Reproductive traits and floral visitors of *Aechmea distichantha* plants growing in different habitats of a South American xerophytic forest

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**Abstract**

*Aechmea distichantha*, a widely-distributed facultative epiphytic bromeliad species, is present from rainforests to xerophytic forests. At its southernmost distribution (Humid Chaco) it grows in the understory and forest edges. This animal-pollinated bromeliad shows high phenotypic plasticity on its vegetative traits, but there is no information about plasticity on its reproductive traits. Infructescences from shade plants were heavier, had longer rachis, more spikelets, higher number of fruits/spikelet and higher number of seeds/fruit than those from sun plants, but they presented similar number of open flowers. The number of visitation events was similar in both habitats, but more flowers were visited in the sun than in the shade. Flowers were visited by seven species (six insects and one hummingbird). In the sun, the carpenter bee was the most frequent visitor and visited almost all flowers, whereas in the shade different species of visitors attained similar proportion of visits and number of visited flowers. Despite visitation events were similar in both habitats, plants growing in the shade set more seeds/fruit than plants growing in the sun. The higher proportion of visits accomplished by carpenter bees compared to hummingbirds is probably a consequence of the climatic conditions in the austral location of these populations.

**Key words:** *Aechmea distichantha*, habitat, pollinators, reproductive characters.

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**Introduction**

The Bromeliaceae is a large monocotyledon Neotropical family that has been traditionally divided into eight subfamilies: Brocchinioideae, Lindmanioideae, Tillandsioideae, Hechtioideae, Navioideae, Pitcairnioideae, Puyoideae, and Bromelioideae (Givnish et al. 2011). Most bromeliad species are mainly pollinated by...
animals, but there are large differences in the relative importance of different pollinators (i.e., hummingbirds, bats, bees, moths, and butterflies) among bromeliad species (Benzing 2000; Kessler & Krömer 2000; Givnish et al. 2014) and even within a bromeliad species (Schmid et al. 2011; Rocca & Sazima 2013).

The pollinator assemblages visiting bromeliads are conditioned by resource availability, such as pollen and nectar (Canela & Sazima 2005), as well as by some morphological traits of flowers, such as corolla color and length (Sazima et al. 1996). According to the bee avoidance hypothesis, these insects are excluded from red flowers due to their blindness for these wavelengths (Bergamo et al. 2016). Likewise, only hummingbirds and those insects whose tongues are long enough to reach the nectary chamber will be able to visit bromeliad flowers with depth-probed floral tube (Taura & Laroca 2001; Vossler et al. 2014). Therefore, hummingbirds are expected to be more important than bees for bromeliads with deep red or pink flowers. For some bromeliad species pollinated by different agents, it is known that bees have lower effectiveness inducing seed set than hummingbirds (Schmid et al. 2011a). However, it is important to assess the contribution of different groups to the reproductive success by analyzing the seed set (Canela & Sazima 2005).

Microclimatic factors are also known to influence pollination. Variability of irradiance associated with environmental heterogeneity has an effect on pollination by affecting foraging responses and thus on the composition of pollinator assemblages (Herrera 1995; Kilkeny Galloway 2008). Microclimatic factors such as irradiance may not influence the floral display, but may have a direct effect on pollinators behavior and consequently on plant reproductive success (Kilkeny & Galloway 2008).

Aechmea distichantha Lem. (Bromeliodeae) is widely distributed in subtropical regions of South America from 14° to 30°S (Smith & Downs 1979). This facultative tank epiphyte lives in the canopy of the rain forests, as well as in the understory of seasonal dry and xerophytic forests from the sea level in the east to 2,200 m above the sea level in the west (Smith & Downs 1979). It reproduces both sexually and asexually (Smith & Downs 1979; Mercier & Guerreiro Filho 1990; Bianchi et al. 2000; Scrok & Varassin 2011). It flowers in the spring and in the summer, and no latitudinal or altitudinal patterns in flowering period have been described (Buzato et al. 2000; Marques et al. 2004; Scrok & Varassin 2011; Vizentin-Bugoni et al. 2014; Vossler et al. 2014). Aechmea distichantha has hermaphrodite flowers, 12.4–18 mm in length (Sazima & Sazima 1999; Krömer et al. 2006), arranged in an inflorescence composed of spikes of spikelets, with centripetal anthesis at the spike and at the spikelet levels. The anthesis extends for only one day (Sazima & Sazima 1999; Scrok & Varassin 2011) and the nectar produced in the septal nectaries accumulates at the bottom of the flower (Bernardello et al. 1991; Buzato et al. 2000; Scrok & Varassin 2011; Oliveira et al. 2016). Flowers are pollinated by hummingbirds, perching birds, bees, and butterflies (Araujo 1996; Sazima et al. 1996; Sazima & Sazima 1999; Taura & Laroca 2001; Krömer et al. 2006; Scrok & Varassin 2011; Bueno 2012; Rosa & Monteiro 2012; Chupil 2013; Vizentin-Bugoni et al. 2014; Vossler et al. 2014), and has been described as partially self-compatible (Scrok & Varassin 2011) or self-incompatible (Bianchi et al. 2000), but a few (5%) fruit set was produced following self-pollination.

Plants of Aechmea distichantha exhibit high phenotypic plasticity; individuals growing in the sun are shorter, have shorter diameter, more leaves, and higher sheath biomass fraction than plants growing in the shade (Cavallero et al. 2009, 2011). The proportion of adult plants in reproductive stage is higher for plants growing in the shade (Barberis, Klekailo & Galetti, unpublished data). Although phenotypic plasticity for vegetative characters has been demonstrated for this species, it is not known how reproductive characters are affected by environmental factors and if these variations in reproductive characters are associated with the composition of the pollinator assemblage.

Therefore, this work has two objectives. Firstly, to evaluate whether there are differences in the characteristics of the inflorescences (length and dry weight of floral rachis, spikelets per rachis, flowers per spikelets) between plants growing in the sun vs. those growing in the shade. As shade plants have larger biomass, it is expected a higher number of flowers in shade vs. sun plants. Secondly, to evaluate whether the activity of pollinators, the pollinator assemblage, and consequently the reproductive success (number of seeds per fruit), are affected by habitat differences and variations in inflorescence traits at different environments. As shade plants have larger inflorescences, a higher number of visits and an increase in the reproductive success are expected.
Material and Methods

Study area

The study was carried out in a stand of Schinopsis balansae forest (‘quebrachal’; Lewis et al. 1997), located at “Las Gamas”, Santa Fe, Argentina (Estación Experimental Tito Livio Coppa, 29°28’S, 60°28’W, 58 m above the sea level). This site at the Cuña Boscosa Santafesina is located at the southernmost range of the distribution of A. distichantha of the Eastern Chaco (Lewis 1991; Barberis et al. 1998, 2002, 2014). The climate is humid temperate to warm, with a mean annual temperature of about 20 °C, and a mean annual precipitation of about 1,000 mm. Rainfall is concentrated in the summer (December–March), and a dry season of variable length occurs during the winter (June–August) (Lewis & Pire 1981). The forest is located on a mosaic of soils with low hydraulic conductivity, high sodium content and high electrical conductivity (Espino et al. 1983). These forests show high environmental heterogeneity, mostly related to topographic variation, with high density of woody species on relatively elevated patches, and open areas with grassy vegetation (Barberis et al. 2002). Aechmea distichantha plants are largely restricted to relatively elevated patches representing an important element of the understory, but may also grow at the edges of the open areas showing a gradient related to light exposure (Barberis et al. 2014). The populations are patchily distributed due to their clonal growth.

In the study area, it mainly flowers in the spring (September–November), even though it is not unusual to find flowering individuals throughout the year, even during the winter (J.L. Vesprini, personal observation).

Inflorescence traits and seed production

Fieldwork was conducted in December 2013. For 10 plants from the sun and 14 plants from the shade, we took one flower and measured the length of their corollas. We chose 30 plants growing in the shade of the understory and 12 plants exposed to full radiation. During fruit development, the infructescences keep all old flowers attached, including the ones that did not develop into fruits. Therefore we used the infructescences to determine the inflorescence traits. The infructescences were harvested, placed in paper bags, labelled and kept in a dark and dry place for two days until its processing. Then, each infructescence was oven-dried at 60 °C for 120 hours and weighed. We also measured the length of its rachis, and counted the number of spikelets. We randomly chose two spikelets from three different positions along the infructescence (bottom, middle, and top), and counted the number of fruits (six spikelets for each infructescence). Then we chose a fruit from the center of each spikelet and counted the number of seeds.

Pollinator sampling

In October 2013, we selected two forest patches separated by 20 m with about 14 flowering plants each. One patch was located in the sun and the other in the shade. We randomly selected six plants at each habitat, and for each inflorescence we counted the number of open flowers. During six days (22–27 October 2013) we carried out eight 10-min sampling periods per day; two sampling periods in the morning (10:00–11:30 h) and two others in the afternoon (16:30–18:00 h) for each habitat. In total, there were 44 sampling periods (22 in the sun and 22 in the shade). At each sampling period we recorded the identity of flower visitors, counted the number of visits and the number of flowers visited at each visit. One morning (26th October 2013), sampling was not carried out due to rainfall.

Data analyses

We applied the Student t-test to analyze differences in infructescence biomass, rachis length and number of spikelets between sun and shade plants. We applied the Bonferroni test to correct for multiple comparisons from a single data set, thus for these tests the significance P-value was set to 0.05/3 = 0.0167. For other data set, we also used the Student t-test to analyze differences in corolla length between sun and shade plants. All these t-tests were carried out with the Rcmdr package in R (Fox & Bouchet-Valat 2016). We performed general linear mixed models with Poisson distribution to analyze differences in fruit number/spikelet and seed number/fruits between spikelets located at different positions, between sun and shade plants, as well as their interactions. Therefore, in these models, the habitat (sun vs. shade) and the spikelet position (bottom, middle, or top) were considered fixed factors, while the random factors considered the following nested structure. Fruits nested within spikelets, and spikelets nested within infructescence. We fitted a model with all terms (i.e., both fixed effects and their interaction) and a model without the interaction. Then we used maximum likelihood tests of nested models (i.e., with and without interactions) to evaluate the effect of the interaction (Crawley 2013). We used
the function glmer from the lme4 package (Bates et al. 2015).

We used Kruskal-Wallis test to evaluate whether there were differences between habitats in the number of open flowers. We also applied this test to analyze if there were differences between habitats in the number of visits/10 min and the number of flowers visited/10 min. These tests were used to analyze if there were differences between habitats in the number of visits/10 min and the number of flowers visited/10 min considering each visitor species separately and for all visitor species together. We used these non-parametric tests because the data sets were small and non-normally distributed. All analyses were carried out with the Rcmdr package in R (Fox & Bouchet-Valat 2016).

We performed a multivariate analysis of variance to evaluate whether there were differences between habitats in the species composition of the assemblage of flower visitors. We used two matrices, one including number of visits/10 min and the other for number of flowers visited/10 min. The sampling periods were used as replicates. This could inflate our sampling due repeated observations on the same plant individuals, but the differences between habitats seems to be consistent (see Results section). These multivariate analyses were carried out with the program MRPP (Multi-Response Permutation Procedure) of the PC-Ord package (McCune & Mefford 2011) using the Sörensen index as a distance measure.

Results

Flowers from plants growing in both habitats presented similar corolla length (Sun: mean = 14.28 mm, SD = 1.32, Shade: mean = 15.23 mm, SD = 0.95; t = –2.05, P = 0.053). There were significant differences in infructescence traits between sun and shade plants (all P < 0.05). The infructescences from plants growing in the shade were heavier (t = 3.06; P = 0.0054), had longer rachis (t = 3.93; P = 0.0003), but had similar spikelets than those from sun plants (t = –1.98; P = 0.0606) (Fig. 1). However, they presented similar number of open flowers in both habitats (Sun: Median = 3.5, Range = 0–14, Shade: Median = 6, Range = 0–9; H = 0.106, P = 0.745).

Shade plants had higher number of fruits/spikelet and higher number of seeds/fruit (P < 0.05) (Fig. 2). In both habitats, spikelets located at the bottom had more fruits than spikelets located at its top (P < 0.05) (Fig. 2). In contrast, there were no differences in number of seeds/fruits for spikelets located at different positions along the infructescences (P > 0.05) (Fig. 2). Interactions were not significant (all P-values > 0.05).

Visits happened in 25 out of the 44 10-min sampling periods (14 in the sun and 11 in the shade). We recorded 101 visitation events (53 in the sun and 48 in the shade) that resulted in 285 flowers visited (167 in the sun and 118 in the shade). Flowers were visited by six species of insects (four Hymenoptera, one Diptera, one Lepidoptera) and one hummingbird species (Chlorostilbon lucidus). In both habitats, most species were recorded only in 20% of the sampled periods (Fig. 3). The exception was the carpenter bee (Xylocopa ordinaria) which was recorded in half of the sampled periods in the sun (Fig. 3). The average visitation rate was 4.53 visits/flower/hour (SE = 1.36) in the sun and 2.57 visits/flower/hour (SE = 0.98) in the shade.

There were no differences between habitats in the number of inflorescences or flowers visited by all floral visitors (Fig. 5; Tab. 1). However, there were differences in the frequency of visit of different species between habitats. The carpenter bees carried out more visits to sun plants than to shade plants, whereas the unidentified Hymenoptera

![Figure 1](image-url) - Boxplots of infructescence dry weight, rachis length and number of spikelets of Aechmea distichantha plants grown in the sun and in the shade. Each boxplot includes the 25th percentile, the median, and the 75th percentile. Whiskers (error bars) above and below the box indicate the 90 and 10th percentiles. For each variable, values with the same letters are not significantly different.
species showed the opposite pattern (Fig. 5; Tab. 1). Even though three species were recorded only at one site (the unidentified Vespidae species and the hummingbird in the shade plants, and Polistes canadensis in the sun) the number of observations was too low to detect significant differences between habitats (Tab. 1). The butterfly (Lepidoptera: Pieridae; one species) and the fly (Diptera: Syrphidae; one species) were recorded in both habitats, but there were no significant differences between habitats (Tab. 1). Many honey bees (Apis mellifera) were observed in the area

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**Figure 2** – Boxplots of flower number/spikelet and seed number/flower for spikelets located at the bottom, middle or top of the infructescence of Aechmea distichantha plants grown in the sun and in the shade. Each boxplot includes the 25th percentile, the median, and the 75th percentile. Whiskers (error bars) above and below the box indicate the 90 and 10th percentiles. For each variable, values with the same letters are not significantly different.

**Figure 3** – Percentage of the sampled periods at each habitat where each animal species was present visiting Aechmea distichantha inflorescences. N denotes the number of sampled periods visited at each habitat.
drinking water in the phytotelmata, but no flower visits were recorded for this species.

There were significant differences between habitats in the community of floral visitors, either measured as number of visits/10 min (MRPP test, A = 0.053, P = 0.016) or as number of flower visited/10 min (MRPP test, A = 0.039, P = 0.035). In the sun, the carpenter bee was the most frequent visitor and visited almost all flowers, whereas in the shade different species of visitors attained similar proportion of visits and number of visited flowers (Fig. 4). We acknowledge that our sampling may be inflated due to repeated observations on the same individuals. However, the differences in assemblages seem to be strong despite this statistical limitation.

Discussion

Plant traits

Infurescences from plants growing in the shade had longer rachis and higher number of fruits than those from sun plants. Nonetheless, infurescences had similar number of open flowers per day in both habitats. A similar pattern was recorded in southern Brazil (Scrok & Varassin 2011). This result indicates that bigger infurescences do not offer a higher number of open flowers each day, and thus probably offer similar amount of nectar at both habitats. Additionally, it should be taken into account that bigger infurescences can offer reward for longer periods (Firmage & Cole 1988; Kilkenny & Galloway 2008).

_Aechmea distichantha_ displays a corolla of 18–20 mm length and the floral tube has a small diameter that allows medium and long-tongued bees to reach the nectar. On the other hand, the flowers seem to hinder the access to short-tongued bees that have no access to nectary chambers because of the combination of the length of their proboscis and the floral tube morphology (length and width) such as _A. mellifera_. There are some reports for medium- or long-tongued bees (Taura & Laroca 2001; Vossler et al. 2014) these reports are supported by our records of honey bees drinking water in the phytotelmata but not visiting the flowers.
Pollinator assemblage

Previous studies on the reproductive biology of *A. distichantha* carried out in areas located at lower latitudes reported hummingbirds and butterflies as its main flower visitors (Bernardello et al. 1991; Araujo 1996; Buzato et al. 2000; Scrok & Varassin 2011; Rosa & Monteiro 2012; Chupil 2013; Vizentin-Bugoni et al. 2014). In contrast, in this study conducted at the southern distribution range of *A. distichantha* we found that carpenter bees are the main pollinators (in terms of visitation rates). Differences in the pollinator assemblages can be explained by different latitudes and climatic conditions (Krömer et al. 2006), but the plant community composition may also play a central role in the assemblage of pollinators due to changes in the floral neighborhood (Lázaro et al. 2009). In the studied community, ornithophilous species are

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**Figure 5** – Boxplots of *Aechmea distichantha* plants in the sun and in the shade visited by each animal species. Each boxplot includes the 25th percentile, the median, and the 75th percentile. Whiskers (error bars) above and below the box indicate the 90 and 10th percentiles.
scarce and only represented by a liana (*Dolichandra cynanchoides*), few epiphytes (*Tillandsia* spp.) and a cactus (*Cleistocactus baumanii*) (Bianchi et al. 2000; Vesprini, personal observation), which mainly flower during the spring and the summer. Although a few individuals of *A. distichantha* were observed flowering in winter, there would not be abundant winter floral resources for large populations of hummingbirds. In contrast, carpenter bees not only are able to forage on a larger number of plant species, but they also collect pollen, and have lower energy requirements (Heinrich 1975). Therefore, it is likely that the high proportion of visits accomplished by carpenter bees is a consequence of the austral location of these *A. distichantha* populations and also of the climatic conditions favoring a large number of bees.

Hymenoptera species have been considered functional pollinators for other bromeliad species (Abrahamovich *et al.* 2001; Fumero-Cabán & Melández-Ackerman 2007; Kamke *et al.* 2011; Schmid *et al.* 2011a, 2011b, 2011c; Guerra *et al.* 2012). For instance, *Xylocopa, Bombus* and some species of Augochlorine bees induced seed set in *Aechmea nudicaulis*, albeit with much lower effectiveness than hummingbirds. However, the low effectiveness of bees may be compensated by their much higher frequency of flower visitation (Schmid *et al.* 2011a). Besides, the low effectiveness of bees may be coupled with low quality pollination with regard to a higher proportion of visits leading to selfing vs. outcrossing (Bergamo *et al.* 2016).

### Table 1 – Results of tests to evaluate differences in number of visits/10 min and number of flowers visited/10 min between sun and shade habitats for all floral visitors and for each visitor species.

<table>
<thead>
<tr>
<th></th>
<th>Visits/10 min</th>
<th>Flower visited/10 min</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>KW-Chi.squared</td>
<td>P</td>
</tr>
<tr>
<td>All floral visitors</td>
<td>0.361</td>
<td>0.5478</td>
</tr>
<tr>
<td><em>Xylocopa ordinaria</em></td>
<td>3.952</td>
<td>0.0468</td>
</tr>
<tr>
<td><em>Hymenoptera</em> sp</td>
<td>5.500</td>
<td>0.0190</td>
</tr>
<tr>
<td><em>Vespidae</em> sp</td>
<td>2.047</td>
<td>0.1526</td>
</tr>
<tr>
<td><em>Polistes canadiensis</em></td>
<td>1.000</td>
<td>0.3173</td>
</tr>
<tr>
<td><em>Syrphidae</em> sp</td>
<td>0.931</td>
<td>0.3347</td>
</tr>
<tr>
<td><em>Chlorostilbon lucidus</em></td>
<td>2.047</td>
<td>0.1526</td>
</tr>
<tr>
<td><em>Pieridae</em> sp</td>
<td>0.031</td>
<td>0.8595</td>
</tr>
</tbody>
</table>

Note: The Kruskal Wallis and P values are shown. Significant results are shown in bold.

### Pollinator assemblage and reproductive success in different habitats

The number of visitation events was similar in sun and shade habitats. However, our results suggest that pollinator assemblages are conditioned by habitat, with carpenter bees showing strong preferences for foraging in the sun. An effect of irradiance on the pollinator assemblage composition was also found in another study: endothermic pollinators (large-size bees) visited flowers independently of habitat while ectothermic insects (small-size bees and butterflies) discriminated between sun and shade patches (Herrera 1995). Interactions between pollinators may also be important in the studied case. In a study on *Impatiens biflora* pollinated by a hummingbird and two large bees, it was found that individuals of one pollinator species can be largely excluded from access to nectar by foragers of other pollinator species with larger tongues. It is likely that hummingbirds would not have problems to forage in sunny habitats, but nectar is depleted at sun a large frequency of visits attained by bees (Laverty & Plowright 1985).

In our study, as carpenter bees have a high visitation rate, more flowers were visited in the sun than in the shade. Conversely there were 2.41 more seeds produced per infructescences in the shade than in the sun (*i.e.*, 1,650 vs. 681 seeds/infructescence). In contrast, Scrok & Varassin (2011) recorded more seeds in the sun...
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produce seeds between plants located at different habitats, but since shade plants have more flowers per inflorescence it is possibly that they remain open flowers at a given time was the same at both habitats. On one hand, this could be related to the higher pollinator richness in the shade, because for other plant species it has been shown that seed set is more affected by the diversity than by the abundance of pollinators (Aizen 2007). It is important to consider that the quality of the visits of bees may be lower than the visits accomplished by hummingbirds, and that sunny habitats may have fewer nutrients which may affect seed set of sun plants. On the other hand, we did not record the length of the flowering period at both habitats. However, when we carried out the survey there was similar number of open flowers between habitats, but since shade plants have more flowers per inflorescence it is possibly that they remain open flowers for longer periods.

At our study site it is rare to observe seedlings originated from seeds. Thus, even though sexual reproduction could be very important for habitat colonization, it is not easy to determine whether these differences found in the ability to produce seeds between plants located at different habitats have an adaptive value. For species, whose reproduction within a colonized habitat is secured by clonality, the importance of sexuality (asexual vs. sexual) would be higher in other processes like clone rejuvenilization or maintenance of genetic variability (Silvertown & Lovett Doust 1993).

Final comments

Habitat influences most inflorescence traits, with shade plants having bigger inflorescences than those in the sun. However, a similar number of flowers per plant are open each day in both habitats, and thus a higher nectar reward for pollinators is not expected between habitats.

At the southernmost portion of its distribution range, this bromeliad species is mainly visited by insects. The pollinator assemblage composition and the visitation rate vary between habitats. The existence of pollinator assemblages between habitats was unexpected. In the sun, the carpenter bee was the most frequent visitor and visited almost all flowers, whereas in the shade different species of visitors attained similar proportion of visits and number of visited flowers.

A higher pollinator richness in the shade may be associated with a higher reproductive success, evidenced by a higher seed set. The observed variation in seed production between habitats has not necessarily an adaptive value because sexual reproduction of this species does not play an important role in the structure and dynamic of its populations.

Finally, factorial experiments would be the most suitable step to test most of the hypotheses here presented, since it would allow to control many environmental variables, as well as biotic interactions.

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