Flood regime and water table determines tree distribution in a forest-savanna gradient in the Brazilian Pantanal

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ABSTRACT

This study aimed to recognize the preferential location of species of the tree sinuaria in response to a moisture gradient in Pantanal Matogrossense, Brazil. We established sampling plots of arboreal sinuaria along a soil moisture and flood gradient. Piezometers were installed, allowing monthly measurements of water table depth and flood height during one year. Detrended Correspondence Analysis, Gradient Direct Analysis, Multi-response Permutation Procedures and Indicator Species Analysis were performed to evaluate the effect of moisture gradient on tree distribution. The annual variation of water table is shallower and similar in Seasonally Flooded Forest and Termite Savanna, with increasing depths in Open Savanna, Savanna Forest and Dry Forest. Circa 64% of the species were characterized as having a preferential location in “terrestrial habitats normally not subjected to inundation”, while 8% preferentially occur in “wet habitats”. Lowest tree richness in flood-affected vegetation types is related to both present-day high climatic seasonality and Late Pleistocene dry paleoclimates in the Pantanal wetland. The tree distribution across different formations in the Pantanal shows a direct relationship with soil moisture gradient.

Key words: floristic composition, flood pulse concept, wetlands, RPPN SESC Pantanal.

INTRODUCTION

The Pantanal Matogrossense is one of largest wetland systems in the world, and the largest seasonally flooded area in South America (Nunes da Cunha and Junk 2015). The great diversity of permanent habitats and seasonally waterlogged and permanent dry areas is the ecological basis for the diversity of species in Pantanal (Junk et al. 2006).

Flood is one of the controlling factors of maintenance of fauna, flora and landscape diversity in Pantanal (Junk et al. 1989, Nunes da Cunha and Junk 1999, 2001). Flood cycle determines the availability of dry and wet areas, which in turn
influences the spatial distribution of many species. Flood provides a permanent exchange of water, sediments, chemical components and organisms between the main river channel and adjacent floodplains, resulting in a great environmental heterogeneity (Lourival et al. 2000).

Based on the concept of flood pulse in floodplain systems (Junk et al. 1989), Nunes da Cunha and Junk (2001) stated that tree vegetation in Pantanal is distributed over a flooding gradient, in accordance to tolerance to either flood or drought. This differential tolerance, with morphological and physiological adaptations of species, leads to formation of peculiar forest communities.

Phillips and Gentry (1994) reported that different disturbances, at different scales, are responsible for the environmental heterogeneity observed in tropical forests, exerting strong influence on structure, dynamics and composition of species of tree communities. Among these disturbances, flood acts in a regional scale, similarly fragmentation, burning and wind-exposure, all affecting the composition and structure of plant communities (Ferreira and Laurance 1997, Cochrane and Schulze 1999).

In tropical forests, rivers are important sources of disturbances due to the annual flooding regime, and also because of river dynamics at landscape scale (Klinge et al. 1990). Some plant communities on floodplains represent the initial phases of primary succession and are, usually, formed by a small group of fast growing species, well adapted to periodic flood (Junk 1989). Permanent water movement and deposition of sediments affect the establishment, growth and reproduction of different species, resulting in instability and a constant reorganization of plant communities (Siebel and Blom 1998).

Flood imposes varying changes on physical, chemical and biological soil processes, affecting soil structure, O₂ diffusion, CO₂ emissions, anaerobic decomposition of organic matter, depletions or toxicity of iron and manganese, all limiting the soil capacity for supporting plant growth (Kozlowski 2002, Bailey-Serres and Voesenek 2008).

Plant tolerance to flood varies according to species, genotype and age of the plant; intensity and duration of the flood, and chemical composition of water (Kozlowski 2002). Flood promotes strong effects on germination and initial development of seedlings because physiological processes needed for germination requires hight O₂ availability (Kozlowski and Pallardy 1997). Adverse effects from flooding are responsible for changes on distribution, species composition and successional processes, as demonstrated by Arieira and Nunes da Cunha (2006) and Amorim and Batalha (2007).

The aim of this work was to evaluate the preferential location of species of the tree sinuas in response to a moisture gradient in the Pantanal. We determined the pattern of water table annual oscillation and flood found in the main tree formations along a “savanna (cerrado)-forest” gradient. Accordingly, we expected to answer the following questions: i) Is species composition variable with different annual water table and flood oscillation patterns? ii) Is the water table oscillation pattern similar in areas free from annual flood (Dry Forests and Savannas)? iii) Which species show preferential distribution in the wet or dry extremes of the moisture gradient? iv) Which tree species present wide spatial distribution across the gradient?

MATERIALS AND METHODS

STUDY SITE

This study was carried out within the Reserva Particular de Patrimônio Natural SESC Pantanal (RPPN SESC Pantanal) (16°34’50”S and 56°15’58”W), in Barão de Melgaço, Mato Grosso State, western Brazil. It is located on the Cuiabá and São Lourenço rivers interfluve (Fig. 1). Average precipitation is from 1,100 to 1,200 mm (Hasenack et al. 2003). We observed the predominance of herbaceous fields, savannas (cerrados) and forests mixed to water bodies and aquatic vegetation.
Three vegetation transects were selected, where tree physiognomies were sampled in plots with 20 m × 100 m, divided in subplots with 20 × 20 m. Transect 1 began with a Termite Savanna (Campo de Murundus) adjacent to the paleo-levée (cordilheira) towards its highest central section, crossing four forested phytophysiognomies. The Cordilheiras are geomorphological units formed by river sandy bars of former high energy sedimentation, now stabilized and forming the highest parts of the landscape, usually with sandy sedimentary material (Nunes da Cunha and Junk 2011). Transect 2 began in a Semi Evergreen Forest, across a Termite Savanna, followed by a cordilheira with a Savanna Forest (Cerradão), ending with a Forest in the sequence. Transect 3 began with a Termite Savanna, followed by Open Savanna (Cerrado stricto sensu) physiognomy. Only individuals with DBH (diameter at 1.30 m above ground level) ≥ 4.77 cm for areas with forest physiognomy were sampled and identified. For Open Savanna physiognomy a inclusion criteria of DSL (diameter at soil level) ≥ 4.77 cm was used (Felfili et al. 2005).

Dry Forest and Savanna facies were identified, each facies named considering the populations of tree species more abundant, based on the structure. The moisture gradient was monitored measuring the seasonal water table depth oscillation, using piezometers at each site, including Termite Savannas, and measuring flood height.

DATA ANALYSIS

Classification and ordination analyses were applied to a matrix of binary data. Species with single occurrences (singleton species) were excluded in order to better understand the possible grouping of the samples, according to their most frequent elements. A Detrended Correspondence Analysis - DCA (Hill and Gaugh 1980) was performed to detect floristic patterns, or gradients.

Multi-response Permutation Procedures - MRPP (Mielke Jr et al. 1981) with the Jaccard index and pairwise comparisons was used to check the differences in species composition between the phytophysiognomies along the moisture gradient. The MRPP reports a test statistic (T) describing the separation between groups, with smaller, or negative values indicating greater separation. The
agreement statistic (A) describes within-group homogeneity compared to random, where values close to zero indicating the degree of heterogeneity expected by chance, and values close to one indicating highly homogeneous groups. Values below zero correspond to highly heterogeneous groups (Faith et al. 1987).

The way in which the moisture gradient affects the preferential localization of plant species was evaluated by Indicator Species Analysis (ISA) (Dufrêne and Legendre 1997) and Direct Analyses of Gradients (Gauch-Jr 1991) with the distribution of the species with greater indicator value (IV) derived from the Indicator Species Analysis (ISA). The statistical significance of species IV was assessed via Monte Carlo randomizations. DCA, MRPP and ISA analyses were performed with the PC-ORD program, version 6.0.

Finally, species were classified according to their preferential habitat (Nunes da Cunha and Junk 1999). These authors classified the Pantanal species based on their preferential localization in: S = terrestrial habitats normally not subjected to inundation, T = habitats inundated during short periods (< two months); A = habitats inundated during long periods (> two months).

RESULTS

VEGETATION FACIES

Fig. 2 shows the phytophysiognomies and their respective facies sampled in the studied transects. Dry Forest, Seasonally Flooded Semi Evergreen Forest (Cambarazal), Savanna Forest (Cerradão de Carvoeiro) and Open Low-tree Savanna (Cerrado de Lixeira) are the arboreal formations, whereas Termite Savanna represents the grassland formation (Fig. 2).

The Dry Forest, located in well-drained habitats (paleo-levée), were divided in five facies:
1. Transitional Dry Forest with Scheelea phalerata; 2. Dry Forest with S. phalerata and Callisthene fasciculata; 3. Dry Forest with S. phalerata and Myracrodruon urundeuva; 4. Dry Forest with S. phalerata, M. urundeuva and C. fasciculata; 5. Dry Forest with S. phalerata, M. urundeuva and Combretum leprosum (Fig. 2).

A Seasonally Flooded Semi Evergreen Forests dominated by Vochysia divergens Pohl (regionally known as Cambarazal) (Arieira and Nunes da Cunha 2006) are located in the hydromorphic lowlands, representing wet areas along the moisture gradient. The Savanna Forest (Cerradão) and the Open Low-tree Savanna (Cerrado sensu stricto) were the two savanna phytosociologies studied, locally called Cerradão de Carvoeiro and Cerrado de Lixeira, respectively. The Cerradão de Carvoeiro were divided in two facies: 1. Cerradão de Carvoeiro with Magonia pubescens and 2. Cerradão de Carvoeiro with Lafoensia pacari. The Cerrado de Lixeira was represented by a single facie: 1. Cerrado de Lixeira with Terminalia argentea (Fig. 2).

VEGETATION VERSUS MOISTURE GRADIENT

The relation between phytosociology and water table depth is illustrated in Table I and Fig. 3.

Dry Forests have the lowest water table during the year, with low seasonal variability. The water table oscillation regime in Cerrados was similar for Cerradão de Carvoeiro and Cerrado de Lixeira, however, with slight higher depths during the dry season in the latter (Table I; Fig. 3). The Cerrado assumes a denser forested physiognomy (Cerradão de Carvoeiro) with accompanying increment in richness (23 species in Cerrado da Lixeira to 33 species in Cerradão de Carvoeiro), as water table remains deeper during the wet season (1.90 m and 3.21 m, respectively). This suggest that the different Cerrado phytosociologies in Pantanal are strongly conditioned by the water table depth in the wet season.
At the other extreme of the moisture regime are Termite Savannas and Cambarazal, where water table oscillation is close to the surface year round, with waterlogged soils during the rainy season. However, Cambarazal differs from the other phytophysiognomies for being subject to annual seasonal flood (Table I; Fig. 3).

Concerning the water table and flood oscillations over the year (Table I, Fig. 3), we ranked the Pantanal phytophysiognomies by increasing water saturation and flooding, as follows: Dry Forests < Cerradões de Carvoeiro (Savanna Forest) < Cerrado de Lixeira (Open Low-tree Savanna) < Termites Savannas < Cambarazal (Semi Evergreen Forest).

Figure 2 - Phytophysiognomies and the respective vegetation facies sampled in transects 1, 2 and 3 in RPPN SESC Pantanal, Brazilian Pantanal, where: 1. Transitional Dry Forest (DF) with Scheelea phalerata; 2. DF with S. phalerata and Callisthene fasciculata; 3. DF with S. phalerata and Myracrodruon urundeuva; 4. DF with S. phalerata, M. urundeuva and C. fasciculata; 5. Cambarazal; 6. Cerradão de Carvoeiro with Lafoensia pacari; 7. Cerradão de Carvoeiro with Magonia pubescens; 8. DF with S. phalerata, M. urundeuva and Combretum leprosum; 9. Cerrado de Lixeira with Terminalia argentea.
We sampled one hundred and eleven tree and shrub species distributed in 77 genus and 35 families. Among the studied formations, we recorded 68 species in Dry Forests, 33 species in the Cerradão de Carvoeiro, 23 species in Cerrado de Lixeira and 15 species in Cambarazal.

Quantitative analyses confirm a floristic differentiation among the studied phytosociological formations. The Dry Forest facies sampled are separated from Cerradão de Carvoeiro on the left of the first axis of DCA (Fig. 4), not forming a isolated floristic group, but rather a gradient. The other two studied phytosociological formations, Cerrado de Lixeira and Cambarazal, formed distinct floristic group.

A moisture gradient was confirmed in the axis 1 of the DCA (Fig. 4). The Seasonally Flooded Semi Evergreen Forest (Cambarazal), situated in hydromorphic lowlands, are on the right position.

**Table I**

<table>
<thead>
<tr>
<th>Phytophysiognomies</th>
<th>Water table average depth (m)</th>
<th>Flood (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Wet season</td>
<td>Dry season</td>
</tr>
<tr>
<td>Dry Forest facies</td>
<td>5.60 ± 0.33</td>
<td>5.30 ± 0.82</td>
</tr>
<tr>
<td>Cerradão de Carvoeiro facies</td>
<td>3.21 ± 1.36</td>
<td>4.00 ± 1.75</td>
</tr>
<tr>
<td>Cerrado de Lixeira</td>
<td>1.90 ± 0.26</td>
<td>4.10 ± 2.10</td>
</tr>
<tr>
<td>Termite Savanna</td>
<td>1.32 ± 0.27</td>
<td>2.75 ± 0.86</td>
</tr>
<tr>
<td>Cambarazal</td>
<td>1.30 ± 0.46</td>
<td>3.30 ± 1.19</td>
</tr>
</tbody>
</table>

**Figure 3** - Mean values (m) of annual range of water table depth and flood height in the studied formations in RPPN SESC Pantanal, Brazilian Pantanal.

**Figure 4** - Two-dimensional plots ordination diagram derived from DCA for arboreal sinusiae along a soil moisture and flood gradient in RPPN SESC Pantanal, Brazilian Pantanal. △ - Dry Forests facies; ○ - Cerradão de Carvoeiro facies (Savanna Forest); □ - Cerrado de Lixeira (Open Low-tree Savanna); ● - Cambarazal (Seasonally Flooded Semi Evergreen Forest).
in axis 1; the *Cerradão de Carvoeiro* and *Cerrado de Lixear*, in the middle of the axis 1, are in an intermediate position in the moisture gradient; finally, the Dry Forests on the left of the axis 1, are in the driest habitats.

There were significant differences in species composition between the four phytophysiognomies (T value = -20.52; observed δ = 0.200; expected δ = 0.500; A = 0.59; p< 0.0000) based on MRPP analysis. The pairwise comparisons (Table II) indicated differences between all phytophysiognomies, in term of the arboreal sinuasiae. In this regard, Dry Forest and *Cambarazal* and Dry Forest and *Cerrado de Lixeira* show greater distinction, while *Cambarazal* and *Cerrado de Lixeira* are less differentiated.

**TABLE II**

Summary statistics for pairwise comparison (MRPP) among the studied phytophysiognomies along a soil moisture and flood gradient in RPPN SESC Pantanal, Brazilian Pantanal. T = degree of separation between the groups, higher negative values indicating greater separation A = level of within-group agreement, comparing the distance within and between the groups, higher values indicating greater separation.

<table>
<thead>
<tr>
<th>Pairwise comparison of phytophysiognomies (MRPP)</th>
<th>T</th>
<th>A</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry Forest facies vs. <em>Cambarazal</em></td>
<td>-14.91</td>
<td>0.36</td>
<td>0.0000</td>
</tr>
<tr>
<td>Dry Forest facies vs. <em>Cerradão de Carvoeiro</em> facies</td>
<td>-12.74</td>
<td>0.29</td>
<td>0.0000</td>
</tr>
<tr>
<td>Dry Forest facies vs. <em>Cerrado de Lixeira</em></td>
<td>-14.24</td>
<td>0.34</td>
<td>0.0000</td>
</tr>
<tr>
<td><em>Cambarazal</em> vs. <em>Cerradão de Carvoeiro</em> facies</td>
<td>-7.63</td>
<td>0.46</td>
<td>0.0002</td>
</tr>
<tr>
<td><em>Cambarazal</em> vs. <em>Cerrado de Lixeira</em></td>
<td>-5.59</td>
<td>0.45</td>
<td>0.0017</td>
</tr>
<tr>
<td><em>Cerradão de Carvoeiro</em> facies vs. <em>Cerrado de Lixeira</em></td>
<td>-6.98</td>
<td>0.40</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

The Indicator Species Analysis and Gradient Direct Analysis can detect ecological preferences of certain tree species in a moisture gradient. The indicator value (IV) of six major indicative species in each phytophysiognomy is illustrated in Fig. 5. ISA results (p<0.05) presented six species indicators of Dry Forest (in descending order of IV), as follows: *Scheelea phalerata* (Mart.) Bur., *Myracrodruon urundeuva* (Engl.) Fr. All., *Tabebuia roseo-alba* (Rid.) Sandw., *Casearia gossypiosperma* Briquet., *Combretum leprosum* Mart. and *Platypodium elegans* Vog.

For *Cerradão de Carvoeiro*, the species with significant IV were *Lafoensia pacari* A. St. Hil., *Qualea parviflora* Mart., *Plathymenia reticulata* Benth., *Callisthene fasciculata* Mart., *Pseudobombax tomentosum* (Mart. & Zuc.) Robyns, *Magonia pubescens* A. St.-Hil. (Fig. 5) and *Cordia glabrata* (Mart.) A. DC. (IV=37.1). For *Cerrado de Lixeira*, the species were *Zanthoxylum hasslerianum* (Chodat) Pirani, *Diospyros hispida* DC., *Byrsonima coccolobifolia* (L.) H. B. K., *Erythroxylum suberosum* A. St.-Hil., *Pouteria ramiflora* (Mart.) Radlk., *Davilla elliptica* A. St.-Hil. (Fig. 5), *Mouriri elliptica* Mart. (IV=60), *Terminalia argentea* Mart. & Zuc. (IV=58), *Bowdichia virgilioides* Kunth (IV=56.3), *Luehea paniculata* Mart. (IV=54.2), *Tabebuia aurea* (Manso) B. & H. ex Moore (IV=52.3), *Curatella americana* L. (IV=41.6), *Hancornia speciosa* Gom., *Kielmeyera coriacea* Mart. and *Andira cuyabensis* Benth., *Qualea grandiflora* Mart. and *Himatanthus obovatus* (Mull. Arg.) Woods (IV=40) and *Simarouba versicolor* A. St.-Hil. (IV=36.1).

For *Cambarazal*, in turn, typical species were *Vochysia divergens*, *Rheedea brasiliensis* (Mart.) Pl. & Tr., *Licania parvifolia* Huber, *Alchornea discolor* Poepp., *Andira inermis* H. B. K., *Eugenia*
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florida DC. (Fig. 5), Mouriri guianensis Aubl. (IV=40) and Brosimum lactescens (Moore) Berg (IV=40).

Although not a single species has been sampled in all the studied phytophysiognomy, Astronium fraxinifolium Schott showed the widest distribution in the moisture gradient, occurring in eight of the nine facies (89% of occurrence), being absent only in Cambarazal. Other four species, Callisthenes fasciculata, Curatella americana, Magonia pubescens and Dipteryx alata Vog. were sampled in seven facies (78%).

In line with Fig. 6, we can observe that the number of species associated with seasonally flood areas (AT) is reduced, compared with species typical of well-drained habitats, normally not subject to inundation (S). Casearia gossypiosperma, Combretum leprosum, Tabebuia roseo-alba, Myracrodruon urundeuva, Astronium fraxinifolium, Anadenanthera colubrina, Pouteria ramiflora, Scheelea phalerata and Qualea parvifolia are some of the 71 species (64%) with a preferential location in terrestrial habitats normally not subject to inundation (S) (Fig. 6).

The other 20 species (18%) are from terrestrial habitats, but can stand flooding during short periods (ST), as for example, Curatella Americana, Cecropia pachystachya Trécul and Simarouba versicolor (Fig. 6).

Only nine species (8%) are typical from “inundated habitats” (AT), for example, Vochysia divergens, Licania parvifolia, Rheedia brasiliensis, Mouriri guianensis and Brosimum lactescens. The remaining species, 11 (10%), have unespecific, wide distribution along the moisture gradient (STA, TAS, TSA, ATS) (Fig. 6). For example, Bromelia balansae Mez. was observed in the understorey of all facies in Dry Forest and in facies in Cerrado

![Graph showing species distribution](image)

**Figure 5** - Gradient direct analysis with distribution of the species identified as significant indicators based on Indicator Species Analysis (ISA) in each phytophysiognomy of Brazilian Pantanal.
de Carvoeiro, but was absent in Cerrado de Lixeira and in Cambarazal.

**DISCUSSION**

**EXAMPLES OF ADAPTATION TO DROUGHT AND FLOODING**

Overall, these results indicated that water table depth and flood height influence the species composition and distribution at each plant community in the vegetation gradient. In line with that, we observed that a given group of species is well adapted to flooding, such as those from Cambarazal. Similarly, another group is typical of high water table level, but not subjected to inundation, in the Cerrado da Lixeira and the Cerradões de Carvoeiro, and finally, the Dry Forest is composed of species with very low tolerance to waterlogging and seasonal flood conditions.

Tree vegetation in Pantanal is distributed along a moisture gradient according to its degree of tolerance to inundation or drought. In some cases, survival during flood periods is due to the existence of physiological and phenological peculiarities of the species. *V. divergens*, a dominant species in Cambarazal, for example, have seedlings and young individuals that can stand inundation and survive until the end of flooding period. On the other hand, adult individuals show deciduity at the end of flooding period, followed by a rapid leaf replacement (Nunes da Cunha and Junk 2001).

Survival to water stress in the dry season is due to phenological, anatomical and physiological adaptations, as observed with *A. colubrina* and *Anadenanthera macrocarpa* (Benth.) Brenan. Both loose virtually all leaves in the dry season, and have tuberous root system, in addition to reduced growth rate during periods of low water availability (Silva and Barbosa 2000, Monteiro et al. 2006). Other species, such as *Curatella americana*, *Bowdichia virgilioides* and *Casearia sylvestris* Sw., have leaf mesophile structure with typical xeromorphic traits which reduce water losses. The main features are reduced intercell space, large vascular bundles, colorless parenchyma, very thick

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*Figure 6* - Distribution of 111 tree species in the Brazilian Pantanal according to their preference along the moisture gradient. S = terrestrial habitats normally not subjected to inundation; T = habitats inundated during short periods (< two months), A = habitats inundated during long periods (> two months). Arrows indicate the supposed direction of expansion of the species from the center of maximum density (according to Nunes da Cunha and Junk 1999).
cuticles and trichomes (Sarmiento et al. 1985), and fire resistance.

WHY ARE THERE MORE SPECIES IN DRIER HABITATS OF PANTANAL?

Recent estimates of the number of plant species for Pantanal show a total richness of 2,000 species distributed in 144 families, in which 104 families are exclusively of terrestrial habitats, 21 families are exclusively aquatic and 19 both terrestrial and aquatic species. Most species (1,656) are terrestrial: 900 are herbs, grasses, vines, epiphytes and parasites, whereas the remaining 756 are woody plants like shrubs, sub shrubs, trees, lianas and palms. Only 247 species are considered as hydrophytes or aquatic macrophytes (Pott et al. 2011, Junk et al. 2006).

In Pantanal there is less flood-resistant trees than species adapted to well-drained lands. Concerning this, Nunes da Cunha and Junk (1999) reported for the Pantanal of Mato Grosso (Brazil), north of Poconé that only four were exclusive from flooded environments, out of 86 tree species sampled. Two possible and complementary explanations for the low richness of flood-resistant tree species in Pantanal are: (1) the highly seasonal moisture regime in the present, and (2) inheritance from prevailing semi-arid paleoclimates in the Late Quaternary of the Pantanal plains (Ab’Sáber 2006).

Before the onset of Late Quaternary wet climates, the Pantanal remained a vast semi-arid lowland, unfavourable to species typical to wetlands (Pott et al. 2011). In fact, the rainfall of the part of Pantanal, towards the borders with Bolivia and Paraguay, is greatly reduced to less than 800 mm/year (Ab’Sáber 2006), while most areas have rainfall ranges from 1,000 mm to 1,400 mm/year (Hasenack et al. 2003), with very high seasonality and long dry season.

The paleoclimatic history of Pantanal is marked by alternating dry spells during the Late Pleistocene and early Holocene, which accounted for the “invasion” of a xeric and mesic flora from the adjacent drylands, like Chaco and Cerrado (Ab’Sáber 2006, Junk et al. 2006). The occurrence of dry paleoclimate spells in the Late Pleistocene (between 23,000 – 13,000 y.b.p.) enabled the expansion of the semi-arid arboreal and shrubry flora on the Pantanal plains (Prado 2000, Ab’Sáber 2006).

The biodiversity of Pantanal has experienced strong climate disturbances in the past. Nonetheless, the expected raise in climate fluctuations will occur with unprecedented levels by human impacts (Úbeda et al. 2013). Typical examples are (1) eutrophication of rivers resulting in cyanobacteria blooms (Cózar et al. 2012), (2) conversion of natural vegetation into pastures for cattle grazing, (3) conversion of Cerrado vegetation into crop fields in the high plateau headwaters, (4) environmental contamination (eg. agricultural chemicals and mercury contamination by gold mining), (5) unregulated tourism, (6) introduction of exotic species and (7) infrastructure and unplanned human occupation (Alho 2011, Nunes da Cunha and Junk 2015). According to Prado and Gibbs (1993), Pennington et al. (2004), Caetano et al. (2008) and Linares-Palomino et al. (2009) Dry Forests occur in South America along an extensive arch named “Pleistocene Arc of Dry Formations”, where Dry Forests expanded during the Late Pleistocene, followed by fragmentation in the Holocene. Its maximum extension occurred in the dry, cool period between 18,000 to 12,000 years ago.

The species Anadenanthera colubrina, Myracrodruon urundeuva, Pterogyne nitens Tul., Combretum leprosum, Cereus peruvianus (L.) Mill., Combretum leprosum, Cordia glabrata and Maclura tinctoria (L.) Engl. have been described as typical of the Pleistocene Arc of Dry Formations of South America (Prado and Gibbs 1993, Prado 2000). Accordingly, the presence of these species
can be interpreted as relicts of a former wider xeric flora which remained in places where current local environmental conditions (microclimate, soils) were favourable to their maintenance.

After the Late Pleistocene dry spells, the newly established trend of wetter conditions in the Holocene allowed elements coming from Cerrado and Pre-Amazon source to invade the former dry space. The Cerrado flora, present in the adjacent high tableland (chapadas, entered the Pantanal through the northeastern and eastern sectors of the depression, mainly along the alluvial fan of Taquari river (Ab’Sáber 2006, Junk et al. 2006).

**EFFECT OF FLOOD ON TREE PHYTOCENOSES IN PANTANAL AND FLOODPLAIN FORESTS IN THE AMAZON REGION**

The number of tree species in Pantanal adapted to long periods of flood is much lower compared with the observed for the Amazon floodplain. It is estimated that on Amazonian floodplains, there are more than 1,000 tree flood-resistant species. For the Pantanal, the estimates account for 355 species tolerating different degrees and intensities of flood (Junk et al. 2006).

Species richness and composition, structure (Ferreira 2000), total area, number and composition of species of canopy gap regenerations (Ferreira and Almeida 2005) and phenological events of tree species of lowland forests in Central Amazon (Ferreira and Parolin 2007) vary with flood susceptibility, where typical plant communities develop in response to their adaptation to flood.

Areas subjected to longer flood periods showed lower richness and diversity in function of selective pressure of adaptation to low oxygen in the rizosphere (Junk 1989). Reduction on species richness and diversity in function of increasing flood duration and intensity was also registered by Ferreira (2000) in flooded Forests in the Jaú and Tarumã-Mirim floodplains, although this author claimed that other factors, such as the distance of the study area from upland forest and class diversity, might also contribute to the differences observed. Concerning the Pantanal, Arieira and Nunes da Cunha (2006) observed a negative correlation between flood height and species diversity in a Seasonally Flooded Semi Evergreen Forest (Cambarazal) on the floodplain of river Cuiabá, in Pantanal of Poconé.

The differences in distribution of dominant species in flooded forests is probably related to their ability to tolerate seasonal floods. However, the role the soil and topographical attributes on the distribution of plant species become significant, especially for less dominant species, locally rare or generalists (Ferreira 2000).

The coexistence of species with very contrasting environmental requirements in Pantanal is only possible because of the proximity of large adjacent phytogeographical spaces (Chaco, Cerrado and Amazonia). Due to a great geomorphological and soil diversity over short distances, the Pantanal landscape offers diverse habitats that, under varying flooding intensity/height, allows the establishment and survival of a peculiar flora.

We concluded that the moisture gradient is determining factor to explain floristic variations of plant communities. Hence, flood and soil water table oscillations are key environmental (abiotic) variables which define plant community patterns in Pantanal.

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RESUMO

O objetivo deste estudo foi conhecer a localização preferencial de espécies da sinuária arbórea em resposta ao gradiente de umidade do solo estudado. Foram estabelecidas parcelas para amostragem com base no gradiente de umidade ao longo dos diferentes tipos vegetacionais no Pantanal. Piezômetros foram instalados para permitir medições mensais da profundidade do lençol freático e da altitude da lâmina de inundação durante um ano. Análise de Correspondência Detrendenciada, Análise Direta de Gradientes, Procedimento de Permutação Multi-resposta e Análise de Espécies Indicadoras foram realizadas para avaliar o efeito do gradiente de umidade sobre a distribuição das plantas. A oscilação do lençol freático é mais superficial e similar na Floresta Sazonalmente Inundada e nos Campos de Murundus, com acréscimo da profunidade do lençol freático e da altura da lâmina de inundação durante um ano. A composição de espécies arbóreas em types vegetacionais sujeitos a inundação está relacionada à alta sazonalidade climática atual e aos paleoclimas secos no Pleistoceno Superior no Pantanal. Cerca de 64% das espécies foram caracterizadas como de localização preferencial em “habitats terrestres normalmente não sujeitos a inundação”, enquanto 8% ocorrem preferencialmente em “habitats inundáveis”. A baixa riqueza de espécies arbóreas em tipos vegetacionais sujeitos a inundação está relacionada à alta sazonalidade climática atual e aos paleoclimas secos do Pleistoceno Superior no Pantanal. A composição de espécies arbóreas ao longo das diferentes formações no Pantanal apresenta uma relação direta com o gradiente de umidade no solo estudado.

Palavras-chave: composição florística, conceito de pulso de inundação, áreas úmidas, RPPN SESC Pantanal.

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