Environmental structure affects taxonomic diversity but not functional structure of understory birds in the southwestern Brazilian Amazon

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ABSTRACT

Changes in environmental characteristics can affect habitat use by birds, impacting the number of individuals, number of species, and changing species composition and functional structure of assemblages. Metrics that evaluate the functional structure of biological assemblages constitute a complementary tool to the traditional taxonomic approach, because they quantify the differences between species by means of functional traits. We assessed the effect of environmental characteristics on the taxonomic diversity (species richness, species composition and number of individuals) and functional structure (functional richness, functional evenness, Rao’s functional diversity, and community-weighted mean traits) of bird assemblages in northern Mato Grosso state, in the southwestern Brazilian Amazon. We sampled birds in 32 plots. At each plot birds were captured using mist nets, and eight environmental variables were measured: canopy openness, leaf litter, elevation, number of trees in three classes of DBH, soil clay content, and distance to nearest stream. To evaluate functional structure, we measured seven morphological traits from individuals of each bird species. Habitat variables had a significant effect on taxonomic diversity. However, the general functional structure was not affected. Elevation and distance to nearest stream were the main variables driving changes in taxonomic diversity and had a minor effect on functional richness. The other metrics of functional structure were not significantly affected by the set of environmental variables. Our results suggest that the sampled bird assemblages exhibit some functional tolerance (redundancy) to small-scale environmental variation, implying certain resilience to ecosystem modification.

KEYWORDS: Aves, assemblage structure, habitat, functional richness

Estrutura ambiental afeta a diversidade taxonômica, mas não a estrutura funcional de aves de sub-bosque no sudoeste da Amazônia brasileira

RESUMO

Mudanças nas características ambientais podem influenciar o uso do habitat por aves, afetando o número de indivíduos, a riqueza de espécies e alterando a composição de espécies e estrutura funcional das assembléias. Medidas que avaliam a estrutura funcional de assembléias biológicas constituem uma ferramenta complementar à abordagem taxonômica, pois quantificam as diferenças entre as espécies através de seus atributos funcionais. Nós acessamos o efeito de variáveis ambientais sobre a diversidade taxonômica (riqueza, composição de espécies e número de indivíduos) e estrutura funcional (riqueza funcional, equabilidade funcional, diversidade funcional e média dos atributos ponderada pelas abundâncias) de assembléias de aves no norte do estado de Mato Grosso. Amostramos aves em 32 parcelas. Em cada parcela as aves foram capturadas usando redes de neblina e oito variáveis ambientais foram mensuradas: abertura de dossel, volume de serapilheira, elevação, número de árvores em três classe de DAP, teor de argila no solo e distância ao riacho mais próximo. Para avaliar a estrutura funcional nós tomamos sete medidas morfológicas de cada espécie de aves. Encontramos um efeito significativo das variáveis do habitat sobre a diversidade taxonômica, porém, a estrutura funcional, de maneira geral, não foi afetada. Elevação e distância ao rio mais próximo foram as variáveis que direcionaram as mudanças na diversidade taxonômica e tiveram um pequeno efeito na riqueza funcional. As demais métricas de estrutura funcional não foram afetadas significativamente pelo conjunto de variáveis ambientais. Nossos resultados sugerem que a assembléia estudada exibe certa tolerância funcional (redundância) à variação ambiental em pequena escala, implicando em alguma resiliência à modificação do ecossistema.

PALAVRAS-CHAVE: Aves, estrutura de assembléias, habitat, riqueza funcional

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INTRODUCTION

The structure of the environment constitutes a crucial component for bird assemblages (Wiens et al. 1987). Environments with greater structural complexity (e.g. rainforests) support high local diversity due to greater opportunities for resource specialization and niche partitioning, enabling the coexistence of species with different ecological features (MacArthur 1972; Tews et al. 2004; Mansor and Mohd Sah 2012). Forest birds are highly dependent on habitat and microhabitat conditions, particularly those related to vegetation structure (Terborgh et al. 1990; Cintra et al. 2006). Changes in the structure of vegetation within forests (e.g. canopy cover, density of trees) affects food availability, nesting sites, microclimate characteristics, vulnerability to predators and bird dispersal ability (Johns 1991; Mason and Thiollay 2001). These conditions can influence the use of habitats by Amazonian birds, affecting the number of individuals, the number of species and compositional change among assemblages (Johns 1991; Cintra et al. 2006; Rodrigues et al. 2016). The depth of leaf litter, for example, can affect the utilisation of the forest floor by many species of insectivorous birds (Pearson 1975), while canopy cover is crucial to the occurrence of birds that depend on a shaded understory (Banks-Leite and Cintra 2008).

Besides forest structure, topographical variation is also broadly considered in ecological studies because it can structure bird assemblages at both large (Quintero and Jetz 2018) and local scales (Cintra et al. 2006; Banks-Leite and Cintra 2008). Locally, variation in elevation is directly related to soil components, which influence the floristic and structural complexity of vegetation in tropical forests (Puig 2008), consequently affecting assemblages (Cintra et al. 2006; Banks-Leite and Cintra 2008; Cintra and Cancelli 2008; Cintra and Naka 2012).

Assemblage diversity is usually measured in terms of taxonomic diversity, which only accounts for species composition and abundance of individuals (Magurran 1988). Functional diversity can also be an effective tool in environmental monitoring studies, constituting a complementary tool to the traditional taxonomic approach (De Bello et al. 2010). This component of biodiversity quantifies the differences between species by means of functional traits, which are morphological, physiological, reproductive or behavioural characteristics of the organisms that affect their fitness (Tilman 2001; Petchey and Gaston 2006). They are often related to their environmental tolerances and competition for resources (Tilman 2001; Petchey and Gaston 2002, 2006). In this way, functional diversity allows to understand the competitive abilities of species and their patterns of co-occurrence, assembly rules, and the role of different functional traits in the ecosystem (Laureto et al. 2015).

Many studies have demonstrated the importance of habitat structure for the composition, richness and guild structure of birds in the Brazilian Amazon (Cintra et al. 2006; Banks-Leite and Cintra 2008; Cintra and Cancelli 2008; Lees and Peres 2010; Menger et al. 2017). Comparatively few studies on Amazonian birds have considered functional diversity, evaluating the effects of monoculture (Almeida et al. 2016), phytophysiognomies and biogeography (Almeida et al. 2018), urbanization (Lees and Moura 2017) and wildfire (Hidasi-Neto et al. 2012) on this trait. However, little is known about the response of forest bird functional diversity to forested habitat structure in the Amazon (e.g. Hidasi-Neto et al. 2012). The functional characteristics of the species determine the impact of biodiversity on ecosystem functioning, ecological patterns and structuring of biological assemblages (Tilman 2004; Laureto et al. 2015).

The purpose of this study was to assess the effects of habitat structure on the taxonomic and functional diversity (as derived from morphological traits) of Amazonian birds. Specifically, we tested the effects of a set of habitat variables (related to vegetation, elevation, soil clay content and distance to nearest stream) on richness, abundance, composition and functional structure of understory bird assemblages. Because changes in habitat conditions influence the structuring of ecological communities, we expected that both taxonomic and functional diversity would be affected by the environmental structure, especially those related to vegetation.

MATERIAL AND METHODS

Study area

The study was conducted at three sites (I, II, III) located in the municipality of Cláudia, in the northern region of the state of Mato Grosso, Brazil (Figure 1). The region is located within the Mato Grosso tropical dry forests ecoregion, which constitutes a transitional area between the Amazon rainforest and savanna-type cerrado vegetation (Carvalho 2006). This ecoregion represents approximately 10% of the Amazon biome and covers the northern part of Mato Grosso and the southern part of Pará state (Carvalho 2006). The climate is of type Am (Köppen 1948), characterised by high temperature (20°C – 33°C) and humidity, and an average annual rainfall of 2,000 mm, with higher rainfall (80%) from September to April, and a dry season from May to September, with precipitation below 60 mm (Vourlitis 2004). The three sampling sites have a history of selective logging, containing clearings that were used for the storage of harvested timber. Site I was logged 29 years, site II 15 years, and site III eight years prior to sampling. However, scarce evidence of impacts from the selective logging can be observed today at the sites, due to the use of reduced-impact logging practices, which resulted in lower environmental degradation.

Sampling design

Sampling was performed in 32 plots distributed in the three sites. Each plot was 250 m in length and located 1 km...
from neighbouring plots. Sites I and II had 12 plots each, distributed in a 1 x 5 km (5 km²) system of trails. Site III contained eight plots comprising a 1 x 3 km (3 km²) system of trails, as the size of the forest area was not large enough for the installation of a 1 x 5 km trail system (Figure 1c). Sampling design was based on the Rapid Assessment model for Long Term Ecological Research (RAPELD) adapted by Magnusson et al. (2005) and adopted by the Research Program on Biodiversity (PPBio) (for more information see http://ppbio.inpa.gov.br/). We captured birds in the 32 plots during two campaigns (March/April 2010 and July/August 2010), so that each plot was sampled twice overall.

**Bird sampling**

Birds were captured using 25 mist nets (36-mm mesh), each 10 m long and 2.5 m height, linearly set within the 250 m, covering an area of 625 m² in each plot. The sampling effort in each plot was 6,250 h m⁻² (sensu Straube and Bianconi 2002). The nets were arranged adjacent to each other along the plots from ground level and remained functioning from 06:00 to 11:00 am, and were checked every hour for the removal of captured birds. Each captured individual received a numbered metal ring provided by CEMAVE (ICMBio/SNA authorisation nr. 3144). Birds not identified in the field were collected (IBAMA/SISBIO license nr. 22923-2) and later identified by experts and deposited in the bird collection of Universidade Federal de Mato Grosso.

**Environmental variables**

Eight environmental variables were measured in each plot during the first sampling campaign (March/April 2010): canopy openness, litter volume, elevation, tree density for three DBH classes, soil clay content, and distance to nearest stream (Table 1). Canopy openness, which is an indirect measurement of the amount of light reaching the understory, was measured at five equidistant points in each plot using a concave spherical densiometer (Robert & Lemmon Forest Densiometer, model C). Measurements were taken along the cardinal points (east-west and north-south) and the average was calculated. The five measurements obtained at each plot were averaged and used in the analysis. Litter volume (in litres) was sampled at five points in each plot (every 50 m), with all litter from a 1 m² area collected at each point. The collected material was pressed three times in a graduated bucket using a wooden press that was released at a distance of 1.5 m from the bucket. The volume of litter measured from the five points was averaged for subsequent analysis. Elevation was measured at the beginning of each plot with a GPS Garmin® model 60CSx MAP altimeter, as the topographic level of each plot was roughly constant along its entire length. Tree density was estimated by counting all trees within three DBH (diameter at breast height at 1.30 m from the ground) classes, using different sub-plot areas for each DBH class: DBH = 1 − 9.9 cm (DBH 1-10) in sub-plots 250 m long x 4 m wide (0.1 ha); DBH = 10 − 29.9 cm (DBH 10-30) in sub-plots 250 m x 20 m (0.5 ha); and DBH ≥ 30 cm (DBH 30) in sub-plots 250

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**Table 1.** Mean, standard deviation (SD) and range (min – max) of values for environmental variables tested as predictors of understory bird assemblage patterns in 32 plots in northern Mato Grosso state, southwestern Brazilian Amazon. See Material and Methods for variable definitions.

| Variables                  | Mean ± SD (min – max) |
|---------------------------|-----------------------|
| Elevation (m)             | 357 ± 20.8 (289 – 388) |
| Canopy openness (%)       | 23.6 ± 2.7 (19.8 – 30.1) |
| Tree DBH 10-30 (ind 0.1 ha⁻¹) | 229.3 ± 38.5 (148 – 310) |
| Tree DBH 1-10 (ind 0.5 ha⁻¹) | 523 ± 116.2 (351 – 850) |
| Tree DBH 30 (ind ha⁻¹)    | 50.6 ± 13.2 (29 – 88) |
| Leaf litter volume (L)    | 3520 ± 355.7 (2880 – 4240) |
| Soil clay content (%)     | 37.81 ± 11.4 (22.7 – 58.3) |
| Distance to nearest stream (m) | 1238.1 ± 635.6 (130 – 2880) |
m x 40 m (1 ha). All previous variable measurements followed PPBio protocols (https://ppbio.inpa.gov.br/manuais). Soil samples were taken from five equidistant points at a depth of 0-10 cm in each plot following the methodology of EMBRAPA (1997) and sent to a laboratory for analysis of clay content. Distance from the mid-point of the 250-m trail to the nearest stream was calculated using georeferenced points in ArcGIS 9.0.

To eliminate highly correlated variables (|r| > 0.70) we analyzed a correlation matrix of the eight environmental variables using the Pearson coefficient (Supplementary Material, Figure S1). All variables were weakly correlated and none was eliminated. We also included the time since last logging in the analyses, to control for any possible effect of past human activities on bird assemblage structure. No model selected this predictor, and none of the habitat variables considered above showed any correlation with time since logging, confirming the absence of influence of this activity in the patterns observed.

**Functional traits**

For each recorded bird species we measured seven morphological traits related to resource use, as birds perform most of their ecological roles via resource acquisition (Sekercioğlu 2006). The following biometric parameters were measured, given their relationship to foraging strategies adopted by the species, and the amount and types of resources they exploit: body mass, wing length, bill length, bill width, bill height, tarsus length and hallux length. Body mass indicates the amount and size of food required for a given individual and strongly relates to metabolic rate, foraging behaviour, longevity and home range size (Luck et al. 2012). Wing length is related to capacity of movement, which in turn influences resource use, seed dispersal and nutrient cycling (Luck et al. 2012). Bill shape is a morphological adaptation related to food access and type of food consumed, influencing pollination effectiveness and handling of fruit and seeds (Luck et al. 2012). The hallux (opposable digit 1) is related to grasping and perching ability, and the tarsus can influence foraging behaviour as well (Luck et al. 2012). These measurements were obtained from the ornithological collection of the Museu Paraense Emílio Goeldi, in Belém, Pará, Brazil (see Supplementary Material, Table S1 for details). Six male specimens of each species were measured using a digital calliper with 0.1 mm precision, and the mean of these six values being used for analysis. The specimens were selected based on the proximity of their collecting localities to our study sites. All measurements were taken by one observer (SMA).

**Data analysis**

To examine the response of bird assemblages to environmental variables, we conducted a redundancy analysis (RDA) using the ‘vegan’ package in the R environment. This method calculates the variation in a set of response variables (e.g. species composition) that can be explained by a set of explanatory variables (e.g. environmental variables). The response variable was the matrix of species composition, and the eight environmental variables measured were used as predictor variables, employing a forward selection (Blanchet et al. 2008). The effect of the environmental gradients (each variable selected by the forward selection) on relative species abundance was shown through a direct gradient analyses (Gauch 1982).

We used different and complementary analytical approaches to describe assemblage functional structure and to understand how biodiversity interacts with habitat variables. We calculated functional richness (FRic), functional evenness (FEve), Rao’s functional diversity (FD), and community weighted means traits (CWM). FRic represents the amount of functional space occupied by all species of an assemblage according to Euclidean distances between them as calculated from the morphological traits (Villéger et al. 2008). FRic is independent of species abundance but the values monotonically increase with species richness, because more species fill a larger volume of trait space. This measure is considered the best performing index, mainly for communities with low richness (Mouchet et al. 2010). FEve quantifies the regularity between species in trait space, weighted by their abundance (Villéger et al. 2008). FEve is not biased by species richness and has a range between 0 and 1, with values close to 1 indicating very constant, and values close to 0 indicating very irregular functional distances between species or a less even abundance distribution among species (Villéger et al. 2008). FRic and FEve were calculated using the “dbFD” function in the ‘FD’ package in R (Laliberté and Legendre 2010). We standardized FRic values for the assemblage in each plot by expressing them as a proportion of the total volume filled by the regional species pool, here represented by all species recorded in this study.

To calculate the functional diversity of assemblages (FD), we calculated a matrix using the Euclidean distance between the morphological trait measures. Then, using the distance matrix with all species, we quantified the functional diversity of the birds in the plots through Rao’s quadratic entropy (Rao 1982; Botta-Dukát 2005). The Rao index is greater when there is a greater number of functionally unique species, because a new species in the community increases the species-abundance based diversity, while it decreases the average dissimilarity among species (De Bello et al. 2016). We calculated FD in the R environment using the ‘melodic’ function, considering abundance data (De Bello et al. 2016).

The community-weighted means (CWM) were calculated for each morphological trait of the assemblage. The CWM can define the functional composition of assemblages, and enables to evaluate the association between trait dominance and environmental gradients for a set of local communities (Duarte et al. 2018). CWM were calculated using the ‘matrix’
function in the R package 'SYNCSA' (Debastiani and Pillar 2012). To analyze the relationship between morphological traits of species and habitat structure we performed simple correlations combining data on environmental variables and the CWM of traits.

Generalised linear models (GLMs) using Gaussian distribution were used to identify the variables related to species richness, number of individuals (log-transformed) and functional richness. For each dependent variable we performed best subset models with all possible combinations of predictors (the eight explanatory variables), and used the Akaike criterion corrected for small sample sizes (AICc) to evaluate the models’ performances. We selected the one with lower AICc as the fittest model (Burnham and Anderson 2002). We ran the GLMs using the ‘bestglm’ package in R.

We tested for the occurrence of spatial autocorrelation in our data, but Moran’s I tests indicated a lack thereof for species richness (Moran’s I = 0.125, p = 0.066) and number of individuals (Moran’s I = 0.127, p = 0.063). Mantel’s statistic also revealed that there was no significant spatial autocorrelation of bird composition among the 32 plots (r = 0.033, p = 0.280). All analyses were conducted in the R environment, version 3.3.2.

## RESULTS

We captured 621 birds belonging to 72 species and 26 families (Supplementary Material, Table S1). Forward selection analysis presented two variables related to species composition of understory birds (elevation and distance to nearest stream), which together explained 14% of variation in species composition. Elevation alone explained 11.4% of the variation (Adj R² = 0.114, F = 4.87, p = 0.001, Table 2). All analyses were conducted in the R environment, version 3.3.2.

Few species were strongly associated with the extremes of the gradients, with most species occurring in the middle of the gradients of elevation and distance to nearest stream. Among the species that occurred more often near the streams were Galbula cyanicollis, Schateria neavea, Automolus subulatus and Corythopis torquatus (Figure 2). Species that occurred predominately in plots at lower altitudes were Corythopis torquatus, Arrermon taciturnus, Willisornis poecillinotus, Manacus manacus, Campylorhamphus procurvvoides, Campylopterus largipennis, Geotrygon montana, Tyrannutes stolzmanni and Phlegopsis nigromaculata. Species that occurred at higher altitudes were Xiphorhynchus guttatus, Lania cristatus, Hylocharis cyanus, Platyrinchus platyrrhynchos, Dacnis albiventris, Myrmotherula longipennis and Monasa morphoeus (Figure 3).

The models that best explained species richness and number of individuals consisted of three variables (dbh1-10 + elevation + distance to nearest stream), and explained 53% and 63% of the variation of these responses, respectively. For functional richness, the best explanatory model contained two variables (elevation + distance to nearest stream), which explained 15% of the variation. However, we failed to find any effect of habitat variables on functional evenness and Rao’s functional diversity in our analyses (Table 3). We also found no significant correlation between the habitat variables and CWM of traits (Table 4).

## DISCUSSION

Our results indicated significant effects of environmental structure on the taxonomic diversity (species composition, species richness and number of individuals) of avifauna found in the dry forests of the southwestern Brazilian Amazon. However, functional structure was not affected by the habitat variables we measured, with the exception of functional richness.

### Table 2. Environmental variables selected to explain bird species composition in 32 plots in northern Mato Grosso, southwestern Brazilian Amazonia.

| Variables                  | R² | R²Cum | AdjR²Cum | F     | p-value |
|----------------------------|----|-------|----------|-------|---------|
| Elevation                  | 0.143 | 0.143 | 0.114 | 4.870 | 0.001 |
| Distance to nearest stream | 0.053 | 0.197 | 0.140 | 1.875 | 0.009 |

### Table 3. Best models for the influence of environmental variables on species richness, number of individuals and functional structure of understory birds in 32 plots in northern Mato Grosso, southwestern Brazilian Amazonia.

|                              | Estimate | Std. Error | Z value | Pr(>|z|)² |
|------------------------------|----------|------------|---------|----------|
| **Species Richness**         |          |            |         |          |
| AICc = 170.6, AdjR² = 0.53   |          |            |         |          |
| (Intercept)                  | 11.77    | 0.62       | 18.96   | < 0.001 |
| dbh1-10                      | -1.28    | 0.63       | -2.02   | 0.053   |
| Elevation                    | -4.47    | 0.74       | -6.01   | < 0.001 |
| Distance to nearest stream   | 2.09     | 0.73       | 2.83    | 0.008   |
| **Number individuals**       |          |            |         |          |
| AICc = 212.11, AdjR² = 0.63  |          |            |         |          |
| (Intercept)                  | 20.03    | 1.19       | 16.73   | < 0.001 |
| dbh1-10                      | -3.085   | 1.29       | -2.38   | 0.02    |
| Elevation                    | -12.12   | 1.86       | -6.49   | < 0.001 |
| Distance to nearest stream   | 4.55     | 1.43       | 3.16    | 0.006   |
| **Functional Richness**      |          |            |         |          |
| AICc = 375.4, AdjR² = 0.15   |          |            |         |          |
| (Intercept)                  | 0.14     | 0.02       | 6.64    | < 0.001 |
| Elevation                    | -0.06    | 0.02       | -2.48   | 0.01    |
| Distance to nearest stream   | -0.02    | 0.02       | -1.01   | 0.32    |
| **Functional Evenness**      |          |            |         |          |
| (Intercept)                  | 0.66     | 0.01       | 50.5    | < 0.001 |
| Rao's Functional diversity   | 1.41     | 0.13       | 10.29   | < 0.001 |
| Best model is null model     |          |            |         |          |
Figure 2. Bird species captured in 32 lines of mist-nets in the Cláudia municipality, northern Mato Grosso state, southwestern Brazilian Amazon, in relation to distance to nearest stream gradient. The bars represent the presence and abundance of each species at each point of the gradient, whereas the height of the bars at the top represent the values of the predictor variable.
Figure 3. Bird species captured in 32 lines of mist-nets in the Cláudia municipality, northern Mato Grosso, southwestern Brazilian Amazon, in relation to elevation gradient. The bars represent the presence and abundance of each species at each point of the gradient, whereas the height of the bars at the top represent the values of the predictor variable.
Table 4. Relationship between CWM of traits and environmental variables from Pearson’s correlation analysis. All correlations were non-significant at α = 0.05.

|                   | dbh1.10 | dbh10.30 | dbh30 | Elevation | Canopy | Litter | Soil clay | Stream |
|-------------------|---------|----------|-------|-----------|--------|--------|-----------|--------|
| CWM-Body mass     | -0.19   | 0.04     | 0.03  | -0.1      | 0.11   | 0.03   | 0.08      | -0.26  |
| CWM-Bill length   | -0.15   | 0.15     | -0.01 | -0.1      | 0.18   | -0.1   | 0.07      | -0.27  |
| CWM-Bill height   | -0.19   | -0.03    | -0.01 | 0.18      | 0.31   | 0.14   | 0.19      | -0.04  |
| CWM-Bill width    | -0.26   | -0.01    | 0.01  | 0.11      | 0.21   | 0.12   | 0.05      | -0.06  |
| CWM-Tarsus length | -0.17   | -0.02    | 0.05  | -0.1      | 0.07   | -0.01  | -0.04     | -0.13  |
| CWM-Wing length   | -0.13   | -0.01    | -0.03 | 0.01      | 0.2    | 0.09   | 0.14      | -0.24  |
| CWM-Hallux length | -0.09   | -0.03    | 0.13  | -0.2      | 0.01   | 0.03   | 0.03      | -0.25  |

(FRic). Topographic variation and distance to nearest stream were more important for changes in bird species composition. Although birds are closely associated with vegetation structure, the variables related to vegetation (e.g. canopy openness) did not have any major influence on the diversity measures, contradicting our initial expectation. Only one variable related to vegetation structure was retained in the generalized linear models, dbh1-10 (number of trees with a diameter at breast height between 1 and 9.9 cm), and it composed the models that explained both species richness and number of individuals. All three models generated (for species richness, number of individuals and functional richness) retained elevation and distance to nearest stream.

In evaluating the influence of landscape features and forest structure, Cintra and Naka (2012) found no significant relationships between forest components (e.g. tree abundance, canopy openness) and species composition and number of bird species and individuals sampled. However, they observed that bird species composition was influenced by elevation and distance to forest streams, similar to what we found in the present study, and also leaf litter depth. We found that streams were important for the occurrence of some species. *Scletaria naevia* and *Corythopis torquatus*, for example, are known to occur along the banks of streams and forage in the humid foliage on the ground or in dense vegetation. In the central Amazon, small variations in topography (~ 75 m) have indirectly affected the distribution of plants (Costa et al. 2009) and animals, including birds (Menin et al. 2007; Cintra and Naka 2012; Menger et al. 2017). This parameter can influence soil characteristics (e.g. soil clay content) which in turn influence the structure of vegetation and floristic composition, ultimately affecting bird assemblages (Castilho et al. 2006, Pomba et al. 2012). We also observed this tendency in our results because, although presenting only moderate values of correlation, elevation influenced other environmental variables measured (canopy openness, leaf litter, distance to nearest stream and soil clay content) (Supplementary Material, Figure S1).

Functional richness was little affected by the habitat variables, indicating that an almost similar number of ecological functions were provided throughout the environmental gradient. In addition, we must consider that FRic increases with species richness (Mouchet et al. 2010). In this way, this index may have been influenced by the number of species, which, in turn, was significantly affected by environmental structure in our study sites. The other measures of functional structure (i.e. FEve, FDq and CWM traits) were not influenced by any environmental variables considered. Changes in FRic without changes in FEve indicate that, despite finding some impact of habitat variables on functional richness, there were no shifts in the range of functional traits as a function of environmental variation (Magnago et al. 2014). If FEve values had been influenced by environmental variation, this would indicate a decline or disappearance of some parts of the functional space (e.g. Mouchet et al. 2010).

The average dissimilarity between species (Rao’s functional diversity, FDQ) and average of each of the seven traits (CWM) were not influenced by the habitat variables. Other studies have found an influence of habitat in functional metrics of forest bird assemblages. However, these studies have evaluated strong gradients of disturbance or environmental changes, for example, the effects of monoculture on Amazonian birds (Almeida et al. 2016) and the reduction of riparian forest width on birds of the Atlantic Forest (Maure et al. 2018). We suggest that environmental differences among our plots were not strong enough to generate major changes in functional structure of the assemblages. This might also be explained by the coexistence of functionally similar species (Cornwell et al. 2006, Tews et al. 2004), which may allow a greater stability in the face of small environmental changes (Petchey and Gaston 2002). In addition, although functional structure can be influenced by the environmental filtering of certain characteristics of the birds, it also can be caused by other ecological mechanisms, such as interaction between species (e.g. interspecific competition; Gomez et al. 2018).

**CONCLUSIONS**

We observed that the species richness, number of individuals and composition of understory birds in 32 sampling plots in tropical dry forest in the southwestern Brazilian Amazon were influenced by local scale environmental variables. In contrast, environmental differences among plots were not enough to cause major alterations in the functional structure of bird assemblages. Thus, because taxonomic diversity changed without a variation in functional structure, we conclude that the Amazonian avifauna in this ecoregion can exhibit some functional redundancy to small-scale environmental variations, which may be important for the resilience of the bird assemblages.
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OLIVEIRA et al. Environmental structure affects taxonomic diversity but not functional structure of understory birds in the southwestern Brazilian Amazon

Table S1. Composition of understory birds from 32 plots sampled in the Cláudia Municipality, Mato Grosso, southern Amazon. Information on family, species, number and proportion of occupied plots (N. plots and % plots, respectively), and the number of individuals with excluded recaptures (N) are shown. The biometric measures are also presented: body mass in grams (live body mass of bird as shown on the specimen label), bill length measured from tip to nostril, bill height measured at proximal end of operculum, bill width measured at proximal end of operculum, mean value of tarsus length measured on specimen’s right-hand side, mean value of wing length measured on specimen’s right-hand side, and halluc length measured on specimen’s right-hand side. Species classification followed the Brazilian Committee of Ornithological Records (Comitê Brasileiro de Registros Ornitológicos, 2015).

| Family            | Species                                      | N (plots) | % plots | N | Body mass | Bill length | Bill height | Bill width | Tarsus length | Wing length | Hallux length |
|-------------------|----------------------------------------------|-----------|---------|---|-----------|-------------|-------------|------------|---------------|-------------|---------------|
| Columbidae         |                                               | 2         | 6.25    | 2 | 109       | 4.69        | 4.19        | 24.3       | 135           | 13.7        |
| Cuculidae         | Dromococcyx pavoninus Pelzeln, 1870           | 1         | 3.12    | 1 | 51.3      | 17.4        | 6.03        | 7.99       | 28            | 126         | 16.6          |
| Caprimulgidae     | Nyctiphrynus ocellatus (Tschudi, 1844)        | 6         | 18.8    | 8  | 37.7      | 6.36        | 3.6         | 6.39       | 13.8          | 120         | 5.97          |
| Trochilidae       |                                               | 4         | 12.5    | 5  | 2.02      | 22.4        | 2.63        | 2.04       | 2.09          | 33.4        | 3.03          |
|                   |                                               | 1         | 3.12    | 1  | 4.8       | 36.1        | 3.21        | 3.47       | 2.7           | 61.8        | 3.62          |
|                   |                                               | 3         | 9.37    | 5  | 7.75      | 25.2        | 2.71        | 3.16       | 2.45          | 74.6        | 4.41          |
|                   |                                               | 7         | 21.9    | 10 | 4.06      | 17.9        | 2.16        | 2.29       | 2.37          | 50.8        | 4.46          |
|                   |                                               | 1         | 3.12    | 1  | 3.48      | 16.7        | 2.17        | 2.49       | 2.5           | 48.8        | 4.13          |
|                   |                                               | 1         | 3.12    | 2  | 5.88      | 16.6        | 2.36        | 3.01       | 2.77          | 64.8        | 3.79          |
| Momotidae         |                                               | 1         | 3.12    | 1  | 118       | 29.3        | 12.1        | 11         | 25.3          | 133         | 15.2          |
| Galbulidae        |                                               | 2         | 6.25    | 2  | 25.1      | 41          | 7.56        | 6.7        | 9.26          | 82          | 5.49          |
| Bucconidae        |                                               | 4         | 12.5    | 6  | 41        | 19.3        | 8.78        | 8.23       | 17.6          | 87          | 7.73          |
|                   |                                               | 2         | 6.25    | 2  | 73        | 25.7        | 11.3        | 9.62       | 19.3          | 117         | 8.6           |
| Falconidae        |                                               | 1         | 3.12    | 1  | 160       | 12          | 13          | 10.8       | 52.5          | 166         | 22.5          |
|                   |                                               | 2         | 6.25    | 2  | 204       | 13.1        | 13.2        | 10.6       | 64           | 175         | 16.2          |
| Thamnophilidae    |                                               | 15        | 46.9    | 27 | 8.1       | 9.34        | 3.8         | 4.82       | 15.7          | 50          | 10.3          |
|                   |                                               | 1         | 3.12    | 1  | 15.6      | 10.1        | 3.85        | 5.44       | 24            | 56.4        | 11.7          |
|                   |                                               | 15        | 46.9    | 26 | 7.32      | 9.3         | 3.17        | 4.66       | 13.9          | 52.4        | 9.11          |
|                   |                                               | 2         | 6.25    | 3  | 7.92      | 9.13        | 3.49        | 4.87       | 13.7          | 61          | 8.29          |
Table S1. Continued.

| Species                              | (N plots) | % plots | N | Body mass | Bill length | Bill height | Bill width | Tarsus length | Wing length | Hallux length |
|--------------------------------------|-----------|---------|---|-----------|-------------|-------------|------------|---------------|-------------|---------------|
| Isleria hauseri (Sclater, 1857)      | 8         | 25      | 13| 9.54      | 8.38        | 3.44        | 4.95       | 18.3          | 50          | 11.1          |
| Thamnomanes caesius (Temminck, 1820) | 4         | 12.5    | 5 | 16.4      | 11.3        | 5.19        | 7.85       | 16.3          | 74          | 11.9          |
| Thamnophilus schistaceus d’Orbigny, 1835 | 2         | 6.25    | 2 | 17.1      | 11.9        | 5.86        | 7.74       | 17.7          | 65          | 11.3          |
| Thamnophilus aestivus Sclater, 1858  | 9         | 28.1    | 13| 21.7      | 12.2        | 6.62        | 7.47       | 20.9          | 69.8        | 14.1          |
| Scateria naevia (Gmelin, 1788)       | 1         | 3.12    | 2 | 19.7      | 14.8        | 4.23        | 6.23       | 22.9          | 69.8        | 14.4          |
| Mymelaestes ruficaudus (Helmiet, 1929)| 1         | 3.12    | 1 | 24        | 13.8        | 4.9         | 6.62       | 24.5          | 69.6        | 15.1          |
| Myrmoborus myotherinus (Spix, 1825)  | 12        | 37.5    | 18| 16.2      | 10.2        | 4.4         | 5.97       | 23.8          | 62          | 14.1          |
| Pygionela leucotona (Spix, 1824)     | 4         | 12.5    | 5 | 33.6      | 12.1        | 5.39        | 6.26       | 30.2          | 79.2        | 17.1          |
| Sciaphylax hemipelmaeoa (Sclater, 1857)| 13        | 40.6    | 23| 14.6      | 9.93        | 3.92        | 5.52       | 21.8          | 57.2        | 13.3          |
| Cercomacra nigrescens (Cabanis & Heine, 1859) | 2         | 6.25    | 2 | 18.3      | 11.6        | 5.11        | 6.94       | 24.3          | 66.2        | 13.9          |
| Hypocnemis striata (Spix, 1825)      | 6         | 18.8    | 15| 12.4      | 9.42        | 3.99        | 5.68       | 19.2          | 54.2        | 12.4          |
| Willasis poecilinotus (Cabanis, 1847) | 5         | 15.6    | 15| 16.9      | 11.2        | 4.7         | 6.89       | 21.8          | 69          | 15.6          |
| Phlegopsis nigromaculata (d’Orbigny & Lafresnaye, 1837) | 3         | 9.37    | 7 | 46.4      | 12.9        | 6.26        | 7.95       | 29.2          | 90.2        | 18.6          |
| Rhegmatanthis gymnonotus Ridgway, 1888 | 10        | 31.3    | 25| 26.2      | 12.3        | 4.87        | 6.71       | 26.5          | 77.8        | 15.9          |

Conopophagidae

Conopophaga australis (Gmelin, 1789)

Formicariidae

Formicarius colma Boddaert, 1783

Dendrocolaptidae

Dendrocincla fuliginosa (Vieillot, 1818)

Dendrocincla merula (Lichtenstein, 1829)

Dendrocolaptes nigricatus (Vieillot, 1818)

Sittaxieus nigricapillus (Vieillot, 1818)

Glyptornis spurus (Vieillot, 1819)

Xiphorhynchus elegans (Pelzeln, 1868)

Xiphorhynchus guttatus (Lichtenstein, 1820)

Camptothorax borealis (Lafrasay, 1850)

Xenopidae

Xenops minutus (Sparrman, 1788)

Furnariidae

Automolus rufipileatus (Pelzeln, 1859)

Automolus subulatus (Spix, 1824)

Automolus ochroloemus (Tschudi, 1844)

Syliaxius rutilans Temminck, 1823

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Table S1. Continued.

|amily           | Species                          | % plots | N plots | N Body mass | Bill length | Bill height | Bill width | Tarsus length | Wing length | Hallux length |
|----------------|----------------------------------|---------|---------|------------|-------------|-------------|------------|---------------|-------------|---------------|
| Pipridae       | Tyranneutes stolzmanni (Hellmayr, 1906) | 1       | 3.12    | 2          | 9.18        | 6           | 3.6        | 4.28          | 12.6        | 50.2          | 8.98          |
|                | Ceratopipra rubrocapilla (Temminck, 1821) | 19      | 59.4    | 38         | 11.4        | 5.95        | 3.57       | 3.93          | 13.8        | 59.8          | 9.24          |
|                | Lepidothrix nattereri (Sclater, 1865) | 27      | 84.4    | 87         | 6.32        | 6.47        | 3.95       | 4.27          | 12.9        | 49.4          | 6.83          |
|                | Manacus manacus (Linnaeus, 1766) | 1       | 3.12    | 1          | 16.1        | 6.92        | 3.76       | 5.6           | 19.3        | 49.6          | 11.6          |
|                | Machaeropterus pyrocephalus (Sclater, 1852) | 9       | 28.1    | 22         | 8.5         | 5.98        | 3.32       | 4.13          | 13          | 53.2          | 9.1           |
| Onychorhynchidae | Onychorhynchus coronatus (Statius Muller, 1776) | 5       | 15.6    | 7          | 15.1        | 15          | 4.71       | 7.07          | 14.1        | 80.8          | 11            |
|                | Terenotriccus erythrurus (Cabanis, 1847) | 2       | 6.25    | 2          | 6.76        | 5.59        | 2.67       | 4.09          | 13.4        | 50.8          | 8.95          |
| Tityridae      | Schiffornis turdina (Wied, 1831) | 11      | 34.4    | 15         | 25.8        | 9.59        | 5.58       | 6.66          | 19.1        | 88.4          | 13.6          |
| Platyrinchidae | Platyrinchus mystaceus Vieillot, 1818 | 1       | 3.12    | 1          | 9.6         | 6.85        | 3.27       | 8.44          | 15.4        | 57.4          | 9.31          |
|                | Platyrinchus platyrhynchos (Gmelin, 1788) | 4       | 12.5    | 4          | 11          | 7.75        | 4.47       | 11.5          | 12.5        | 63.8          | 8.58          |
| Rhynchocyidae  | Mionectes oleagineus (Lichtenstein, 1823) | 5       | 15.6    | 7          | 10.8        | 8.19        | 3.27       | 4.56          | 13.8        | 66.4          | 9.82          |
|                | Corythops torquatus Tschudi, 1844 | 1       | 3.12    | 1          | 15.3        | 9.17        | 3.86       | 5.56          | 24.1        | 65            | 12.2          |
|                | Hemitriccus minor (Snethlage, 1907) | 15      | 46.9    | 21         | 6.7         | 7.09        | 3.2        | 5.06          | 14.8        | 49.6          | 8.63          |
| Tyrannidae     | Myiopagis vindicata (Vieillot, 1817) | 1       | 3.12    | 1          | 10.6        | 6.75        | 3.65       | 4.76          | 14.4        | 58.6          | 9.12          |
|                | Ramphothrix ruficauda (Spix, 1825) | 2       | 6.25    | 3          | 19.6        | 10          | 5.03       | 8.2           | 15          | 76.8          | 9.62          |
| Vireonidae     | Hylophilus ochraceiceps (Sclater, 1860) | 1       | 3.12    | 1          | 10.1        | 8.86        | 4.22       | 5.08          | 14.5        | 58.8          | 10.4          |
| Troglodytidae  | Pheugopedius genibarbis (Swainson, 1838) | 4       | 12.5    | 4          | 19.4        | 12.4        | 4.38       | 5.25          | 21.1        | 62.6          | 15.4          |
| Turdidae       | Catharus fuscescens (Stephens, 1817) | 1       | 3.12    | 1          | 34          | 10.4        | 4.44       | 6             | 28.6        | 100           | 14.1          |
| Passerellidae  | Ammoman taedatus (Hermann, 1783) | 2       | 6.25    | 2          | 25.4        | 10.3        | 8.06       | 6.77          | 24          | 75            | 14.2          |
| Thraupidae     | Lanius cristatus (Linnaeus, 1766) | 3       | 9.37    | 3          | 21.7        | 10.8        | 6.5        | 7.28          | 18.1        | 81.2          | 11.4          |
|                | Dacnis albiventris (Sclater, 1852) | 1       | 3.12    | 1          | 11          | 6.85        | 3.06       | 4.8           | 10.9        | 61.5          | 8.52          |
| Cardinalidae   | Habia rubra (Vieillot, 1819) | 8       | 25      | 13         | 29.6        | 11.7        | 8.43       | 7.39          | 21.8        | 93.2          | 12.8          |
|                | Cyanoloxia rothschildi (Bartlett, 1890) | 3       | 9.37    | 3          | 24.3        | 13.5        | 12.2       | 11.7          | 18.9        | 81            | 12.4          |
Figure S1. Results of the correlation between habitat variables at the 32 sampling sites in the southwestern Brazilian Amazon. The diagonal presents the data distribution of the variables (identified by their acronyms). Squares below the diagonal present the scatterplots between variables, and squares above the diagonal present the values of the correlation coefficients (r) and their statistical significance (*p < 0.05, **0.05 > p > 0.001, *** p < 0.001).