

# The role of deterministic factors and stochasticity on the trophic interactions between birds and fish in temporary floodplain ponds

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**Abstract** Trophic interaction between contiguous habitats is an understudied area despite their ecological importance. We tested four hypotheses related to the visit of foraging birds in temporary ponds inhabited by fish: the local characteristics of ponds influence the (1) abundance and (2) composition of foraging birds; (3) the number of foraging bird visits increases in higher fish abundance; and (4) pond characteristics lead to a non-random spatial structure of fish assemblage. We studied 18 temporary ponds in the

floodplain area of Pantanal (center-west Brazil), where we measured environmental variables (pond size, depth, macrophyte coverage, and forest canopy coverage), recorded the number of foraging birds and sampled fish. Foraging birds' abundance and composition were mainly influenced by forest canopy coverage and pond size, corroborating our first and second hypotheses. The hypotheses 3 and 4 were rejected. Fish abundance was not correlated with higher number of visits of foraging birds and fish distribution was random. Local environmental variables did not affect significantly fish richness and composition. Our results suggest that the abundance of foraging birds and fish presence are determined by different assembly processes (deterministic vs.

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random), which may limit ponds selection by birds due to the uncertainty in fish distribution and ephemeral nature of temporary ponds.

**Keywords** Intermittent wetland · Predator–prey dynamic · Metacommunity · Pantanal

## Introduction

The community assembly is governed by a complex network of processes that widely vary in the spatiotemporal scale (HilleRisLambers et al., 2012). In general, a given community structure is limited by colonization of species from a regional pool (Ricklefs, 2004). The success to colonize local habitat depends on the availability of resources, abiotic conditions, and the interaction with resident organisms (HilleRisLambers et al., 2012). This deterministic process, generally called environmental filtering, suggests that only species with particular phenotypes that tolerate these local limitations will establish and persist in a given habitat (Kraft et al., 2014). Consequently, environmental gradients influence the distribution of species on space, shaping particular metacommunity structures (e.g., checkerboard, nestedness or evenly spaced; Leibold & Mikkelsen, 2002). On the other hand, stochasticity and randomness may play an important role in short timescales (Tilman, 2004), and, currently, it is a subject of intense debate in ecology (Rosindell et al., 2012). Despite most studies indicating that deterministic factors drive ecological community structures (Gotelli & McCabe, 2002), random species associations are not uncommon (Hubbel & Foster, 1986; Bellay et al., 2011; Ortega et al., 2015).

Trophic interactions are recognized as one of the strongest biotic interactions that may also shape community structure (Stevens, 2010). For instance, studies have suggested that predators may influence the number of rare species (Spiller & Schoener, 1998), patterns of co-occurrence (Maly & Maly, 1997) and behavior of preys (Fortin et al., 2005). However, the intensity and importance of predation depends on the nature where the interaction takes place. Complex and large environments generally could provide more refuges, weakening the antagonistic interaction strength (McCann, 2011; Bellmore et al., 2015) and, consequently, influencing population dynamics

(Cooper & Frederick, 2007) and diversity of preys (Hixon & Menge, 1991). In this sense, landscape heterogeneity generates asynchrony in resource availability, which turns out to affect predator's behavior (Eveleigh et al., 2007). Under these conditions, it is expected that large predators, which have higher mobility and capability to respond to complex environmental information, will move and explore more profitable habitats (Rooney et al., 2008).

Waterbirds are highly mobile organisms with high basal metabolic rates (Bicudo et al., 2010). To sustain the high energy requirements, waterbirds need to consume high levels of food. Searching for food, waterbirds generally explore a large area, connecting different habitats and ecosystems (Kameda et al., 2006; Bauer & Hoyer, 2014; Kloskowski & Trembaczowski, 2015). The decision to where forage may depend on several factors linked with historical past events (Salewski & Bruderer, 2007), the foraging area detectability (Butler & Gillings, 2004), distribution and availability of prey type (Eveleigh et al., 2007; Benoit-Bird et al., 2013; Stanton et al., 2014), and the presence of predators (e.g., wild cats; Yasué et al., 2003). Since most waterbirds rely on visual cues to detect prey, environments with complex structure (e.g., tall grass, forest canopy coverage) may reduce prey accessibility and detectability in potential foraging areas (e.g., Nystrand & Granström, 1997; Frone-man et al., 2001; Butler & Gillings, 2004; Whittingham et al., 2004; Pérez-García et al., 2014). Therefore, attack rate and prey handling could vary widely among environments and may also depend on prey type and thus may have strong effects on waterbirds' foraging behavior and on waterbirds' distribution (Benoit-Bird et al., 2013).

Fish are rich in proteins and lipids (Bone & Moore, 2008), being an important source of food for many waterbird species. However, the contrast between terrestrial and aquatic habitats lead waterbirds to face a complex task to catch fish. For instance, the foraging success of long-legged wading birds (e.g., *Ardeidae* and *Ciconiidae* members) depends on prey availability, which is limited by water depth as they cannot forage at waters deeper than their leg length (Powell, 1987; Bancroft et al., 2002; Gawlik, 2002). Then, shallow lentic water bodies, such as temporary ponds and wetlands, become an attractive foraging habitat for waterbirds due to their high abundance of resources and high capture rates (Weller, 1999).

Shallow lakes, ponds and wetlands are abundant in floodplain areas (Mendonça et al., 2009), which are subjected to seasonal floods responsible for the input of nutrients, sediments, and organic matter that make floodplains one of the most productive habitats in the world (Junk et al., 1989). Numerous species of fish move to floodplains in the high water season seeking for food or suitable areas for spawning or protection (Fernandes, 1997; Winemiller & Jepsen, 1998; Osorio et al., 2011). In the falling water period, most fish return to the river main channel, while some get trapped by the desiccation and stay retained in lateral lakes or temporary ponds (Lowe-McConnell, 1987). The rapid hydrological dynamic made some authors to postulate that the species association and diversity in floodplains are generated randomly (Lowe-McConnell, 1987; Goulding et al., 1988). However, recent studies have recognized the effect of several variables on the structure of fish communities in temporary ponds, such as hydroperiod, depth, aquatic macrophytes occurrence, and connection to perennial water bodies (Baber et al., 2002; Pazin et al., 2006; Fernandes et al., 2010; Tondato et al., 2013).

Here, we verified the deterministic role of local characteristics of temporary ponds (forest canopy coverage, aquatic macrophyte coverage, depth, and size of ponds) on the foraging birds' abundance and on the distribution of fish. Moreover, we examined the correlation between foraging birds and fish abundance. We tested four hypotheses: the local characteristics of ponds determine the abundance (hypothesis 1) and composition (hypothesis 2) of foraging birds; the number of foraging bird visits increases in higher fish abundance (hypothesis 3); and local characteristics of ponds determine a non-random spatial structure of fish assemblage (hypothesis 4). We also quantified the influence of ponds characteristics over fish richness and composition as a way to determine the main modelers of metacommunity structure.

## Methods

### Study area

The study was carried out in temporary ponds of the floodplain area of Miranda River, middle-west of Brazil (Fig. 1). This region is part of the Pantanal, a savanna biome recognized as one of the largest

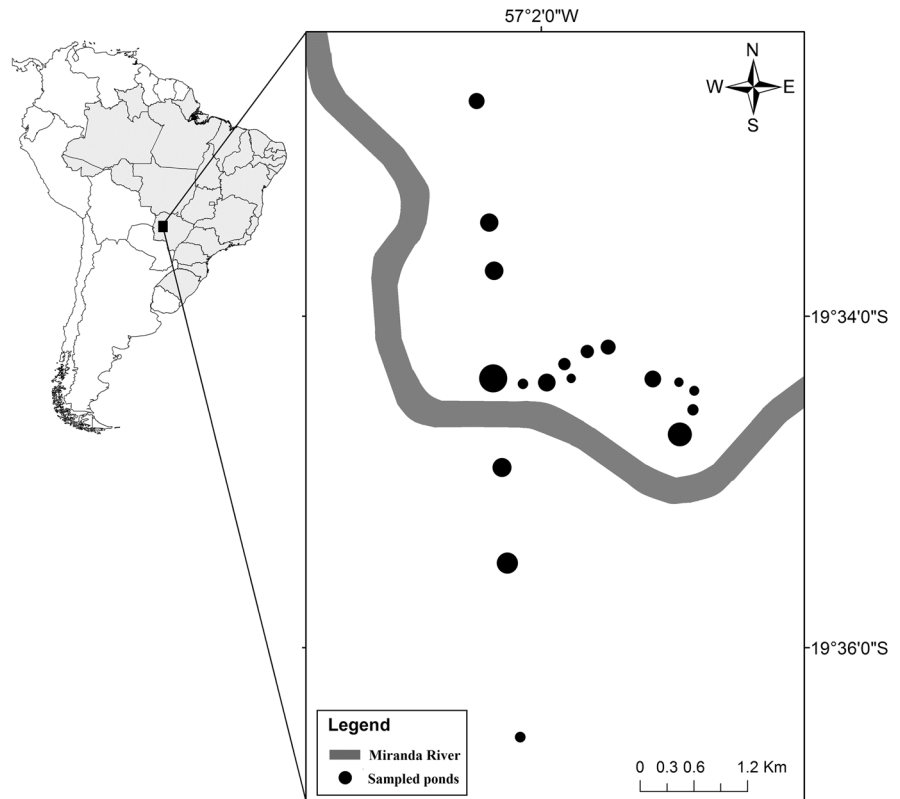
floodplain areas in the world (Junk et al., 2011). The climate in the Miranda sub-basin is classified as tropical humid, with annual temperatures varying between 22.5 and 26.5°C (Gonçalves et al., 2011). The annual rain average varies between 1000 and 1200 mm, with pronounced seasonal variation (abundant rains between November–March). Consequently, the river water level varies significantly throughout the year. The highest water level occurs between December and February (Gonçalves et al., 2011), when the main river channel overflows to the adjacent floodplain. Then, the water level starts to decline up to September–October (Gonçalves et al., 2011), forming several temporary ponds in the floodplain. These temporary ponds are highly heterogeneous, varying in size, forest canopy coverage, depth, and the presence of aquatic macrophytes. In most cases, aquatic macrophytes cover the pond diffusely as a single multi-specific stand, which include floating, emergent, and submerged species. The terrestrial vegetation in the ponds' surroundings varies typically between open grasslands and riparian forest (~4–15 m high). Additionally, this region has one of the highest concentrations of bird individuals in the world (Bini, 2014). Several resident and migratory birds use temporary ponds and other shallow waters to feed on fish, especially in the beginning of the dry season (Donatelli et al., 2014).

### Sampling

We carried out fish samplings and foraging birds records in 18 temporary ponds (Fig. 1) in September of 2013 during four consecutive days. The temporary ponds studied ranged in size from 17.29 to 1500 m<sup>2</sup> and in depth from ~7.4 to ~31 cm (Supplementary Material A). Fish were sampled along each temporary pond using a circular sieve (50 cm of diameter and mesh size of 0.5 cm). The sampling effort for fish assemblages was standardized in 20 min. The sampled fish were anesthetized with a lethal dose of benzocaine and fixed in a 10% formalin solution. Later in the laboratory, fish were identified (Britski et al., 2007), and the standard length (SL; mm) was measured.

Bird record was performed through 3 observations of 10 min in each pond, amounting 54 observations throughout the study. We kept a minimum distance of 20 m between the temporary pond and the observer. Two observations were realized in the morning,

**Fig. 1** Study area location with the 18 temporary ponds' position in the floodplain area of Miranda River, middle-west of Brazil. For better visualization, the ponds were illustrated larger than their natural size. The difference in size between ponds is represented in a square root scale



between 7:00 a.m. and 10:00 a.m., and one in the afternoon, between 4:00 p.m. and 5:30 p.m. Bird observations over time were conducted in different days to ensure sampling independency. Besides, fish sampling and bird observation were conducted in different days to avoid influence over birds' behavior. Birds were identified in the field to species level using specialized literature (Erize et al., 2006; van Perlo, 2009). Later, these recorded birds were classified according to their feeding habit (Sick, 1997; Antas & Palo Jr., 2004), and only those species considered piscivores or potentially piscivores were included in further analyses. We also classified the foraging birds' species according to foraging technique (wading, diving, still hunting and unconventional; Sick, 1997; Antas & Palo Jr., 2004; Bergmann et al., 2013) and main foraging habitat (terrestrial or aquatic; Sick, 1997) for descriptive purposes. It is worth noting that the annual flood pulse creates an open area in the understory close to the soil level, which allows the visual detection of birds independently of the forest structure (authors' personal observation). Moreover,

with the exception of *Jabiru mycteria*, which has a parental care longer than a year, the bird species found in the area were not breeding or feeding their offspring during the studied period.

Depth, macrophyte coverage, and forest canopy coverage were measured once in each temporary pond. Depth was estimated using a graduated stick through 3 measurements (near shores and center), whereas the percentage of macrophyte coverage (0–100) was estimated visually considering the whole pond. Forest canopy coverage was measured through 3 digital pictures mounted vertically (sky direction), located in the center and in the two extreme shores of each temporary pond, standardized in the breast height (1.6 m). These images were then analyzed in the Image J software (Rasband, 2015) to estimate the percentage of canopy coverage over the whole temporary pond. Since the studied ponds resembled to square and rectangular shapes, we opted to estimate the area of each system through the multiplication of its length (m) by its width (m). The width and length of each pond were determined through the mean of 3

measures using measuring tape. Detailed information about each studied ponds is shown in supplementary material A.

### Data analysis

The frequency of occurrence (FO) and relative abundance (RA) of foraging birds and fish were calculated according to Amundsen et al. (1996). The FO and RA was used to construct diagrams according to Costello (1990); modified by Amundsen et al. (1996), which allows distinguishing the dominance of different bird and fish species in the studied temporary ponds. In this graphic analysis, the x axis represents the frequency of occurrence (FO), and the y axis represents the relative abundance (RA) of a specific taxa.

We estimated the importance of each predictor (forest canopy coverage, aquatic macrophyte coverage, pond depth, and size) to the foraging birds' abundance (hypothesis 1) and to the fish richness by the model averaging approach (Burnham & Anderson, 2002). We also included fish and foraging birds' abundances as predictors of each other in the analysis to test our hypothesis 3. Before conducting the model averaging analysis, we verified multicollinearity in our data using the variance inflation factor (VIF; Nakazawa, 2014), which indicated low correlation among the predictors (VIF < 2.2). The model averaging method consisted of a sequence of steps. First, we carried out all possible models with a maximum of two predictors per model to each dataset (foraging birds and fish). For the foraging birds' dataset, we conducted generalized linear mixed models (GLMM) using the ponds ID as a random variable, while for the fish dataset, we used generalized linear models (GLM). Considering that fish richness and foraging birds' abundance are discrete variables and do not have normal error distribution, we used Poisson distribution using the inverse link function. Second, the second-order Akaike information criterion with a correction for small sample sizes (AICc) was used to measure the plausibility of each candidate model. Third, the Akaike weight was calculated for each model (WI) normalized across the set of candidate models to sum to one. Fourth, the Akaike weights were used to obtain averaged estimates for each parameter. Finally, the relative importance of each predictor variable was calculated by summing all Akaike weights over all models that included each predictor. The relative importance (I) ranges from 0 to 1, and the larger

the value of the relative importance of a predictor, the more important it is compared to the others (Burnham & Anderson, 2002). We also evaluated the significance of models with delta AICc lower than 2 using a parametric bootstrap comparison (1000 randomizations; Halekoh & Højsgaard, 2014) with a null model (without variables). For the forest canopy coverage, we also carried out quantile regressions (5th, 25th, 50th, 75th, and 95th) to deal with an over-dispersion of foraging birds in areas with open canopy. In this last case, we think that the forest canopy coverage may set the upper limit for the foraging bird abundance rather than affecting the mean values, driving a relationship with triangular shape in graphs (Gotelli & Ellison, 2004). The model averaging procedure was conducted in the R package "MuMIn" (Barton, 2015), the parametric bootstrap comparison was conducted in the R package "pbkrtest" (Halekoh & Højsgaard, 2014), and the quantile regression was conducted in the R package "quantreg" (Koenker, 2015).

The non-metric multidimensional scaling (NMDS) ordination based on Bray–Curtis dissimilarity was used to assess the variation in fish and birds structure among the 18 sampled ponds. Both the fish and bird data (abundance values) were square root transformed before NMDS ordinations. The pond characteristics (depth, size, forest canopy coverage, fish abundance, foraging birds' abundance and aquatic macrophyte coverage) were correlated onto the birds (hypothesis 2) and fish (hypothesis 4) NMDS ordination using the vector fitting procedure (Oksanen, 2009; Keppel et al., 2010). The significance (*P* value) between the ordination and the predictors was assessed after 1000 permutations. Only the forest canopy coverage was correlated with bird species composition (see Results), and in this case, we also carried out a similarity percentage breakdown procedure (SIMPER) to investigate the difference in foraging birds' composition along the forest canopy coverage gradient. Given that SIMPER analysis discriminates species among levels of categorical variables, we separated the temporary ponds in two main groups: ponds with low forest canopy coverage (<20% of forest canopy coverage) and ponds with high forest canopy coverage (>30% of forest canopy coverage). The probability of getting equal or larger average contribution for each species (*P* value) was accessed through 1000 random permutations of the group factor. To guarantee that both groups (ponds with low forest canopy coverage and

high forest canopy coverage) are legitimate, we also tested their difference regarding foraging birds composition through a Permutational Multivariate Analysis of Variance (PERMANOVA; 1000 randomizations). The vector fitting, NMDS, SIMPER, and PERMANOVA were carried out in the R package “vegan” (Oksanen et al., 2009).

We tested the non-random distribution of fish in temporary ponds (hypothesis 4) using the elements of metacommunity structure (EMS). This method allows classifying metacommunities through a unique framework according to multiple patterns of species distribution, such as checkerboards, nestedness, and Gleasonian distribution (Henriques-Silva et al., 2013). The EMS is based on the evaluation of three main metrics: coherence (level in which species respond to the same environmental gradient), turnover (number of times a species replaced another between sites), and boundary clumping (dispersion of species occurrences among sites) (Leibold & Mikkelsen, 2002). The coherence, turnover, and boundary clumping were calculated from a presence/absence matrix (sites as rows and species occurrences as columns) ordinate via reciprocal averaging (Gauch, 1982). More specifically, the coherence was calculated by counting the number of embedded absences, while the turnover was measured by calculating the number of times one species replaced another between sites (Dallas, 2014). The coherence and turnover were then compared with an appropriated null model to identify non-random patterns (positive or negative). In this study, we used the algorithm “swap” (sum of rows and columns is maintained) to create the null models due to its best performance on the reduction of type I and II errors (Gotelli & Entsminger, 2001; Ulrich & Gotelli, 2007). On the other hand, we calculated the boundary clumping using the Morisita’s index, which is a measure of the dispersion of species occurrences among sites (Morisita, 1971). Boundaries are considered not clumped if the Morisita’s index is one, clumped when values are greater than one and hyperdispersed boundaries when the values are less than one. We expected that coherence should be different from the null model (1000 simulations,  $P < 0.05$ ), which indicates a non-random distribution of species independently from turnover and boundary clumping results (Leibold & Mikkelsen, 2002; Presley et al., 2010). The EMS procedure was carried out in the R package “Metacom” (Dallas, 2015).

## Results

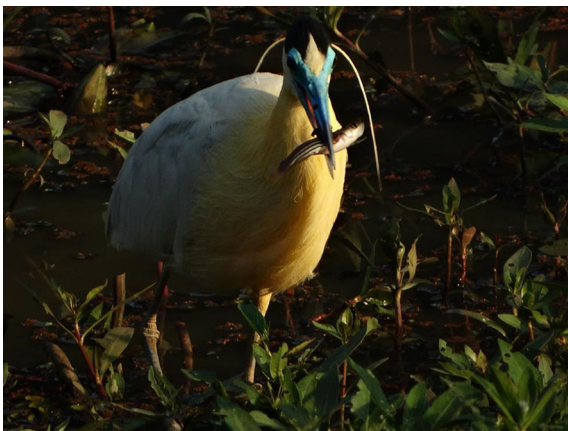
Overall, foraging birds were abundant in the study area, being absent in just three ponds. A total of 230 birds distributed among 14 families and 37 species were observed foraging in the temporary ponds. From these birds species, 13 (147 individuals) were known as piscivores (25.1%) or facultative piscivores (74.9%) and were used in data analysis (Table 1). The majority (79.6%) of the foraging birds recorded were wading foragers (Fig. 2), while 17% were opportunistic terrestrial birds with unconventional foraging methods, 2% were diving birds, and 1.4% were still hunting birds (Table 1). *Phimosus infuscatus* (Lichtenstein, 1823) (FO = 50, RA = 21.7; Fig. 3a), *Jacana jacana* (Linnaeus, 1766) (FO = 27.8, RA = 14.9; Fig. 3a), and *Pitangus sulphuratus* (Linnaeus, 1766) (FO = 61.1, RA = 12.9; Fig. 3a) were the most frequent bird species in the studied temporary ponds.

Local characteristics influenced the numerical abundance and composition of foraging birds, suggesting a deterministic effect of local characteristics on the visit of birds in temporary ponds. The numerical abundance of foraging birds was mainly influenced by pond size ( $I = 0.72$ ; Tables 2, 3; Fig. 4a) and forest canopy coverage ( $I = 0.62$ ; Table 2, 3; Fig. 4b). Besides, the models with pond size and forest canopy coverage were significant ( $P < 0.05$ ; Table 3) in the best models selected by AICc. On the other hand, fish abundance (hypothesis 3), pond depth, and macrophyte coverage had low values of relative importance (Table 2) and were related to models with high delta AICc ( $>2$ ; Table 3), suggesting that these variables are weak predictors. We also found significant negative relationship between the numerical abundance of foraging birds and forest canopy coverage for the 50° ( $P = 0.04$ ), 75° ( $P = 0.009$ ), and 95° ( $P < 0.001$ ) quantiles, while the 5° and 25° were not significant ( $P > 0.05$ ; Fig. 4b). These results from the quantile regressions indicated that forest canopy cover caused an upper limit for the abundance of foraging birds rather than affecting the mean values. In relation to foraging birds composition, forest canopy coverage was the only variable that was significant ( $R^2 = 0.40$   $P = 0.04$ ; Fig. 5a). Large birds, such as members of *Ardeidae* and *Ciconiidae* family, were more related to open areas (Fig. 5a). Similarly, the PERMANOVA test also showed that ponds with high forest canopy

**Table 1** Foraging bird species recorded in 18 temporary ponds located in Pantanal, middle-west of Brazil

Family	Species	Total abundance (min–max)	Foraging technique	Feeding habit	Main foraging habitat
Ciconiidae	<i>Jabiru mycteria</i>	2 (0–2)	Wading	Piscivorous	Aquatic
Ardeidae	<i>Egretta thula</i>	15 (0–10)	Wading	Piscivorous	Aquatic
	<i>Ardea alba</i>	14 (0–11)	Wading	Piscivorous	Aquatic
	<i>Tigrisoma lineatum</i>	2 (0–2)	Wading	Piscivorous	Aquatic
	<i>Nycticorax nycticorax</i>	1 (0–1)	Wading	Piscivorous	Aquatic
	<i>Phimosus infuscatus</i>	32 (0–7)	Wading	Facul. piscivorous	Aquatic
Threskiornithidae	<i>Theristicus caerulescens</i>	7 (0–2)	Wading	Facul. piscivorous	Aquatic
	<i>Mesembrinibis cayennensis</i>	3 (0–3)	Wading	Facul. piscivorous	Aquatic
	<i>Phalacrocorax brasilianus</i>	1 (0–1)	Diving	Piscivorous	Aquatic
Phalacrocoracidae	<i>Phalacrocorax brasilianus</i>	1 (0–1)	Diving	Piscivorous	Aquatic
Rallidae	<i>Aramides cajanea</i>	14 (0–4)	Wading	Facul. piscivorous	Aquatic
Aramidae	<i>Aramus guarana</i>	4 (0–2)	Wading	Facul. piscivorous	Aquatic
Jacaniidae	<i>Jacana jacana</i>	22 (0–10)	Wading	Facul. piscivorous	Aquatic
Recurvirostridae	<i>Himantopus melanurus</i>	1 (0–1)	Wading	Facul. piscivorous	Aquatic
Sternidae	<i>Phaetusa simplex</i>	1 (0–1)	Diving	Piscivorous	Aquatic
Cuculidae	<i>Guira guira</i>	4 (0–4)	Unconventional	Facul. piscivorous	Terrestrial
Alcedinidae	<i>Chloroceryle</i> sp.	1 (0–1)	Diving	Piscivorous	Aquatic
Falconidae	<i>Caracara plancus</i>	2 (0–1)	Unconventional	Facul. piscivorous	Terrestrial
Accipitridae	<i>Rostrhamus sociabilis</i>	2 (0–1)	Still hunting	Facul. piscivorous	Aquatic
Tyrannidae	<i>Pitangus sulphuratus</i>	19 (0–5)	Unconventional	Facul. piscivorous	Terrestrial

*Facul. piscivorous* Facultative piscivorous. Minimum (Min) and maximum (Max) abundance values with regard to single observations

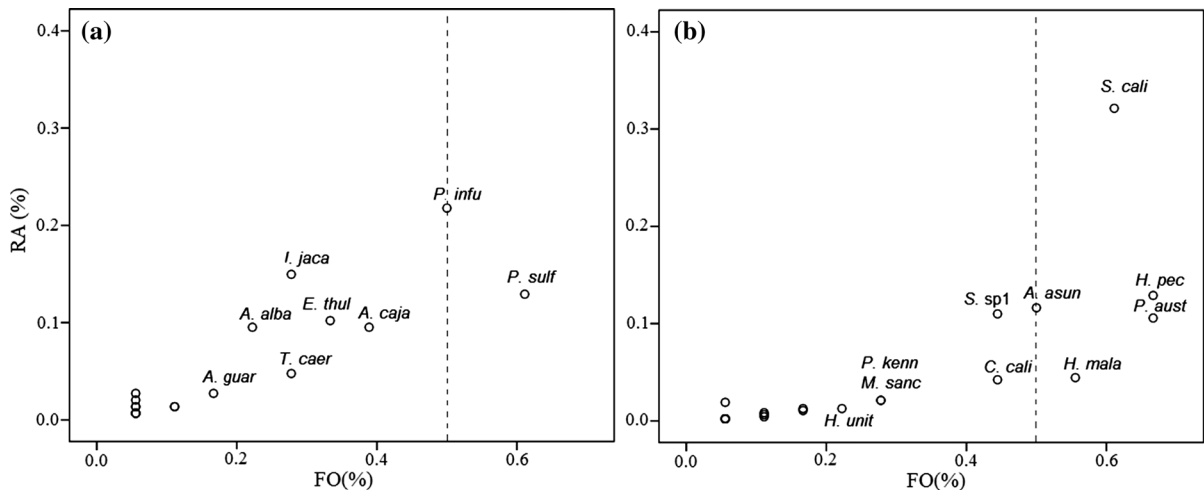


**Fig. 2** Heron (*Pelecanus erythrorhynchos*), a wading forager, preying the annual fish *Trigonectes balzanii* in one of the temporary ponds studied in the Pantanal, middle-west of Brazil. To our knowledge, this is the first picture evidencing the role of birds as a predator of annual fish, fact attested indirectly by several authors (Costa, 1998; Lanés et al., 2014)

coverage differed in terms of foraging bird composition from ponds with low forest canopy coverage ( $F_{1,12} = 3.43$ ,  $P = 0.007$ ,  $R^2 = 0.22$ ). Seven

foraging birds species accounted for a little more than 70% of the dissimilarities among ponds with high forest canopy coverage and ponds with low forest canopy coverage (Table 4), but *Aramides cajanea* ( $P = 0.04$ ), *Theristicus caerulescens* ( $P = 0.09$ ), and *Egretta thula* ( $P = 0.08$ ) were the only species that contributed significantly or marginally significantly to distinguish between both pond groups. *Aramides cajanea* was more related with ponds with high forest canopy coverage, while *T. caerulescens* and *E. thula* were more related with ponds with low forest canopy coverage (Table 4; Fig. 5a).

A total of 473 fish, divided into 8 families and 23 species, were found in the studied temporary ponds (Table 5). Fish abundance varied from 5 to 72 among the temporary ponds, and *Characidae* (59%), *Callichthyidae* (18%), and *Lebiasinidae* (10%) were the most abundant families. According to the EMS, the coherence (embedded absences observed = 106, embedded absences expected =  $129.91 \pm SE 17.53$ ,  $z = 1.36$ ,  $P = 0.17$ ) and turnover (replacements observed = 2139, replacements expected =  $1816.9 \pm SE 556.58$ ,  $z = -0.57$ ,  $P = 0.56$ ) were not affected



**Fig. 3** Modified Costello Diagram showing the frequency of occurrence (FO) and relative abundance (RA) of foraging birds (a) and fish (b) found in the temporary ponds of Pantanal,

middle-west of Brazil. Dashed lines indicate where frequency of occurrence is equal to 0.5 (50%)

**Table 2** Model-averaged importance of predictors (I) for foraging bird abundance and fish richness in temporary ponds of Pantanal

Parameter	Foraging birds' abundance	
	Relative importance	Parameter estimation (95% CI)
Pond size	0.72	0.0014 (0.0003 to 0.0025)
Forest canopy coverage	0.62	−0.0209 (−0.0404 to −0.0014)
Pond depth	0.16	0.0308 (−0.0545 to 0.1162)
Fish abundance	0.1	0.0064 (−0.0140 to 0.0269)
Aquatic macrophyte coverage	0.1	0.0016 (−0.0125 to 0.0159)
Parameter	Fish richness	
	Relative importance	Parameter estimation (95% CI)
Aquatic macrophyte coverage	0.44	−0.0061 (−0.0139 to 0.0016)
Pond depth	0.23	−0.0179 (−0.0571 to 0.0212)
Pond size	0.18	−0.0001 (−0.0008 to 0.0004)
Bird abundance	0.17	−0.0062 (−0.0350 to 0.0224)
Forest canopy coverage	0.15	0.0007 (−0.0071 to 0.0085)

CI confidence interval

negatively from what was expected by the null models, while the boundary clumping was significantly higher than expected (Morisita's index = 1.90,  $DF = 20$ ,  $P = 0.0002$ ). These results suggest a random distribution of fish species in the studied temporary ponds. Similarly, we did not find any interpretable pattern in fish ordination (Fig. 5b) nor significant influence of pond characteristics over fish composition ( $P > 0.05$ ; Fig. 5b). In relation to fish richness, aquatic macrophyte coverage

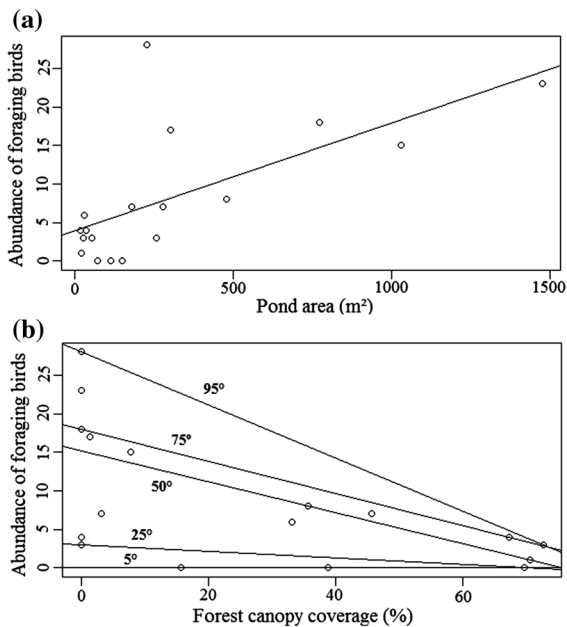
was the most important predictor ( $I = 0.44$ ; Table 2) and was present in the best model according to AICc criterion (Table 3). However, all models, including the one with aquatic macrophyte coverage parameter, were not significant ( $P > 0.05$ , Table 3). Besides, the second best model, which had a delta AICc of 0.41, was the one without any predictor, indicating that in general the pond variables measured were weak predictors for fish richness.



**Table 3** Best models for foraging bird abundance and fish richness according to AICc value

Model (foraging birds' abundance)	AICc	Delta AICc	AICc Weight	Sig. ( <i>P</i> value)
1 + Forest canopy coverage + pond size	222.04	0.00	0.36	0.002
1 + Pond size	223.40	1.36	0.18	0.003
1 + Forest canopy coverage	224.66	2.62	0.10	–
1 + Forest canopy coverage + pond depth	224.79	2.75	0.09	–
1 + Pond size + fish abundance	225.58	3.54	0.06	–
Model (fish richness)	AICc	Delta AICc	AICc Weight	Sig. ( <i>P</i> value)
1 + Aquatic macrophyte coverage	87.09	0.00	0.21	0.830
1	87.51	0.41	0.17	1.000
1 + Pond depth	88.88	1.78	0.09	0.480
1 + Aquatic macrophyte coverage + pond depth	89.28	2.18	0.07	–
1 + Pond size	89.29	2.20	0.07	–

Generalized linear mixed models (GLMM) were used to fit the foraging bird abundance, while Generalized Linear Models (GLM) were used to fit fish richness (more details in Data Analysis section). All possible models with a maximum of 2 predictors were considered in these comparisons. The constant 1 included in all models is the intercept. The significance was calculated using a parametric bootstrap comparison for models with Delta AICc lower than 2

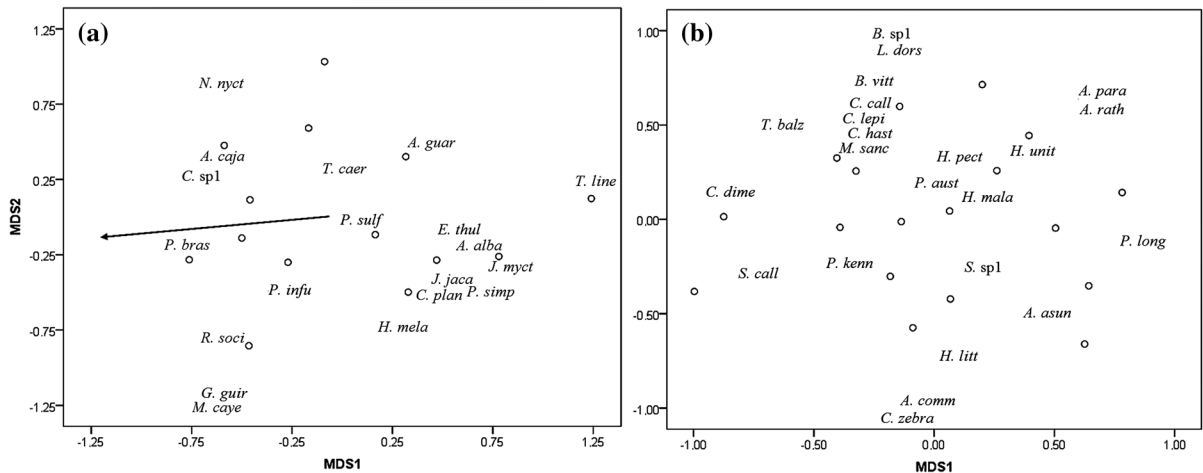


**Fig. 4** Relationship between pond size (a) and forest canopy coverage (b) with foraging bird abundance in temporary ponds of Pantanal, center-west of Brazil. A linear regression was performed for pond size (a), and quantile regressions (95°, 75°, 50°, 25°, 5°) were fitted for forest canopy coverage (b)

## Discussion

Deterministic processes drive bird abundance and composition in temporary ponds as expected in our

first and second hypotheses, respectively. Forest canopy coverage and pond size affected numerical abundance of foraging birds. Open large ponds have higher foraging bird visitation than small ponds with high forest canopy coverage. These results reflect the detectability and accessibility of the ponds by birds. Prey detectability and accessibility decrease with the increasing habitat structure for birds that rely on visual cues to detect their prey (Butler & Gillings, 2004). All the birds found foraging in the studied temporary ponds are first oriented by their visual system (Sick, 1997). Therefore, small ponds in forest areas with large canopy cover should affect negatively pond detectability and also the mobility of birds, decreasing the number of visits and the attack rate over aquatic organisms. It is noteworthy that the forest canopy coverage created a limiting response in bird abundance instead of a classic linear relationship. The over-dispersion found in ponds with low forest canopy coverage, which created a triangular shape in the relation between forest canopy coverage and foraging bird abundance, was probably a consequence of the heterogeneity in pond sizes. Even in open areas, the chance of find birds in small ponds is much lower than in the large ones. Indeed, we observed foraging birds in all ponds that were located in open areas with no forest canopy coverage (foraging birds were absent in three ponds under high forest canopy coverage).



**Fig. 5** Ordinations generated through a NMDS analysis of species composition of **a** foraging bird and **b** fish found in temporary ponds of the floodplain of Miranda River. Only forest

canopy coverage (*black arrow*) significantly affected the composition of foraging birds' species

**Table 4** Average contribution of foraging birds species to overall dissimilarity (Bray–Curtis) between ponds with high forest canopy coverage and ponds with low forest canopy coverage in the floodplain of Miranda River, middle-west of Brazil

Foraging bird species	Average contribution (SD)	Average abundance in ponds with high forest canopy coverage	Average abundance in ponds with low forest canopy coverage
<i>Phimosus infuscatus</i>	0.11 (0.10)	1.28	1.09
<i>Aramides cajanea</i> *	0.09 (0.06)	1.13	0.44
<i>Jacana jacana</i>	0.07 (0.08)	0.00	1.04
<i>Egretta thula</i> <sup>o</sup>	0.07 (0.07)	0.00	0.90
<i>Pitangus sulphuratus</i>	0.06 (0.06)	0.88	1.05
<i>Theristicus caerulescens</i> <sup>o</sup>	0.06 (0.06)	0.00	0.64
<i>Ardea alba</i>	0.05 (0.08)	0.00	0.70

Only foraging birds that accounted for at least 0.5 of the average contribution were shown. The abundance of foraging birds was square root transformed prior to the analysis. <sup>o</sup>  $P < 0.1$  (marginally significant), \*  $P < 0.05$  (significant)

Foraging decisions, such as the choice of where to feed, are usually portraiture of the balance between energetic costs and gains (Pyke, 1984). Following this idea, several studies have suggested that prey availability is the main constraint of the numerical abundance of wading birds in wetlands (Kushlan, 1976; Butler, 1994; Hafner, 1997; Gawlik, 2002; Master et al., 2005). Kushlan (1976) observed that wading birds forage more intensively when the wetland is shallow and with high density of fish, increasing their success of capture. Our results suggest that pond depth and fish abundance were weak predictors of foraging

birds' abundance, which led us to reject our third hypothesis. The shorter period of water permanence in temporary ponds compared to wetlands may have affected this lack of relation between fish concentration and bird abundance. During the dry season, the high solar intensity in the region increase water temperature causing fast evaporation (Junk et al., 2011). Two days after the end of the study, 3 ponds had already dried completely. Therefore, we suggest that the decreasing water level was fast enough to create a short window of time, which could prevent bird's learning and foraging adaptation. On the other hand, the

**Table 5** Length and abundance of fish found in temporary ponds located in floodplain of Miranda River, middle-west of Brazil

Family	Species	Total abundance (Min–Max)	Mean length (SD)
Lebiasinidae	<i>Pyrrhulina australis</i>	50 (0–20)	2.53 (0.50)
Characidae	<i>Moenkhausia sanctaefilomenae</i>	10 (0–3)	2.57 (0.38)
	<i>Serrapinnus</i> sp.1	52 (0–22)	1.71 (0.27)
	<i>Serrapinnus calliurus</i>	152 (0–39)	1.84 (0.27)
	<i>Aphyocharax rathbuni</i>	1 (0–1)	3.20
	<i>Aphyocharax paraguayensis</i>	2 (0–1)	2.18 (0.12)
	<i>Psellogrammus kennedy</i>	10 (0–5)	3.15 (0.13)
	<i>Astyanax asuncionensis</i>	55 (0–12)	3.90 (0.98)
	Erythrinidae	<i>Hoplias malabaricus</i>	21 (0–7)
<i>Hoplerythrinus unitaeniatus</i>		6 (0–2)	4.33 (1.87)
Crenuchidae	<i>Characidium zebra</i>	1 (0–1)	2.89
Hypopomidae	<i>Brachyhypopomus</i> sp. 1	1 (0–1)	11.00
Rivulidae	<i>Trigonectes balzanii</i>	1 (0–1)	3.95
	<i>Pterolebias longipinnis</i>	1 (0–1)	4.00
Callichthyidae	<i>Corydoras hastatus</i>	4 (0–2)	1.45 (0.06)
	<i>Hoplosternum pectorale</i>	61 (0–24)	2.99 (0.64)
	<i>Hoplosternum littorale</i>	1 (0–1)	6.50
	<i>Callichthys callichthys</i>	20 (0–10)	2.90 (1.03)
Cichlidae	<i>Bujurquina vittata</i>	6 (0–4)	2.21 (0.18)
	<i>Laetacara dorsigera</i>	9 (0–9)	2.76 (0.20)
	<i>Crenicichla lepidota</i>	3 (0–2)	4.42 (0.89)
	<i>Cichlasoma dimerus</i>	5 (0–2)	4.14 (1.18)
	<i>Apistogramma commbrae</i>	2 (0–1)	2.44

Minimum (Min) and maximum (max) abundance values with regard to single pond samplings

similarity in depth and fish abundance among the ponds may have caused the absence of the expected variation in foraging birds' abundance. The maximum difference found in terms of depth was ~23.6 cm, while for fish abundance was 67 individuals. Although these differences are significant for small temporary ponds, the limitations of depth and fish abundance over the foraging of birds may not apply, leading to the absence of pond's selection by birds. Either way, studies in large spatial areas, comparing ponds with different hydroperiods, depths, and fish abundances are needed to clarify these hypotheses.

Interestingly, Haas et al. (2007) observed that the presence of fish in ponds influence negatively the occurrence of waterbirds. More specifically, fish compete with birds for invertebrate preys, and thus, birds select ponds without fish (Haas et al., 2007). It seems unlikely that this process occurred in our studied sites, since fish species inhabiting the

temporary ponds were small sized (see Table 3) and are also preys of birds, as attested by several attacks witnessed by us. Indeed, in our study, we captured, to our knowledge, the first picture evidencing the role of birds as a predator of annual fish (Fig. 2). However, many bird species found are not strictly piscivorous and their diet also consists of macroinvertebrates (see Table 1). For instance, *P. infuscatu*s, *J. jacana*, and *P. sulphuratus* represent almost half of the foraging birds recorded and all of them are facultative piscivorous. Therefore, it is possible that macroinvertebrates availability also influenced the numerical abundance of bird's foraging, which was not measured in our study.

Foraging birds' composition varied among the temporary ponds as expected in our second hypothesis. Large birds, such as herons and storks, were not found in pond covered by forest vegetation. On the other hand, ponds in open areas included a wide range of species, such as the small *Jacana jacana*

(~23 cm) and the large *J. mycteria* (Linchtenstein, 1819) (~1.5 m). This result supports the modeler role of habitat structural complexity on the composition and morphology of biological assemblages through filtering effect (Gibb & Parr, 2013). The structure and spatial grain of vegetation cover may determine the presence of species with certain body sizes over others (Gunnarsson, 1992; Telleria & Carrascal, 1994; McAbendroth et al., 2005). A structural complex habitat may affect large body birds, reducing their mobility and, consequently, increasing their energetic demands (Brodmann et al., 1997). Alternatively, large slow birds may avoid forested areas due to a higher risk of predation (Whittingham & Evans, 2004). Either way, in a savanna biome, such as the Pantanal, where the landscape is flat and composed by a mosaic of forest and grassland, the vegetation cover play an important role on bird predation dynamics in ponds.

Unlike the deterministic structure of foraging birds' assemblage, fish varied between temporary ponds randomly. This result rejects our fourth hypothesis and supports the importance of flood pulse to the random colonization and distribution of fish in temporary ponds. In general, harsh abiotic conditions strongly limit the number of species that can inhabit temporary ponds (Jocque et al., 2010). With the exception of those fish that are adapted to live in seasonal habitats, such as annual fish (Costa, 1998), the recruitment of species in temporary ponds occur by those species that are incapable to return to perennial habitats and end up retained in these temporary habitats (Pazin et al., 2006). In the 23 fish species sampled, just 2 are strictly adapted to survive the dry period via drought-tolerant embryos (annual fish from *Rivulidae* family; Wourms, 1972). Some species, such as members from the *Callichthyidae* family, are capable to bury into the mud or move small distance creeping or leaping to others water bodies (Magoulick & Kobza, 2003). However, most species occurring in these temporary habitats are fated to desiccation or aerial predation without produce offspring for the next generation.

Although the harsh conditions and low life expectancy for fish in temporary ponds, recent studies suggest that fish assemblages may be deterministic and highly predictable in these habitats (Baber et al., 2002; Pazin et al., 2006; Fernandes et al., 2010; Tondato et al., 2013). Pazin et al. (2006) observed a

nested subset structure and the influence of local factors (e.g., hydroperiod, canopy cover, depth) on fish richness and composition in ponds from Amazon. However, our results indicate that fish species do not exhibit a coherent range along the sampled environmental gradient, which breach an important assumption of all non-random metacommunities distributions (nestedness, checkboard, evenly spaced, Gleasonian, Clementsian, and other quasi structures; Presley et al., 2010). Furthermore, none local variable measured was correlated to fish richness and composition, suggesting a random species assembly in temporary ponds from Pantanal. Despite these strong evidences, we should be cautious and do not assume that fish assemblage structure in temporary ponds is purely random. Community structures that are identified as random may be caused by the interaction among deterministic processes (Boschilia et al., 2008; Caruso et al., 2011). Our data are snapshots of fish community composed by small sized individuals (less than 7 cm), and therefore, deterministic process of long-time scale may have been not identified. Long-time deterministic process could be more prone to be observed during the falling water period, when large body fish may be less likely to be trapped into temporary ponds due to their high mobility and higher capacity to respond to complex environmental information (Rooney et al., 2008).

Predator–prey interactions are strongly affected by how individuals distribute themselves into space (Hammond et al., 2007). Our study suggests that the uncertainty in fish distribution may have limited pond selection by foraging birds. More specifically, the random organization of fish in temporary ponds and the absence of pond characteristics that serve as surrogate for resources availability may have hampered bird recognition and selection of profitable sites. In this scenario, physical characteristics of ponds, such as accessibility and detection, may become the strongest factors influencing the intensity of visits of foraging birds. Additionally, the cost of diet specialization in animals that forage in temporary ponds may be high and disadvantageous due to the wide inter-annual and spatial variation in prey availability (Polacik & Reichard, 2010). Indeed, most wetland birds are generalist predators (Sick, 1997; Eastman, 1999), changing diet according to season and availability of preys (Weller, 1999; Jakubas & Maniowska, 2011). Thus, the dynamic of bird and fish

assemblage in temporary ponds is another good example that food webs are dynamical systems subjected to predators' behavior and preys' fluctuation, instead of just static entities (Polis & Winemiller, 1996; McCann, 2011).

In summary, trophic interaction between contiguous habitats is still an understudied area despite their importance for the maintenance of communities in ecological landscapes (Nakano & Murakami, 2001). In this study, we provide information about the predator–prey relationship between foraging birds and fish in temporary ponds from Pantanal, highlighting the roles of deterministic factors and stochasticity in the structure of both assemblages. These results could help elucidate the energy pathways between aquatic and terrestrial environments in savanna like biomes subjected to seasonal floods, providing basic information for ecological modelling and conservation management.

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