

## SHORT COMMUNICATION

# Flowering as a key factor in ant–*Philodendron* interactions

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With the spread of angiosperms some lineages of ants, originally ground-dwellers and predators, adapted to arboreal life. Ground-nesting worker ants probably constituted the first case of biotic plant protection through their predatory activity while foraging on plant foliage. Then, ants developed tight evolutionary bonds with plants varying from facultative diffuse relationships to obligatory specific associations, necessary to the survival of both partners. In diffuse relationships plants induce different ant species to patrol their foliage by producing energy-rich food rewards such as extra-floral nectar (EFN) and/or food bodies (FBs) (Dejean *et al.* 2007, Heil 2008, Heil & McKey 2003). First described by Janzen (1966), myrmecophytes, or plants that provide specialized plant-ants a nesting place in hollow structures called domatia, represent a good example of a strict association as, in return, they are protected from several kinds of enemies, particularly defoliating insects (Heil & McKey 2003, Hölldobler & Wilson 1990).

The Araceae, a monocotyledon family of 3000 mainly tropical species, has only a few known cases of associations with ants. In a few genera (i.e. *Alocasia*, *Culcasia*, *Philodendron*), ants can exploit nectar produced by young leaves and sometimes tend Aleyrodidae (Bastien & Belin-Depoux 1998, Blüthgen *et al.* 2000). *Philodendron insigne*, a trash-basket epiphyte with suspended soil, is inhabited by a variety of ant species and particularly the Ponerinae *Odontomachus hastatus* (Gibernau *et al.* 2007); while *Philodendron megalophyllum* (= *P. myrmecophyllum*) and *Anthurium gracile* develop in ant-gardens (Orivel & Dejean 1998). Hence, excluding these few examples, no

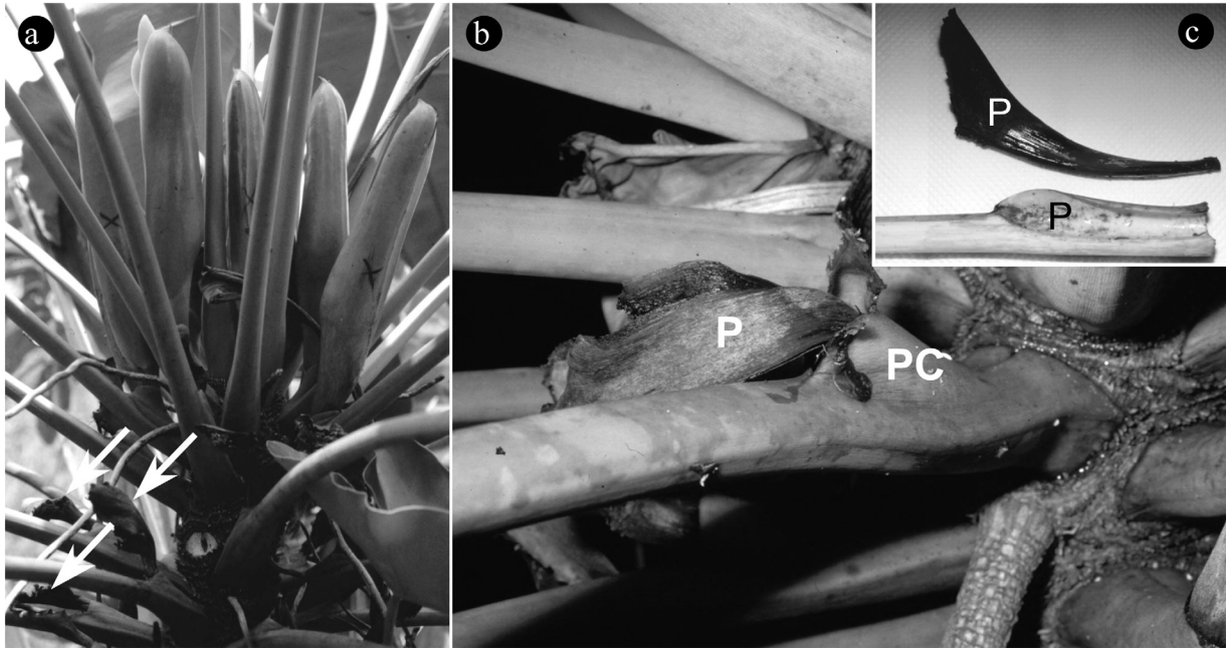
aroid is known to have strict relationships with ants or to be a myrmecophyte.

*Philodendron solimoense* A. C. Smith is a hemi-epiphytic aroid species found in upland or riverine forests where it develops on sandy and flooded soils, as was the case in our study site. Like other species of the genus, *P. solimoense* produces EFN on bud bracts, the petiole, the base of the leaf lamina and the spathe of the inflorescence, suggesting a diffuse relationship with ants (Bastien & Belin-Depoux 1998). Solitary inflorescences, arising vertically from the petiolar sheath at the base of the leaves (Figure 1a), are thermogenic and pollinated by dynastine beetles (Gibernau *et al.* 1999). At maturity the closed spathe abscises from the base, exposing ripe fruits to frugivorous animals. After fruit dispersal, the spadix falls, breaking above the peduncle, which is hollow and remains inserted in the petiolar sheath (Figures 1b-c).

In this study we hypothesized that the *P. solimoense* petiolar sheaths can serve as durable shelters for ant colonies, making this plant species a possible myrmecophyte.

During July 1998 and 1999 we monitored 326 *P. solimoense* growing at 12 sites in French Guiana; each individual was marked, and its developmental stage noted. Seven sites were situated along Route N°1 (RN1) between kilometeric points 87 and 97 (219 plant individuals) and the five others were located along the road leading to the Petit Saut dam (107 plant individuals). The sites along RN1 were situated at the limit between the secondary forest and the savanna, while those along the dam road were at the edge of the rain forest. All of these populations were established from hemi-epiphytic plants growing on trees cut down during the construction of the road in 1989. Due to long-lived leaves (at least 3 y;

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**Figure 1.** Flowering in *Philodendron solimoense*. At the top of the stem, several inflorescences developing at the base of the leaves coming out of their petiolar sheath (a). Note below dry peduncles of fallen inflorescences from previous flowering events (white arrows). Detail of a dry peduncle (P) inserted within the petiolar sheath forming a petiolar cavity (PC) after the inflorescence fell off (b). Dry peduncle (P) removed from the petiolar cavity (PC) (c).

M. Gibernau, pers. obs.) the presence of the hollow, dry peduncles (Figure 1b) indicates that the plant has already flowered.

Each individual plant, as well as every petiolar sheath, was screened 2–6 times at different hours of the day, and ants were collected from the leaves. Voucher specimens were deposited in the CPDC collection (Laboratório de Mirmecologia, Cocoa Research Center, Itabuna, Bahia, Brazil) after identification. We distinguished three groups of ant species. (1) Visitors (i.e. non-resident) only represented by workers patrolling on the plant in order to gather EFN; (2) inhabitants with colonies located in carton pavilions built on the trunk, under dead leaves fallen from large trees, or in dry bracts trapped by lower petioles; and (3) colonies nesting in the petiole and/or peduncle cavities with queen(s), workers and brood.

Using an analysis of variance and a binomial error (GLIM 3.77 software, 1986; Royal Statistical Society, London), we analysed the frequency at which *P. solimoense* was inhabited by ants according to the habitat, the plant's developmental stage, or both factors combined. The number of inhabiting and visiting ant species (count data) per plant was analysed in the same way using a Poisson error (GLIM 3.77 software). The covariate for the number of inhabiting ant species was the logarithm(number of petiole cavity + 1) on each plant individual, since when the number of cavities increases, the number of potential nesting sites for ant species also increases. The covariate for the number of visiting ant

species was the logarithm(number of inhabiting species + 1), as inhabiting ants may become locally dominant and partly exclude visitor ants.

Two developmental stages were distinguished: immature individuals had not yet flowered and did not yet have cavities, and their petiolar sheaths are closed; whereas mature individuals bore several cavities from previous flowerings (Figure 1b).

The proportion of mature individuals was similar in the two habitats: 58% of all *Philodendron*, but the proportion of flowering individuals differed greatly between habitats (Table 1). On average, a plant had  $5 \pm 3.4$  (mean  $\pm$  SD) petiolar sheaths (e.g. cavities available for ants). More than 48 ant species were found on the 326 studied *P. solimoense*, with 27 species nesting on the plants, and more than 40 diurnal ant species were visitors (Appendix 1).

The frequency of ant-visited plants was not affected by the combination of habitat and developmental

**Table 1.** Total number of studied *Philodendron solimoense*, number of individuals inhabited by ants and number of *P. solimoense* belonging to each developmental stage: immature and mature individuals in the two studied habitats (along the Route N°1 and the road leading to the Petit Saut dam). The last column indicates the number of mature individuals which were flowering during the study.

Habitat	Total	Inhabited	Immature	Mature	Flowering
Route N°1	219	114	93	126	87
Dam road	107	62	45	62	14

**Table 2.** Frequencies ( $\pm$  SD) of *Philodendron solimoense* visited by ants according to its habitat (along the Route N°1 or the road leading to the Petit Saut dam) and its developmental stage (immature or mature individuals).

Habitat	Immature	Mature
Route N°1	0.96 $\pm$ 0.20	0.86 $\pm$ 0.34
Dam road	0.78 $\pm$ 0.42	0.69 $\pm$ 0.47

stage ( $\chi^2_1 = 1.3$ ;  $P = 0.25$ ), but this frequency varied significantly according to the developmental stage alone ( $\chi^2_1 = 5.4$ ;  $P = 0.02$ ) and between the two habitats ( $\chi^2_1 = 16.2$ ;  $P < 0.001$ ; Table 2).

The number of ant species visiting a plant was not affected by the combination of habitat and developmental stage ( $\chi^2_1 = 0.02$ ;  $P = 0.88$ ), nor the developmental stage alone ( $\chi^2_1 = 0.35$ ;  $P = 0.55$ ) and was negatively correlated with the number of inhabiting ant species (slope =  $-0.21$ ,  $\chi^2_1 = 4.5$ ,  $P = 0.034$ ). On the other hand, *Philodendron* plants along RN1 were visited by twice as many ant species as those growing along the dam road ( $2.3 \pm 1.4$  versus  $1.2 \pm 1$ ;  $\chi^2_1 = 46.4$ ;  $P < 0.001$ ).

The frequency of plant inhabited by ants was not affected by the combination of habitat and developmental stage ( $\chi^2_1 = 1.7$ ;  $P = 0.19$ ), nor the habitat alone ( $\chi^2_1 = 1.2$ ;  $P = 0.27$ ). Yet the frequency of plants inhabited by ants differed significantly according to the developmental stage alone ( $\chi^2_1 = 118$ ;  $P < 0.001$ ). Immature individuals of *P. solimoense* were less frequently inhabited than mature individuals (ratios:  $0.20 \pm 0.40$  vs.  $0.80 \pm 0.41$ ). In fact, ant colonies were rarely present on immature individuals. When present, they nested only in pavilions, under dead leaves or in dry bracts since the cavities formed by petiolar sheaths were not yet available. In contrast, mature individuals of *P. solimoense* were inhabited by ant colonies nesting in their dry hollow peduncles and in their petiolar sheaths.

The number of ant species inhabiting a plant was not affected by the combination of habitat and developmental stage ( $\chi^2_1 = 1.9$ ;  $P = 0.17$ ), nor of the habitat alone ( $\chi^2_1 = 0.02$ ,  $P = 0.88$ ). Yet, the number of ant species inhabiting a plant varied significantly according to the developmental stage alone ( $\chi^2_1 = 16.2$ ,  $P < 0.001$ ;  $0.23 \pm 0.49$  for immature versus  $1.44 \pm 0.64$  for mature individuals) and increased with the number of petiolar cavities present on the plant (slope =  $0.10$ ;  $\chi^2_1 = 4.41$ ;  $P = 0.036$ ). Interestingly the ant species found in pavilions, under dead leaves or in dry bracts on immature plants were also able to colonize the petiolar sheaths of mature individuals.

Finally, the *P. solimoense*–ant interaction appeared to be non-specific and to be mainly established with ant species present in the forest edge. However, there were few species (1–2) nesting on each plant (Appendix 1).

One important factor in the ant–plant interaction is the habitat or the ecological zone (King *et al.* 1998,

Majer *et al.* 1997, Vasconcelos 1999). In fact, a greater diversity in ant species was found along RN1 than along the dam road. Firstly, the *P. solimoense* located along RN1 grew in more open habitats than those growing in the shade of pioneer trees in the forest edges along the dam road. Consequently, *P. solimoense* individuals along RN1 were exposed to a greater light intensity than the other populations. This light exposure favours flowering in *P. solimoense* (unpubl. data). Hence, there was a greater number of flowering specimens growing along RN1 (40%) than in populations growing along the road leading to the dam (13%), while the same proportion (58%) of individuals were mature at both sites. A population of *P. solimoense* with a greater number of flowering individuals may attract a greater number of ant species (RN1 vs. dam road) due to the EFN produced on the spathes.

Immature and first-flowering individuals of *Philodendron solimoense* are visited by workers from various opportunistic ant species looking for EFN or searching for prey (Appendix 1). This is considered to be a diffuse relationship. On the contrary, ant colonies generally occupied several petiolar cavities on most mature individuals, which can be considered to be a stable association as they provide ants with a permanent residence (see also Way & Bolton 1997 for coconut palms). Moreover, their presence was associated with fewer visitor ants, probably because their territoriality leads to the exclusion of the foraging workers of other species.

Such colonies can remain on a plant for a long time due to long-lived leaves and because new cavities appear at each flowering. Flowering thus appears to be an important factor in the relationships between *P. solimoense* and ants. Consequently, *P. solimoense* falls somewhere between plants attracting ants with EFN and/or FBs (Dejean *et al.* 2007, Heil 2008, Heil & McKey 2003) and true myrmecophytes (Hölldobler & Wilson 1990, Janzen 1966). Hence *P. solimoense* is not a myrmecophyte as its petiolar sheaths, non-specialized structures linked with flowering, are not real domatia. Ant-domatia have been defined by Beattie & Hughes (2002) as ‘plant structures that appear to be specific adaptations for ant occupation, often formed by the hypertrophy of internal tissue at particular locations in the plant, creating internal cavities attractive to ants’. Thus, the interaction between *P. solimoense* and ants is in between a non-specific diffuse relationship and a specific obligatory interaction, and still needs to be properly defined.

Further studies are needed to prove the positive impact ant presence has on this plant to determine if the relationship is a true mutualism. The presence of petiolar cavities resulting from flowering is a general characteristic in the genus *Philodendron*. This suggests that there are numerous possibilities for this genus and

ants to interact. Moreover the cavity present within the petiole is also a common trait in Araceae, but varies in size and degree of inclination based on the genus and, thus, can result in the selection of different kinds of inhabiting ant species. It would be interesting to determine if these cavities promote ant–plant interactions in this plant family, as well as in other monocotyledon families.

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## APPENDICES

Appendices are available as supplementary material on-line at <http://journals.cambridge.org/tro>