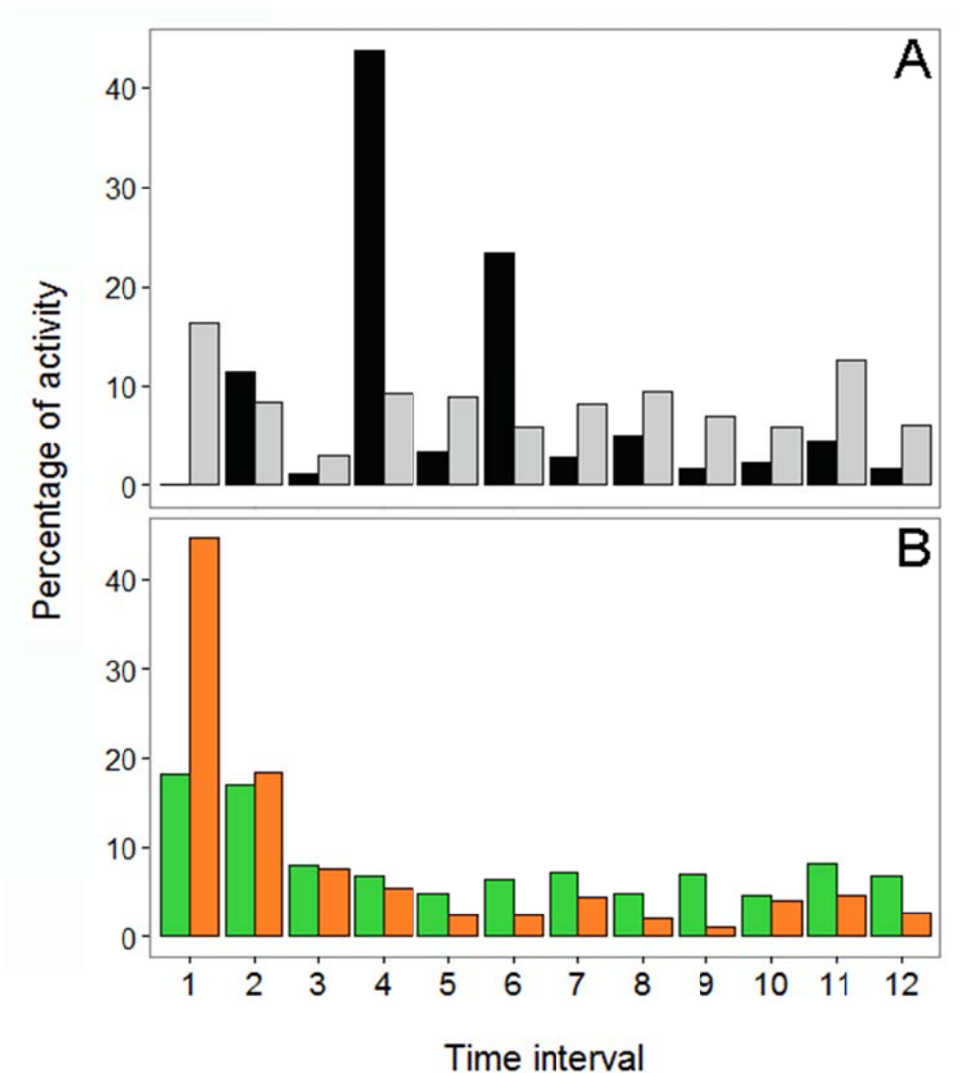
	Pianka index		Czechanowski index	
	Mean overlap	<i>P</i> -value	Mean overlap	<i>P</i> -value
<b>Frugivores</b>				
<i>2011-2013</i>				
Old-growth forest	0.778	<b>0.01</b>	0.675	<b>0.008</b>
Second-growth	0.799	<b>0.01</b>	0.713	<b>0.004</b>
<i>2014</i>				
Old-growth forest	0.572	<b>0.022</b>	0.513	<b>0.013</b>
Forest clearings	0.824	<b>0.033</b>	0.715	<b>0.048</b>
<b>Gleaning animalivores</b>				
<i>2011-2013</i>				
Old-growth forest	0.701	<b>0.002</b>	0.544	<b>0.001</b>
Second-growth	0.551	<b>0.041</b>	0.453	<b>0.022</b>



**Figure 1.** A) Schematic representation of the study area in 2011-2013, before the creation of the experimental clearings by the felling of 100 m of second-growth vegetation. B) Schematic representation of the study area in 2014, following the creation of the experimental clearings by the felling of 100 m of second-growth vegetation. C) Picture of an experimental forest clearing formed by clearing of second-growth vegetation surrounding a 1 ha old-growth forest fragment (picture taken in November 2013 by Adrià Lopéz-Baucells).



**Figure 2.** Temporal activity patterns for the two species that exhibited significant differences in activity between old-growth forest (dark grey) and (A) second-growth or (B) clearings (light grey) based on Kolmogorov–Smirnov 2-sample tests.



**Figure 3.** Temporal activity patterns for the pair of species and guilds that exhibited significant differences in activity patterns: A) *Artibeus obscurus* (black) and *Rhinophylla pumilio* (grey) in second-growth; B) frugivores (green) and gleaning animalivores (orange) in old-growth forest (before the creation of the experimental clearings).