Demography and Structures Population of Actinostemon concolor (Spreng.) Müll. Arg. (Euphorbiaceae) in Flooded Areas in Southern Brazil

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ABSTRACT

The aim of this work was to analyze the size and spatial structures and demography of Actinostemon concolor (Spreng.) Müll. Arg. (Euphorbiaceae) in the flooded areas of Mata dos Godoy State Park. Fifty plots of 100 m² were established and three surveys were conducted. All the individuals were measured, mapped and divided into categories as seedlings, juveniles and adults. The coefficient of skewness, L-function of Ripley, and the matrix population models were used in the study. The population showed positive coefficient of skewness and the individuals were distributed in groups. The population growth rates (λ) were equal or lower than 1.0 and the adult survival and recruitment of juvenile to adult were the vital rates that most affected the values of λ. Considering these results and the unpredictability of the environment, it could be concluded that the individual numbers of A. concolor would fluctuate over time, but the species would remain as one of the most important in the Park.

Key words: elasticity, matrix models, spatial distribution, structure size, tropical tree

INTRODUCTION

Due to agriculture expansion, the Seasonal Semideciduous Forest in the Southern and Southeast Brazil was deforested (Medri et al. 2002), resulting in a highly fragmented landscape. In this devastated scenario, the Riparian Forests were also degraded, although protected by law (Medri et al. 2002). These degradation processes were harmful to the riparian and water ecosystems. Currently, there are several projects aiming to recover the riparian degraded areas and ecosystem function. However, the riparian forest conservation depends on the knowledge about the tree population dynamics and structures, as suggested by van Groenendaal et al. (1996) for the tropical forest. These studies are considered essential to the understanding of the processes that regulate the dynamics and structure of natural communities and to the management and conservation programs (Santos et al. 1998). However, the life histories are unknown for the majority of tropical species (Wright et al. 2003). The structures of tropical tree populations are influenced by the environmental heterogeneity (Cielo-Filho et al. 2007; Bianchini et al. 2010). The heterogeneity can be promoted by the properties of soil, water availability and light intensity (Clark and Clark 1987; Condit et al.

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For *Geonoma macrostachys* Mart., an understory palm, Svenning (2002) detected greater recruitment and fecundity because of canopy openness that increased light availability in the understory. Fonseca et al. (2004) observed that the seedlings of *Aspidosperma polyneuron* Müll. Arg. tended to be concentrated in one region of the area and not distributed in scattered clumps, as reported in other two studied areas. These authors associated the results with slope and its related edaphic factors.

In this work, the population structures and demography of *Actinostemon concolor* (Spreng.) Müll. Arg. (Euphorbiaceae), a common understory tree of the Brazilian Seasonal Semideciduous Forest (Smith et al. 1988) and Riparian Forest was studied. This species is more abundant in flooded areas of Riparian Forest (Bianchini et al. 2003). The following aspects were looked into: the size distribution, the population size, the parameters (fecundity, survival or recruitment) that most affect the species demography, the spatial pattern of the population and the abiotic factors related to this spatial pattern.

**MATERIAL AND METHODS**

This study was carried out in the Mata dos Godoy State Park - MGSP (23°27’S, 51°15’W, visitors center), near Londrina town, Southern Brazil. The MGSP has 680 ha and is surrounded by agricultural fields, pasture and secondary forest (Vicente 2006). The South limit of this Park is the Apertados Stream that in the periods of high precipitation, floods several sites in the riparian forest (Bianchini et al. 2001). At these sites (hereafter referred as flooded areas), some species were very abundant, such as *A. concolor* (Bianchini et al. 2003). The flooded areas in the MGSP were restricted and discontinuous. These areas showed small microtopographic variations and the vegetation cover was heterogeneous (Bianchini et al. 2001) due to the occurrence of small gaps.

The climate is Cfa according to Köppen’s classification system (Bianchini et al. 2003), with a yearly rainfall of about 1600 mm. The rainfall during the summer is three times higher than in the winter. The average annual temperature is around 21°C, and January (23.9°C) and June (16.3°C) are the hottest and the coldest month, respectively. The climate data for the period 1976-2004 were obtained from the Instituto Agronômico do Paraná. *Actinostemon concolor* is a small evergreen tree, reaching 10 m in height and 10 cm in diameter at breast height (dbh) (Smith *et al.*, 1988). It is a shade tolerant species and occurs specially in humid soils (Bianchini et al. 2003). In the Riparian Forest of Paraná State, Southern Brazil, *A. concolor* is among the principal species (Soares-Silva *et al.* 1992; Dias *et al.* 1998; Soares-Silva *et al.* 1998; Bianchini *et al.* 2003) and it can be among the species with great value of importance (IV) in the flooded areas (Silva *et al.* 1992; Nakagima *et al.* 1996; Bianchini *et al.* 2003). Flowering and fruiting occurs from August to September and August to November, respectively (Alberti and Morellato 2008). The fruits are capsules dehiscent and autochoric (Smith *et al.* 1988; Morellato and Leitão-Filho 1992). Fifty 10 m x 10 m plots (5000 m²) were established in the flooded sites of the Park. As flooded areas were discontinuous, these plots were divided into three groups (A, B, C), distant at least 20 m from each other. In May 2000, all *A. concolor* individuals were tagged, mapped to the nearest meter and their height and stem base diameter were measured. The population was grouped into three size categories: seedling – presence of cotyledon; juvenile – without cotyledon and height ≤ 1.50 m; adults – height > 1.5 m. The plots were monitored in January 2003 and January 2004. When individual height was reduced to less than 1.5 m because of break, it was considered that the individual returned from adult to juvenile.

To evaluate the height and diameter distribution in the population, the coefficient of skewness was used. The height and diameter growth rates were calculated using the first and last survey data.

Demographic data were analyzed using Lefkovitch stage-classified transition matrix (Caswell 2000). The matrix model is described by the equation $n_{t+1} = A \times n_t$, where $A$ is a square matrix containing transition probabilities among categories during one time-step and $n$ is a vector containing the individuals’ number in each category at time $t$ and
In this equation, the population structure at time $t + 1$ is obtained by multiplying the structure at time $t$ with the transition matrix, which contains information on the demography of the population, assuming environmental factors to be constant in time. Repeated multiplication of the transition matrix with the current population produces the dominant latent root or dominant eigenvalue of the transition matrix ($\lambda$), which is equal to the intrinsic population growth rate and the stable stage population structure (the right eigenvector $w$) (Caswell 2000). The elements of transition matrix (recruitment, survival and fecundity) were calculated directly from the counts of individuals that grew to the next category or stayed in the same category involved in each transition. Fecundity was expressed as the average number of seedlings produced in the study period per adult in the plots (Alvarez-Buylla 1994). The intrinsic population growth rate ($\lambda$) and stable stage distribution were calculated using the RAMAS Stage Software (Ferson 1990). The confidence interval for $\lambda$ was calculated using the bootstraps techniques following Caswell (2000). The sensitivity and elasticity analyses were carried out to determine the sensitivity of $\lambda$ to the changes in different elements of the transition matrix. Sensitivity analysis examines the effect on $\lambda$ of small changes of fixed size in the demographic parameters (Caswell 1978) while the elasticity analysis shows the proportional contribution of each matrix element to the $\lambda$ (De Kroon et al. 1986).

Maps were used to analyze the spatial distribution. They were made using the individuals’ Cartesian coordinates sampled in 2004. We used Ripley's L-function, which was based on the variance (second-order analysis) of all point-to-point (plant-to-plant) distances in a two-dimensional space (Ripley 1977) to confirm the spatial distribution patterns of species. When L value is zero, the points are randomly distributed; values less than zero indicate clumping while values greater than zero indicate regular spacing (Dale 1999). All the calculations (L-function, randomization, and edge effect) were performed in the Passage software (Rosenberg 2001).

To evaluate that environmental characteristic could explain the spatial distribution of the individuals, the canopy cover, litter and herbaceous covers in the soil and the plots height in relation to the level of the river in normal conditions (avoiding the dry and the rainy period) were measured. The canopy cover was assessed with a spherical densiometer (Lemmon 1956). In January 2003, four readings of canopy cover were made in the centre of each of the 10 m x 10 m plots and the average of the four readings was used as a measure of canopy cover in each plot (Bianchini et al. 2001). The plots were grouped into four categories of canopy cover: $>$90%, $>$85-90%, $>$80-85%, $\leq$80%. The litter and herbaceous covers were visually estimated in four categories: low, medium, high and very high. The plot height was determined in the centre of each plot using a transparent water hose and a ruler. The plots were grouped into five height categories: very low, low, medium, high and very high.

The correspondence analysis (CA) was applied to group the plots according to its environmental characteristics. The CA is a technique of multivariate analysis indicated for cases in which data are grouped in categories (Guedes et al. 1999). The CA allows getting a graphical representation through the distribution of the categories score of lines and/or columns in a system of coordinates. The eigenvalues are extracted from the tables of contingencies in which the original values are transformed in order to be interpreted as conditional probabilities. Because of the nature of this transformation, the relations between columns and lines of the transformed table are the same that those of the original matrix of data (Guedes et al. 1999). The CA was performed with PCord (McCune and Mefford 1999), and Statistica 6.0 (StatSoft 2004), softwares.

The CA groups the plots in a bidimensional graphical space, defined for the two most important eigenvectors in accordance with its environmental characteristics. In the figure produced by CA, the 15 plots with the highest (more than 50 individuals) and lowest (0 - 7 individuals) densities were highlighted.

**RESULTS**

The species presented high population density (above 2900 individuals ha$^{-1}$) with gradual increase in time (Table 1). There was high density of young for three surveys, corresponding to more than 60% of the population. The maximum and minimum number of individuals per plots, in each category, showed the heterogeneity among them.
The highest standard deviations were detected for the juvenile (Table 1). The population size varied with time and this numerical fluctuation must be mainly due to seedling and juvenile (Fig. 1).

**Table 1 -** Number of *Actinostemon concolor* individuals sampled in a flooded area (0.5 ha), Southern Brazil. SD - Standard deviation; CV – coefficient of variation.

|                      | May/2000 | January/2003 | January/2004 |
|----------------------|----------|--------------|--------------|
| Total number of seedling | 81       | 58           | 484          |
| Total number of juvenile | 946      | 1521         | 1414         |
| Total number of adults   | 446      | 423          | 418          |
| Population size             | 1473     | 2002         | 2316         |
| Maximum number of seedling plot$^1$ | 13       | 13           | 96           |
| Minimum number of seedling plot$^1$ | 0        | 0            | 0            |
| Mean (± SD) seedling plots$^1$ | 5.0 ± 3.9| 3.8 ± 3.6    | 12.8 ± 18.1  |
| CV                      | 4.9      | 6.9          | 3.8          |
| Maximum number of juvenile plot$^1$ | 104      | 133          | 125          |
| Minimum number of juvenile plot$^1$ | 0        | 0            | 0            |
| Mean (± SD) juvenile plots$^1$ | 24.9 ± 25.6| 36.9 ± 33.5  | 33.5 ± 31.2  |
| CV                      | 2.7      | 2.2          | 2.2          |
| Maximum number of adult plot$^1$ | 45       | 46           | 46           |
| Minimum number of adult plot$^1$ | 0        | 0            | 0            |
| Mean (± SD) adult plots$^1$ | 12.4 ± 11.0| 12.1 ± 10.8 | 11.9 ± 10.7  |
| CV                      | 2.5      | 2.6          | 2.6          |

**Figure 1** - Life table diagram for *Actinostemon concolor* population in a flooded area (0.5 ha), Southern Brazil. Data from two period of study: 2000-2003 (A) and 2003-2004 (B). I – number of new seedling recorded in the period; $N_0$ – number of individuals in the first survey; $N_1$ – number of survivors that stayed at the same class; M – number of deaths; Arrows to the right (solid) – number of recruitment to next class; Arrows to the left (broken) – number of regression to previous class. * Values resulting from the vital rates between the surveys.
A. concolor population showed a predominance of small individuals. The skewness coefficient ($g_1$) for the height and diameter were $g_1 = 2.3$ and $g_1 = 3.2$, respectively, confirming the distribution with a long tail to the right. The individuals grew slowly. The height and diameter mean growth rates were $0.7 \pm 8.0$ and $0.03 \pm 0.16 \text{ cm year}^{-1}$, respectively for all the population.

The population was characterized by very high survival rates and low recruitment rates between the categories. The largest proportional numerical loss in the population was in the seedlings, mainly in 2000/2003 (Fig. 1, Table 2). In this category, there was recruitment for the juvenile, especially in 2003/2004 (Fig. 1, Table 2).

The juvenile presented high survival probability in all the surveys (Table 2, Fig. 1). About 80% of the juvenile that died were 15 cm of height or minus. Although the recruitment of young for adults had occurred, the registered maximum value was only 1.3% (Fig. 1, Table 2).

Adults’ abundance reduction was observed throughout the study. The mortality was higher than the recruitment. But the main cause of numerical reduction was the return to juvenile (Fig. 1, Table 2), due to breaks in the shoots. The reproduction was registered in both time intervals: however, the fecundity was higher in 2003/2004 (Fig. 1, Table 2).

The intrinsic population growth rate ($\lambda$) was $0.921 \pm 0.026$ in 2000/2003 and $0.996 \pm 0.028$ in 2003/2004. The 2000/2003 $\lambda$ value indicated that the population of A. concolor would reduce in size. However, 2003/2004 $\lambda$ value indicated that the population size of A. concolor would remain stable.

The observed stage distribution was different from the stable stage distribution for both time periods considered (2000-2003: $\chi^2$-test = 12.1, $p = 0.002$; 2003-2004: $\chi^2$-test = 12.7, $p = 0.002$). If the observed transition rates were constant in time, the proportion of seedlings would have increased. The greater differences between the observed and stable stage distributions were recorded in adult’s category.

The intrinsic population growth rates were strongly influenced by the survival elements (principal diagonal) (Table 2). The highest elasticity was obtained for the adult survival (Fig. 2). Survival elasticity (0.97) was higher than the elasticity of recruitment (0.02) and fecundity (0.01), indicating the importance of individual survival, especially adults, for the self-maintenance of the population (Fig. 2).

The highest sensitivity was observed for the recruitment of juvenile to adult (mainly in 2003/2004), followed by the survival of adults (Table 3). Fecundity was less important, considering the impact of numerical changes on $\lambda$ (Table 3).

The population and all tree-classes size of A. concolor showed a patchy spatial pattern (Fig. 3) in all the distances considered (Fig 4). The seedling and juvenile mean density in the plots with the adults were $12.8 \pm 18.7$ and $37.4 \pm 31.8$, respectively. In the plots without adults, the numbers were $1.8 \pm 3.9$ seedlings and $4.4 \pm 8.0$ juvenile. These differences were significant (seedling: $t = 2.2$, df = 48, $P = 0.02$; juvenile: $t = 3.8$, df = 48, $P < 0.001$). The seedling and juvenile mortalities did not alter the patches pattern.
Figure 2 - Elasticity of $\lambda$ to changes in stasis (S) and growth (R - recruitment) for *Actinostemon concolor* population in a flooded area (0.5 ha), Southern Brazil.

### Table 3 - Sensitivity matrices for *Actinostemon concolor* population in a flooded area (0.5 ha), Southern Brazil.

|        | 2000 - 2003 |           | 2003 - 2004 |           |
|--------|-------------|-----------|-------------|-----------|
|        | seedling    | juvenile  | adult       | seedling  |
| Seedling| 0.005       | 0.000     | 0.035       | 0.008     |
| Juvenile| 0.017       | 0.117     | 0.119       | 0.011     |
| Adult   | 0.000       | 0.865     | 0.878       | 0.000     |

Figure 3 - Spatial distribution of mapped *Actinostemon concolor* individuals in a flooded area (0.5 ha), Southern Brazil. Data from 2004 survey.
Figure 4 – Spatial pattern of population (a), adults (b), juvenile (c) and seedling (d) of *Actinostemon concolor* in a flooded area (0.5 ha), Southern Brazil, contrasted to the null model of complete spatial randomness (value 0), using $L$-function of Ripley. The lateral lines to the null model (envelope) indicate the confidence interval.
The first two axes of CA explained 81.5% of the total data variance (Fig. 5). The height of plots \( r = 0.29 \), canopy cover \( r = -0.26 \) and litter covers \( r = -0.26 \) were the mainly variables associated to axis one. In relation to axis two, the variables were herbaceous cover \( r = 0.38 \) and plots height \( r = -0.20 \).

Despite not having the separation of groups, the plots of higher and lower densities of \( A. \) concolor had been situated mainly on the left and on the right of axis one, respectively (Fig. 5). The plots with higher densities showed minor height and covering of herbaceous, more closed canopy and dense litter cover. In contrast, the plots with lower densities showed major height and the soil covered by the herbaceous plants, more opened canopy and low covering of litter.

**DISCUSSION**

The high density and a juvenile-dominated size structure observed for \( A. \) concolor population were expected as for an understory species. The same trend was observed for the shade-tolerant tree species in temperate Japanese forest (Manabe et al. 2000). De Steven (1994) observed high density and predominance of seedling and sapling for two shade-tolerant canopy tree species in Panama. In Mallotus Lour. species, an understory trees of Bornean rain forests, there were approximately ten times more saplings than the adults (Primack and Lee 1991).

The \( A. \) concolor population presented a strong hierarchy among the individuals, with a small number of large individuals that contributed with the majority of biomass population. This size distribution was typical to shade-tolerant species (Wright et al. 2003). Size hierarchy was also observed for the populations of Mallotus species (Primack and Lee 1991), Trichilia tuberculata (Triana and Planch.) C. DC. and Tetragastris panamensis (Engl.) Kuntze (De Steven 1994).

The high density of seedlings and juveniles reflected the occurrence of constant and intense reproductive events (Ramirez and Arroyo 1990). This seems to be the case of \( A. \) concolor because the reproduction occurs every year and apomixy is common (Alberti and Morellato 2008). Mortality in \( A. \) concolor followed the pattern of tree populations, in which the highest mortality rates occurred among the smallest individuals (Alvarez-Buylla and Martínez-Ramos 1992; Gomes et al. 2003). Specific causes of mortality could not be determined for \( A. \) concolor, but litter fall was an important source of mortality because several dead individuals were buried under the litter. Martínez-Ramos et al. (1989) also related the litter fall as an important cause of death to Astrocaryum mexicanum Liebm. ex Mart., a shade-tolerant palm, in Mexican tropical forests. Although \( A. \) concolor tolerated flooding (Colli unpublished data), there was an increase in the seedling and
juvenile mortality after the overflow in January 2003. Sediment deposition at some sites, soil erosion at other sites and dragging of dead trunks and twigs by the water were the main causes of mortality in this period. These results were also observed for Chrysophyllum gonocarpum (Mart. and Eichler ex Miq.) Engl. in the same area (Bianchini unpublished data). Marques and Joly (2000) found that the burial of smaller individuals by the sediment was the main cause of mortality for Calophyllum brasiliense Cambess population in a swamp forest in the southeastern Brazil. Standing dead adult of A. concolor was not observed and the main cause of mortality of these individuals was branches and tree falls. Deaths provoked by litter fall were observed for Cecropia obtusifolia Bertol. (Alvarez-Buylla and Martínez-Ramos 1992) and Dipteryx panamensis (Pittier) Record and Mell (Clark and Clark 1987).

The individuals of A. concolor grew slowly in the understory. As the probability of death of small individuals is larger (De Steven 1994; Olmsted and Alvarez-Buylla 1995), the slow growth of this species contributes to an increase in the mortality among the seedlings and juveniles. Gomes et al. (2003) related that species with low growth rates could present high mortality.

The low fecundity values observed in this study could be related to the mortality of seedlings after the germination. Changes in the stasis rates of A. concolor among the years promoted by stochastic events such as flooding, could be related to the differences in the fecundity values mentioned. The slow growth of A. concolor was also related to the low adult recruitment and the accumulation of juveniles. According to Gomes et al. (2003), species with low growth rates showed low recruitment of individuals. Gomes et al. (2003) described higher values of $\lambda$ to understory species of mountain secondary forest than A. concolor. But A. concolor showed larger death probability for the adults than the species studied by Gomes et al. (2003). This result could explain the lower values of $\lambda$ in A. concolor because its values were very sensitive to the alterations in adults’ survival probabilities.

For A. concolor, the transition probabilities were not constant in time, indicating that the population size changed, mainly because of stochastic factors that acted on the population. The $\lambda$ elasticity and sensitivity of demographic process followed the pattern described in the literature (Barot et al. 2000; Svenning 2002; Guedje et al. 2003; Pulido et al. 2007; Kouassi et al. 2008). Thus, the model would provide higher values of $\lambda$ in the years when environmental conditions favored the survival of adults and/or the recruitment from juvenile to adults.

Considering the different vital rates between the periods, the high $\lambda$ sensitivity for the survival of adults and recruitment from the juvenile to adult, the unpredictability of the environment (flooding for example) and the position of plant species in several phytosociological studies carried out in the region (Silva et al. 1992; Soares-Silva et al. 1992; Nakagima et al. 1996; Dias et al. 1998; Soares-Silva et al. 1998; Bianchini et al. 2003), it could be concluded that the population size of A. concolor could vary over the time, but the species would remain as one of the most important in the community.

A patchy spatial pattern was observed for the A. concolor population in the MGEP’s flooded areas, similar to observed by Bianchini et al. (2010) in other forest fragment in the region. In the MGEP, plots lower in height (longer time of flooding) and with higher canopy cover (lower light intensity) were the main sites of species occurrence. This suggested the occurrence of favorable microenvironmental conditions to the regeneration of A. concolor, as suggested by Manabe et al. (2000) and Miyamoto et al. (2003) for the species studied by them. However, this interpretation required some care because this study was conducted at small scales.

The patchy spatial pattern could be also related to low dispersal seed (Law and Dieckmann 2000). As A. concolor is an autochoric species, the majority of their seeds fall under the parent plant. For the shade-tolerant species, adults crown provide a shaded place suitable for the species establishment (Richards 1997).

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