Congeneric mutualist ant symbionts (Tetraponera, Pseudomyrmecinae) differ in level of protection of their myrmecophyte hosts (Barteria, Passifloraceae)

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Congeneric mutualist ant symbionts (*Tetraponera*, Pseudomyrmecinae) differ in level of protection of their myrmecophyte hosts (*Barteria*, Passifloraceae)

**Running title:** Differing plant protection by congeneric plant-ants

**Keywords:** ant–plant symbiosis, chemical mediation, defence, protection mutualism, specificity

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Abstract

*Barteria fistulosa* and *B. dewevrei*, central African rain-forest trees, provide nesting cavities for *Tetraponera aethiops* and *T. latifrons* ants, respectively, which protect them against herbivores. To compare protection efficiency between these two symbioses, for 20 plants of each species in two sites in Gabon we measured the time elapsed before ants reached a focal leaf, for host leaves that were undisturbed, damaged (cut with scissors) or subjected to slight vibration (mimicking such damage), and for damaged leaves of the non–host *Barteria* species. *Tetraponera aethiops* displayed stronger protective behaviour than did *T. latifrons*. Time to reach a damaged host leaf (4.5 ± 2.6 min, mean ± SD) did not differ significantly from time to reach a leaf subjected to slight vibration (5.2 ± 3.0 min) for *T. aethiops*, but response to a leaf subjected to slight vibration (9.5 ± 1.9 min) was significantly slower than that to a damaged leaf (7.8 ± 1.9 min) for *T. latifrons*. The faster response of *T. aethiops* to slight vibration may have masked a response of this species to chemical signalling. Both ants reached damaged host leaves faster than damaged leaves of the non–host *Barteria* sp., indicating host plant specificity in ant responses.
Introduction

Ant–plant interactions, particularly protection mutualisms (Heil & McKey 2003), are widespread in tropical rain forests. In addition to numerous non-symbiotic interactions mediated by plant-provided food sources such as extrafloral nectar (Heil 2015), symbiotic mutualisms also occur. In these, myrmecophytic plants provide nesting space in the form of hollow twigs, petioles, stipules, leaf pouches, etc.—more or less specialized structures termed domatia—and food in the form of extrafloral nectar, food bodies, secretions of plant-nourished hemipteran trophobionts, or combinations of these, to their host ants (Davidson & McKey 1993, McKey et al. 2005). In return, ants defend their host plant against herbivores and pathogens (Letourneau 1998, Rosumek et al. 2009). About 700 plant and 110 ant species are involved in this type of symbiosis (Chomicki & Renner 2015, Davidson & McKey 1993). In some cases, the interaction is obligatory for one or even both partners (Gaume & McKey 1999, Moog 2009, Yu & Davidson 1997).

Protection of myrmecophytic plants by ants has been thoroughly studied (Janzen 1966, Rosumek et al. 2009). Like many other plants, myrmecophytic plants release volatile compounds, to which their resident ants are attracted (reviewed in Blatrix & Mayer 2010). Whereas the role of chemical signalling in ant–plant symbioses has been demonstrated by several studies, a smaller number of studies show that vibrations produced by herbivore movement also induce ant patrolling (Dejean et al. 2008, 2009, Federle et al. 1998, Lapola et al. 2003, Madden & Young 1992).

Protection level often differs among systems depending on the capacity of the ants to counter plant enemies (Bruna et al. 2004, Lapola et al. 2003). This capacity may
partly depend on colony size (Duarte Rocha & Godoy Bergallo 1992), which in turn depends strongly on plant investment in the resident ant colony. Both protective behaviour and the level of plant investment in ants are driven by coevolution. However, studies comparing the efficiency with which different ants protect the same plant species have so far compared ants belonging to different genera that have independently evolved associations with the same host plant (Bruna et al. 2004, Lapola et al. 2003). These studies thus cannot distinguish differences that may have resulted from coevolutionary interactions with the host plant from those that already existed prior to their host associations.

In this study we compare the level of protection conferred on host plants by two closely related species of Tetraponera (Pseudomyrmecinae), *T. aethiops* Smith, 1877 and *T. latifrons* (Emery, 1912), that both live in symbiotic associations with myrmecophytes of the genus *Barteria* (Passifloraceae), suggesting that their common ancestor was also a symbiont of the plants’ common ancestor. Differences in their protective behaviour thus evolved in the context of this symbiosis. Comparison of their protective behaviour can thus shed light on differences driven by coevolutionary interactions. As association specificity seems weaker for *B. dewevrei* than for *B. fistulosa*, coevolution between *B. fistulosa* and *T. aethiops* might be more intensive than between *B. dewevrei* and *T. latifrons*. We postulated that more intensive coevolution of ant and plant may have resulted in better protection of *B. fistulosa* by its ant associate *T. aethiops*. We tested the following hypotheses: (1) protective behaviour is elicited by vibration and/or chemical signals, (2) protective behaviour of each ant species is more pronounced
toward its main host plant than toward the other *Barteria* species, (3) *T. aethiops* shows a more pronounced protective behaviour than *T. latifrons*.

**Methods**

*Study species and study sites*

The genus *Barteria* (Passifloraceae) comprises four tree species (Breteler 1999) endemic to central Africa. *Barteria solida* Breteler does not host ants. *Barteria fistulosa* Mast., *B. dewevrei* De Wild. & T. Durand and *B. nigritana* Hook. f. are myrmecophytes with hollow lateral branches. *Barteria nigritana* hosts various opportunistic species of ants, some of which provide some protection to their host (Djiéto-Lordon *et al.* 2004).

*Barteria dewevrei* and *B. fistulosa* have three potential ant symbionts: *Tetraponera aethiops*, *T. latifrons* and *Crematogaster* sp. However, *B. dewevrei* preferentially hosts *T. latifrons* or *Crematogaster* sp., whereas *B. fistulosa* hosts *T. aethiops* preferentially, even when both *Barteria* and both *Tetraponera* species occur in syntopy (Kokolo *et al.* 2016).

*Barteria fistulosa* is restricted to the tropical rain-forest environment, whereas *B. dewevrei* occurs in a wider range of habitats: rain forest, forest gallery and the forest/savanna ecotone. Protection of *B. dewevrei* by its ants has never been studied. The vigorous protection of *B. fistulosa* by *T. aethiops* is well-known by local people and has been the subject of several studies (Dejean *et al.* 2008, Janzen 1972, McKey 1974). The association between the two *Barteria* and their two *Tetraponera* ants shows a high degree
of specificity: *Tetraponera aethiops* and *T. latifrons* have never been found nesting outside either *B. fistulosa* or *B. dewevrei*. Moreover, *B. fistulosa* does not grow well if not occupied by one or the other of these two *Tetraponera* species. *Barteria dewevrei* seems to be less dependent upon *Tetraponera* ants, at least in some habitats, as many individual trees in some savanna sites are occupied by *Crematogaster*.

The experiments were conducted on 20 individuals of *B. dewevrei* occupied by *T. latifrons* in Souba (01°36’S, 14°03’E) and 20 individuals of *B. fistulosa* occupied by *T. aethiops* in Bongoville (1°37’S; 13°54’E). Souba is located at ~550 m asl on the Batéké Plateau, a highland covered by grassland and wooded savanna, criss-crossed by gallery forest. Bongoville is located at the foot of the Batéké Plateau, at ~400 m asl, along a stream in a tropical rain-forest environment.

*Behavioural response of the ants to host plant disturbance*

Behavioural tests were conducted in three steps. The first step aimed at measuring the spontaneous level of ant patrolling activity on leaves, i.e. without any disturbance of the tree, as a control. Two leaves were randomly selected among the young leaves of two upper branches on each *Barteria* individual (one leaf per branch). Starting at an arbitrary time, we recorded the time lapse before each leaf was reached by a resident ant, once per leaf. We averaged the time lapse over the two leaves for each tree. This average was meant to temper the effect of heterogeneity expected with low levels of patrolling, and was used as a single value for each tree (undisturbed leaf = control leaf). The second step aimed at testing the effect on ant reaction of physical damage to a leaf
(simulating an attack by a small herbivore, such as a phytophagous insect) and of slight vibration, which inevitably accompanies such experimental infliction of damage. Two leaves were randomly selected among the young leaves of two upper branches on each Barteria individual (one leaf per branch). The apex of one of the two leaves was cut with scissors (damaged leaf), and the other leaf was simply gently scraped superficially with the tip of scissors without inflicting damage (‘scraped leaf’). For each of the two leaves simultaneously we recorded the time lapse before it was reached by a resident ant, and the number of ants that touched the site of damage (damaged leaf only) over 30 min. The two focal ant species, *T. latifrons* and *T. aethiops*, are very sensitive to physical disturbance of their host tree by an experimenter. They literally pour out of the hollow branches as soon as an experimenter manipulates, or even touches, the leaves. Any vibration, even weak, such as cutting a leaf with scissors, may thus induce such a reaction. Damaging the leaf necessarily implies also provoking vibrations. The level of vibration we applied to the leaf by scraping it with the tip of the scissors was meant to mimic that experienced by the leaf damaged with scissors. We recognize that scraping a leaf does not exactly reproduce the same physical disturbance caused by cutting the leaf blade with scissors. However, slightly scraping the leaf was the best proxy we could use to control for a possible effect of the physical disturbance of cutting with scissors to test for the effect of emission of volatile compounds after such damage. Thus, the effect of damaging the leaf was tested by comparing the scraped and the damaged leaves (because the damage treatment corresponds to vibration + physical damage). The effect of scraping the leaf was tested by comparing the undisturbed and the scraped leaves. The third step aimed at testing specificity of the behavioural response of the ant species
to physical damage of the two plant species. On a leaf of each *Barteria* individual we placed a damaged leaf (apex cut with scissors) of the other *Barteria* species and recorded the time lapse before it was contacted by a resident ant. The host leaf upon which the damaged non-host leaf was placed was gently scraped to mimic the level of vibration of a leaf damaged with scissors, so we could compare time lapse of response to the damaged host leaf and to the damaged non-host leaf. We also compared responses of the two ant species to each treatment.

Statistical tests were performed with R 3.1.0. P-values were adjusted using Holm’s method in the case of multiple comparisons.

**Results**

The time lapse before the ants reached the focal leaves varied significantly across treatments (host leaf undisturbed, host leaf scraped, host leaf damaged, non–host leaf damaged) for *T. aethiops* (Friedman test, statistic = 38, $P = 0.000000036$) and for *T. latifrons* (Friedman test, statistic = 36, $P = 0.000000050$).

*Tetraponera latifrons* ants reached the scraped host leaf significantly more quickly than the undisturbed leaf (mean ± SD, scraped: 9.5 ± 1.9 min, undisturbed: 16.3 ± 5.7 min; Wilcoxon signed-rank test, $V = 8$, $P_{adj} = 0.0025$, Figure 1), the damaged host leaf significantly more quickly than the scraped host leaf (damaged: 7.8 ± 1.9 min, scraped: 9.5 ± 1.9 min; $V = 96$, $P_{adj} = 0.020$, Figure 1), and the damaged host leaf (*B. dewevrei*) significantly more quickly than the damaged non–host leaf (*B. fistulosa*)
(damaged host: 7.8 ± 1.9 min, damaged non-host: 10.8 ± 2.1 min; V = 171, $P_{adj} = 0.0014$, Figure 1). Moreover, the number of ants of this species that touched the damaged site was significantly greater for the host leaf ($B. dewevrei$) than for the non–host leaf ($B. fistulos$) (damaged host: 9.1 ± 3.5 ants, damaged non-host: 6.6 ± 2.1 ants; $V = 152, P_{adj} = 0.045$, Figure 1).

*Tetraponera aethiops* ants reached the scraped host leaf significantly more quickly than the undisturbed leaf (mean ± SD, scraped: 5.2 ± 3.0 min, undisturbed: 16.6 ± 5.3 min; Wilcoxon signed-rank test, $V = 1, P_{adj} = 0.00087$, Figure 1). The time lapse before *T. aethiops* ants reached the host leaf did not differ significantly between scraped and damaged leaves (scraped: 5.2 ± 3.0 min, damaged: 4.5 ± 2.6 min; $V = 114, P_{adj} = 0.46$, Figure 1). *Tetraponera aethiops* ants reached the damaged host leaf ($B. fistulosa$) significantly more quickly than the damaged non–host leaf ($B. dewevrei$) (damaged host: 4.5 ± 2.6 min, damaged non-host: 7.15 ± 2.5 min; $V = 178, P_{adj} = 0.0036$, Figure 1). Moreover, the number of ants of this species that touched the damaged site was significantly greater for the host leaf ($B. fistulosa$) than the non–host leaf ($B. dewevrei$) (damaged host: 15.2 ± 4.0 ants, damaged non-host: 13.3 ± 3.3 ants; $V = 162, P_{adj} = 0.045$, Figure 1).

The time lapse before the ants reached the undisturbed leaf did not differ significantly between the two ant species (mean ± SD, *T. latifrons*: 16.3 ± 5.7 min, *T. aethiops*: 16.6 ± 5.3 min; Mann–Whitney U-test, $W = 191, P_{adj} = 0.81$), indicating no significant difference in the spontaneous level of patrolling activity. *Tetraponera aethiops* ants reached their host leaf ($B. fistulosa$) significantly more rapidly than *T.
*latifrons* reached theirs (*B. dewevrei*), whether leaves were simply scraped (*T. aethiops*: 5.2 ± 3.0 min, *T. latifrons*: 9.5 ± 1.9 min; $W = 344, P_{adj} = 0.00084$) or damaged (*T. aethiops*: 4.5 ± 2.6 min, *T. latifrons*: 7.8 ± 1.9 min; $W = 336, P_{adj} = 0.0014$). Moreover, the number of *T. aethiops* ants that contacted the damaged host leaf during 30 min was significantly larger than for *T. latifrons* (*T. aethiops*: 15.2 ± 4.0 ants, *T. latifrons*: 9.1 ± 3.5 ants; $W = 345, P_{adj} = 0.0000029$). *Tetraponera aethiops* ants were significantly faster and more numerous than *T. latifrons* in contacting the damaged non–host leaf (*T. aethiops* on *B. dewevrei*: 7.15 ± 2.5 min, 13.3 ± 3.3 ants; *T. latifrons* on *B. fistulosa*: 10.8 ± 2.1 min, 6.6 ± 2.1 ants; time lapse: $W = 55, P_{adj} = 0.00078$; number of ants: $W = 383, P_{adj} = 0.0000029$).

**Discussion**

*Behavioural response of Tetraponera ants to disturbance of their Barteria host plants*

Most ants associated with myrmecophytes respond to leaf damage in less than five minutes (reviewed in Blatrix & Mayer 2010). The time lapse before *T. aethiops* and *T. latifrons* reached a damaged leaf appeared comparatively long (4.5 ± 2.6 and 7.8 ± 1.9 min, respectively), when we consider that the *Tetraponera* ants associated with *Barteria* are well known for their particularly prompt aggressive reaction to human intruders (Janzen 1972). As we were aware of this particularity, we took special care to limit physical disturbance as much as possible when manipulating experimental leaves (“scraped” and “damaged” treatments). The treatments we applied were probably more representative of the disturbance induced by small insects than that induced by
mammals. Comparing the African Barteria/Tetraponera symbiosis with the American Acacia/Pseudomyrmex symbiosis, Janzen (1972) noted that the protection behaviour displayed by Tetraponera aethiops is particularly adapted to protect its host plant against large herbivores such as mammals. For instance, the number of workers of Tetraponera aethiops patrolling Barteria fistulosa is much lower than in the Acacia/Pseudomyrmex symbiosis. The relatively slow reaction time measured in our experiments with Tetraponera on Barteria likely reflects the low spontaneous patrolling activity of these ants and their weaker reaction to insect herbivory than to mammal herbivory.

Our experiments showed that both ant species, T. latifrons and T. aethiops, reached the leaves more rapidly when these were scraped than when they were undisturbed, demonstrating their response to vibrations. Dejean et al. (2008) already showed that T. aethiops responds to vibration induced by herbivory. However, only T. latifrons responded more strongly to physical damage than to scraping. Behavioural response of T. aethiops did not differ between a scraped and a damaged (thus also vibrated) leaf of its host plant, B. fistulosa, but T. aethiops proved to react more rapidly than T. latifrons in both cases. This difference between the two ants cannot be due to varying levels of spontaneous patrolling, as these did not differ significantly. These results suggest that perception of chemical signals (volatile compounds emitted from the damaged site) improves protection of B. dewevrei by its host ant T. latifrons. Given that damaged leaves were also vibrated by the investigator and that T. aethiops reacts very strongly to scraping, the effect of chemical signals on T. aethiops may have been masked by the effect of scraping in our experimental design. Thus, we cannot conclude
on the effect of chemical signalling on *T. aethiops*. We hypothesize that the responses of *T. aethiops* to chemical signals and vibration could have evolved to target respectively small insects feeding on the leaves (which are unlikely to induce detectable vibration) and large herbivores such as mammals. Response of *T. aethiops* to chemical signals should be tested in a vibration-free context. Similar chemical signalling of leaf damage has been demonstrated in many ant–plant symbioses (reviewed in Blatrix & Mayer 2010). Perception of chemical signals indicating herbivory should improve protection efficiency because it allows the ants to locate sites of herbivory quickly and accurately (Agrawal 1998, Gonçalves-Souza 2016, Pacheco & Del Claro 2018). In various myrmecophytic interactions, ant patrolling is focused on young leaves and shoots, even when these are not disturbed (reviewed in Blatrix & Mayer 2010). Such directed patrolling is most likely driven by chemical signalling to provide constitutive protection of the most vulnerable plant parts. Plant chemical signalling of herbivore damage and ant behavioural response to it are expected to coevolve because both partners benefit from deterrence of herbivores.

**Specificity of ant behavioural response to host plant species**

Our study revealed a certain level of specificity in the behavioural response of the two ant species to disturbance of the plant. We showed that both ant species, *T. aethiops* and *T. latifrons*, responded more strongly (faster recruitment and more ants recruited) to damage to leaves of their usual host plant species (*B. fistulosa* and *B. dewevrei* for *T. aethiops* and *T. latifrons*, respectively) than to damage on leaves of the non–host plant
species (*B. dewevrei* and *B. fistulosa* for *T. aethiops* and *T. latifrons*, respectively). As there is a preferential association between *T. aethiops* and *B. fistulosa* on the one hand and between *T. latifrons* and *B. dewevrei* on the other hand (Kokolo et al. 2016), specificity of the behavioural response might be a product of ant–plant coevolution within each pair of species. Alternatively, specificity of ant response in the *Barteria–Tetraponera* system might result from ants acquiring familiarity with the signals of the plant species they presently live on. For instance, arboreal ants have been shown to nest preferentially on plant species to which they have been exposed during development (Djiéto-Lordon & Dejean 1999), probably because of a pre–imaginal imprinting process directed on plant odour.

Notwithstanding the specific responses of the ants to damage of their host plant, we showed that *T. aethiops* responded more strongly to leaf disturbance (scraping and damage), a signal of potential herbivory, than *T. latifrons*. This difference is expected to translate into better protection conferred by *T. aethiops* than by *T. latifrons*. As the association between *T. aethiops* and *B. fistulosa* is more specific than the association between *T. latifrons* and *B. dewevrei* (Janzen 1972, Kokolo et al. 2016), we speculate that the difference in response level might be the result of a tighter coevolution between *T. aethiops* and *B. fistulosa*. *Tetraponera aethiops* and *T. latifrons* are sister species (Ward 1991) and are the only African species of their species-group (Ward 2006). The differences we observed in their protective behaviour have thus likely evolved after their divergence from a common ancestor that was also in symbiosis with *Barteria*. This situation is different from those studied by Bruna et al. (2004) and Lapola et al. (2003), where comparisons involved phylogenetically distant ant species that
independently colonized the same plant. The differences between the two species of *Tetraponera* much more likely reflect coevolution with their *Barteria* hosts. From a mechanistic point of view, our study demonstrates that these differences are not due to different levels of spontaneous patrolling activity. However, the shorter time lag before reaching the disturbed leaves and the larger number of ants recruited for *T. aethiops* than for *T. latifrons* could result from colonies of *T. aethiops* being larger, leading to more ants involved in defending the plant. It would be worth investigating the level of food rewards provided by the two *Barteria* species to their ant symbionts to test the hypothesis that *B. fistulosa* invests more resources into its symbiont to sustain a higher level of protection, and to see whether such a difference has translated into larger, denser colonies of *T. aethiops* in its host than of *T. latifrons* in its host.

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Figure caption

Figure 1. Behavioural response of the ants *Tetraponera aethiops* and *T. latifrons* to treatments on leaves of their host plants, *Barteria fistulosa* and *B. dewevrei*, expressed as differences in time lapse (“T”, min) before the ants reached the focal leaves and in number of ants (“N”) that touched the damaged site. Treatments were tested in a paired design as follows: host leaf undisturbed vs host leaf scraped with the tip of scissors, host leaf scraped vs host leaf damaged, non-host leaf damaged vs host leaf damaged. Damage was performed with scissors, and thus damaged leaves were inevitably also subjected to slight vibration that we mimicked by gently scraping the leaf with the tip of scissors. Experiments were conducted in Gabon. Vertical lines represent median, boxes represent interquartile range, and whiskers extend to the data extremes. n.s.: not significant, *: P < 0.05, **: P < 0.01, ***: P < 0.001.
Figure 1

 Difference in time lapse of ant reaction (T) or in number of ants recruited (N) between leaf treatments.