

Acta Scientiarum. Biological Sciences

ISSN: 1679-9283 eduem@uem.br

Universidade Estadual de Maringá

Brasil

Gomes da Silva, André Luiz; Ramos Chaves, Samyra; Mesquita Brito, Jéfferson Reproductive biology of Bowdichia virgilioides Kunth (Fabaceae) Acta Scientiarum. Biological Sciences, vol. 33, núm. 4, 2011, pp. 463-470 Universidade Estadual de Maringá .png, Brasil

Available in: http://www.redalyc.org/articulo.oa?id=187121352012



Complete issue

More information about this article

Journal's homepage in redalyc.org



Reproductive biology of Bowdichia virgilioides Kunth (Fabaceae)

André Luiz Gomes da Silva*, Samyra Ramos Chaves and Jéfferson Mesquita Brito

¹Centro de Ciências Agrárias e Ambientais, Universidade Federal do Maranhão, BR-222, km 74, 65500-000, Boa Vista, Chapadinha, Maranhão, Brazil.*Author for correspondence. E-mail: andrebotanico@gmail.com

ABSTRACT. The aim of this study was to investigate the reproductive biology of *Bowdichia virgilioides* in an area of Cerrado (Brazilian Savanna). The study was carried out in Chapadinha, State of Maranhão in northeastern Brazil. *Bowdichia virgilioides* has cornucopia flowering and annual pattern. Flowering occurred at the beginning of the dry season, between June and August; and fruiting in the middle of this season, between July and October. The anthesis of *B. virgilioides* is diurnal, and the main flower resource is the nectar. During the flower opening, the anthers becomes dehiscent and thus pollen grains are deposited at the stigma, promoting automatic self-pollination. The species is self-compatible and apomitic. In the study area, they were visited by nine species of bees, four butterflies, two wasps, and two hummingbirds. The visits began in early morning and persisted throughout the day, with a peak of activity between 8:00 and 9:00 hours. Species of *Centris* were the main visitors of *B. virgilioides*. The butterfly and wasp species were considered resource robbers, and other floral visitors were considered occasional visitors. Although the flowers of *B. virgilioides* are widely visited, the species shows a high rate of inbreeding because of automatic self-pollination.

Keywords: breeding system, Cerrado, pollination, reproductive success.

RESUMO. Biologia reprodutiva de Bowdichia virgilioides Kunth (Fabaceae). Este trabalho teve como objetivo estudar a biologia reprodutiva de Bowdichia virgilioides Kunth (Fabaceae) e avaliar o papel dos visitantes florais em seu sucesso reprodutivo. Bowdichia virgilioides apresenta padrão de floração anual e do tipo cornucópica. A floração ocorre no início da estação seca, entre os meses de junho e agosto e a frutificação no meio desta estação, entre os meses de julho e outubro. Suas flores são de antese diurna e apresentam néctar como recurso floral. Durante a abertura floral, as anteras se encontram deiscentes e por esta razão há a adesão dos grãos de pólen no estigma da própria flor, promovendo o processo de autopolinização passiva. A espécie é autocompatível e apomitica. As flores foram visitadas por nove espécies de abelhas, quatro espécies de borboletas, duas espécies de vespas, e por duas espécies de beija-flores. As visitas iniciam logo ao amanhecer e perduram por todo o dia, com um pico de atividades de todos os insetos entre 8 e 9h. Em virtude do seu comportamento intrafloral, por sua abundância e frequência, Centris spp. são os principais visitantes de B. virgilioides. As borboletas e as vespas foram consideradas pilhadoras e os outros visitantes florais foram considerados visitantes ocasionais. Embora as flores de B. virgilioides sejam intensamente visitadas, a espécie pode apresentar alta taxa de endogamia pela autopolinização passiva.

Palavras-chave: sistema de reprodução, Cerrado, polinização, sucesso reprodutivo.

Introduction

Bowdichia virgilioides has a wide geographical distribution, occurring naturally from the States of Rio de Janeiro to Roraima, mainly in central and northern Brazil (SILVA JÚNIOR; SANTOS, 2005). Commonly known as "sucupira", "sucupiraroxa" or "sucupira-preta", it is economically value because of its timber, which is used for various purposes; the medicinal properties of its stem bark; its landscaping potential; and its use in programs for management and restoration of degraded areas (LORENZI, 1992; SILVA JÚNIOR; SANTOS, 2005; SMIDERLE; SOUZA, 2003). For these

reasons, there is a need for conservation and promotion of the spread of this species. Studies on seed germination of *B. virgilioides* have been conducted for this purpose (ALBUQUERQUE; GUIMARÃES 2007; RODRIGUES; TOZZI, 2007; SILVA et al. 2001; SMIDERLE; SOUZA, 2003).

The reproductive phenology of *B. virgilioides* differ in different places where it occurs (FUNCH et al. 2002; LOCATELLI; MACHADO, 2004). Thes variations may be linked to climate and soil condition in different regions.

The reproductive biology of native species is an important parameter in maintaining the ecological balance of many biological systems, and leads to

various plant-animal interactions such as the pollination mechanism. *B. virgilioides* is adapted for bee pollination, because its nectariferous flowers show mellitophilic characteristics (SILVA JUNIOR; SANTOS, 2005). However; Rojas and Ribbon (1997) described a guild of nine species of birds, including six hummingbirds and three passeriform that visited its flowers, which indicates a variety of pollination mechanisms.

Many self-compatible species have special adaptations to prevent the automatic self-pollination, such as species herkogamous, protandrous and protogynous. However, in many other species not there is a morphological or temporal explicit barriers to prevent the transfer of pollen grains of the same flower, as observed in several species of Fabaceae (ARROYO, 1981; NOGUEIRA; ARRUDA, 2006) as well as in species of families not related (SILVA; PINHEIRO, 2009). This mechanism can favor the reproductive efficiency of a species in a fragmented environment where pollinator availability is limited (MOTTEN; ANTONOVICS, 1992).

The Self-compatibility and automatic selfpollination is a common phenomenon in many species of Fabaceae (BORGES, 2006; JACOBI et al., 2005; NOGUEIRA; ARRUDA, 2006; VIEIRA et al., 2002). Although flowers with a wide availability of resources are heavily visited, many self-compatible species have a high degree inbreeding caused by geitonogamous pollination, apomixis, and also automatic selfpollination (BAWA, 1979). These processes can drastically decrease the genetic variability of biological populations, but may also increase the chances of producing fruits and seeds, even with a low rate of (CHARLESWORTH; floral visits CHARLESWORTH, 1995). Self- versus crosspollination can occur concurrently in many plant species, and only specific studies for each group can infer any specific explanation for the prevalence of one or another mechanism (BAWA, 1979).

This study aimed to describe the phenological aspects, floral biology, floral visitors, reproductive system and reproductive success of *B. virgilioides* in an area of Savanna in the city of Chapadinha, Maranhão State, Brazil. In general, it is intended to answer the following question: Floral visitors are important for fruit set of *B. virgilioides*?

Material and methods

Study area

The city of Chapadinha is located in the microregion of Chapadinha (03° 44'17" S and 43° 20'29" W), also known as the microregion of "Alto

Munim" (IBAMA, 2006), in the southwestern part of the state of Maranhão, Brazil. The region is characterized by Cerrado *sensu lato*, formed by a mosaic of plant communities. *Bowdichia virgilioides* occurs in savanna, mainly in "campo sujo", grassland with scattered trees and shrubs.

The climate type of the region, according Köppen (1948) is wet tropical. Precipitation is high during the months of January to July and very low from August to November/December. The highest temperatures occur in the dry period.

Species studied

Bowdichia virgilioides Kunth (Fabaceae), Clade Genistoids (WOJCIECHOWSKI et al., 2004) is an arboreal species that is typical of the Cerrado sensu stricto, "Campo Sujo" (grassland with scattered trees and shrubs), and "Cerradão" (woodland with a welldeveloped tree canopy) (SILVA JUNIOR; SANTOS, 2005). However, it occurs predominantly in open areas such as "Campo Sujo", because the establishment and development of its seedlings is strongly influenced by shading, which is greater in areas of "Cerradão" (KANEGAE et al., 2000). According to LORENZI (1992), B. virgilioides occurs mainly in dry, nutrient-poor soils. However, Miranda et al., (2002) reported that this species is typical of sodium-rich soils in savannas in the state of Roraima, Brazil.

Flowering and fruiting phenology

Flowering and fruiting phenology were investigated from January 2007 until December 2008. Ten individuals previously selected, adults and healthy, were observed twice a month. During reproductive period, observations were weekly.

Floral biology

Floral biology of *B. virgilioides* was verified in natural populations. Buds and flowers were collected and analyzed in the laboratory. The stigma receptivity was tested although hydrogen peroxide (H₂O₂) 3% (KEARNS; INOUYE, 1993) in flowers in natural conditions. Anthesis was observed over a period of five days in five adults and healthy focal individuals, registering time of flower opening, flower longevity, and process of anther dehiscence.

Two samples of 15 flowers were collected in natural conditions to verify nectar production. These flowers were previously bagged one day prior to the anthesis. The first sample was collected at 8:00 hours and the second at 15:00 hours, on two consecutive days. Fifteen Flowers previously bagged were dissected to collect the nectar. Concentration

of the total volume of 15 flowers from each sample was analyzed with a portable refractometer with a scale from 0 to 32% Brix. The volume of nectar from each flower was measured with the aid of micropipettes of 0.2 mL.

Floral visitors

Floral visitors were observed, collected and identified during flowering period of five preindividuals, adults and Observations were from 6:00 to 17:00 hours, during ten consecutive days, totaling about 100 hours of observation. Floral visitors were classified as effective when the behavior of visitors promotes the contact of anthers with stigma and frequency of visits is high, more than 10 visits per day, occasional when the behavior of visitors promotes the contact of anthers with stigma and frequency of visits is low, less than 10 visits per day or floral resources robbers when visitors do not touch the stigma nor the anthers.

Breeding system

The following tests were carried out to evaluate the breeding system: 1-Manual self-pollination (autogamy); 2-Cross-pollination (xenogamy). These tests were conducted with previously bagged flowers one day prior to the anthesis; 3-Control - Unbagged flowers; 4-Automatic self-pollination - calculated from the proportion of fruits formed from bagged buds.

The pollen load on the stigmas of open (naturally pollinated) and bagged (automatic self-pollinated) flowers was determined by counting the pollen grains on each stigma, using an optical microscope. The styles of some buds were removed to test for the occurrence of autonomous apomixis (RICHARDS, 1986).

Significance difference between the percentages obtained from xenogamy, autogamy and the control experiments were evaluated by comparison formula between two percentages, with a 5% significance difference level (PAGANO; GAUVREAU, 2004). The obtained values of "t" were then compared with tabulated theoretical "t" values.

Reproductive success

To estimate the fecundity rate, we followed Cruden (1972) procedure, which is the product of two ratios (seed/ovule and fruit/flower).

Results and discussion

Flowering phenology

Bowdichia virgilioides has cornucopia flowering and a annual pattern, according to the classification proposed by Gentry (1974) and Newstron et al.

(1994), at the population level. Although, individual plants exhibit a supra-annual pattern. In 2007 flowering occurred at the beginning of the driseason, between June and August; and fruiting in the middle of this season, between July and October. In 2008, only three individuals of the population flowered between June and August, and none of the individuals that flowered in 2008 flowered in 2008. These observations agree with those of Bulhão and Figueiredo (2002) and Figueiredo (2008), studying a natural plan community in the Cerrado of Maranhão State According to these same authors, this pattern of flowering early in the dry season is common in many species of the Cerrado.

The abscission of the leaves occurs before the beginning of flowering and increases the display of the flowers. Flowering in the dry season may be characteristic of low latitudes, because in a Cerradoregion in the state of São Paulo, the tree specie flower mainly in the beginning of the rainy season (BATALHA; MANTOVANI, 2000).

The flowering of *B. virgilioides* can occur a different times in different localities, but alway in the dry periods of the year. Funch et al. (2002 observed flowering between June and October in the "Chapada Diamantina", Bahia State and Locatelli and Machado (2004) observed flowering between November and January in a montant forest in Pernambuco State, Brazil. These differences may be related to climate and soit conditions in each area.

Floral biology

Bowdichia virgilioides has a paniculate inflorescence with 23.7 flowers (\pm 18.3, N = 18) The large number of flowers per inflorescence is common feature among the species of Fabaceae (KIILL; DRUMOND, 2001). Flowers are hermaphroditic, with 1.75 cm length (± 0.17, N = 20), pedicel short, calyx campanulate with five ciliate teeth, 0.5 cm in diameter (\pm 0.05, N = 20 and 0.8 cm in depth (\pm 0.1, N = 21). The inne part of the calyx forms a nectariferous chamber The color of the petals varies along the anthesis purple at the beginning and purplish toward the end. This feature showed no relationship with pollination mechanism. The flowers ar papilionacea type with vexillus subunguiculate transverse-oblong, deeply emarginate, unguiculate wings, obovate, keels free and elliptical, a

suggested by Silva Júnior and Santos (2005).

The ovary is unicarpellate, unilocular, with 8.2 ovules (± 1.32, N = 35), the stigma is wet and papillose and remains receptive during the first day

of anthesis. The androecium is formed by ten stamens with orbicular anthers with rimose dehiscence.

The floral resource is the nectar that is produced inside the nectariferous chamber, in the base of the floral receptacle around the ovary. The flowers produce nectar only the first day of anthesis. In two samples, the first at 8:00 hours and the second at 15:00 hours, the volume of nectar per flower was about 0.1 mL and the concentration of nectar was 13% (Brix scale). Nectar with a low concentration of sugar cannot satiate the floral visitor on the first visit, and only prolongs their foraging to collect more food. In many species of Fabaceae, the concentration of nectar does not exceed 28% sugar (ETCHEVERRY; ALEMÁN, 2005; KUDO; HARDER, 2005; MENDONÇA; ANJOS, 2006).

The anthesis was diurnal and began soon after sunrise. The flowers remained open only on the first day of anthesis, on the second day the petals wilted, became whitish, and the anthers had no more pollen grains. At the moment of floral opening, only part of the style and stigma receptive were exposed (Figure 1A-1); during the day, the petals reflexed and exposed the anthers (Figure 1A-2). This mechanism can act as herkogamic, temporally separating the receptive female and male structures (FAEGRI; VAN DER PIJL, 1979). However, before opening the anthers are already dehiscent, and therefore the pollen grains adhere on the stigma, causing automatic self-pollination.

The self-pollination mechanism is common in other species of Fabaceae (KILL; DRUMOND, 2001; NOGUEIRA; ARRUDA, 2006; BORGES, 2006) and may to favor a significant increase in the rate of inbreeding in self-compatible species (BAWA; WEBB, 1984). The automatic self-pollination can hinder cross-pollination pollen tube growth, owing to competition with the autogamic pollen tubes, thus reducing xenogamy rate (MAHORO, 2003; SILVA; PINHEIRO, 2009).

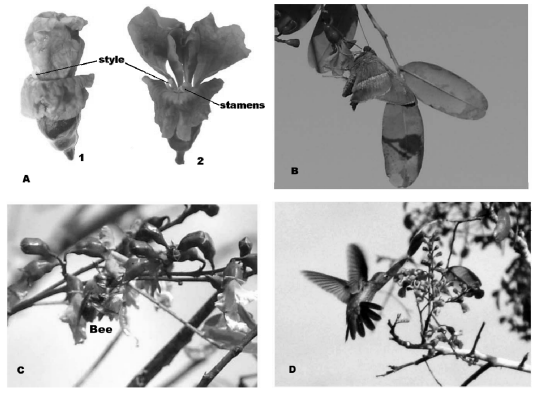


Figure 1. A- Flower of Bowdichia virgilioides Kunth (Fabaceae): A1- Flower at the beginning of anthesis, showing part of estyle; A2- Flower with stamen and style exposed; B- Visit of Lepidoptera; C- Visit of Centris sp.; D- Visit of Chlorostilbon mellisugus.

Floral visitors

Flowers of *B. virgilioides* were visited by nine species of bees (Apidae), four species of butterflies (Hesperidae); two species of wasps (one of Braconidae and the other Vespidae) and two species of hummingbirds (Trochilidae) (Table 1).

Table 1. Floral visitors of *Bowdichia virgilioides* Kunth (Fabaceae) in an area of Cerrado in Chapadinha, Maranhão, Brazil: mymain visitors; ov- occasional visitors, ro- robber.

Order/Family	Species	Pollination efficiency	
Himenoptera			
Amidaa	Centris (Centris) aenea	mv	
Apidae	Lepeletier		
	Centris (Trachina) grupo		
	fuscata Lepeletier	mv	
	Xylocopa sp.	ov	
	Apis mellifera L	ov	
	Trigona sp. 1	ov	
	Trigona sp. 2	ov	
	Partamona sp.	ov	
	Geotrigona sp.	ov	
Vespedae	sp. 1	ro	
Braconidae	sp. 2	ro	
Lepidoptera			
Hesperidae	Urbanus dorantes dorantes	ro	
	Chioides catillus catillus	ro	
	sp. 1	ro	
	sp. 2	ro	
Trochiliformes			
Trochilidae	Chlorostilbon mellisugus	ov	
	Amazilia fimbriata	ov	

The bees visited the flowers throughout the day, especially from 7:30 to 8:30 hours. The bees foraged more intensely intra-plant because of the abundance of flowers on one individual, which guarantees an unlimited supply of resources throughout the day. The species of Apidae were the most frequent and abundant visitors, and of these, *Centris (Centris) aenea* Lepeletier, 1841 and *Centris (Trachina)* group *fuscata* Lepeletier, 1841 were the most common species. Species of Apidae are the main pollinators of other species of Fabaceae (VIEIRA et al., 2002; GUEDES et al., 2009), and bees are the main pollinators in the Cerrado biome (MARTINS; BATALHA, 2007).

During foraging, each bee remained around 10 seconds on each flower. They were positioned in the flower, seizing the wings and keels, and introduced the proboscis inside the nectariferous chamber to collect nectar (Figure 1C). All visiting bees showed the same intra-floral behavior. Large amounts of pollen grains were found, mainly in Corbicula and on the feet of these bees. Because of their intra-floral behavior, its abundance and frequency, species of *Centris* are the main visitors of *B. virgilioides*.

Apis mellifera showed low frequency of visits to flowers of *B. virgilioides*; however, Carvalho and Marchini (1999) reported that this species is frequent in the flowers of *B. virgilioides* in the municipality of

Castro Alves, Bahia State, Brazil. Although species of *Xylocopa* are considered efficient pollinators in species of Fabaceae (KIILL; DRUMOND, 2001; BORGE, 2006; GUEDES et al., 2009), they were infrequent in this study. Two species of *Trigona* observed visiting the flowers of *B. virgilioides* showed intra-floral behavio similar to other bees, although they were also uncommon, as were *Geotrigona* sp. and *Partamona* sp. Because of the low frequencies of these species of bees they were considered occasional visitors.

The butterflies visited the flowers intensely throughout the day, and because of the wide availability of flowers in one individual, the exhibited a foraging behavior like bees, visiting the flowers of the same plant for long periods. The land on the flower and introduce their probosci into the nectariferous chamber to collect the nectar which according to Proctor and Yeo (1972) is the principal food of Lepidoptera adults. During the visit, only the proboscis touches the flora reproductive structures (Figura 1B), and few or no pollen grains from this structure were observed in specimens collected. For this reason, the butterflie were considered only as robbers of nectar. These insects also act only as nectar robbers from othe species of Fabaceae (KIILL; DRUMOND, 2001 NOGUEIRA; ARRUDA, 2006), and also from species of other families (FAEGRI; VAN DEF PIJL, 1979; MARTINS; BATALHA, 2007) According to Arroyo (1981), the morphology o flowers of Fabaceae does not correspond to the Lepidoptera pollination syndrome.

Wasps were rarely observed, and it was not possible to determine whether their visits were to seek for pollen grains or nectar. Because of the low frequency of visits and because they did not enter the full flower reproductive structures, these insects were considered as robbers of floral resource. In *Centrosema pubescen* Benth. (Fabaceae), wasps can act as pollinators as much as pillagers (BORGES, 2006), which shows the great diversity of pollination systems among species of Fabaceae (ARROYO, 1981).

Hummingbirds were also rarely observed visiting flowers of *B. virgilioides*. On each visit to a plant, a bird visited up to five flowers and left soon after The purpose of the intra-floral visits is to search for nectar. Because the flower tube is short, only part of the beak of this animal touches the flower (Figure 1D). These animals were observed more frequently around 6:30 and 8:30 hours. Because of their low frequency, these birds were considered occasional visitors. However, Rojas and Ribbon (1997) observed that in a Cerrado remnant in Minas Gerais State, Brazil, *B. virgilioides* was heavily visited by nine species of birds, including three species of

passariforms and six species of hummingbirds (Trochilidae), which visited the flowers to look for nectar and small insects. Hummingbirds are considered effective pollinators of many species of Fabaceae (GALETTO et al., 2000; ETCHEVERRY; ALEMÁN, 2005).

Breeding system

Bowdichia virgilioides is self-compatible, because it showed a rate of fruit production of 36% (N = 25) in an experiment of manual self-pollination (Table 2). The xenogamy test showed an index of 36% (N = 25), and the fruit/flower ratio under natural conditions was 13.2% (N = 114). Comparing the results of the experiments of autogamy and xenogamy, which were identical to each other, to the result for the control, found that there was a significant difference between them $(t_{obs} = 32.85 <$ $t_{teor} = 2.06$). These data indicate that there is a significant increase in the production of fruit when there is deposition of grains in the stigma, which may represent inefficiency in the mechanism of natural pollination. The low production of fruit in natural conditions is a common phenomenon in many species of Fabaceae, and may be related to several factors, such as the low rate of pollination, herbivores, and the effect of climate, among other factors (GALETTO et al., 2000; KIILL; DRUMOND, 2001; ETCHEVERRY; ALEMÁN, 2005). However, mass flowering of B. virgilioides can counterbalance the low fruit/flower ratio.

Flowers previously bagged in bud had approximately 29 pollen grains on the stigma (N = 25), whereas unbagged control flowers had 36 (N = 25); these means are not significantly different $(t_{obs} = 1.04 < t_{teor} = 2.06)$. Natural biotic pollination did not significantly increase the amount of pollen grains on the stigma, and therefore automatic selfpollination is possible in this species. The fruit set in an experiment of automatic self-pollination was 20.4% successful (N = 201). This index, although higher than the fruit set in natural conditions, was not significantly different from it ($t_{obs} = 1.75 < t_{tcor}$ = 2.06). Automatic self-pollination is also common in other species of Fabaceae (BORGES, 2006; NOGUEIRA; ARRUDA, 2006), as well as in species of unrelated families, and can promote fruit set if there is a decrease in rates of natural pollination (BAWA; WEBB, 1984; CHARLESWORTH; CHARLESWORTH, 1995, FAEGRI; VAN DER PIJL, 1979; SILVA; PINHEIRO, 2009).

Self-compatibility may be responsible for a higher production of fruit in natural conditions caused by the process of automatic self-pollination in *B. virgilioides*. The reproductive system is

diversified in Fabaceae, which includes both self-compatible (ARROYO, 1981; VIEIRA et al., 2002; KUDO; HARDER, 2005; NOGUEIRA; ARRUDA, 2006; BORGES, 2006) and self-incompatible species (ETCHEVERRY; ALEMÁN, 2005; GALETTO et al., 2000; GUEDES et al., 2009; KIILL; DRUMOND, 2001).

The rate of apomixis was 8% (N = 25), significantly different from that found in natural pollination, 13.2% ($t_{\rm obs} = 10.0 < t_{\rm tcor} = 2.06$). Self-compatible species of Fabaceae usually have some incidence of apomixis (BORGES, 2006; NOGUEIRA; ARRUDA, 2006), whereas in self-incompatible species this phenomenon does not occur (ETCHEVERRY; ALEMÁN, 2005; GALETTO et al., 2000; KIILL; DRUMOND, 2001). As with automatic self-pollination, apomixis can also promote fruit set when there is a low rate of natural pollination (BAWA; WEBB, 1984; FAEGRI; VAN DER PIJL, 1979).

Table 2. Breeding system of *Bowdichia virgilioides* Kunth (Fabaceae) in an area of Cerrado in Chapadinha, Maranhão State, Brazil.

	Flowers	Fruits
Control	114	15 (13%)
Xenogamy	25	9 (36%)
Self-pollination	25	9 (36%)
Automatic self- pollination	201	41 (20.4%)
Apomixis	25	2 (8%)

Reproductive success

In natural conditions the number of seeds per fruit was 3.5 (\pm 1.63, N = 26), the seed/ovule was 0.43, and the resulting fecundity rate (CRUDEN, 1972) was 0.056 (Table 3), a value considered low for self-compatible species (WIENS et al., 1987). In an automatic self-pollination experiment with bagged flowers in bud, the number of seeds was 3.9 (\pm 1.96; N = 25), the seed/ovule was 0.47, and the resulting fecundity rate was 0.096 (Table 3). These results show that *B. virgilioides* is primarily autogamous, which is a common feature among many species of Fabaceae (BORGES, 2006; NOGUEIRA; ARRUDA, 2006).

Table 3. Reproductive success of *Bowdichia virgilioides* Kunth (Fabaceae) in an area of Cerrado in Chapadinha, Maranhão State, Brazil, in natural conditions.

	Control	Bagged flowers (automatic self-pollination)
Ovule/flower	$8.2 \pm 1.32 \mathrm{N} = 25$	
Pollen grains/stigma	36.0 N = 25	28.4 N = 25
Seed/fruit	$3.5 \pm 1.63 \text{ N} = 26$	$3.9 \pm 1.96 \mathrm{N} = 25$
Seed/ovule	0.43	0.47
Fruit/Flower	0.13 N = 114	0.2 N = 201
Fecundity rate	0.056	0.096

Conclusion

Although the flowers of B. virgilioides are visited by a variety of animals with a great potential to pollinate, the dynamics of flower's opening allows the transfer of pollen grains to the stigma through of the automatic self-pollination. This process promotes the fruit set, since this species is self compatibility. Natural biotic pollination did not significantly increase the amount of pollen grains on the stigma in relation to the amount of pollen deposited by automatic self-pollination. All these data show that B. virgilioides is independent of pollinators for fruit set. However, only studies on the genetic diversity in natural populations will aid in understanding the balance between self- and cross-pollination and its consequent genetic structure.

Acknowledgements

We wish to thank FAPEMA for financial support, and we particularly thank Prof. Fernando Silveira (UFMG) for the identification of Hymenoptera species, Prof. Dr. Olaf Mielke (UFPR) for the identification of Lepidoptera species, and Prof. Dr. Carlos Martinez (UFMA) for identifying the hummingbirds. Dr. Janet W. Reid revised the English text.

References

ALBUQUERQUE, K. S.; GUIMARÃES, R. M. Comportamento fisiológico de sementes de *Bowdichia virgilioides* Kunth. sob diferentes temperaturas e condições de luz. **Cerne**, v. 13, n. 1, p. 64-70, 2007.

ARROYO, M. T. K. Breeding systems and pollination biology in Leguminosae. In: POLHILL, R. M.; RAVEN, P. H. (Ed.). **Advances in Legume Systematic, part 2**. London: Royal Botanic Gardens Kew, 1981. p. 723-769.

BATALHA, M. A.; MANTOVANI, W. Reproductive phenological patterns of cerrado plant species at the Pé-de Gigante Reserve (Santa Rita do Passa Quatro, São Paulo, Brazil): a comparation between the herbaceus and woody floras. **Revista Brasileira de Biologia**, v. 60, n. 1, p. 129-145, 2000.

BAWA, K. S. Breeding systems of trees in a tropical wet forest. **New Zealand Journal of Botany**, v. 17, p. 521-524, 1979.

BAWA, K. S.; WEBB, C. J. Flower, fruit and seed abortion in tropical forest trees: implications for the evolution of paternal and maternal reproductive patterns. **American Journal of Botany**, v. 71, n. 5, p. 36-751, 1984.

BORGES, H. B. N. Biologia reprodutiva de *Centrosema pubescens* Benth. (Fabaceae). **Boletim do Museu Paraense Emilio Goeldi**, v. 1, n. 1, p. 31-38, 2006.

BULHÃO, C. F.; FIGUEIREDO, P. S. Fenologia de leguminosas arbóreas em uma área de Cerrado marginal

no nordeste do Maranhão. **Revista Brasileira de Botânica**, v. 25, n. 3, p. 361-369, 2002.

CARVALHO, C. A. L.; MARCHINI, L. C. Planta visitadas por *Apis mellifera* L. no vale do rio Paraguaçu Município de Castro Alves, Bahia. **Revista Brasileira de Botânica**, v. 22, n. 2, p. 333-338, 1999.

CHARLESWORTH, D.; CHARLESWORTH, E Transposable elements in inbreeding and outbreeding populations. **Genetics**, v. 140, n. 1, p. 415-417, 1995.

CRUDEN, R. W. Pollinators in high elevation ecosystems: relative effectiveness of birds and bees **Science**, v. 176, p. 1439-1440, 1972.

ETCHEVERRY, A. V.; ALEMÁN, C. E. T. Reproductive biology of *Erythrina falcata* (Fabaceae-Papilionoideae) **Biotropica**, v. 37, n. 1, p. 54-63, 2005.

FAEGRI, K.; VAN DER PIJL, L. **The principles o pollination ecology**. 3rd ed. Oxford: Pergamon Press 1979.

FIGUEIREDO, P. S. Fenologia e estratégias reprodutiva das espécies arbóreas em uma área marginal de Cerrado na transição para o semi-árido no nordeste do Maranhão Brasil. **Revista Trópica - Ciências Agrárias Biológicas**, v. 2, n. 2, p. 8-21, 2008.

FUNCH, L. S.; FUNCH, R.; BARROSO, G. M. Phenology of Gallery and Montane Forest in the Chapad Diamantina, Bahia, Brazil. **Biotropica**, v. 34, n. 1 p. 40-50, 2002.

GALETTO, L.; BERNADELLO, I. C.; ISELE, J. VESPRINI, G.; SPERONI, G.; BERDUC, A Reproductive biology of *Erythrina crista-galli* (Fabaceae) **Annals of the Missouri Botanical Garden**, v. 87, n. 2 p. 127-145, 2000.

GENTRY, A. H. Flowering phenology and diversity in tropical Bignoniaceae. **Biotropica**, v. 6, n. 1, p. 64-68 1974.

GUEDES, R. S.; QUIRINO, Z. G. M.; GONÇALVES E. P. Fenologia reprodutiva e biologia da polinização d *Canavalia brasiliensis* Mart. ex Benth (Fabaceae) **Biotemas**, v. 22, n. 1, p. 27-37, 2009.

IBAMA-Instituto Brasileiro do Meio ambiente. **Laude** sócio-econômico e biológico para criação de Reserva de Chapada Limpa. Maranhão: Centre Nacional de Populações Tradicionais-CNPT/MA, 2006.

JACOBI, C. M.; RAMALHO, M.; SILVA, M. Pollination biology of the exotic rattleweed *Crotalaria retusa* I (Fabaceae) in NE Brazil. **Biotropica**, v. 37, n. 3 p. 357-363, 2005.

KANEGAE, M. F.; BRAZ, V. S.; FRANCO, A. C. Efeito da seca sazonal e disponibilidade de luz na sobrevivência crescimento de *Bowdichia virgilioides* em dua fitofisionomias típicas dos Cerrados do Brasil. **Revist: Brasileira de Botânica**, v. 23, n. 4, p. 459-468, 2000.

KEARNS, C. A.; INOUYE, D. W. **Techniques fo Pollination Biologists**. Colorado: University Press
1993

KIILL, L. H. P.; DRUMOND, M. A. Biologia floral e sistema reprodutivo de *Gliricidia sepium* (Jacq.) Steud (Fabaceae - Papilionoidae) na região de Petrolina

Pernambuco. Ciência Rural, v. 31, n. 4, p. 597-601, 2001

KÖPPEN, W. **Climatologia**: con un estudio de los climas de la tierra. México: Fondo de Cultura Econômica, 1948.

KUDO, G.; HARDER, L. D. Floral and inflorescence effects on variation in pollen removal and seed production among six legume specie. **Functional Ecology**, v. 19, n. 2, p. 245-254, 2005.

LOCATELLI, E.; MACHADO, I. C. Fenologia das espécies arbóreas de uma mata serrana (brejo de altitude) em Pernambuco, Brasil. In: PORTO, K. C.; CABRAL, J. P.; TABARELLI, M. (Org.). **Brejos de altitude em Pernambuco e Paraíba**. História natural, ecologia e conservação. Brasília: Ministério do meio Ambiente, 2004. p. 255-276.

LORENZI, H. **Árvores brasileiras**: manual de identificação e cultivo de plantas arbóreas nativas do Brasil. Nova Odessa: Plantarum, 1992.

MAHORO, S. Effects of flower and seed predators and pollinators on fruit production in two sequentially flowering congeners. **Plant Ecology**, v. 166, n. 1, p. 37-48, 2003.

MARTINS, F. Q.; BATALHA, M. A. Pollination systems and floral traits in Cerrado woody species of the upper taquari region (Central Brazil). **Braziliam Journal of Biology**, v. 66, n. 2, p. 543-552, 2007.

MENDONÇA, L. B.; ANJOS, L. Feeding behavior of hummingbirds and perching birds on *Erythrina speciosa* Andrews (Fabaceae) flowers in an urban area, Londrina, Paraná, Brasil. **Revista Brasileira de Zoologia**, v. 23, n. 1, p. 43-49, 2006.

MIRANDA, I. S.; ABSY, M. L.; REBÊLO, G. H. Community Structure of Woody Plants of Roraima Savannahs, Brazil. **Plant Ecology**, v. 164, p. 109-123, 2002.

MOTTEN, A. F.; ANTONOVICS, J. Determinants of outcrossing rate in a predominantly self-fertilizing weed, Datura stramonium (Solanaceae). **American Journal of Botany**, v. 79, n. 4, p. 419-427, 1992.

NEWSTRON, L. G.; FRANKIE, G. W.; BAKER, H. G. A new classification for plant phenology based on flowering patterns in lowland tropical rain forest at La Selva, Costa Rica. **Biotropica**, v. 26, n. 2, p. 141-159, 1994.

NOGUEIRA, E. M. L.; ARRUDA, V. L. V. Fenologia reprodutiva, polinização e sistema reprodutivo de *Sophora tomentosa* L. (Leguminosae – Papilionoideae) em restinga da praia da Joaquina, Florianópolis, sul do Brasil. **Biotemas**, v. 19, n. 2, p. 29-36, 2006.

PAGANO, M.; GAUVREAU, K. **Princípios de Bioestatística**. São Paulo: Pioneira Thomson Learning, 2004.

PROCTOR, M.; YEO, P. **The pollination of flowers**. New York: Taplinger Publishing Company, 1972.

RICHARDS, A. J. **Plant Breeding Systems.** London: Publishers George Allen and Unwin Ltda., 1986.

RODRIGUES, R. S.; TOZZI, A. M. G. A. Morfologia de plântulas de cinco leguminosas genistóides arbóreas do Brasil (Leguminosae - Papilionoideae). **Acta Botânica Brasilica**, v. 21, n. 3, p. 599-607, 2007.

ROJAS, R.; RIBON, R. Guilda de aves em *Bowdichia virgilioides* (Fabaceae – Faboidae) em área de Cerrado de Furnas- Minas Gerais. **Ararajuba**, v. 5, n. 2, p. 189-194, 1997.

SILVA, A. L. G.; PINHEIRO, M. C. B. Reproductive success of four species of *Eugenia* L. (Myrtaceae). **Acta Botanica Brasilica**, v. 23, n. 2, p. 526-534, 2009.

SILVA, L. M. M.; AGUIAR, I. B.; TERESINHA. J. D.; RODRIGUES, T. J. D. Seed germination of *Bowdichia virgilioides* Kunth, under water stress. **Revista Brasileira de Engenharia Agrícola e Ambiental**, v. 5, n. 1, p. 115-118, 2001.

SILVA JÚNIOR, M. C.; SANTOS, G. C. **100 Árvores do Cerrado**: guia de campo. Brasília: Ed. Rede de Sementes do Cerrado, 2005.

SMIDERLE, O. J.; SOUSA, R. C. P. Dormência em sementes de Paricarana (*Bowdichia virgilioides* Kunth-Fabaceae-Papilionidae). **Revista Brasileira de Sementes**, v. 25, n. 2, p. 48–52, 2003.

VIEIRA, R. E.; KOTAKA, C. S.; MITSUI, M. H.; TANIGUCHI, V. A. A. T.; RUVOLOTAKASUSUKI, M. C. C.; TERADA, Y.; SOFIA, S. H.; COSTA, F. M. Biologia floral e polinização por abelhas em siratro (*Macroptilium tropurpureum* Urb.). Acta Scientiarum. Animal Sciences v. 24, n. 4, p. 857-861, 2002.

WIENS, D.; CALVIN, C. L.; WILSON, C. A.; DAVERN, C. I.; FRANK, D.; SEAVEY, S. R. Reproductive success, spontaneous embryo abortion, and genetic load in flowering plants. **Oecologia**, v. 71, n. 4, p. 501-509, 1987.

WOJCIECHOWSKI, M. F.; LAVIN, M. F.; SANDERSON, M. J. A phylogeny of legumes (Leguminosae) based on analysis of the plastid *matk* gene resolves many well-supported subclades within the family. **American Journal of Botany**, v. 91, n. 11, p. 1846-1862, 2004.

Received on December 8, 2009. Accepted on April 24, 2010.

License information: This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.