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1. Introduction

Tropical forests fragmentation often reduces the tree regeneration process as a result of ruptures in biological interactions such as seed dispersal, with subsequent effects on seedling recruitment (Benítez-Malvido, 1998; Cordeiro and Howe, 2003; Tabarelli et al., 2008; Benítez-Malvido et al., 2018). This impact is particularly severe in forest fragments lacking large-bodied seed vectors due to the decline in dispersal and recruitment of large-seeded trees species, mainly at the edges (Benítez-Malvido et al., 2018). These changes are, at least, partly driven by edge effects, the diverse physical and biotic changes associated with the abrupt artificial boundaries of habitat fragments (Laurance et al., 2006; Melo et al., 2007). Changing plant-frugivore interactions will probably change forest community characteristics, which may exacerbate the long-term species diversity loss in fragmented and disturbed tropical forests (Wright and Duber, 2001; Benítez-Malvido et al., 2018).
Palm recruits are one of the most abundant and diverse groups of animal-dispersed plants in the Americas (Henderson, 2002). Indeed, most palm species produce large fruits, widely consumed by large frugivores (Zona and Henderson, 1989; Muñoz et al., 2019). In this respect, studies have shown mammals as important seed vectors of Neotropical palms (Muñoz et al., 2019; Dracxler and Kissling, 2021). Although some mammals may function as both seed predators and dispersers, associations with mammalian frugivores benefit palms in terms of invertebrate seed predation avoidance, seed germination, and seedling survival by seed dispersal by these vectors (Zona and Henderson, 1989; Dracxler and Kissling, 2021). Medium and large mammals are usually scarce in the forest edges, implying in reduced seed dispersal at this modified habitat (Asquith and Mejia-Chang, 2005; Wright et al., 2007). Consequently, edges of forest fragments typically experience significant changes in their palm community composition (Baez and Balslev, 2007; Wright et al., 2007). However, few studies focused on the responses of palm species to habitat loss and fragmentation at the population level (Souza and Martins, 2003; Mariano and Christianini, 2016). Worth note, Galetti et al. (2006) pointed out the deleterious consequences of the loss of seed-dispersing agents for the long-term sustainability of palm populations in forest remnants.

Studies in fragmented landscapes suggest that the edge effect affects palms differently (Scariot, 1999; Baez and Balslev, 2007; Browne and Karubian, 2016). Some palm species may experience an increase in their abundance in modified habitats, as forest edges (Souza and Martins, 2003; Pimentel and Tabarelli, 2004), while others decline (Arroyo-Rodriguez et al. 2007; Baez and Balslev, 2007). Basically, some palm species benefit from increased light availability (Souza and Martins, 2003) and changes in dispersal-predation dynamics, which favor palm species in terms of recruitment rates, becoming dominant after forest fragmentation (Wright et al., 2000; Pimentel and Tabarelli, 2004). On the other hand, in Amazonia, for example, the density of two species of palms and the overall species richness of the palm community tended to decrease toward the edges within forest fragments (Baez and Balslev, 2007). In view of those different patterns, desert attention the importance of understanding palm recruitment in fragmented vegetation in terms of population dynamic changes. Importantly, the Brazilian Cerrado has been severely fragmented (Carvalho et al., 2009; Beuchle et al. 2015), but no study compared palm recruitment simultaneously in the edge and the interior of remnants. In this respect, I predict the distance from the edge affects positively both seedling abundance and far distribution from parent palms due to their dependence on medium-large animal seed-vectors, which are scarce or absent along forest edges (Wright et al., 2007). Thus, in this study, I compared both Syagrus flexuosa (Mart.) Becc. seedling abundance and distribution (according to the distance from adult palms) at the edges with the interiors of Cerrado remnants.

2. Materials and Methods

2.1. Study sites

I developed this study in three large remnants of dense Cerrado, which, as a habitat type, resembles the dry forests due to a low and almost continuous canopy, besides the deciduous pattern during the dry season (Ribeiro and Walter, 1998). The remnants were: Matão 20°25′16″S, 51°51′32″W, 1200 ha in area, São Matheus 20°27′33″S, 51°38′36″O, 753 ha; Bom Jardim 20°30′40.8″S, 52°31′49″O, 667 ha, all of which are private reserves located in the Municipality of Três Lagoas, State of Mato Grosso do Sul, Brazil; 51°48′W, 20°46′S, elevation 340 m. The clearing of Cerrado in this region has produced landscape that are mosaics of pastures with isolated trees, riparian forest remnants, palm (Mauritia L.f.) stands remnants, and Cerrado fragments. Cerrado remnants mentioned above were spaced by at least 23 km. Some common trees present in the remnants are Qualea grandiflora Mart., Q. parviflora Mart., Copaifera langsdorfii Desf., Dipteryx alata Vog., Pterodon emarginatus Vogel, Annona crassiloba Mart., Anadenanthera falcata (Benth.) Speg. and Vatairea macrocarpa (Benth.) Deke. The canopy in these remnants was 6–10 m tall, while emergent trees may reach 12–14 m. From June to September, most tree species shed their leaves, contributing to a deciduous pattern. Annual rainfall is around 1400 mm, most of which (70–80%) occurs from October to March (wet season). During the wet season average temperature is 24 °C, whereas, during the dry season (April to September) an average of 17 °C is recorded. Frosts may occur in the coldest months.

2.2. Study species

Syagrus flexuosa (Mart.) Becc. is present in the Cerrado of São Paulo, Minas Gerais, Mato Grosso do Sul, Mato Grosso, Goiás, Tocantins, Maranhão, and Bahia states (Lorenzi et al., 2010). This palm reaches 1–5 m high, and the crown has 7–15 leaves 0.9–1.1 m long, in which leaflets (40–80 at each side) are regularly arranged in different planes (Lorenzi et al., 2010). It occurs both in preserved and disturbed Cerrado, as well as in anthropized areas as pastures. It produces one to four infructescences per stem, each with 1 to 60 ovoid fruits (3.0–6.0 cm in length; 2.0–3.0 cm in diameter) with a fleshy edible pulp, and a highly-nutritious, coconut-like endosperm (Lorenzi et al., 2010). Syagrus flexuosa fruiting occurs all year round, and seeds are dispersed by mammals (Henderson, 2002). Mature individuals often produce fruits once a year (Henderson, 2002). At the study sites, mammals as Tapirus terrestris (Linnaeus, 1758), Dasyprocta sp. (Illiger, 1811), Thrichomys aperioideus (Lund, 1839), and Cynomys sp. (Thomas, 1916), are putative S. flexuosa seed dispersers (Zona and Henderson, 1989; Marinho-Filho et al., 2002). Except for T. terrestris, and in addition to Cuniculus paca (Linnaeus, 1776), all other species also prey on S. flexuosa seeds.

2.3. Sampling

At the edge and in the interior (habitat types) of Cerrado remnants, I sampled seedling abundance (August 2018) according to the distance from adult palms. To do...
that, in each Cerrado remnant, I delimited one transect in the edge at 25 m from the Cerrado boundary and one transect in the interior at 600 m away from the edge. Initially, I searched for adult palm individuals, identified by the presence of infructescence or vestiges of previous fruiting events. Then, I randomly selected at each habitat type 10 adult individuals (height ≥ 3 m) located in flat areas at least 40 m away from the nearest conspecific adult. For each adult individual, I counted seedlings along four 20-m transects, which were randomly set starting from the base of the palm stem. To count seedlings, in each transect, five plots of 3 m² (rectangle of 1 m [width] x 3 m [length]) were positioned at every 5 m from the adult palm (presumably parent; 0, 5, 10, 15, and 20 m). Non-dispersed fruits may be found on the soil surface under palm crowns (commonly from 0.5 to 2-3 m radius from the base of the palm stem) after falling from the infructescence. Thus, seedlings found up to 3 m from a given focal adult palm (i.e., under the palm crown) were considered as originating from non-dispersed seeds. On the other hand, those located between > 3 m and 23 m away from palms were assumed as originating from dispersed seeds (Draczler et al., 2011; Andreazzi et al., 2012). Seedlings had simple leaves and no stem, or had pinnate leaves and also no stem. In both cases, I counted, only seedlings smaller than 25 cm in height. I defined this ontogenetic criterion in order to evaluate early seedling establishment resulting from recent animal dispersal events, in face of the focus on the edge effect on seedling recruitment. Both in the edge and interior, seedlings were easily detected due to the scarce understory.

2.4. Data analysis

To compare the abundance of seedlings between the edge and interior of each remnant, I used a t-test in which I took as a replicate the number of seedlings sampled per parent palm; these data were log-transformed to achieve normality. To compare seedling distribution along distance classes from parent palms [edge versus interior], I used a Chi-square contingency analysis. In each edge or interior of every remnant, I took the accumulated number of seedlings sampled at each distance class to compare their distribution at edges with at interiors. In a contingency table, columns (distance class [0, 5, 10, 15, and 20 m]) represent the different states of one nominal variable, while rows (habitat type [edge or interior]) represent the states of another nominal variable. The cells include the counts of occurrences (accumulated number of seedlings) of that specific state (distance class, habitat type) of the two variables. The significance of association between the two variables (based on chi-squared) is then given, with p values from the chi-squared distribution. The contingency table analysis then gives information on whether the two variables are associated.

3. Results

Along the Matão edge, seedlings around parent palms ranged from 2 to 29 (average ± se: 8.2 ± 2.5, N = 82 seedlings), while in the interior ranged from 4 to 27 (13.4 ± 2.2, N = 134). This difference was significant (t-test, t = 2.18, P < 0.050). In the São Matheus the number of seedlings was also higher in the interior (from 2 to 27; 9.6 ± 2.4, N = 96, versus [at the edge] from 1 to 11; 4.7 ± 1.1, N = 47, t = 2.02, P < 0.050), as well as in the Bom Jardim remnant (interior: from 2 to 13; 8.4 ± 1.2, N = 84, versus edge: from 1 to 11; 5.7 ± 1.3, N = 57, t = 2.05, P < 0.050; Figure 1). Seedling distribution across distance classes differed between habitat types. For palms located at the edges, most seedlings were found under parent crowns (Matão: 69.5%, São Matheus: 53.2%, and Bom Jardim: 56.6%), while the contrary occurred in the interiors, where most seedlings were found away (> 3m) from palm individuals (Matão: 30.6%, São Matheus: 32.3%, and Bom Jardim: 19.0%; contingency, Matão: $\chi^2 = 40.27$, $P < 0.0002$, São Matheus: $\chi^2 = 10.48$, $P < 0.030$, and Bom Jardim: $\chi^2 = 21.68$, $P < 0.0003$; Figure 2).

![Figure 1](image-url). Number of Syagrus flexuosa seedlings (log transformed) found around parent palms at the edge and in the interior of Cerrado remnants (namely Matão, São Matheus, and Bom Jardim; Três Lagoas, MS, Brazil, August 2018). Bars show mean ± standard errors.
4. Discussion

It is well known that seed dispersal can decrease the spatial seedling aggregation by spreading seeds over a larger area (Howe et al., 1985; Galetti et al., 2006; Dracxler and Forget, 2017). Moreover, studies have demonstrated that seed removal depends positively on disperser abundance (Alcántara et al., 1997). In the interior of Cerrado remnants, the higher number of seedlings found far from parent palms may reflect the persistence of the seed dispersal process (Howe et al., 1985; De la Peña-Domene et al., 2016). Scatter-hoarding rodents, for example, commonly move palm seeds of distances between 5-10 m from the seed source (Jansen et al., 2014; Dracxler and Kissling, 2021), which, in the interiors, is coherent to the observed distribution of seedlings around parent palms. On the other hand, the high seedling concentration under parent palms suggests that large-seeded dispersers interact less with *Syagrus flexuosa* fruits present in the edges (Silva and Tabarelli, 2000; Ragusa-Netto, 2017). Presumably, once on the soil, most *S. flexuosa* fruits continued untouched, as documented in other anthropized areas (Wright and Duber, 2001; Ragusa-Netto, 2017). Indeed, in sites where key seed dispersers were extirpated, large-seeded palm species suffer limitations on long-distance seed dispersal (Wright and Duber, 2001). In these conditions, important to consider the potential density-dependent seed mortality caused by natural enemies under parent palms (Janzen, 1971; Johnson et al., 1995; Terborgh, 2012; Ragusa-Netto, 2019). *Syagrus flexuosa* produce fruits asynchronously (Henderson, 2002), and, during the dry season curculionid larvae (*Revena sp* Casey, 1922) prey on average, only 9% of their seeds (Ragusa-Netto, J, unpublished results). Then, the diaspores accumulated under palms during the diapause of bruchid beetles (dry season), might experience a less severe mortality rate (Wright, 1990). Starting the wet season, those intact seeds might germinate (Gandolfi et al., 2007). The reduced seed dispersal added to a decreased seed mortality during the dry season is coherent with seedling concentration under adult trees (Silva and Tabarelli, 2000). According to that, in sites where large-seeded dispersers declined, an increase of seedling patches has been documented due to seed accumulation beneath parent palms (Wright and Duber, 2001). Even

Figure 2. Proportions of *Syagrus flexuosa* seedlings found at different distance classes from parent palms at the edge and interior of Cerrado remnants (namely Matão, São Matheus, and Bom Jardim; Três Lagoas, MS, Brazil, August 2018). Bars show mean ± standard errors.
then, despite the initial development, it is important to consider that seedlings may become unable to growth close to parents, incurring in mortality process (Gandolfi et al., 2007). Many studies have shown that seedling mortality rates are higher near conspecific adults (Wright, 2002). In the studied edges, no late-stage seedling or sapling was present close to adult palms, suggesting a reduced recruitment process under adults S. flexuosa.

Herbivores as peccaries, tapirs, and ants often feed on palm seedlings (Silman et al., 2003; Wirth et al., 2008). All of them occurred in the studied fragments (Ragusano-Netto, J., pers. obs.), and it is unlikely that, only in the interior, those herbivores consumed seedlings close to adult palms. Actually, the edge vegetation often favors generalist herbivores, which may exert an important regulatory role on seedling abundance at these areas (Wirth et al., 2008). Syagrus flexuosa seedlings were less abundant at edges but common close to adult individuals, suggesting a decline of interactions with medium-large seed vectors (Wright and Duber, 2001; Muñoz et al., 2019). Although some palm species can exhibit increased abundance along forest edges (Souza and Martins, 2003; Pimentel and Tabarelli, 2004), I documented here that S. flexuosa individuals found at Cerrado fragment edges show patterns of seedling abundance and distribution unfavorable to an intense regeneration process (Benítez-Malvido et al., 2018). Then, because S. flexuosa is a widely distributed palm that serves as an important resource for several animals across Cerrado habitats (Lorenti et al., 2010), changes in their regeneration process due to edge effects can further impact frugivore populations. In this respect, as the recruitment of large-seeded tree species arises as a priority to tropical vegetation conservation (Melo et al., 2007), S. flexuosa progressive rarity at fragments’ edges implies substantial changes that may compromise the persistence of other species and ecological processes, as well (Peters et al., 2008; Arroyo-Rodríguez et al. 2007; Benítez-Malvido et al., 2018). Moreover, the differences between edges and interiors in the seedling recruitment process observed here may be widespread among other large-seeded tree species (Melo et al., 2007; Ragusano-Netto, 2017). Thus, Cerrado fragments might have similar problems of species impoverishment, as documented for tropical forests (Silva and Tabarelli, 2000; Melo et al., 2007; Tabarelli et al., 2008; Browne and Kubkiyan, 2016). In this way, with only half of its original area left, the accelerated clearing of Cerrado needs urgent intervention (Carvalho et al., 2009; Beuchle et al., 2015). Conservation actions to re-establish S. flexuosa populations at edges may require the management of Cerrado remnants to imitate undisturbed sites, thereby preventing mortality of adult palms and promote the recruitment of juveniles.

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