Fly pollination and pollinator sharing in two synchronopatric species: *Cordia multispicata* (Boraginaceae) and *Borreria alata* (Rubiaceae)

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(Received: August 11, 1999; Accepted: April 17, 2000)

**Abstract** — The floral biology of *Cordia multispicata* (Boraginaceae) and *Borreria alata* (Rubiaceae) was studied in natural populations in a fragment of the Atlantic forest in Pernambuco, northeastern Brazil. Both species flower during almost the whole year. *Cordia multispicata* is a shrubby species with white, distylous and tubular flowers. *Borreria alata* is a herbaceous species. Its flowers are whitish, tubular and have a polymorphism in relation to the size of the style. Floral anthesis in both species begins at 6:00 a.m. Sugar concentration in the nectar was about 16% in *C. multispicata* and 30% in *B. alata*. Nine species of flies, mainly of the genus *Palpada* (Syrphidae), were observed visiting flowers of the two species. Seven of them were observed visiting and pollinating flowers of both *C. multispicata* and *B. alata*. Two species visited only flowers of *C. multispicata*, whereas no fly was exclusive to *B. alata* flowers. Both species have similar flower morphology, flowering time, habitats in the forest and establish populations very close to each other. These facts can favour the pollinators’ sharing and increase pollinator attraction.

**Introduction**

Many dipterans are flower visitors and show a great variety of modes of pollination, which characterize the syndromes of myophily and sapromyophily (Faegri & van der Pijl 1979). The family Syrphidae is considered the most important family of Diptera, regarding pollination (Endress 1994, Proctor et al. 1996), but there are still few studies concerning the effective role of these flies as pollinators, especially in tropical regions (Arruda & Sazima 1996).

The genus *Cordia* (Boraginaceae) has species adapted to different pollinators. Bees and butterflies are their main visitors, although there are records of pollination by hummingbirds, beetles and flies (Opler et al. 1975). *Borreria* (Rubiaceae) is apparently an entomophilous genus, however there is almost no information in the literature about the pollination biology of its species.

*Cordia multispicata* Cham. and *Borreria alata* DC. have similarities in the size, form and colour of their flowers. In some cases, the similarity of floral characteristics in species of different genera and families may be explained as a convergence towards utilization of the same type of pollinator (Brown & Kodric-Brown 1979).

In this study we present comparative data about floral biology and pollination by flies in *Cordia multispicata* and *Borreria alata* discussing aspects related to floral features, breeding system and polli-
nator sharing between these two synchronopatric species.

**Material and methods**

Field observations were carried out in the Forest Reserve of Dois Irmãos (8°7'30"S and 34°52'30"W), in Recife – Pernambuco State, northeastern Brazil. It’s one of the few remnants of Atlantic forest in the State. Observations went from August 1992 to July 1993, with additional observations in November and December 1997. The annual rainfall at Dois Irmãos is more than 2000 mm and mean annual temperature is 25 °C (20-30 °C). Both species were studied in natural populations distributed at the edges of the forest.

The morphology and size of the flowers, time, sequence and duration of anthesis, colour, emission of odour, sugar concentration in the nectar, as well as time of visits, frequency, and behaviour of the visitors were registered. Measurements of floral parts were based on 10 flowers of *B. alata* and 20 of *C. multispicata* (10 short- and 10 long-styled). Sugar concentration in nectar from both species was determined from flowers previously isolated by paper bags on the day prior to anthesis. The sugar concentration was recorded for 20 flowers of *C. multispicata* and 39 of *B. alata* with a compensated temperature refractometer (Atago). Pollen viability was assessed using acetocarmine (Radford et al. 1974) on 10 flowers of *Borreria alata* and 20 of flowers of *Cordia multispicata* (10 of each morph), from 10 individuals from each species. In order to examine the reproductive system of *C. multispicata*, controlled pollinations were performed in the field, according to the methods described by Radford et al. (1974). Treatments included tests for manual self- and cross-pollination. The crosses included flowers of both morphs. Buds were tagged one day before anthesis and the inflorescences were bagged with semi-permeable paper. Treatments were performed at anthesis. To estimate fruit production under natural conditions (control) some, inflorescences were tagged and observed for fruit development. Vouchers of both species are in the herbarium of the Federal University of Pernambuco (UFP 8064 and 8065).

Flowers were observed for visitors between 5:00 a.m. to about 4:00 p.m., during 23 days, totalizing 96 hours of field observations. The behaviour of the visitors was analysed visually and through photographic records. Frequency of the visitors to flowers of both species was registered over three days during the flowering peak, in five plants of each species. Some insects were captured, sent to specialists for identification and stored in the collection of the Laboratory of Floral and Reproductive Biology - Department of Botany, Federal University of Pernambuco.

**Results**

*Cordia multispicata* is a shrub with flowers disposed in dense inflorescences, usually positioned outside the leaves. Inflorescence position varies from erect to hanging, depending on the size of the peduncle, which varies from 2 to 10 cm. The spikes are 2-9 cm long, with a variable number of flowers (\( \bar{X} = 62 \)). The inflorescences occur from 10 cm up to 3-4 m from the ground. *Cordia multispicata* is heterostylous, with short- and long-styled flowers, white, pentamerous and tubular (figure 1A). The corolla is about 4-5 mm long, the stamens are alternipetalous and inserted in the corolla, the pollen is white and powdery and the stigma is four-lobed. The nectary is yellow, annular and is located at the base of the superior ovary. The corolla length in long- and short-styled flowers were not significantly different. The length of the filaments is 2.2 mm in short-styled and 1.6 mm in long-styled flowers, and the length of the style is 3.0 and 5.0 mm, in short- and long-styled flowers, respectively.

Anthesis begins at 6:00 a.m., and the flowers last until 3:00-4:00 p.m. of the same day. The flowers emit a strong odour, slightly sweet. The sugar concentration in the nectar is 15-16.6% (\( \bar{X} = 16\% \); \( n = 20 \)). The nectar volume is very low (ca.1 µL per flower). Pollen viability is 96% in both long- and short-styled flowers.

Results of manual pollination experiments (table 1) show that only crosses between different floral types set fruit. Self- and intramorph pollinations were completely incompatible. The fruit set under natural conditions was ca. 48%.

*Borreria alata* is a herbaceous species, also with tubular flowers (figure 1B) disposed in glomerulous inflorescences. Each glomerule has ca. 65 flowers, and opens 2-4 flowers per day. The flowers are white with a purple corolla tip and are covered with glandular hairs at the tube entrance. The corolla is about 6 mm long and 4 mm wide, the stamens are alternipetalous and inserted in the corolla, and the pollen is white and powdery. The nectary is a discon-

![Figure 1.](image-url)
tinuous ring located at the top of the inferior ovary. There is a floral polymorphism in relation to the length of the style and, in some flowers, also in relation to the length of the filaments, which can vary from 0.6 to 1.6 mm. The length of the style varies from 2.5 to 6.0 mm, with some intermediate sizes in the same inflorescence (figure 2).

Anthesis begins at 6:00 a.m. and the flowers last until 2:00 p.m. The flowers are odourless and the mean sugar concentration in the nectar was 30%, varying from 29 to 31% (n = 39). The volume of nectar per flower was ca. 2-3 µL. Under natural conditions the fruit set was 78% (n = 100).

The flowering period of both species extends almost throughout the year, with peaks in the rainy season (from May to July) as well as during the dry season (from October to December).

Nine species of flies, mainly of the genus Palpada (Syrphidae), were recorded on flowers of Cordia multispicata and Borreria alata (table 2). Seven of these fly species were observed pollinating flowers of both C. multispicata and B. alata. The same fly can visit flowers of both species, where plants were very close. Two fly species of Dipterans visited exclusively flowers of C. multispicata, whereas none visited exclusively flowers of B. alata (table 2). The visiting behaviour was similar for all the fly species. During a visit the flies land on the corolla (figure 3), moving towards the nectary and inserting the mouth parts into the central region of the flower, thus drinking nectar. At this time, they touch anthers and stigma with the head, torax, abdomen and/or with the legs. The flies walk on the inflorescences visiting different flowers before moving to another plant. They remain on each flower from 3 to 50 seconds. The visits occur from 6:00 a.m. to 2:00 p.m., mainly from 8:00 a.m. to 12:00 in C. multispicata (figure 4).

Two species of Palpada (P. vinetorum and Palpada sp.) were considered as the main visitors and pollinators (figure 4), of both C. multispicata and B. alata flowers. The former was most frequent in C. multispicata and the latter in B. alata.

Bees (Apis mellifera, Trigona spinipes and Augochloropsis sp.) and butterflies were also observed visiting flowers of the two species. They can pollinate the flowers since they touch the anthers and stigma, but due to their low visiting frequency they were considered as occasional pollinators.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Flowers (number)</th>
<th>Fruits (number)</th>
<th>Success (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Self-pollination</td>
<td>180</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Cross-pollination</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Long-styled* X long-styled</td>
<td>50</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Short-styled* X short-styled</td>
<td>60</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Long-styled* X short-styled</td>
<td>60</td>
<td>30</td>
<td>50</td>
</tr>
<tr>
<td>Short-styled* X long-styled</td>
<td>50</td>
<td>28</td>
<td>56</td>
</tr>
<tr>
<td>Control</td>
<td>240</td>
<td>115</td>
<td>48</td>
</tr>
</tbody>
</table>

*Pollen donor.

Figure 2. Flowers of Borreria alata without the corolla tubes and the stamens, showing the different sizes of the styles in the same inflorescence (x1.6).
Discussion

*Cordia multispicata* and *Borreria alata* have diurnal, whitish, actinomorphic tubular flowers with exposed sexual organs. These attributes together with the easy access to nectar and the high number of flowers per inflorescence attract, in general, a great number of floral visitors, especially those with short mouth parts, as most of the dipterans (Faegri & van der Pijl 1979, Proctor et al. 1996). Most of the flies of the family Syrphidae have proboscides varying from 2 to 4 mm (Proctor et al. 1996), which are compatible with the lengths of the floral tubes of the studied species. Although the pollination carried out by small flies can be, in many cases, inefficient due to their size (Proctor et al. 1996), in the case of small flower species, such as *C. multispicata* and *B. alata*, these insects seem to be very efficient as pollinators.

The sugar concentration in the nectar varied between the two species, being highest in *B. alata* flowers. According to Vogel (1983), flies and bees prefer a more concentrated nectar than the ones found in *B. alata* and *C. multispicata*. On the other hand, Percival (1965) refers to fly-pollinated blossoms with sugar concentration varying from 5 to 15%. Baker (1975) indicates a mean value of 20% for fly-pollinated flowers, based on the information of Percival (1974). In this same work, Percival found a concentration of 16% in *Borreria laevis*, a species visited by many Syrphidae. In turn, Pombal & Morellato (1995) found a very low sugar concentration (2%) in the nectar of *Dendropanax cuneatum* (Araliaceae), a species which is also pollinated by flies. Our data show mean sugar concentrations of 16% in *Cordia multispicata* and 30% in *Borreria alata*, which reinforce the suggestion of Pombal & Morellato (1995) that sugar concentrations in the nectar of fly-pollinated flowers may vary widely and does not seem to be a determining factor in the searching for flowers by the flies. The low amount of nectar offered per flower in both species is compensated by the great number of available flowers in each species per day. Flies are known to use variable feeding resources in their diet (Faegri & van der Pijl 1979), however nectar is an important source of carbohydrates to their metabolism (Vogel 1983).

The visits to the flowers of both species occurred during the morning, mainly between 8:00 a.m. and 12:00, in the case of *C. multispicata*. In fact, many species of Syrphidae feed only at specific hours of the day (Weems 1953). Regarding specificity, most of them forage on many types of flowers, but they are generally observed on white, greenish, inconspicuous flowers (Faegri & van der Pijl 1979, Proctor et al. 1996, Weems 1953), such as *C. multispicata* and *B. alata* flowers.

The flowers of *B. alata* in contrast to those of *C. multispicata*, emit a strong sweet odour, but this

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**Table 2. Size and frequency of the Diptera visitors to flowers of *Cordia multispicata* and *Borreria alata***

<table>
<thead>
<tr>
<th>Diptera</th>
<th>Length (mm)</th>
<th><em>C. multispicata</em></th>
<th><em>B. alata</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Syrphidae</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Ornidia obesa</em> (Fabricius 1775)</td>
<td>10</td>
<td>f</td>
<td>F</td>
</tr>
<tr>
<td><em>Palpada furcata</em> (Wiedemann 1830)</td>
<td>10</td>
<td>f</td>
<td>F</td>
</tr>
<tr>
<td><em>Palpada geniculata</em> (Fabricius 1805)</td>
<td>12</td>
<td>f</td>
<td>F</td>
</tr>
<tr>
<td><em>Palpada vinetorum</em> (Fabricius 1798)</td>
<td>12</td>
<td>f</td>
<td>F</td>
</tr>
<tr>
<td><em>Palpada sp. 1</em></td>
<td>10</td>
<td>f</td>
<td>F</td>
</tr>
<tr>
<td><em>Palpada sp. 2</em></td>
<td>10</td>
<td>f</td>
<td>F</td>
</tr>
<tr>
<td>Calliphoridae</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Chrysomya megacephala</em></td>
<td>9</td>
<td>r</td>
<td>-</td>
</tr>
<tr>
<td>Sarcophagidae</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Sarcophagula sp</em></td>
<td>4</td>
<td>r</td>
<td>-</td>
</tr>
<tr>
<td>Tachinidae</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not identified</td>
<td>10</td>
<td>r</td>
<td>R</td>
</tr>
</tbody>
</table>

vf- very frequent > 70% of the total of visits, f- frequent 30-70%, lf- less frequent 10-30%, r-rare < 10%.
attribute as in the case of nectar sugar concentration does not seem to determine the searching for flowers by the flies. Arruda & Sazima (1996) found that flowers of many plant species visited by Syrphidae emit a perceptible odour. The fact that the flowers of *B. alata* emit a strong sweet odour could be an

Figure 3. *Palpada* spp. (Syrphidae) visiting flowers of *Cordia multispicata* (a- x2.8, b- x2.0, c- x2.6) and *Borreria alata* (d-f x2.2). a,d- *Palpada* sp.; b, c- *Palpada geniculata*; c- *Palpada vinetorum*; f- *Palpada furcata*. 
advantage to that species in attracting flies over a long distance. However, this situation does not seem to occur, since the diversity and frequency of visitors to the flowers of *C. multispicata* and *B. alata* are similar, and they share seven species.

The frequency of flies visiting flowers of a species or an individual plant may be related to climate conditions (Arruda & Sazima 1996, Vieira & Silva 1997) and to the presence of other species of plants flowering at the same time (Arruda & Sazima 1996). Under some conditions, co-occurring species may mutualistically favour or improve their reproduction (Brown & Kodric-Brown 1979, Schemske 1981). Some studies, on the other hand, provide evidence that the convergence of floral characters in co-occurring species induces a competition for the same pollinators (Armbruster & Herzig 1984, Yanagizawa & Gottsberger 1982/83) and loss of pollen through interspecific pollen flow (Armbruster & Herzig 1984). The floral similarity between *C. multispicata* and *B. alata*, the overlap in flowering time and the occupation of the same environment in the forest, in glades or edges, seem to be advantageous for both, resulting in benefits to them, since they may increase pollinator attraction. In a study carried out with *C. multispicata* in grazing areas in Amazon, Vieira & Silva (1997) observed that, during the dry season, when the number of pollinators was low, the fruit set under natural conditions was also low, varying from 11 to 25%, while in the rainy season it increased to 48 to 61%. According to Vieira & Silva (1997), the pollen flow between the two floral morphs of *C. multispicata* is small when the activity of the insects is reduced during the dry season, and it is one of the determinant factors of the low fruit set. Our data show that the fruit set of 48% under natural conditions in the forest of Dois Irmãos is very near to the maximum value obtained in the interform manual crossing (see table 1) in *C. multispicata* and indicates that activity and efficiency of pollinators are high in the study area.

**Figure 4.** Average number of Dipteran visitors to flowers of *Cordia multispicata* (A) and *Borreria alata* (B) during three consecutive days.

Regarding the reproductive system, *C. multispicata* is self-incompatible and heterostylic with an isoplectic frequency of the two floral morphs, similar to many other distylic species of Boraginaceae or other families (Ornduff 1971, Sobrevila et al. 1983, Barrett & Shore 1985, Richards & Barrett 1992). The occurrence of heterostyly in *Cordia* is known since Darwin (1877) and it is considered the most common type of reproductive system in the genus (Opler et al. 1975). In *C. multispicata* the existing mutual correspondence related to the size of stamens and styles makes easier the intermorph pollen flow, similar to other species of *Cordia* (Opler et al. 1975).

*Borreria alata*, in turn, is probably a self-compatible species, with a high fruit set under natural conditions. The occurrence of apomixis was not tested. In the family Rubiaceae heterostyly also occurs widely. It was mentioned for 13 studied genera (Barrett & Richards 1990), from the 91 heterostylistous genera listed for this family (see Ganders 1979, Bir Bahadur 1968, Vuilleumier 1967). Barrett & Richards (1990) did not include the genus *Borreria* in their list (even though it is mentioned in the lists of Bir Bahadur (1968) and Ganders (1979)) once the genus had not been effectively studied. The flowers of *B. alata* show a polymorphism related to the size of the style. Some species of Rubiaceae show a
characteristic distyly, associated with a strong mechanism of self-incompatibility, as in *Palicourea fendleri* (Sobrevila et al. 1983). Other species show polymorphisms of the style that differ from those typically distylos, as occurred in *Guettarda scabra* (Richards & Koptur 1993), a self-compatible species. It is possible that *Borreria alata* shows an "anomalous heteromorphism", as defined by Barret & Richards (1990). Further studies would be interesting in order to analyse this hypothesis.

Acknowledgments - We thank Dr. A. X. Linhares for the identification of the flies and the researchers of the IPA Herbarium (Empresa Pernambucana de Pesquisa Agropecuária) for the identification of the plants. To Dr. S. Vogel, Msc. A. V. Lopes, Dr. C. Westerkamp and two anonymous reviewers for their critical comments and improving on the manuscript. The COMPESA for permission to work in lands under their care; the CNPq and CAPES for essential financial support.

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