


Diet specialisation reduces the occupancy of cocoa agroforests by diurnal raptors

JULIANA MONTEIRO DE ALMEIDA-ROCHA ,
JULIO AMARO BETTO MONSALVO and
LEONARDO DE CARVALHO OLIVEIRA

Summary

Habitat loss and fragmentation, especially due to agriculture expansion, pose the main threats to biodiversity conservation. While some species are able to survive and proliferate in human-modified habitats (winner species), others are highly dependent on well-preserved habitats, being more vulnerable to extinction (losers). Many raptors can be considered loser species due to their high trophic position, large home ranges, and low reproductive rates. Consequently, this group is frequently used to reflect the environmental quality of habitats and the biodiversity status of communities. Here we describe the diurnal raptor assemblages found in shaded-cocoa agroforests (*cabruças*) of Southern Bahia, Brazil, and the determinants of raptor occupancy in this system. We systematically surveyed diurnal raptors in 16 *cabruça* sites using complementary sampling methods – active search, playback, and point-count – and modeled species occupancy using covariates related to vegetation structure, landscape, management intensity, and biological traits (body mass, trophic level, and diet specialisation). We found a high number of species (at least 18) as able to use *cabruças* and a negative relationship between species occupancy and the degree of diet specialisation. Thus, our results suggest that *cabruças* have a high potential to mitigate the effects of forest loss and fragmentation on diurnal raptors, but this system alone may not support complete assemblages of this group. It highlights the need for the maintenance of forest remnants in the landscapes to increase species diversity at a regional scale and to assure the conservation of this group in Southern Bahian Atlantic Forest.

Introduction

Anthropogenic habitat loss and change are rapidly expanding worldwide (Venter *et al.* 2016, Watson *et al.* 2016), causing the loss, fragmentation and structural modification of natural habitats of many forest species. Species tolerance to habitat changes depends on their ecological flexibility to deal with the new conditions, such as alterations to the microclimate and vegetation structure, and this flexibility will strongly determine their ability to survive in human-modified habitats (Swihart *et al.* 2003, Tabarelli *et al.* 2012).

Among birds, raptors are expected to be highly affected by habitat disturbance due to their high trophic position in food webs, large home ranges, and low reproductive rates (Carrete *et al.* 2009). For example, raptors have shown to be underrepresented in agricultural lands where high inputs of pesticides are applied, since bioaccumulation and magnification of such substances impact on top predators (Espín *et al.* 2016). Consequently, this group is often used to indicate the biodiversity status of biological communities (Sergio *et al.* 2006, 2008) and the quality of the surrounding environment (Rodríguez-Estrella *et al.* 1998).

Three orders of diurnal raptors (Cathartiformes, Accipitriformes, and Falconiformes), encompassing 102 species, are found in the Neotropics (Remsen *et al.* 2017). Seventy-five species occur in Brazilian territory (Piacentini *et al.* 2015) and about 76% of these inhabit the Atlantic Forest (Ferguson-Lees and Christie 2006), one of the most threatened biomes in the world which is currently reduced to about 12% of its original extent (Ribeiro *et al.* 2009). The Corredor Central da Mata Atlântica (Central Atlantic Forest Ecological Corridor) is a region within this biome that combines a large extent of forest remnants with a high level of biodiversity (Galindo-Leal and Câmara 2003). Within this corridor, there are some Important Bird and Biodiversity Areas (IBAs), such as the Southern Bahia region (Bencke *et al.* 2006), where three of the six species of Accipitriformes that are threatened in Brazil can be found: the Crested Eagle *Morphnus guianensis*, the Harpy Eagle *Harpia harpyja* and the White-necked Hawk *Buteogallus lacernulatus* (MMA/ICMBio 2016). In Southern Bahia, much of the original forest was converted into cocoa *Theobroma cacao* plantations, mostly cultivated under the “cabruca” system (Landau *et al.* 2008). In *cabrucas*, the understorey is replaced by cocoa trees that grow under the canopy of predominantly native forest trees partly retained when cocoa is planted, plus trees that regenerated or have been planted later (Alves 1990).

Cabruca has been considered a wildlife-friendly system (Schroth *et al.* 2011) because it supports a significant portion of the native fauna (Cassano *et al.* 2009). Despite the vertical structural simplification compared to forests, *cabrucas* are much more complex than other farmlands such as annual crops (Alves 1990), thus acting as a suboptimal habitat for species from different taxonomic groups (Cassano *et al.* 2009). To our knowledge, there is no study specifically developed to investigate the use of *cabrucas* by raptors. Understanding how raptors are being affected in agricultural lands - particularly in conservation priority areas such as Southern Bahia - is a research priority in Brazil's National Action Plan for the Conservation of Birds of Prey (Soares *et al.* 2008). Also, the specific response of raptor species to environmental degradation gradients is still poorly understood in the Neotropics (Carrete *et al.* 2009), calling for new field data.

Since the carnivorous mammal fauna is very depauperate in Southern Bahia region (Cassano *et al.* 2012), raptors end up being the major (and sometimes the only) predators regulating populations of many species, including some native endangered species such as the Golden-headed Lion Tamarin, *Leontopithecus chrysomelas* (Oliveira and Dietz 2011). Predator-prey relationships can be unbalanced and sometimes lead to the local extinctions of prey in human-modified habitats (Irwin *et al.* 2009). Therefore, understanding the abundance and diversity of raptors using *cabrucas* is important to evaluate the trophic structure and the stability of biological communities in this agrosystem. This study aimed to describe the diurnal raptor assemblages of *cabrucas* from Southern Bahia and to identify features of this system that could determine occupancy by raptors. We expected a positive relationship between raptor occupancy and habitat structural complexity (Thiollay 1985, Jullien and Thiollay 1996), specifically: *cabrucas* with better-connected canopy, high-density, taller and large-diameter shade trees, as well as a lower density of cocoa trees and a low management intensity, would have a higher chance of being occupied by raptors. We also expected the percentage of vegetation cover in the landscape to positively affect raptor occupancy in *cabrucas* (Jullien and Thiollay 1996).

We added some biological traits such as body mass, dietary requirements, and use of vertical strata, to predict raptor occupancy in *cabruca* sites. We expected a lower occupancy of (1) species with higher dependence on intermediate forest strata, since these are replaced by cocoa trees in *cabrucas* (Johns 1998); (2) larger species, since they usually occur at low population densities and have large home ranges, being more sensitive to habitat disturbance (Gaston and Blackburn 1995); and (3) species with higher energetic requirements and dietary specialisation, since they are expected to be less tolerant to changes in the availability of food resources (Terraube *et al.* 2011).

Methods

Study area

The surveys occurred between August 2014 and May 2015, a period that includes the breeding season for most raptor species in Atlantic Forest (Mañosa *et al.* 2003, Zorzin 2011), and when these predators are expected to behave more conspicuously, increasing their detectability (Seymour *et al.* 2010, Monsalvo 2012). Also, according to field records, migratory species such as the Swallow-tailed Kite *Elanoides forficatus* and the Plumbeous Kite *Ictinia plumbea* can be found in the study region during the major part of this period (<https://en.wikiaves.com/>).

Sixteen *cabruca* sites separated by at least 11 km (mean 52 km; range 11–114 km) were surveyed, covering 11 municipalities (~ 4,000 km²) in the cocoa-growing region of Southern Bahia, Brazil (Figure 1). The dominant vegetation is tropical lowland rainforest (Oliveira-Filho and Fontes 2000), the mean annual temperature is 24°C and rainfall averages 2,500 mm/yr, with no marked seasonality (Mori *et al.* 1983).

Data collection

Species dataset

A subset of 46 species potentially occur in the study region (<http://www.listavermelhahabia.org.br/>; <http://www.iucnredlist.org/>; Del Hoyo *et al.* 2017). Taxonomy follows Remsen *et al.* (2017). For all of them, we compiled information in the literature about (1) diet preferences (proportional consumption of each food category: vertebrates, invertebrates, and fruits); (2) body mass (in grams); (3) percentage of foraging time in each stratum (ground, understorey, mid-high, canopy, and air); (4) degree of sensitivity to habitat disturbance (low, medium or high); (5) relative abundance throughout the range (uncommon, fairly common, common, or patchily distributed); and (6) conservation status according to IUCN ('Vulnerable', 'Near Threatened', 'Endangered', or 'Critically Endangered'). We derived two indexes based on diet preferences: (1) trophic level – the sum of the proportional consumption of each food category weighted by its energetic value, assuming a decreasing energy content from vertebrates to invertebrates to fruits; and (2) dietary specialisation – the number of different food categories in species dietary. Species list, ecological information, details about indexes calculation, and literature references are available in the online supplementary material (Tables S1 and S2).

Field survey

The field methodology was adapted from Granzinolli and Motta-junior (2008). A combination of active search, playback and point-counts were performed in each *cabruca* site by the same observer with the help of a field assistant. All sampling was carried out in two visits between 06h00 and 12h00, a period during which most species of diurnal raptors are active (Thiollay 1989, Mañosa *et al.* 2003), avoiding days with rain and strong wind due to a possible decrease in species detectability (Jones 2000).

Using Landsat images with 30 m resolution obtained from Google Earth (Google Inc. 2016), we designed two parallel linear transects of 800 m separated by 400 m in each *cabruca* site (Figure 2). Between 06h00 and 09h00, the active searches and playbacks were carried out in these transects, focusing on species which predominantly occur in forested habitat, such as the Crested and Harpy Eagles and the Forest-falcons *Micrastur* spp., plus soaring species that have not started their flight activities yet, such as the Grey-headed Kite *Leptodon cayanensis* and the Hawk-eagles *Spizaetus* spp. (Thiollay 1989). The active search was performed by walking through these transects at a constant velocity while identifying species using 10 x 50 binoculars and a digital voice recorder to record vocalisations whenever possible.

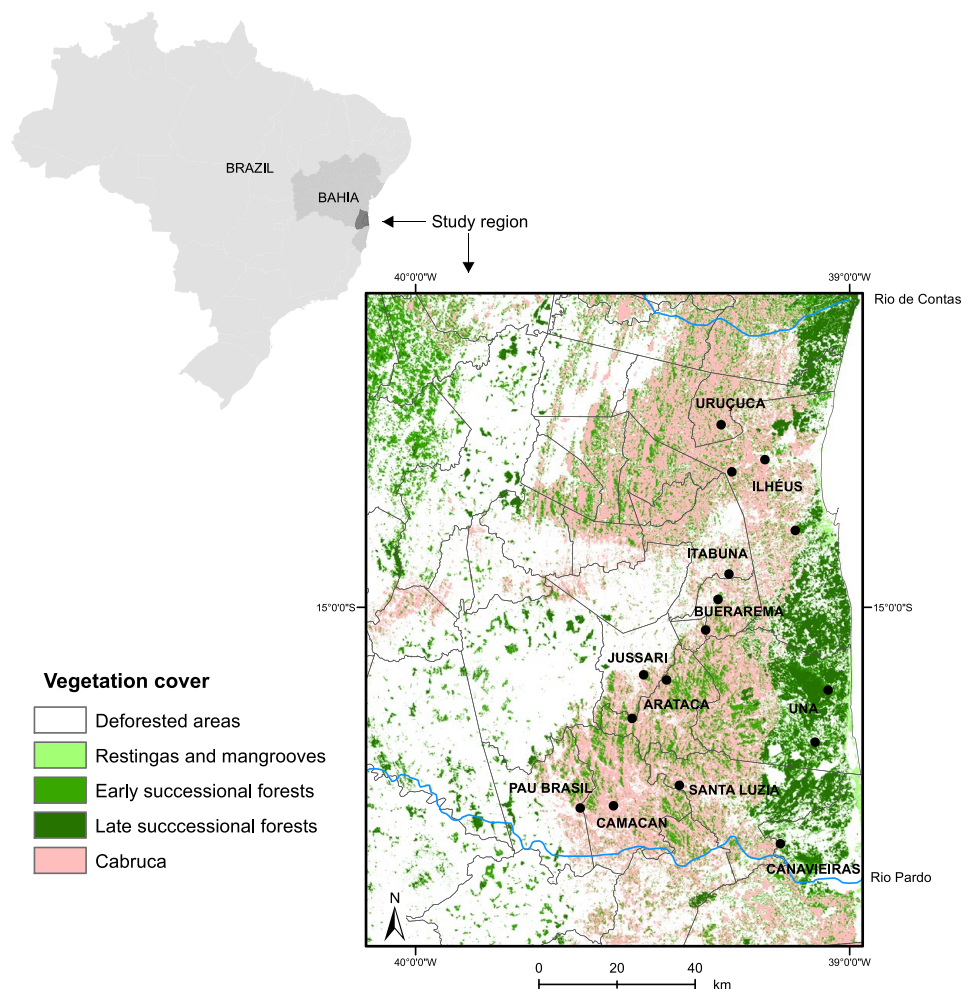


Figure 1. Vegetation cover in the study region and the location of the 16 *cabruca* sites surveyed in this study (black circles). Vegetation classification followed Landau *et al.* (2008).

Playback was performed to a set of species that are known to respond well to this technique: the Grey-headed Kite, Barred Forest-falcon *Micrastur ruficollis*, Collared Forest-falcon *M. semitorquatus*, Bicolored Hawk *Accipiter bicolor* and Black Hawk-eagle *Spizaetus tyrannus* (Zorzin 2011, JABM pers. obs.). This method consists of playing a recording of the species' vocalisation to attract individuals and stimulate intra-specific responses. Often playback resulted in inter-specific responses as well. Playback was performed at two opposite points of the sampling grid (Figure 2), separated by a mean linear distance of $665 \text{ m} \pm 160 \text{ SD}$, a placement that is consistent with those used by previous studies (Carvalho Filho *et al.* 2008/2009, Vázquez-Pérez *et al.* 2009). We used recordings available on Wiki Aves (<http://www.wikiaves.com/>), preferentially those made in the study region and avoiding aggressive vocalisations and pair duets. Using the aforementioned recorder coupled with a portable speaker, the recordings of all focal species were played in a pre-established order that took into account both body size and aggressive behavior, considering that large and fierce species could repel smaller ones (Mosher *et al.* 1990). Each vocalisation was played continuously for 3 min, holding the speaker at

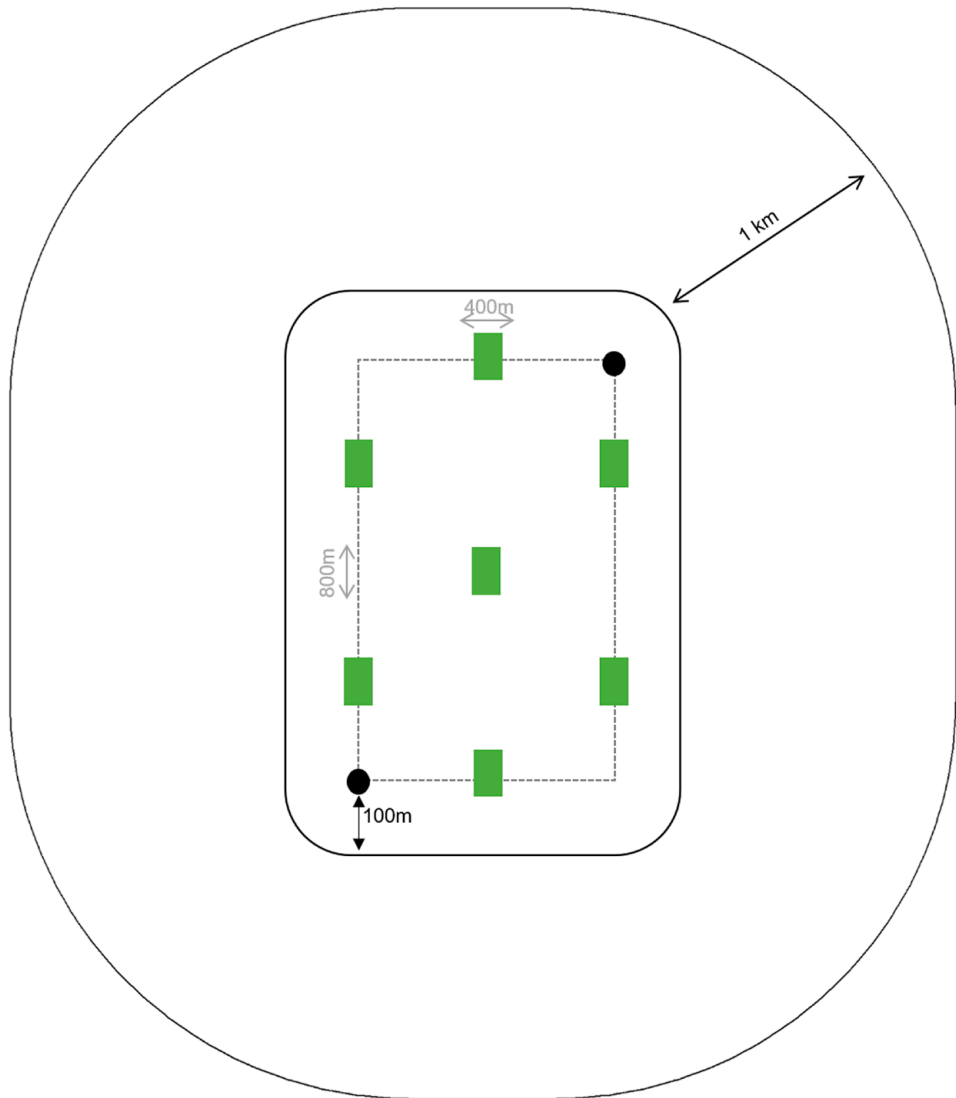


Figure 2. Scheme of the sampling grid designed in each *cabruca* site showing the linear transects (dashed gray line) where the active searches were performed, the location of the playback points (black circles) and the vegetation plots (gray rectangles). The transects are located at least 100m distant from fragment's edge and the vegetation cover was estimated within the area delimited by the 1-km radius.

approximately 2 m above the ground and rotating it over 360° at a constant speed, followed by a 3-min on-site wait.

Most raptors, such as the hawks *Buteogallus* spp. and the Mantled Hawk *Pseudastur polionotus*, often start soaring when thermals are well-formed, so the best period to perform the point-count methodology is between 09h00 and 12h00 (Thiollay 1989, Mañosa *et al.* 2003). During this period, we recorded all individuals visually or through their vocalisations from a fixed location. Point-counts for raptors are typically located in high places that offer a wide view of the canopy,

such as tops of hills adjacent to the study area (Mañosa *et al.* 2003). Such an approach was used whenever possible, but in six *cabruca* sites the relief was very flat so we located two complementary point-counts at the edge of the *cabruca* (~100 m from the edge and 690 ± 170 m apart from each other) and remained for 1h30 min at each point.

Except for single point-counts, the starting sampling point of each method was alternated on the second visit to ensure that species with different peaks of activity could be detected at all points (Jones 2000). Since most Atlantic Forest fragments are smaller than the mean territory sizes of Neotropical raptors (Thiollay 1989, Zorzin 2011), repeated detections of the same species at the same site were attributed to the same individual unless when more than one individual was observed simultaneously.

Habitat characterisation

In each *cabruca* site, seven 200-m² plots were placed at interspersed points of the sampling grid (Figure 2). Six features related to vegetation structure and *cabruca* management were collected in these plots, resulting in eight explanatory variables to be used in data analysis: (1) shade tree density; (2) mean shade tree diameter; (3) mean canopy height; (4) mean canopy connectivity; (5) heterogeneity of the vertical strata; (6) cocoa tree density; (7) shading level and (8) management intensity (Table S3). For the statistical analysis, we created an abundance index for each variable so that each of them was represented by a unique value per area. This index is the sum of the values obtained for each variable per plot.

Responses of diurnal neotropical raptors to forest fragmentation usually occur between the scales of 500 m and 1 km (Zorzin 2011). Based on the Landsat images, we created a buffer defined by a 1-km radius from the extreme points of the sampling grid (Figure 2) where we estimated the percentage of vegetation. Clearings within the buffers were visually identified and their area was subtracted from the total area of the buffer. This estimate included both *cabruca* and native forest due to the difficulty of accurately differentiating it in the images. Spatial analyses were performed on Quantum GIS 2.18.2 (www.qgis.org) using its interface with Google Earth (Google Inc. 2016). The distance measurements were made with the Raster Package (Hijmans *et al.* 2016) in R 3.3.1 (R Core Team 2016).

Data analysis

Occupancy modeling was performed in the program Mark version 8.x (White and Burnham 1999) to evaluate the influence of several variables on raptor occupancy. We used the single-season multi-species method to model the occupancy (Ψ) and detectability (p) of all species simultaneously, assuming an imperfect detection (Mackenzie *et al.* 2002). In this approach, 'species' and 'visits' represent the sample units and occasions, respectively (see chapter 9 of Mackenzie *et al.* [2006] for more details). For each species, a detection history was created based on the two independent visits per sampling method, which resulted in six occasions per species/site. This analysis included only the subset of species detected at least once at one of the sampling areas.

Single-season modeling assumes that the population is closed to changes in occupancy during the season (Mackenzie *et al.* 2006). Given that raptor home ranges are probably larger than all of our sampling areas (Thiollay 1989), this assumption may not have been achieved. To deal with this problem, we interpreted the estimated occupancy as the proportion of the area 'used' by the species rather than the true occupancy (Mackenzie *et al.* 2006). The detectability was interpreted as the probability of detecting the species when it is present in the area *and* using the sampling unit during the survey, assuming that movement of species through their home range is random (see similar interpretations in Keane *et al.* 2012 and Kalan *et al.* 2015).

Habitat features, landscape metrics, and biological traits were used as covariates to model Ψ and p in a series of competing models. Prior to analyses, we assessed their pairwise correlations through a Spearman correlation test using the R Stats package (R Core Team 2016), and excluded

those with correlation ≥ 0.60 , that is: percentage of foraging in intermediate strata (negatively correlated with body mass; $r = -0.63$); percentage of foraging in air (positively correlated with percentage of foraging in the canopy; $r = 0.67$); percentage of shade (positively correlated with canopy connectivity; $r = 0.60$ and vertical stratification; $r = 0.66$); and canopy height (positively correlated with canopy connectivity; $r = 0.84$) and diameter of shade trees ($r = 0.64$). We assessed the multicollinearity among the remaining variables through the Variance Inflation Factor (VIF) using the CAR package of R (R Core Team 2016). Since no variable had $VIF > 4$, we kept all of them for the analyses.

We finished with seven covariates to model p : (1) sampling method; (2) body mass; (3) percentage of foraging in the canopy; (4) canopy connectivity; (5) vertical strata heterogeneity; (6) shade tree density; and (7) cocoa tree density. We expected that: sampling methods would differ in species detectability; larger species would be more easily detected than smaller ones; species could be more or less easily detected depending on its preferred foraging strata; canopy connectivity, vertical heterogeneity, shade tree density, and cocoa density could interfere with visual obstruction and sound propagation, thus affecting species detectability during active searches and playbacks. We modeled Ψ as a function of 10 covariates: (1) shade trees density; (2) shade trees diameter; (3) canopy connectivity; (4) vertical strata heterogeneity; (5) management intensity; (6) cocoa tree density; (7) vegetation cover; (8) body mass; (9) trophic level; and (10) specialisation degree. We also included models where p and Ψ were held constant.

Since we were more interested in determining the importance of covariates rather than in the final estimates of Ψ and p , we built a model set based on all possible additive covariate combinations (Doherty *et al.* 2012), totalling 3,215 competing models. Then we calculated the cumulative AIC_c weight (w_+) for each covariate to interpret their relative importance, and the final estimate of Ψ was averaged among all competing models (Burnham and Anderson 2002). We investigated the fit of the most general model (the model with the greatest number of parameters) by estimating the overdispersion parameter c -hat using 10,000 bootstrap samples (Mackenzie and Bailey 2004) in the program PRESENCE 11.7 (<http://www.mbr-pwrc.usgs.gov/software/presence.shtml>).

Results

Raptor assemblages in cabrucas

A total sampling effort of 64 playback points (four per area), 91h11min of active search (mean effort: 5h40 per area; range: 4h48–6h45) and 96h of point-count (6h per area), resulted in the record of 18 species plus unconfirmed records of three species (Table S4). Some species for which we did not perform playbacks, as *Buteo* hawks, responded to the vocalisations of other species, therefore increasing their detectability (Table S5). Although we did not systematically survey for the smaller New World vultures – the Turkey Vulture *Cathartes aura*, Lesser Yellow-headed Vulture *C. burrovianus*, and Black Vulture *Coragyps atratus* – they were observed in all *cabruca* sites with all survey methods. Considering their constant presence in almost all sampling units and the difficulty in estimating their abundance, we decided not to include them in the analyses.

Among all recorded species, 44% (11 spp.), 48% (12 spp.), and 8% (2 spp.) are considered to be slightly, moderately and highly sensitive to habitat disturbance, respectively (Parker III *et al.* 1996). The most common species in our survey – ordered from the most to the less abundant – were the Southern Caracara *Caracara plancus*, the Zone-tailed Hawk *Buteo albonotatus*, the Laughing Falcon *Herpetotheres cachinnans*, and the Roadside Hawk *Rupornis magnirostris*. With the exception of the Zone-tailed Hawk – which is classified as uncommon, patchily distributed, and moderately sensitive to habitat disturbances – the aforementioned species are all considered to be common and slightly sensitive to habitat disturbance. The highly sensitive species detected in this study were the Black-and-white Hawk-eagle *Spizaetus melanoleucus* (recorded in four sites), and the Mantled Hawk *Pseudastur polionotus* (recorded in 10 sites).

The recorded species vary a lot in their dependence on forested habitats. The Collared Forest-falcon *Micrastur semitorquatus*, a forest-dependent species that uses mainly the low forest strata to forage, was recorded in two *cabruca* sites. Both records were made near regenerating forest plots, in places with a high density of shade trees. At the opposite extreme, the Savanna Hawk, *Buteogallus meridionalis*, a species that is typical of open areas, was observed during point counts in pastures near *cabruca* sites.

Some evidence of reproductive activity were observed during surveys, such as breeding pairs from six species (Black-and-white Hawk-eagle, and Black Hawk-eagle, Great Black-hawk *Buteogallus urubitinga*, Mantled Hawk, Southern Caracara, and Zone-tailed Hawk), juveniles of three species (Gray-lined Hawk, Roadside Hawk and King Vulture), and a family group of the Zone-tailed Hawk (two adults and one juvenile) foraging in the canopy near the edge of a *cabruca*.

Habitat characterisation

The structure of the *cabruca*s varied significantly among sites, with a mean \pm SD density of 623 ± 182 cacao trees/ha, and 182 ± 60 shade trees/ha (Table S6). Mean \pm SD diameter of shade trees is 37.2 ± 30.7 cm, mean \pm SD canopy height is 15.6 ± 2.6 m and mean \pm SD shading level is $73 \pm 10\%$. Hunting signs (traps, hunters and/or firearm blows) and logging signs (chainsaw noises and stumps) were recorded in 10 and nine of the 16 areas, respectively. The vegetation cover ranged between 73% and 96%.

Occupancy modeling

The most parsimonious model had a low AIC_c weight ($w_+ = 0.08$), reflecting the high number of competing models and a high degree of uncertainty about the best-ranked models (Table 1). There is no evidence of overdispersion ($\chi^2 = 73.37$; $P = 0.12$; $c\text{-hat} = 1.28$). Although the detectability of diurnal raptors was higher with the point-count method ($P = 0.24$; 95% CI: 0.19, 0.32) compared to the others ($P_{\text{both}} = 0.13$; 95% CI: 0.09, 0.18), the sampling method did not significantly affect detectability (Table 2). Detectability decreases as both the percentage of foraging in the canopy ($w_+ = 0.97$) and the density of shade trees ($w_+ = 0.82$) increase (Table 2; Figures 3 and 4), and there is a significantly negative correlation between occupancy and dietary specialisation ($w_+ = 0.60$; Table 2; Figure 5). The model-averaged estimate of Ψ was 0.66 (95% CI: 0.42; 0.84).

Discussion

In this study we described the diurnal raptor assemblages found in *cabruca*s of Southern Bahia, Brazil. We used occupancy modeling to investigate the role of vegetation structure, management intensity, landscape context, and biological traits in determining raptor occupancy in this agroecosystem. A significant percentage of the species expected to occur in the study region (~40%) was recorded in our study sites, and they have a high probability of occurrence in *cabruca* sites. It does not necessarily mean that *cabruca*s can assure the conservation of these species, but considering the high degree of deforestation and habitat loss in this region, *cabruca*s can act as a suboptimal habitat and assist in the persistence of diurnal raptor populations. Despite this promising perspective, there is a tendency towards impoverishment of communities in this system from the decrease of specialists.

Raptor assemblages in *cabruca*s

At least 18 (possibly 21) diurnal raptor species occur in the *cabruca*s of Southern Bahia. Most of these are considered to be naturally common, but species such as the Grey-headed Kite, Mantled Hawk, and Zone-tailed Hawk, which are classified as 'uncommon' by Parker III *et al.* (1996),

Table 1. Results for the 10 top-ranked models of occupancy (Ψ) and detection (p) probabilities of diurnal raptors in 16 *cabruças* of Southern Bahia, Brazil. 'AIC_c' is the Akaike information criterion corrected for small samples, 'ΔAIC_c' is the difference between the AIC_c value of each model and the top-ranked model, 'AIC_cW' is the Akaike weight, and 'Dev' is the deviance (model adjustment).

Model	AIC _c	Δ AIC _c	AIC _c W	Dev
{Ψ(DIE+TRO) p (FOR+DEN)}	1013.39	0.00	0.08	1001.05
{Ψ(BOD) p (FOR+DEN+BOD)}	1013.72	0.33	0.07	1001.38
{Ψ(DIE) p (FOR+DEN)}	1014.08	0.68	0.06	1003.84
{Ψ(DIE+MAN) p (FOR+DEN)}	1014.12	0.73	0.06	1001.78
{Ψ(DIE+BOD) p (FOR+DEN)}	1015.12	1.73	0.03	1002.78
{Ψ(DIE+DEN) p (FOR+DEN)}	1015.14	1.75	0.03	1002.80
{Ψ(BOD+MAN) p (FOR+DEN)}	1015.35	1.96	0.03	1003.01
{Ψ(DIE+COC) p (FOR+DEN)}	1015.49	2.09	0.03	1003.15
{Ψ(DIE+CON) p (FOR+DEN)}	1015.57	2.18	0.03	1003.24
{Ψ(DIE) p (FOR+DEN+BOD)}	1015.61	2.21	0.03	1003.27

DIE = dietary specialisation; DEN = shade tree density; FOR = percentage of foraging in the canopy; BOD = body mass; MAN = management intensity; COC = cocoa tree density; and TRO = trophic level.

were actually very frequent in our surveys. It is important to emphasise that such a classification is being used to allow comparison with other studies since we believe this is the only classification available for all Neotropical raptors to date. We recognise that it may suffer some limitations since it does not consider regional variations in factors that strongly determine species abundance. Anyway, since no systematic surveys of raptor assemblages have been made in forests from this region, we cannot conclude if the aforementioned species are particularly common in *cabruça*, or in the study region as a whole. However, as they are

Table 2. Cumulative AIC_c weight for covariates used to model occupancy (Ψ) and detection (p) probabilities of diurnal raptors in 16 *cabruças* of Southern Bahia, Brazil. The covariate effects (β parameters) were derived from the most parsimonious model including each covariate. LL and UL represent the lower and upper limits of the confidence interval (95%), respectively.

Covariate	Cumulative	β parameters		
	AIC _c Weights	Estimate	LL	UL
Detection (p)				
% of canopy foraging	0.97	-0.02	-0.03	-0.01
density of shade trees	0.82	-5.99	-8.43	-3.55
body mass	0.17	-0.72 x 10 ⁻³	0.00	-0.41 x 10 ⁻³
canopy connectivity	0.11	-0.01	-0.04	0.03
vertical stratification	0.10	-0.83 x 10 ⁻³	-0.02	0.03
method – active search	0.07	-0.24	-0.90	0.43
method - playback	0.07	-0.03	-0.68	0.63
density of cocoa trees	0.05	0.10	-1.01	1.20
Occupancy (Ψ)				
dietary specialisation	0.60	1.12	0.29	1.95
body mass	0.43	0.00	-0.83 x 10 ⁻³	0.01
management intensity	0.18	0.11	-0.03	0.26
canopy connectivity	0.16	-0.03	-0.11	0.05
trophic level	0.11	1.42	-0.31	3.16
density of shade trees	0.10	-3.7	-10.75	3.34
vertical stratification	0.07	0.02	-0.04	0.09
dbh of shade trees	0.07	0.00	-0.04 x 10 ⁻³	0.01
density of cocoa trees	0.06	1.40	-1.99	4.78
% of vegetation cover	0.04	0.82	-0.17	1.81

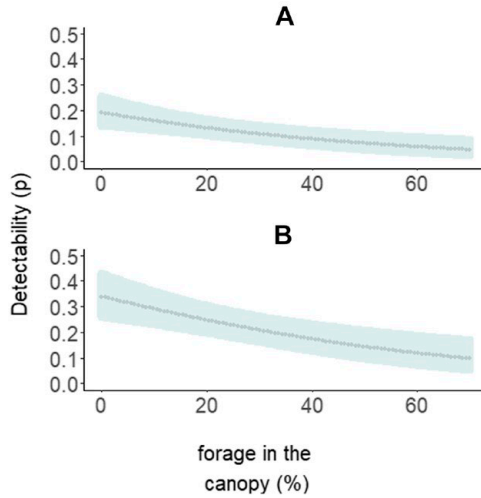


Figure 3. Relationship between diurnal raptor detectability and species' percentage forage in the canopy using three sampling methods: active search and playback (both represented in graph A), and point-count (B). Dotted lines and shaded areas represent the predicted values and the 95% confidence intervals, respectively.

frequently sighted throughout this region (even foraging in urban areas, in the case of the Zone-tailed Hawk; JABM pers. obs.), we believe that they can indeed be fairly common in Southern Bahia.

More than half of the recorded species are classified as moderately or highly sensitive to habitat disturbance, such as the Black-and-white Hawk-eagle and the Mantled Hawk. In fact, some authors have been questioning this classification by claiming that species-specific responses to habitat alterations depend on the disturbance context (Sergio *et al.* 2008; Alexandrino *et al.* 2016).

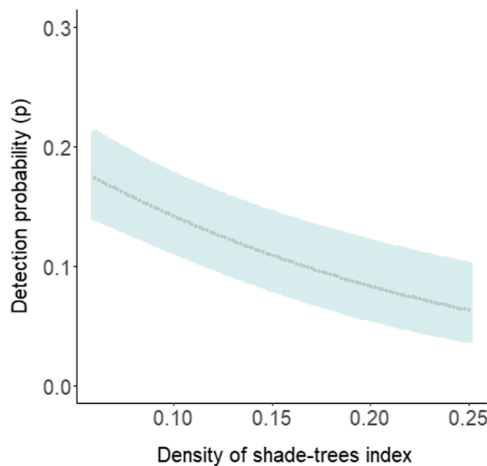


Figure 4. Diurnal raptor occupancy probability as a function of the density of shade trees in *cabruca* sites. The dotted line represents predicted values and the shaded areas the 95% confidence intervals.

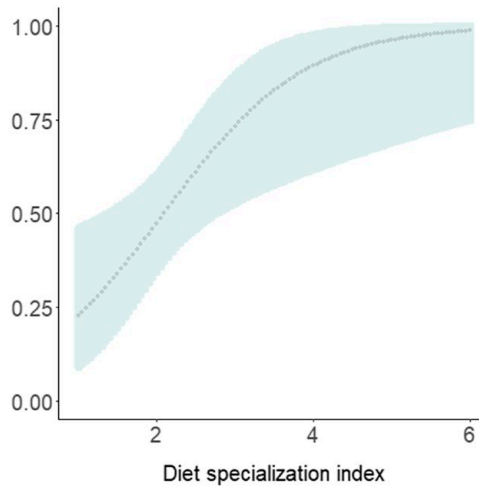


Figure 5. Diurnal raptor occupancy probability as a function of the diet specialisation index (= the number of food items in species diet). A high index value means a low degree of dietary specificity. The dotted line represents predicted values and the shaded areas the 95% confidence intervals.

As the classification proposed by Parker III *et al.* (1996) did not specify the type and intensity of disturbance under consideration, conclusions may have been based on habitats with levels of disturbance more intense than those observed in *cabruças*.

We did not record any globally threatened species, and only one 'Near Threatened' species, the Mantled Hawk, in the *cabruça* sites. The commonness of this species in *cabruça* is supported by unpublished reports from the Serra Bonita reserves complex in Camacan, located within the study region (JABM pers. obs.). An earlier study has found the Mantled Hawk to be extremely affected by forest fragmentation (Zorzini 2011), thus a high tolerance to *cabruças* may reveal an important role of this agroecosystem on mitigating the detrimental effects of forest fragmentation.

The observation of juveniles and breeding pairs from many species in *cabruça* sites was a very interesting and promising finding because it suggests they may be breeding there. However, specific studies on this topic are needed to confirm if these species are actually nesting in *cabruças* since it would contradict the high-quality requirements proposed for raptor breeding sites (Joenck *et al.* 2011, Canuto *et al.* 2012). There are some nesting records of Neotropical accipitrids such as the Short-tailed Hawk and the Harpy Eagle in areas that do not match such high-quality demands (Silva 2007, Monsalvo 2012), suggesting that a re-evaluation of this topic is necessary at least for some hawks and eagles (Monsalvo *et al.* 2018).

The failure to detect some species from the potential dataset may be partly explained by their high dependence on specific habitat types. The Snail Kite *Rostrhamus sociabilis* for example, is most commonly associated with wetlands, and the Long-winged Harrier *Circus buffoni* prefers open areas (Thiollay 2007). An adult pale-morph male of the latter species was occasionally observed while flying above a *restinga* (vegetation on the sandbanks) confirming their presence in the region despite the absence of records in the *cabruça* sites. A group of Swallow-tailed Kites, a migratory species that reproduces in the Atlantic Forest, was also occasionally recorded foraging under a pasture in the proximities of a *cabruça* site. To our knowledge, this is the first published record of this species in this region, and although we did not detect any individual within *cabruça* sites, such record suggests an ability to forage in landscapes totally dominated by this system. The same conclusion applies to our records of the Savanna Hawk, a common species in altered landscapes (Thiollay 2007).

The largest species from our potential dataset (the Crested and Harpy Eagles) are among the rarest Neotropical raptors (Thiollay 2007) and were also not recorded in our surveys. However, there are recent records of both species near and within our study region (Sánchez-Lalinde *et al.* 2011, Araújo *et al.* 2015, Suscke *et al.* 2016), including the observation of Harpy Eagles foraging in *cabruças* from the Serra Bonita reserves complex between 2012 and 2013, and the recent record of nesting in that same area (JABM unpublished data). Considering the rarity of such species, an increased survey effort would be necessary to best estimate their occupancy in *cabruca* sites.

Determinants for raptor occupancy in cabruças

Our analysis suggests that the availability of food resources may play a key role in determining the assembly of raptor communities in *cabruças*. Despite the certain degree of habitat complexity and heterogeneity that is preserved in *cabruças*, this system is indeed simplified compared to forests, so the overall prey availability is expected to be lower (August 1983). Consequently, specialists may be less likely to occupy *cabruças* than species with more diversified diet. Previous studies have already shown that specialist species are strongly affected by habitat disturbance and may be excluded from altered habitats, which tend to be dominated by generalists (Ferrer-Sánchez and Rodríguez-Estrella 2014). Thus, although *cabruças* may represent a suboptimal habitat for a significant number of diurnal raptor species, this system alone may not support complete assemblages of raptors.

Finally, although we failed to find any significant effect of habitat and landscape features on raptor occupancy, our results should be interpreted under the regional context. The *cabruca* sites surveyed in this study preserve a high density of shade trees, which is typical of traditional *cabruças* in this region (Schroth *et al.* 2015). There may be a critical density of trees, below which significant impacts on raptor occupancy may be observed, and it is therefore possible that the study did not include *cabruças* with such a low density of trees. Assuming this possibility is particularly important due to the current scenario in Southern Bahia of encouragement of *cabruca* management intensification. A state decree published by the Bahia Government in 2014 (N° 15.180; article 19) has sanctioned the management intensification of *cabruças* by allowing the reduction in shade tree density to very low levels (40 trees per ha). If implemented, it will promote a huge change and will likely decrease habitat heterogeneity in *cabruca* (Benton *et al.* 2003). Unfortunately, we cannot predict how raptors will respond to the changes that will occur in habitat features such as canopy connectivity or the diameter of shade trees. However, since a minimum level of habitat complexity and heterogeneity is necessary to support species-rich bird communities in altered habitats (Abrahamczyk *et al.* 2008, Philpott *et al.* 2008), raptor assemblages may not be sustained in very intensified *cabruças*, as already shown in other types of farmland (Donald *et al.* 2001, Carrete *et al.* 2009).

Conservation implications and recommendations

This study is a first attempt to understand the role of *cabruças* for diurnal raptor conservation in Southern Bahia. We found a significant number of diurnal raptor species are able to use *cabruças*, and evidence that some species may even be breeding in these areas. However, there is a tendency towards impoverishment of communities through the reduction in dietary specialists' occupancy. These findings emphasise the importance of considering human-modified habitats in conservation planning but highlight the need for the maintenance of forest remnants in the landscape to increase the regional species diversity.

Finally, a complete understanding of the role of *cabruças* in assisting raptor conservation depends on a better knowledge about raptor assemblages in the forests of this region. More surveys using appropriate sampling methods, as well as ecological and behavioural studies in both forests and *cabruca* sites, are crucial to better understand how raptors are dealing with habitat changes. Such knowledge would clarify the extent to which *cabruças* can contribute to raptor conservation at a

regional scale, and how different species can be specifically affected by *cabruca* intensification. Ultimately, since raptors play a key role as top predators, understanding such issues would allow a better understanding of how stable these communities are, and also of the long-term persistence probability of many species (including endangered species) in *cabruças* of Southern Bahia.

Supplementary Material

To view supplementary material for this article, please visit <https://doi.org/10.1017/S0959270919000017>

Acknowledgements

We are grateful to all landowners, especially Juliana Torres and Elizabeth Torres from Fazenda Almada, and to the Instituto Chico Mendes de Conservação da Biodiversidade (ICMBio) for the permission to work in the Biological Reserve of Una. We also thank Gastón Giné for the help with spatial analysis; Rodrigo Massara and Ana Maria Paschoal for the help with occupancy analysis; Victor Arroyo-Rodrigues and Stephen Ferrari for helpful comments on the manuscript; and all people who helped in field work, especially Edivaldo Francisco de Jesus and Jiomário dos Santos Souza. JMAR's doctoral studentship was funded by the Fundação de Amparo ao Pesquisador do Estado da Bahia (FAPESB) and the Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPQ) through the "Ciência sem Fronteiras" Program. JABM's master studentship was funded by CNPq. This study received funds from CNPq/CAPES through the Casadinho/PROCAD Project UESC-UFRJ (N°:552198/2011-0) and had the support of the Idea Wild (<http://www.ideawild.org>) through field equipment donations.

References

- Abrahamczyk, S., Kessler, M., Dwi Putra, D., Waltert, M. and Tschardt, T. (2008) The value of differently managed cacao plantations for forest bird conservation in Sulawesi, Indonesia. *Bird Conserv. Interantn.* 18: 349–362.
- Alexandrino, E. R., Buechley, E. R., Piratelli, A. J., Ferraz, K. M. P. M. B., Moral, R. de A., Sekercioglu, Ç. H., Silva, W. R. and Couto, H. T. Z. (2016) Bird sensitivity to disturbance as an indicator of forest patch conditions: An issue in environmental assessments. *Ecol. Indic.* 66: 369–381.
- Alves, M. C. (1990) The role of cocoa plantations in the conservation of the Atlantic Forests of Southern Bahia, Brazil, Thesis, University of Florida.
- Araújo, R. C., Silveira, L. F. and Luz, D. E. (2015) Rediscovery of the Crested Eagle *Morphnus guianensis* (Daudin, 1800) in the fragmented Atlantic Forest of Bahia, Brazil. *Rev. Bras. Ornitol.* 23: 25–28.
- August, P. V. (1983) The role of habitat complexity and heterogeneity in structuring tropical mammal communities. *Ecology* 64: 1495–1507.
- Bencke, G. A., Maurício, G. N., Develey, P. F. and Goerck, J. M. (2006) *Áreas importantes para a conservação das aves no Brasil. Parte I - Estados do domínio da Mata Atlântica.* São Paulo: SAVE Brasil.
- Benton, T. G., Vickery, J. A. and Wilson, J. D. (2003) Farmland biodiversity: is habitat heterogeneity the key? *Trends Ecol. Evol.* 18: 182–188.
- Burnham, K. P. and Anderson, D. R. (2002) *Model selection and multimodel inference: A practical information-theoretic approach.* 2nd edition. New York: Springer.
- Canuto, M., Zorzin, G., Carvalho-Filho, E. P. M., Carvalho, C. E. A., Carvalho, G. D. M. and Benfic, C. E. R. T. (2012) Conservation, management and expansion of protected and non-protected tropical forest remnants through population density estimation, ecology and natural history of top predators: case studies of birds of prey (*Spizaetus taxon*). Pp. 359–388 in P. Sudarshana,

- M. Nageswara-Rao and J. R. Soneji, eds. *Tropical Forests*. InTech. <http://www.intechopen.com/books/tropical-forests>
- Carrete, M., Tella, J. L., Blanco, G. and Bertellotti, M. (2009) Effects of habitat degradation on the abundance, richness and diversity of raptors across Neotropical biomes. *Biol. Conserv.* 142: 2002–2011.
- Carvalho Filho, E., Zorzini, G., Canuto, M., Carvalho, A. and Carvalho, G. D. M. (2008/2009) Aves de rapina diurnas do Parque Estadual do Rio Doce, Minas Gerais, Brasil. *MG Biota* 1: 4–43.
- Cassano, C. R., Barlow, J. and Pardini, R. (2012) Large mammals in an agroforestry mosaic in the Brazilian Atlantic Forest. *Biotropica* 44: 818–825.
- Cassano, C. R., Schroth, G. G., Faria, D., Delabie, J. H. C. and Bede, L. (2009) Landscape and farm scale management to enhance biodiversity conservation in the cocoa producing region of southern Bahia, Brazil. *Biodivers. Conserv.* 18: 577–603.
- Del Hoyo, J., Elliott, A., Sargatal, J., Christie, D. A. and De Juana, E. (2017) Handbook of the birds of the world alive. Available at: <http://www.hbw.com/>.
- Doherty, P. F., White, G. C. and Burnham, K. P. (2012) Comparison of model building and selection strategies. *J. Ornithol.* 152: S317–S323.
- Donald, P. F., Green, R. E. and Heath, M. F. (2001) Agricultural intensification and the collapse of Europe's farmland bird populations. *Proc. R. Soc. Lond. B* 268: 25–29.
- Espín, S., García-Fernández, A. J., Herzke, D., Shore, R. F., van Hattum, B., Martínez-López, E., Coeurdassier, M., Eulaers, I., Fritsch, C., Gómez-Ramírez, P., Jaspers, V. L. B., Krone, O., Duke, G., Helander, B., Mateo, R., Movalli, P., Sonne, C. and van den Brink, N. W. (2016) Tracking pancontinental trends in environmental contamination using sentinel raptors - what types of samples should we use? *Ecotoxicology* 25: 777–801.
- Ferguson-Lees, J. and Christie, D. A. (2006) *Raptors of the world: a field guide*. Princeton: Princeton University Press.
- Ferrer-Sánchez, Y. and Rodríguez-Estrella, R. (2014) Man-made environments relationships with island raptors: endemics do not cope with habitat changes, the case of the island of Cuba. *Biodivers. Conserv.* 24: 407–425.
- Galindo-Leal, C. and Câmara, I. G. (2003) Atlantic Forest Hotspot: An overview. Pp. 3–11 in C. Galindo-Leal and I. G. Câmara, eds. *The Atlantic Forest of South America: Biodiversity status, threats, and outlook*. Washington, DC: Island Press.
- Gaston, K. J. and Blackburn, T. M. (1995) Birds, body size and the threat of extinction. *Philos. Trans. R. Soc. London B* 347: 205–212.
- Google Inc. (2016) *Google Earth*. Mountain View, CA.
- Granzinolli, M. A. M. and Motta-junior, J. C. (2008) Aves de rapina: levantamento, seleção de habitat e dieta. pp. 169–187 in S. V. Matter F. Straube I. Accordi V. Piacentini and J. F. Cândido Jr., eds. *Ornitologia e conservação: Ciência aplicada, técnicas de pesquisa e levantamento*. Rio de Janeiro: Technical Books.
- Hijmans, R., van Etten, J., Cheng, J., Mattiuzzi, M., Sumner, M., Greenberg, J. A., Lamigueiro, O., Bevan, A., Racine, E. and Shortridge, A. (2016) Raster: Geographic data analysis and modeling. Available at: <http://cran.r-project.org/package=raster>.
- Irwin, M. T., Raharison, J.-L. and Wright, P. C. (2009) Spatial and temporal variability in predation on rainforest primates: do forest fragmentation and predation act synergistically? *Anim. Conserv.* 12: 220–230.
- Joenck, C. M., Zilio, F. and Mendonça-Lima, A. (2011) First record of breeding of the Ornate Hawk-Eagle (*Spizaetus ornatus*) in southern Brazil. *Hornero* 26: 163–166.
- Johns, N. D. (1998) Conservation in Brazil's chocolate forest: The unlikely persistence of the traditional cocoa agroecosystem. *Environ. Manage.* 23: 31–47.
- Jones, M. (2000) Study design. In C. Bibby M. Jones and S. Marsden, eds. *Expedition field techniques: Bird survey*. Cambridge, UK: BirdLife International.
- Jullien, M. and Thiollay, J.-M. (1996) Effects of rain forest disturbance and fragmentation: Comparative of the raptor and human-made changes community along natural gradients in French Guiana. *J. Biogeogr.* 23: 7–25.

- Kalan, A. K., Mundry, R., Wagner, O. J. J., Heinicke, S., Boesch, C. and Kühl, H. S. (2015) Towards the automated detection and occupancy estimation of primates using passive acoustic monitoring. *Ecol. Indic.* 54: 217–226.
- Keane, A., Hobinjatovo, T., Razafimanahaka, H. J., Jenkins, R. K. B. and Jones, J. P. G. (2012) The potential of occupancy modelling as a tool for monitoring wild primate populations. *Anim. Conserv.* 15: 457–465.
- Landau, E., Hirsch, A. and Musinsky, J. (2008) Vegetation cover and land use in the Atlantic forest of southern Bahia, Brazil, based on satellite imagery: a comparison among municipalities. Pp. 221–244 in W. Thomas, ed. *The Atlantic Coastal Forest of north-eastern Brazil*, New York: The New York Botanical Garden Press.
- Mackenzie, D. I. and Bailey, L. L. (2004) Assessing the fit of site-occupancy models. *J. Agric. Biol. Environ. Stat.* 9: 300–318.
- Mackenzie, D. I., Nichols, J. D., Lachman, G. B., Droege, S., Andrew, J. and Langtimm, C. A. (2002) Estimating site occupancy rates when detection probabilities are less than one. *Ecology* 83: 2248–2255.
- Mackenzie, D. I., Nichols, J. D., Royle, J. A., Pollock, K. K., Bailey, L. L. and Hines, J. E. (2006) *Occupancy estimation and modeling: inferring patterns and dynamics of species occurrence*. Burlington, MA: Elsevier Academic Press.
- Mañosa, S., Mateos, E. and Pedrocchi, V. (2003) Abundance of soaring raptors in the Brazilian Atlantic rainforest. *J. Raptor Res.* 37: 19–30.
- MMA/ICMBio (2016) *Brazil Red Book of Threatened Species of Fauna (Executive Summary)*. Brasília/DF: Instituto Chico Mendes/Ministério do Meio Ambiente.
- Monsalvo, J. A. B. (2012) Reprodução de *Buteo brachyurus* em um parque urbano de São Paulo, sudeste do Brasil. *Atualidades Ornitológicas* 170: 33–40.
- Monsalvo, J. A. B., Heming, N. M. and Marini, M. Â. (2018) Breeding biology of Neotropical Accipitriformes: current knowledge and research priorities. *Rev. Bras. Ornitol.* 26: 151–186.
- Mori, S. A., Boom, B. M., de Carvalho, A. M. and dos Santos, T. S. (1983) Southern Bahian moist forests. *Bot. Rev.* 49: 155–232.
- Mosher, J. A., Fuller, M. R. and Kopeny, M. (1990) Surveying woodland raptors by broadcast of conspecific vocalizations. *J. Field Ornithol.* 61: 453–461.
- Oliveira-Filho, A. and Fontes, M. (2000) Patterns of floristic differentiation among Atlantic Forests in southeastern Brazil and the influence of climate. *Biotropica* 32: 793–810.
- Oliveira, L. C. and Dietz, J. M. (2011) Predation risk and the interspecific association of two Brazilian Atlantic forest primates in Cabruca agroforest. *Am. J. Primatol.* 73: 852–860.
- Parker III, T. A., Stotz, D. F. and Fitzpatrick, J. W. (1996) Ecological and distributional databases. Pp. 111–410 in D. F. Stotz, J. W. Fitzpatrick, T. A. Parker III and D. K. Moskovits, eds. *Neotropical birds: Ecology and conservation*. Chicago: The University of Chicago Press.
- Philpott, S. M., Arendt, W. J., Armbrrecht, I., Bichier, P., Diestch, T. V., Gordon, C., Greenberg, R., Perfecto, I., Reynoso-Santos, R., Soto-Pinto, L., Tejada-Cruz, C., Williams-Linera, G., Valenzuela, J. and Zolotoff, J. M. (2008) Biodiversity loss in Latin American coffee landscapes: Review of the evidence on ants, birds, and trees. *Conserv. Biol.* 22: 1093–1105.
- Piacentini, V. de Q., Aleixo, A., Agne, C. E., Maurício, G. N., Pacheco, J. F., Bravo, G. A., Brito, G. R. R., Naka, L. N., Olmos, F., Posso, S., Silveira, L. F., Betini, G. S., Carrano, E., Franz, I., Lees, A. C., Lima, L. M., Pioli, D., Schunck, F., Amaral, F. R. do, Bencke, G. A., Cohn-Haft, M., Figueiredo, L. F. A., Straube, F. C. and Cesari, E. (2015) Annotated checklist of the birds of Brazil by the Brazilian Ornithological Records Committee/Lista comentada das aves do Brasil pelo Comitê Brasileiro de Registros Ornitológicos. *Rev. Bras. Ornitol.* 23: 91–298.
- R Core Team (2016) R: A language and environment for statistical computing. Available at: <https://www.R-project.org/>.
- Remsen, J. V. J., Areta, J. I., Cadena, C. D., Claramunt, S., Jaramillo, A., Pacheco, J. F., Pérez-Emán, J., Robbins, M. B., Stiles, F. G., Stotz, D. F. and Zimmer, K. J. (2017) A classification of the bird species of South America. American Ornithologists' Union.

- Available at: <http://www.museum.lsu.edu/~Remsen/SACCBaseline.htm>.
- Ribeiro, M. C., Metzger, J. P., Martensen, A. C., Ponzoni, F. J. and Hirota, M. M. (2009) The Brazilian Atlantic Forest: How much is left, and how is the remaining forest distributed? Implications for conservation. *Biol. Conserv.* 142: 1141–1153.
- Rodríguez-Estrella, R., Donazar, J. A. and Hiraldo, F. (1998) Raptors as indicators of environmental change in the Scrub Habitat of Baja California Sur, Mexico. *Conserv. Biol.* 12: 912–925.
- Sánchez-Lalinde, C., Vélez-García, F., Cornélio, A. C., Silveira, L. F. and Alvarez, M. R. (2011) Records of the Harpy Eagle (*Harpia harpyja*) in the Serra Bonita reserves complex, Camacan, Bahia, with evidence of breeding. *Rev. Bras. Ornitol.* 19: 436–438.
- Schroth, G., Bede, L. C., Paiva, A. O., Cassano, C. R., Amorim, A. M., Faria, D., Mariano-Neto, E., Martini, A. M. Z., Sambuichi, R. H. R. and Lôbo, R. N. (2015) Contribution of agroforests to landscape carbon storage. *Mitig. Adapt. Strateg. Glob. Chang.* 20: 1175–1190.
- Schroth, G., Faria, D., Araujo, M., Bede, L., Van Bael, S. a., Cassano, C. R., Oliveira, L. C. and Delabie, J. H. C. (2011) Conservation in tropical landscape mosaics: the case of the cacao landscape of southern Bahia, Brazil. *Biodivers. Conserv.* 20: 1635–1654.
- Sergio, F., Caro, T., Brown, D., Clucas, B., Hunter, J., Ketchum, J., McHugh, K. and Hiraldo, F. (2008) Top predators as conservation tools: ecological rationale, assumptions, and efficacy. *Annu. Rev. Ecol. Evol. Syst.* 39: 1–19.
- Sergio, F., Newton, I., Marchesi, L. and Pedrini, P. (2006) Ecologically justified charisma: Preservation of top predators delivers biodiversity conservation. *J. Appl. Ecol.* 43: 1049–1055.
- Seymour, A. S., Hatherley, G., Contreras, F. J., Aldred, J. and Beeley, F. (2010) Hatching synchrony, green branch collecting, and prey use by nesting Harpy Eagles (*Harpia harpyja*). *Wilson J. Ornithol.* 122: 792–795.
- Silva, F. H. A. (2007) Dieta do gavião-real *Harpia harpyja* (Aves: Accipitridae) em florestas de terra firme de Parintins, Amazonas, Brasil. Manaus: Instituto Nacional de Pesquisas da Amazônia/Universidade Federal do Amazonas.
- Soares, E. S., Amaral, F. S. R., Granzinolli, M. A. M., Albuquerque, J. L. B., Lisboa, J. S., Azevedo, M. A. G., Moraes, W., Sanaïotti, T. and Guimarães, I. G. (2008) *Plano de ação nacional para a conservação de aves de rapina/Instituto Chico Mendes de Conservação da Biodiversidade, Coordenação-Geral de Espécies Ameaçadas*. Brasília: ICMBio.
- Suscke, P., Verderane, M., de Oliveira, R. S., Delval, I., Fernández-Bolaños, M. and Izar, P. (2016) Predatory threat of harpy eagles for yellow-breasted capuchin monkeys in the Atlantic Forest. *Primates* 58: 141–147.
- Swihart, R. K., Gehring, T. M. and Kolozsvary, M. B. (2003) Responses of “resistant” vertebrates to habitat loss and fragmentation: the importance of niche breadth and range boundaries. *Divers. Distrib.* 9: 1–18.
- Tabarelli, M., Peres, C. A. and Melo, F. P. L. (2012) The “few winners and many losers” paradigm revisited: Emerging prospects for tropical forest biodiversity. *Biol. Conserv.* 155: 136–140.
- Terraube, J., Arroyo, B., Madders, M. and Mougeot, F. (2011) Diet specialisation and foraging efficiency under fluctuating vole abundance: A comparison between generalist and specialist avian predators. *Oikos* 120: 234–244.
- Thiollay, J.-M. (1985) Composition of falconiform communities along successional gradients from primary rainforest to secondary habitats. *ICBP Tech. Publ.* 5: 181–190.
- Thiollay, J.-M. (1989) Censusing of diurnal raptors in a primary rain forest: comparative methods and species detectability. *J. Raptor Res.* 23: 72–84.
- Thiollay, J.-M. (2007) Raptor communities in French Guiana: distribution, habitat selection, and conservation. *J. Raptor Res.* 41: 90–105.
- Vázquez-Pérez, J. R., Enríquez-Rocha, P. L. and Rangel-Salazar, J. L. (2009) Diversidad de aves rapaces diurnas en la Reserva de la Biosfera Selva El Ocote. *Rev. Mex. Biodivers.* 80: 203–209.
- Venter, O., Sanderson, E. W., Magrath, A., Allan, J. R., Beher, J., Jones, K. R., Fekete, M., Levy, M. A., Possingham, H. P., Laurance, W. F., Wood, P. and Watson, J. E. M. (2016)

- Sixteen years of change in the global terrestrial human footprint and implications for biodiversity conservation. *Nat. Commun.* 7: 12558.
- Watson, J. E. M., Shanahan, D. F., Marco, M. Di, Sanderson, E. W. and Mackey, B. (2016) Report catastrophic declines in wilderness areas undermine global environment targets. *Curr. Biol.* 26: 2929–2934.
- White, G. C. and Burnham, K. P. (1999) Program MARK: survival estimation from populations of marked animals. *Bird Study* 46: 120–139.
- Zorzín, G. (2011) Os efeitos da fragmentação da mata atlântica sobre a riqueza e abundância de accipitriformes e falconiformes na zona da mata de Minas Gerais. Universidade Federal de Viçosa.

JULIANA MONTEIRO DE ALMEIDA-ROCHA*

Applied Ecology and Conservation Lab, Programa de Pós-Graduação em Ecologia e Conservação da Biodiversidade, Universidade Estadual de Santa Cruz, Ilhéus, BA, 45662-900, Brazil.

JULIO AMARO BETTO MONSALVO

Laboratório de Ecologia e Conservação de Aves, Programa de Pós-graduação em Ecologia, Universidade de Brasília, DF, 70910-900, Brazil.

LEONARDO DE CARVALHO OLIVEIRA

Faculdade de Formação de Professores, Universidade do Estado do Rio de Janeiro, São Gonçalo, RJ, 24435-005, Brazil; and Bicho do Mato Instituto de Pesquisa, Belo Horizonte, MG, 30360-082, Brazil.

*Author for correspondence; e-mail: almeidarocha.jm@gmail.com

Received 29 April 2018; revision accepted 4 January 2019;
Published online 5 March 2019