Widespread army ant aversion among East African jumping spiders (Salticidae)

Ximena J. Nelson · Samuel Aguilar-Arguello · Robert R. Jackson

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Abstract
Jumping spiders (Salticidae) typically prey on a variety of arthropods of similar size to themselves, but rarely on ants. Using 28 salticid species from East Africa, we first investigated vision-based aversion to ants by recording latency to enter a transparent sealed chamber flanked by chambers containing living army ants (Dorylus sp.) or tsetse flies (Glossina pallidipes) of comparable size. For all species, entry latency was significantly longer when the stimuli were ants. In another experiment, we used dead ants and tsetse flies mounted in a life-like posture as stimuli; except for Goleba puella, a species with unusual retinal ultrastructure, we again found significantly longer entry latency when the stimuli were ants. Our findings imply that these salticids express an aversion specifically to ants even when restricted to using vision alone and, except for G. puella, even when relying on solely the static appearance of the insects. Having used salticids from laboratory cultures with no prior experience with ants, our findings are consistent with vision-based aversion to army ants being innate.

Keywords Object discrimination · Vision · Formicidae · Movement cues · Innate aversion · Myrmecomorphy

Introduction
Despite a tendency to think of insectivorous birds as the primary predators of arthropods (e.g., Gunnarsson 2007; Gunnarsson and Wiklander 2015), through sheer biomass it seems likely that the dominant predators of arthropods are actually other arthropods. For example, in temperate regions (King et al. 2013), and especially in the tropics (Davidson 1997; Davidson et al. 2003), there is evidence that the biomass of ants, which can constitute over 80% of the total biomass of arthropods, exceeds that of all vertebrates combined. Many ants also have characteristics that make them particularly dangerous for similar-sized arthropods (Sanders and Platner 2007; Mestre et al. 2012; Ramesh et al. 2016), including formidable weapons such as powerful mandibles and potent venom, along with a capacity for mobbing attacks on would-be predators and prey (Eisner 1970; Hölldobler and Wilson 1990). In response to selection pressures from particular predators, prey species may acquire predator-specific defenses. Knowing that ants can have strong impacts on the abundance of other arthropods (Halaj et al. 1997; Piñol et al. 2012) suggests that it might also be common for ant-vulnerable arthropods to acquire, by natural selection, acute capacity to detect, identify and then avoid ants.

Jumping spiders (family Salticidae) provide especially interesting case studies of how other arthropods adapt to ants (Huang et al. 2011; Nelson and Jackson 2011a, 2014; Jackson and Nelson 2012). Salticidae is the largest family of spiders (more than 6100 described species in 640 genera) and, like ants, a major predatory group which has diversified especially in the tropics (Maddison 2015; Prószyński 2017; Platnick 2019). However, the most distinctive characteristic of salticids is their extraordinarily good eyesight.

The salticid visual system consists of a pair of large forward-facing ‘principal’ eyes and three pairs of smaller eyes, collectively called the secondary eyes, situated to the side and behind the principal eyes. To discern visual detail pertaining to the shape and form of objects, an animal needs especially good spatial acuity (Land and Nilsson 2012) and...
salticids stand out by having the best spatial acuity known for terrestrial arthropods (Goté et al. 2019). To achieve this, the salticid relies primarily on its principal eyes (Land 1969; Blest et al. 1990), with the secondary eyes serving other functions, such as motion-detection (Land 1972; Zurek et al. 2010; Zurek and Nelson 2012; Jakob et al. 2018). This visual system underpins some of the most intricate vision-based predatory strategies known for animals in general (Nelson and Jackson 2011b) and enables salticids to discriminate between different kinds of prey at a distance, even in the absence of cues from other sensory modalities (Harland et al. 1999; Nelson and Jackson 2012; Cerveira et al. 2019).

That encounters with ants tend to be exceptionally dangerous for salticids is the rationale for a hypothesis that, for salticids, proficiency at identifying ants from a distance is dangerous to terrestrial arthropods (Huseynov et al. 2008; Schöning et al. 2008), and determines the effect of movement on the potential aversion that they cause to a wide variety of salticid species. Thus, here we consider whether aversion to the static appearance of army ants specifically might also be considerable variation between ant species, but candidates might include the ant’s characteristically narrow body form, a slender pedicel where the thorax joins the abdomen, lack of wings, and thin, bare, antennae (Nelson 2010). However, instead of considering salticid ability to generalize the characteristics of ants across multiple species, here our objective was to consider only one group of ants (Dorylus sp. army ants, also known as driver ants), that are well-known to be dangerous to terrestrial arthropods (Huseynov et al. 2008; Schöning et al. 2008), and determine the effect of movement on the potential aversion that they cause to a wide variety of salticid species. Thus, here we consider whether aversion to the static appearance of army ants specifically might also be prevalent in a different salticid fauna found in East Africa, where army ants are a keystone species (Schöning et al. 2008). Here, in addition to testing with stationary mounts, we also tested salticids with live ants. This is of interest because, besides their distinctive static appearance, ants have an erratic style of locomotion, characterized by quick, fast-paced changes in direction (Shamble et al. 2017). There are a significant minority of spiders, including salticids, that specialize on eating ants and have a prey preference for ants (Jackson and Nelson 2012; Pekár and Toft 2015), and there is recent evidence that an ant-like style of moving is salient to ant-eating as well as ant-averse salticids (Nelson and Jackson 2014; Nelson and Card 2016; Shamble et al. 2017).

**Materials and methods**

Experiments were carried out in Mbita Point, Western Kenya, between 800 and 1400 h (laboratory photoperiod 12L:12D, lights on 700 h). For test spiders, we used 28 salticid species (Table 1) from laboratory cultures (F2 or F3 generation). The cultures for Hyllus deyrollei, Parajotus cinereus, and Schenkelia modesta originated from Uganda; all others originated from Kenya. Spiders were housed individually in cages made from 250 ml transparent plastic containers with water available through a cotton wick submerged in water which protruded into the container. Each spider was kept on a mixed diet of mosquitoes (Anopheles gambiae), from an insectary, and non-biting midges (Chao-boridiae and Chironomidae), collected from the field. As Cyba ocellata, Holocalaethis vellarea, Meleon solitaria, and Portia africana are known to prefer spiders to insects as prey, we included field-collected spiders in their diets (Argyrodes spp., Oecobius amboseli, Tetragnatha spp.). For more information about maintenance procedures, see Cerveira and Jackson (2013). Our testing apparatus consisted of a stack of three glass chambers (Fig. 1): a ‘testing chamber’ in the middle with a ‘top stimulus chamber’ above and a ‘bottom stimulus chamber’ below. There was also a glass ‘holding chamber’, which housed the spider at the beginning of each trial and opened into the testing chamber. The 5-mm walls of all chambers were opaque, with their 2-mm tops and bottoms being transparent. For each chamber, the top was removable. During each trial, the spider could move out of the holding chamber into the testing chamber. Because the chambers were airtight, the spider could see, but not smell or touch, insects which were in the stimulus chambers. The insects used during a trial were either army ants (Dorylus sp.), which, due to their swarm-raiding foraging style, are known to be especially dangerous to salticids and other arthropods inhabiting leaf-litter (Huseynov et al. 2008; Schöning et al. 2008), or tsetse flies (Glossina pallides), which are comparatively harmless. The ants were collected as needed from the field and the flies were from laboratory culture (body length of ants 9–11 mm; body length of flies 12 mm).

The holding chamber had a ‘resting area’ and an ‘entry area’ (Fig. 1). Before a trial began, the spider was taken into a glass tube (diameter 8 mm; length 45 mm) plugged with stoppers. With the spider quiescent in the tube, we removed the lid of the holding chamber and the stopper from the end of tube farthest from the spider’s location within. The tube was then laid on the resting area of the holding chamber with
the open end facing the entry area, and the other end against the distal end of the resting area. Next, we closed the holding chamber lid and the spider was now free to move out of the tube and into the entry area.

Trials began when the test spider went into the entry area and ended when it entered the testing chamber. The elapsed time between arriving in the entry area and entering the testing chamber was the spider’s ‘entry latency’. The apparatus was specifically designed to investigate the salticid’s disinclination to being in the company of army ants. This is why we had a holding chamber to the side of the apparatus that forced the salticids to move into the wider ‘entry area’ if they wanted to venture into the vicinity of the ants (Fig. 1). We used entry latency as the metric that expressed the level to which spiders were deterred by what they saw in the stimulus chambers, with longer latencies implying stronger aversion. Once a trial began, the spider was allowed 15 min to enter the testing chamber. Whenever a test spider began a trial and then failed to enter the testing chamber within the allowed 15-min interval, we recorded an entry latency of 900 s. Testing was dismissed on the rare occasions when spiders failed to enter the entry area within 15 min. Sample sizes for each test are stated in Table 1.

As our objective was to detect evidence of innate aversion to army ants, no spiders were tested twice and no test spiders or their parents had prior experience with ants or tsetse flies. All spiders were of standard size (body length 4–6 mm) and about half the size of the insects used in tests (9–12 mm). For the salticid species with large adult body length, we used juveniles; otherwise, we used adult males, which are usually the smaller sex. With some variation, salticids tend to prey on arthropods that are similar in size to themselves (Jackson and Pollard 1996), suggesting that, owing to the size disparity we had between test spider and stimulus insect, we minimized the likelihood that test spider would respond to the stimuli as prey.

In each trial, we had four stimulus insects of the same type in both stimulus chambers. Depending on the experiment, these were either living individuals or else mounted dead individuals. Mounts (see Jackson et al. 2005 for details)

Table 1  Sample sizes for aversion experiments in five conditions using 28 salticid species

| Species                        | Control | Live ants | Live flies | Dead ants | Dead flies |
|--------------------------------|---------|-----------|------------|-----------|------------|
| Asemonea murphyae1             | 27      | 22        | 23         | 23        | 20         |
| Cyba ocellata2                 | 27      | 24        | 20         | 24        | 22         |
| Dendryphantes hewitti3         | 23      | 23        | 20         | 21        | 21         |
| Evarcha culicivora3            | 25      | 24        | 24         | 25        | 25         |
| Evarcha ignea3                 | 24      | 20        | 20         | 24        | 23         |
| Goleba puella1                 | 26      | 22        | 23         | 20        | 20         |
| Harmochirus bianoriformis3     | 22      | 22        | 22         | 20        | 20         |
| Hasarius adansoni3             | 20      | 20        | 20         | 22        | 22         |
| Heliophanus jacksoni3          | 23      | 25        | 23         | 25        | 25         |
| Holocolaetis vellerea2         | 23      | 23        | 23         | 24        | 23         |
| Hyllus deyroelii5              | 22      | 20        | 20         | 25        | 20         |
| Icasis mbitaensis3             | 24      | 22        | 23         | 23        | 25         |
| Meleon solitaria2              | 23      | 22        | 20         | 24        | 23         |
| Menemera congobensis3          | 25      | 20        | 23         | 25        | 24         |
| Natta horizontalis3            | 20      | 20        | 20         | 20        | 20         |
| Pachyballus cordiformis3       | 24      | 20        | 20         | 26        | 25         |
| Parajotus cinereus3            | 22      | 22        | 22         | 22        | 23         |
| Pellenes rufoclypeata3         | 22      | 20        | 20         | 23        | 22         |
| Phintella aequipes3            | 24      | 22        | 22         | 25        | 22         |
| Phlegra sp.3                   | 22      | 20        | 20         | 20        | 20         |
| Plecius auberti3               | 24      | 24        | 23         | 26        | 23         |
| Pseudicius roberti3            | 25      | 25        | 22         | 23        | 21         |
| Portia africana2               | 25      | 24        | 25         | 26        | 20         |
| Portia schulzi2                | 24      | 24        | 21         | 25        | 22         |
| Rhene sp.3                     | 25      | 24        | 23         | 23        | 20         |
| Schenkelia modesta3            | 22      | 20        | 21         | 21        | 20         |
| Thyene inflata1                | 29      | 26        | 24         | 25        | 24         |
| Tusitala byrata1               | 22      | 26        | 20         | 26        | 22         |

Subfamilies: 1Asemoneinae, 2Spartaeinae, 3Salticinae (from Maddison 2015)
were made by immobilizing the insect under carbon dioxide, placing it in 70% ethanol for 60 min, gluing it in place on a small cork disc, and then spraying it with an aerosol plastic adhesive (Crystal Clear Lacquer, Atsco Australia Pty). The four mounts were situated facing the center, about 15 mm from the edge, along two rows in each stimulus box (see Fig. 1) and were held in place with double-sided tape stuck on the underside of the cork disc. The mounts were on the bottom of the bottom stimulus chamber and on the top glass sheet of the upper stimulus chamber. There were also control trials, in which we used empty stimulus chambers, for each salticid species. If spiders can determine the morphology of an ant and correctly identify it in the absence of its characteristic movements, we would predict longer entry latencies when stationary ant mounts are visible instead of stationary tsetse flies, or empty chambers.

We used a linear mixed effects (LME) model implemented under restricted maximum likelihood (REML) in R v 3.3.3 to broadly analyze the data. As the dataset contained censored data (trials ending at 900 s) and mixed...
models cannot adequately process censored data, 900 s latencies (n = 535 out of 3170 data points) were treated as missing data and were omitted from this model (known as the Complete-Case method, Wu 2010; see Supplementary Material for the LME model applied to the complete dataset). The selected model accounted for entry latency (log-transformed) as the response variable, the five treatments (control, dead flies, live flies, dead ants, and live ants) as fixed factors, and spider species as the random factor. We then applied contrast analysis (Crawley 2007) to make specific comparisons between treatments. We used the R packages ‘nlme’ (Pinheiro et al. 2018) and ‘gmodels’ (Warnes et al. 2015) for analyses and ‘ggplot2’ (Wickham 2009) and ‘ggpubr’ (Kassambara 2018) to draw the figures.

To investigate differences between species, we used two-tailed Mann–Whitney U tests, as the data were not normally distributed. When multiple comparisons were made using the same dataset, Bonferroni corrections were applied. In the one instance where correction caused a qualitative change, this was noted. Detailed results of this analysis can be found in Table S2.

Results

Results from the LME model showed that spiders did not differ in their latency to enter the testing chamber when exposed to dead and live ants in the stimulus chambers. Spiders also did not differ in their latency to enter the testing chamber when exposed to dead and live flies, but these latencies were significantly shorter than when spiders were flanked by ants (dead or alive). The shortest latencies were when spiders were flanked by empty, control treatment, chambers (Fig. 2, Table 2).

When species-specific differences were investigated, we found the same overall pattern as above (see detailed results in Table S2). However, in these analyses, with the exception of Thyene inflata as an outlier, there was no significant difference for any salticid species in entry latency between control tests and in tests with live flies within the stimulus boxes (Table S1; Fig. 3). Using Mann–Whitney tests, we also could not distinguish between the effect of ant movement and the static appearance of dead ants, such that test spiders were averse to ants even in the absence of movement as a variable. As found in the LME model, these analyses confirmed that entry latencies were significantly longer when flanked by mounted ants as opposed to mounted flies, with the single exception of Goleba puella (Table S1). We also found no significant differences in entry latencies when flanked by active, living ants compared with stationary ant mounts for 27 of the 28 species (Table S1). Again, G. puella was the exception, having a significantly longer entry latency when flanked by living, compared with dead, ants (Fig. 1, Table S2).

Discussion

We found evidence of a strong innate aversion to army ants among East African salticids. As in other research on salticid predatory behavior (Nelson and Jackson 2011b), we ruled out prior experience by test spiders and their parents with the prey used in our experiments here by ensuring that no test spider or its parents had prior experience with ants or tsetse flies. It is on this basis that our findings support the hypothesis that these salticids express an aversion to ants that is innate with respect to experience with ants (see O’Neil 2015). We also found evidence that this was specifically a vision-based aversion because aversion to ants was evident despite the apparatus being designed to rule out the use of chemical cues.

When stimulus chambers contained living ants instead of being empty, the entry latencies of all 28 salticids were longer when entering the testing chamber with ants. While these differences were sometimes not large, these are biologically-relevant, as salticids typically only have a few seconds in which to decide whether to attack a potential prey item. Furthermore, our results make it unlikely that we were
dead ants/live effect = species. ‘Species’ as a random effect explains 2.11% of the overall variance of the data. Presented estimates and 95% confidence intervals (CI) were back-transformed from the log scale.

| Comparison               | Estimate | CI       | t   | df | P       |
|--------------------------|----------|----------|-----|----|---------|
| Dead ants/live ants      | 1.065    | 0.951–1.194 | 1.118 | 2603 | 0.263   |
| Dead ants/empty          | 2.817    | 2.554–3.107 | 21.143 | 2603 | <0.0001 |
| Dead ants/dead flies     | 2.439    | 2.208–2.695 | 17.909 | 2603 | <0.0001 |
| Dead ants/live flies     | 2.324    | 2.103–2.568 | 16.894 | 2603 | <0.0001 |
| Live ants/empty          | 2.643    | 2.376–2.941 | 18.229 | 2603 | <0.0001 |
| Live ants/dead flies     | 2.289    | 2.054–2.551 | 15.313 | 2603 | <0.0001 |
| Live ants/live flies     | 2.181    | 1.957–2.430 | 14.389 | 2603 | <0.0001 |
| Empty/dead flies         | 0.865    | 0.790–0.948 | −3.153 | 2603 | 0.001   |
| Empty/live flies         | 0.825    | 0.752–0.904 | −4.204 | 2603 | <0.0001 |
| Dead flies/live flies    | 0.952    | 0.867–1.045 | −1.039 | 2603 | 0.298   |
| Ants (dead + live)       | 2.306    | 2.143–2.483 | 22.701 | 1301.5 | <0.0001 |
| /flies (dead + live)     |          |          |      |     |         |

Response variable = latency (log(s)), fixed effect = treatment, random effect = species. ‘Species’ as a random effect explains 2.11% of the overall variance of the data. Presented estimates and 95% confidence intervals (CI) were back-transformed from the log scale.

Table 2: Results of the LME model omitting censored data (latency = 900 s)

seeing a general aversion to seeing moving ant-sized objects instead of specifically ants because, when flanked by moving flies instead of ants, entry latencies were not significantly different from those in trials with dead flies. This conclusion is further supported by finding that entry latencies of spiders surrounded by live ants were significantly longer than when surrounded by live tsetse flies. However, the fact that empty stimulus chambers caused the shortest entry latencies suggests that the mere presence of some stimulus does affect their approach behavior, possibly because they take more time to visually inspect their surroundings.

It is unlikely that our findings pertained primarily to prey-choice behavior instead of aversion to ants because, although all salticid species may be willing to prey on flies, the ants and flies that we used as stimuli were considerably larger than the test spiders and salticids generally prey on insects comparable to or smaller than themselves. One of the 28 salticid species that we used, Natta horizontalis, is known to express a preference for ants as prey (Jackson and van Olphen 1992), but not army ants, which are substantially larger than the small ant species typically targeted by these rather timid salticids (pers. obs.). In addition, trials with flies could be expected to have motivated the spiders to approach faster than an empty chamber, but salticids in our experiments were not responding to the tsetse flies that we used as prey: for 27 of the 28 salticid species, entry latencies in trials with tsetse flies were not significantly different from entry latencies in trials with empty stimulus boxes (Table S1). The exception was Thyene inflata. Although T. inflata, like the other 27 species, expressed an aversion to ants, this salticid’s entry latency when tsetse flies were in the stimulus chamber was shorter than when the stimulus chamber was empty. However, T. inflata is unusual in that it often preys on tsetse flies and other similar-size insects in the field (unpublished observations).

Not all spiders are averse to ants, and, indeed, spiders that specialize at preying on ants are attracted both to their movement and static appearance (Jackson and van Olphen 1992; Pekár and Toft 2015; Nelson and Jackson 2006b; Nelson and Card 2016). However, despite their distinctive pattern of locomotion, this did not seem to be a major contributing factor to ant identification in our experiments. This corroborates previous suggestions that ant-averse salticids appear to be sufficiently deterred by the shape of the ants (Nelson and Jackson 2006a; Nelson and Card 2016) that motion, if used, is only of secondary importance—perhaps reinforcing or speeding-up identification, rather than being used as an unambiguous cue to discriminate ants. That salticids use multiple visual cues in some form of hierarchical order for the identification of arthropods is well-attested (Harland and Jackson 2000, 2002; Nelson and Jackson 2012; Dolev and Nelson 2014). While motion is an important prey-recognition cue (Bartos and Minias 2016), it appears redundant when recognizing ants as a potential threat. This finding parallels recent results in which many species of birds were found to mob cuckoo dummies with similar intensity and frequency as they mob real cuckoos, and that mobbing was more intense against cuckoos that parasitized the specific mobbing bird species (Tryjanowski et al. 2018). This suggests that selection has been exerted in birds to discriminate against this potential parasite, even in the absence of movement.

There was only one species, Goleba puella, for which we found a significant difference when we compared entry latencies from using stationary ant mounts to entry latencies from using living insects (Table S1). This species also was the only one that showed no differential response between dead ants and dead flies. Part of the explanation for G. puella being the only species for which making ant-fly discriminations seemed to require active, living individuals of these insects may pertain to the ultrastructure of the principal-eye retinas of this species having relatively poor capacity for spatial acuity (Blest et al. 1990). It is possible that comparatively poor capacity for spatial acuity may constrain G. puella to rely more strongly on motion-related cues.
Fig. 3 Boxplot (median, 1st and 3rd quartile) of entry latencies (s) for salticids to enter the testing chamber flanked by different stimuli. Whiskers depict minimum and maximum latencies. See Table 1 for sample sizes. Where medians are not visible in the boxes, these are all 900 s.
Fig. 3 (continued)
Our findings for 28 East African salticids, along with our earlier findings for 12 salticid species from the Philippines (Nelson and Jackson 2006a) suggest that an innate capacity to visually identify and avoid ants is a geographically widespread salticid characteristic. This capacity may also be phylogenetically widespread. Seven salticid subfamilies are currently recognized (Maddison 2015), with the large majority of species belonging to the Salticinæ. We now have experimental evidence of vision-based aversion to ants from three of these subfamilies: 32 salticines, six spartaeinæ and two asemoneinæ. Additionally, there is experimental evidence of innate vision-based capacity to identify ants by salticids that specialize on, and express an active preference for, ants as prey (Jackson and Nelson 2012), and by salticids that affiliate with ants for protection from their own predators (Nelson and Jackson 2014). Taken together, these studies suggest that ants have exerted especially strong selection pressures in the evolutionary shaping of salticid perceptual and decision-making processes.

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