Evidence that olfaction-based affinity for particular plant species is a special characteristic of

Evarcha culicivora, a mosquito-specialist jumping spider

Running title: Plant affinity in a jumping spider

Ximena J. NELSON^{1*}, Robert R. JACKSON^{1,2}

¹School of Biological Sciences, University of Canterbury, Private Bag 4800, Christchurch, New

Zealand.

²International Centre of Insect Physiology and Ecology, Thomas Odhiambo Campus, P.O. Box

30 Mbita Point, Kenya.

*Email: ximena.nelson@canterbury.ac.nz

Phone: 64-3-3642987 extn. 4050

Fax: 64-3-3642590

Key words: Plant-arthropod interactions, *Evarcha culicivora*, *Lantana camara*, Salticidae,

olfaction

ABSTRACT.

Evarcha culicivora, an East African jumping spider (family Salticidae), was shown in an earlier study to have an affinity for the odor from two particular plant species, namely Lantana camara and Ricinus communis. The olfactometer used in the earlier study was designed for choice testing. Here we focus on L. camara and, by using a second olfactometer method (retention testing), add to the evidence that the odor of this plant is salient to E. culicivora. Another 17 East African salticid species, all from different genera, were investigated using the same two olfactometer designs as used when investigating E. culicivora. The number of individuals of each of these 17 species that chose L. camara odor was not significantly different from the number that chose a no-odor control and, for each species, the latency to leave a holding chamber (retention time) in the presence of L. camara odor was not significantly different from retention time in the presence of a no-odor control. Based on these findings, we conclude that, rather than being a widespread salticid characteristic, an affinity for the odor of L. camara is a special characteristic of E. culicivora.

Many insects that specialize at feeding on nectar and pollen associate with particular plant species (e.g., Chittka et al. 1999; Waser & Ollerton 2006; Díaz et al. 2007), and many insects are known to rely on specific blends of plant-derived volatile compounds for identifying the particular plant species they exploit as sites for feeding or oviposition (e.g., Pichersky & Gershenzon 2002; Bruce et al. 2005; Anfora et al. 2009; Karlsson et al. 2009). There are also examples of spiders that associate with particular types of plants, especially pitcher plants (Cresswell 1993) and bromeliads (Romero & Vasconcellos-Neto 2004, 2005). Besides offering opportunity for nectar and pollen meals (Vogelei & Greissl 1989; Pollard et al. 1995; Jackson et al. 2001; Taylor & Pfannenstiel 2008, 2009; Taylor & Bradley 2009), associating with plants may reward spiders with opportunity to feed on other plant products (Meehan et al. 2009) and on insects that land on the plants (Ruhren & Handel 1999; Whitney 2004). In some instances, the benefits of associating with plants may include opportunity to feed on insects ensnared by the plant's sticky glandular hairs (Vasconcellos-Neto et al. 2007).

Little is known about the chemical cues by which spiders might identify specific plant species, but spiders are known to make use of chemosensory information regarding the sex, maturity, virgin-mated status, and fighting ability of conspecific individuals (Pollard et al. 1987; Clark et al. 1999; Roberts & Uetz 2005). Chemical cues are also known to be used by some spiders for detecting prey (Blanke 1972; Persons & Rypstra 2000; Clark et al. 2000a,b; Jackson et al. 2002, 2005) and predators (Persons et al. 2002; Li & Lee 2004; Li & Jackson 2005), and for determining the individual attractiveness of potential mates (Searcy et al. 1999; Roberts & Uetz 2005; Cross et al. 2009).

Two studies in particular suggest that further research is needed on how spiders might make use of plant-derived volatile compounds when identifying particular plant species. One of these studies showed associative learning by 'ghost spiders' (*Hibana futilis* Banks 1898, Anyphaenidae) when artificial odor was paired with artificial nectar (Patt & Pfannenstiel 2008). The other study showed that *Evarcha culicivora*, an East African salticid, responds in olfactometer experiments to the odor of two particular plant species on which it is commonly found, *Lantana camara* and *Ricinus communis* (Cross & Jackson 2009). That these two plant species might have a role in the mating system of *E. culicivora* has been suggested by other research (Cross et al. 2008) in which it was shown that, when on these plants, the courtship behavior of *E. culicivora* is more variable in display sequencing, more active, and more persistent. These effects are not evident during intraspecific interactions on a variety of other plant species (RRJ unpubl. data).

Here we focus on *L. camara* in particular and test 17 additional East African salticids with the odor of this plant species. Our hypothesis is that an affinity for the odor of *L. camara* is a special characteristic of *E. culicivora*. The rationale for testing other salticids is to consider, as an alternative hypothesis, the possibility that having an affinity for *L. camara* odor is a widespread salticid characteristic. Besides using choice-test olfactometers with these 17 salticid species (as adopted in Cross & Jackson's (2009) study on *E. culicivora*), we also use retentiontest olfactometers in experiments with these 17 salticids and with *E. culicivora*. Retention-test olfactometers are used for determining how long a test spider will remain in a small holding chamber when exposed to specific odors. This type of testing has been used in earlier research with *E. culicivora* (Cross et al. 2009), but never before specifically for examining response to plant odor.

METHODS

General.- Olfactometer testing was carried out using salticids from laboratory cultures (F2 and F3 generation). Rearing methods, as well as the basic procedures used in olfactometer experiments, were as in earlier research (Cross & Jackson 2009; Cross et al. 2009) and only essential details are provided here.

For rearing and maintenance, each spider was fed to satiation three times a week on blood-carrying female mosquitoes (*Anopheles gambiae s.s.* from laboratory culture) and 'lake flies' (*Nilodorum brevibucca*, Chironomidae; collected as needed from field). Hunger level was standardized by subjecting each test spider to a 7-day pre-trial fast.

Two olfactometer methods were used (choice testing and retention testing), with the odor source being a plant cutting held in an odor chamber (glass-cube box). Each cutting was two *Lantana camara* umbels (clusters of flowers) with accompanying leaves and stems (no flowers senescent) taken from the field immediately before setting up for an experiment; median weight/umbel (1st and 3rd quartiles) = 364 (329 and 384) mg, (n=10). Disposable surgical gloves were worn while collecting and handling plant material.

Testing was carried out between 0800 h and 1400 h (laboratory photoperiod 12L-12D, lights on at 0700 hr). Between trials, olfactometers were dismantled and cleaned with 80% ethanol followed by distilled water and then dried in an oven. Airflow in the olfactometers was adjusted to 1500 ml/min (Matheson FM-1000 airflow regulator) and there was no evidence that this setting had any adverse effects on the salticid's locomotion or other behavior. The spiders used in choice tests were different from the spiders used in retention tests, but no test spiders had prior experience with plants. No spider was used in more than one choice test or more than one pair

(experimental one day, control another day) of retention tests. All test spiders were adults that matured 2-3 weeks before being tested and none had mated. Both sexes of all species were used in choice testing, but only males were used in retention testing.

Choice testing.- Y-shaped glass olfactometers were used for choice testing (Fig. 1a). The two ends of the Y were the 'choice arms', with each choice arm being connected to an odor chamber. Which of the two odor chambers contained the plant cutting was determined at random. Air was pumped separately into the two odor chambers and then through the choice arms before converging at the stem of the Y ('test arm').

Before testing began, the test spider (N = 70 per sex and species) was confined for 2 min in a holding chamber at the far end of the test arm. While in the holding chamber, the test spider's access to the test arm was blocked by a removable metal grill that fit within a slit in the chamber roof. Testing began by lifting the grill. When the spider entered a choice arm and remained there for 30 sec, we recorded the arm entered as the test spider's choice. The spider was allowed 30 min within which to make a choice and the number of spiders that failed to make a choice was, for each species, always fewer than 5% of the spiders tested.

Retention testing.- During retention testing, air was pushed successively through an odor chamber, a holding chamber and an exit chamber (Fig. 1b). The holding chamber was a glass tube (rubber stopper in one end, other end open). The open end of the holding chamber fit securely in the hole in the glass cube that formed the exit chamber, flush with the inner wall of the exit chamber. At the other end of the holding chamber, there was a hole in the stopper with a glass tube going through to the odor chamber, which was identical in size to the exit chamber (see Fig. 1b for dimensions). A

nylon-netting screen over the stopper (new netting for each test) ensured that the test spider could not enter the odor chamber, the only way out of the holding chamber being via the opening into the exit chamber. The exit chamber was another glass cube identical to the odor chamber.

The test spider (N = 20 for each species) was first kept in the holding chamber for 2 min, with the holding chamber not yet connected to the stimulus and exit chambers. The end of the holding chamber that would go into the exit chamber was plugged with a rubber stopper. To begin a test, this stopper was removed and the holding chamber was positioned between the stimulus and exit chamber, but with a prerequisite being that the test spider had to be in the half of the holding chamber distal to the exit chamber. If this prerequisite was not met at the end of the 2-min pre-test period, the beginning of the test was delayed until the spider moved on its own accord into the distal half of the chamber and remained there for 2 min. Testing was aborted if this criterion was still not met after waiting 15 min, but aborted tests were rare (< 5% for any given species).

No-odor control tests and odor tests were randomized. Once testing began, we recorded retention time (i.e., the test spider's latency to leave the holding chamber, defined as the time elapsing between the beginning of a test and departure by the spider into the exit chamber; maximum time allowed, 60 min). By default, the spider's retention time was recorded as 60 min whenever the 60-min test period ended with the test spider still in the holding chamber.

Data analysis.- Choice-test data were analyzed using tests for goodness of fit (H_0 =50:50) and retention-testing data were analyzed using non-parametric Wilcoxon tests for paired comparisons (null hypothesis: latency to leave holding chamber when tested with odor source matched latency to leave holding chamber when tested with no-odor control). Retention testing data are shown based on each test spiders' calculated absolute difference score (subtracting its latency to leave

holding chamber when tested with control from latency to leave holding chamber when tested with odor), resulting in positive scores when the spider spent more time in the holding chamber when tested with odor, and resulting in negative scores when spider spent more time in the holding chamber when tested with no odor.

Voucher specimens of all species have been deposited in the Florida State Collection of Arthropods, Gainesville, Florida, USA.

RESULTS

Choice-test data from males and females of each species did not differ in any case, so these data were pooled for simplification. In the earlier study (Cross & Jackson 2009), *Evarcha culicivora* chose *Lantana camara* odor significantly more often than the no-odor control in choice-test olfactometers (Fig. 2) and, in the present study, *E. culicivora* had a significantly longer latency to leave the holding chamber when in the presence of *L. camara* odor than when in the presence of a no-odor control. However, for the other 17 salticid species, the number of individuals that chose *L. camara* odor was not significantly different from the number that chose the no-odor control in the choice-test olfactometers. In the retention-test olfactometers the retention time in the presence of *L. camara* odor was also not significantly different, for these 17 species, from retention time in the presence of the no-odor control (Table 1, Fig. 3; note non-significant trend for *Natta rufopicta* to display greater retention in the control tests). In concordance with results from olfactometer choice tests, *L. camara* odor did induce a significant retention in *E. culicivora* (Table 1).

DISCUSSION

The earlier study (Cross & Jackson 2009) demonstrated that the odor of *Lantana camara* is salient to *Evarcha culicivora*, but left unresolved the question of whether responsiveness to *L. camara* odor by *E. culicivora* is an unusual characteristic of this particular salticid species or, alternatively, a characteristic that is widespread in the Salticidae. Here we investigated another 17 salticid species, all from different genera, from East Africa. For each of these species, when we used the same odor-based choice-testing methods and achieved the same sample sizes as in the earlier experiments with *E. culicivora*, the number of individuals that chose *L. camara* odor was not significantly different from the number that chose the no-odor control. Using the retention-testing olfactometers, we again found evidence that the odor of *L. camara* is salient to *E. culicivora* and, for retention tests, as for choice tests, the response to *L. camara* odor by each species other than *E. culicivora* was not significantly different from response to no-odor controls.

As there are more than 5,000 described species in the family Salticidae (Platnick 2010), our findings should not be construed as proving that *E. culicivora* is absolutely unique, but it seems unlikely that responsiveness to *L. camara* odor is widespread within the Salticidae.

The precise role of *L. camara* in the biology of *E. culicivora* is poorly understood. Earlier research (Cross et al. 2008) suggested that we need a better understanding of the role plants might play in the mating strategy of *E. culicivora*, but plants may also have a role in its feeding strategy. As *E. culicivora* is known to feed on nectar (RRJ unpubl. data), one hypothesis that should be considered is that responding to *L. camara* odor is related to visiting this plant species for nectar meals. This would make *E. culicivora* comparable to *Heliconius melpomene*, a butterfly that, by responding to the odor of *L. camara*, locates and feeds on the nectar of this plant (Andersson et al. 2002; Andersson & Dobson 2003). However, there is a complication with any hypothesis concerning *E. culicivora* having evolved mechanisms of exploiting specifically *L.*

camara. This plant species is native to the Americas and is an introduced weed in many parts of the world, including East Africa (Day et al. 2003). We need a better understanding of how *E. culicivora* responds to a wider range of plant species, including native species with which it has shared a longer evolutionary history, in addition to the primary volatile components of various plants, before we can tease apart the basis of this affinity. This large topic is the subject of ongoing research.

Evarcha culicivora has an unusual predatory strategy, as its preferred prey are bloodcarrying mosquitoes (Jackson et al. 2005). An alternative hypothesis is that this mosquitospecialist spider locates its prey by visiting L. camara or other plants. Only female mosquitoes feed on blood (Clements 1999). Male mosquitoes feed primarily on nectar, but E. culicivora is proficient at discriminating between males and females, has an active preference for female mosquitoes as prey and chooses *Anopheles* in preference to other mosquitoes (Nelson & Jackson 2006). However, it is now well established that visiting plants for nectar meals is important not only for the male but also for the female of a variety of mosquito species, including *Anopheles* species (McCrae et al. 1969, 1976; Gujral & Vasudevan 1983; Clements 1999; Foster & Takken 2004; Impoinvil et al. 2004; Manda et al. 2007a,b). However, it is unlikely that encounters between E. culicivora and female mosquitoes, including Anopheles, often happen on L. camara or other plants, as E. culicivora, like most salticids (Richman & Jackson 1992), appears to be active as a predator during daylight hours (RRJ unpubl. data) while its prey, the female mosquito, probably feeds from plants primarily at night. This problem notwithstanding, E. culicivora might find mosquitoes during the daytime resting post-feeding in the vicinity of the plants to which the spider and the mosquito have been attracted, albeit at different times.

ACKNOWLEDGMENTS

We thank Godfrey Otieno Sune, Stephen Abok Aluoch and Jane Atieno Obonyo for their assistance at ICIPE. We also gratefully acknowledge support from the Royal Society of New Zealand (Marsden Fund (M1096, M1079) and James Cook Fellowship (E5097)), the National Geographic Society (8676-09, 6705-00) and the US National Institutes of Health (R01-AI077722).

REFERENCES

- Andersson, S. & H. E. M. Dobson. 2003. Behavioral foraging responses by the butterfly Heliconius melpomene to Lantana camara floral scent. Journal of Chemical Ecology 29:2303-2318.
- Andersson, S., L. A. Nilsson, I. Groth, & G. Bergström. 2002. Floral scents in butterfly-pollinated plants: possible convergence in chemical composition. Botanical Journal of the Linnean Society 140:129-153.
- Anfora, G., M. Tasin, A. De Crostofaro, C. Ioriatti & A. Lucchi. 2009. Synthetic grape volatiles attract mated *Lobesia botrana* females in laboratory and field bioassays. Journal of Chemical Ecology 35:1054-1062.
- Blanke, R. 1972. Untersuchungen zur ökophysiologie und ökethologie von *Cyrtophora citricola* Forskål (Araneae, Araneidae) in Andalusien. Forma et Functio 5:125–206.
- Bruce, T. J. A., L. J. Wadhams & C. M Woodcock. 2005. Insect host location: a volatile situation. Trends in Plant Science 10:269-274.
- Chittka, L., J. D. Thomson & N. M. Waser. 1999. Flower constancy, insect psychology, and plant evolution. Naturwissenschaften 86:361-377.

- Clark, R. J., R. R. Jackson & J. R. Waas. 1999. Draglines and assessment of fighting ability in cannibalistic jumping spiders. Journal of Insect Behavior 12:753-766.
- Clark, R. J., D. P. Harland & R. R. Jackson. 2000a. Speculative hunting by an araneophagic salticid spider. Behaviour 137:1601-1612.
- Clark, R. J., R. R. Jackson & B. Cutler. 2000b. Chemical cues from ants influence predatory behavior in *Habrocestum pulex* (Hentz), an ant-eating jumping spider (Araneae, Salticidae). Journal of Arachnology 28:299-341.
- Clements, A. N. 1999. The biology of mosquitoes. CABI Publishing, Wallingford, England.
- Cresswell, J. E. 1993. The morphological correlates of prey capture and resource parasitism in pitchers of the carnivorous plant *Sarracenia purpurea*. American Midland Naturalist 129:35-41.
- Cross, F. R., R. R. Jackson & S. D. Pollard. 2008. Complex display behaviour of *Evarcha culicivora*, an East African mosquito-eating jumping spider. New Zealand Journal of Zoology 35:151-187.
- Cross, F. R. & R. R. Jackson. 2009. Odour-mediated response to plants by *Evarcha culicivora*, a blood-feeding jumping spider from East Africa. New Zealand Journal of Zoology 36:75-80.

- Cross, F. R., R. R. Jackson & S. D. Pollard. 2009. How blood-derived odor influences mate-choice decisions by a mosquito-eating predator. Proceedings of the National Academy of Science USA 46:19416-19419.
- Day, M. D., C. J. Wiley, J. Playford & M. P. Zalucki. 2003. *Lantana*: current management status and future prospects. CABI Publishing, Canberra.
- Díaz, P. C., C. Gruter & W. M. Farina. 2007. Floral scents affect the distribution of hive bees around dancers. Behavioral Ecology & Sociobiology. 61:1589-1597.
- Foster, W. A. & W. Takken. 2004. Nectar-related vs. human-related volatiles: behavioural response and choice by female and male *Anopheles gambiæ* (Diptera: Culicidae) between emergence and first feeding. Bulletin of Entomological Research 94:145-157.
- Gujral, G. S. & P. Vasudevan. 1983. *Lantana camara* L., a problem weed. Journal of Scientific and Industrial Research 42:281-286.
- Impoinvil, D. E., J. O. Kongere, W. A. Foster, B. N. Njiru, G. F. Killeen, J. I. Githure, J. C. Beier, A. Hassanali & B. G. J. Knols. 2004. Feeding and survival of the malaria vector *Anopheles gambiæ* on plants growing in Kenya. Medical and Veterinary Entomology 18:1-8.

- Jackson, R. R., R. J. Clark & D. P. Harland. 2002. Behavioural and cognitive influences of kairomones on an araneophagic spider. Behaviour 139:749-775.
- Jackson, R. R., X. J. Nelson & G. O. Sune. 2005. A spider that feeds indirectly on vertebrate blood by choosing female mosquitoes as prey. Proceedings of the National Academy of Science USA 102:15155-15160.
- Jackson, R. R., S. D. Pollard, X. J. Nelson, G. B. Edwards & A. T. Barrion. 2001. Jumping spiders (Araneae: Salticidae) that feed on nectar. Journal of Zoology 255:25-29.
- Karlsson, M. F., G. Birgersson, A. M. C. Prado, F. Bosa, M. Bengtsson & P. Witzgall. 2009.

 Plant odour analysis of potato response of Guatemalan moth to above- and belowground potato volatiles. Journal of Agricultural and Food Chemistry 57:5903-5909.
- Li, D. & R. R. Jackson. 2005. Influence of diet-related chemical cues from predators on the hatching of egg-carrying spiders. Journal of Chemical Ecology 31:333-342.
- Li, D. & W. S. Lee. 2004. Predator-induced plasticity in web-building behaviour. Animal Behaviour 67:309-318.
- Manda, H., L. C. Gouagna, W. A. Foster, R. R. Jackson, J. C. Beier, J. I. Githure, & A. Hassanali. 2007a. Effect of discriminative feeding from different plant species on the survival and fecundity of *Anopheles gambiæ* s.s. Malaria Journal 6:e113.

- Manda, H., L. C. Gouagna, E. Nyandat, E. W. Kabiru, R. R. Jackson, W. A. Foster, J. I. Githure, J. C. Beier & A. Hassanali. 2007b. Discriminative feeding behaviour of *Anopheles gambiæ* s.s. on endemic plants in western Kenya. Medical and Veterinary Entomology 21:103-111.
- McCrae, A. W. R., Y. Ssenkubuge, P. Manuma, C. Mawjje & A. Kitama. 1969. Mosquito and tabanid activity at plant sugar sources. Repository of the East African Virus Research Institute 18:96-102.
- McCrae, A. W. R., P. F. L. Boreham & Y. Ssenkubuge. 1976. The behavioural ecology of host selection in *Anopheles implexus* (Theobald) (Diptera, Culicidae). Bulletin of Entomological Research 66:587-631.
- Meehan, C. J., E. J. Olson, M. W. Reudink, T. K. Kyser & R. L Curry. 2009. Herbivory in a spider through exploitation of an ant-plant mutualism. Current Biology 19:R882-R883.
- Nelson, X. J. & R. R. Jackson. 2006. A predator from East Africa that chooses malaria vectors as preferred prey. PLoS ONE 1(1): e132. doi:10.1371/journal.pone.0000132.
- Patt, J. M. & R. S. Pfannenstiel. 2008. Odor-based recognition of nectar in cursorial spiders.

 Entomologia Experimentalis et Applicata 127:64-71.

- Persons, M. H. & A. L. Rypstra. 2000. Preference for chemical cues associated with recent prey in the wolf spider *Hogna helluo* (Araneae: Lycosidae). Ethology 106:27-35.
- Persons, M. H., S. E. Walker & A. L. Rypstra. 2002. Fitness costs and benefits of antipredator behavior mediated by chemotactile cues in the wolf spider *Pardosa milvina* (Araneae: Lycosidae).

 Behavioral Ecology 13:386-392.
- Pichersky, E. & J. Gershenzon. 2002. The formation and function of plant volatiles: perfumes for pollinator attraction and defense. Current Opinion in Plant Biology 5:237-243.
- Plantick, N. I. 2010. World Spider Catalogue, version 11.0 American Museum of Natural History, http://research.amnh.org/iz/spiders/catalog/COUNTS.html
- Pollard, S. D., A. M. MacNab & R. R. Jackson. 1987. Communication with chemicals:

 Pheromones and spiders, pp. 133-141, in W. Nentwig (ed.). Ecophysiology of Spiders.

 Springer-Verlag, Berlin.
- Pollard, S. D., M. W. Beck & G. N. Dodson. 1995. Why do male crab spiders drink nectar.

 Animal Behaviour 49:1443-1448.
- Richman, D. B. & R. R. Jackson. 1992. A review of the ethology of jumping spiders (Araneae, Salticidae). Bulletin of the British Arachnological Society 9:33-37.

- Roberts, J. A. & G. W. Uetz. 2005. Information content of female chemical signals in the wolf spider, *Schizocosa ocreata*: male discrimination of reproductive state and receptivity.

 Animal Behaviour 70:217-223.
- Romero, G. Q. & J. Vasconcellos-Neto. 2004. Beneficial effects of flower-dwelling predators on their host plant. Ecology 85:446-457.
- Romero, G. Q. & J. Vasconcellos-Neto. 2005. The effects of plant structure on the spatial and microspatial distribution of a bromeliad-living jumping spider (Salticidae). Journal of Animal Ecology 74:12-21.
- Ruhren, S. & S. N. Handel. 1999. Jumping spiders (Salticidae) enhance the seed production of a plant with extrafloral nectaries. Oecologia 119:227-230.
- Searcy, L. E., A. L. Rypstra & M. H. Persons. 1999. Airborne chemical communication in the wolf spider *Pardosa milvina*. Journal of Chemical Ecology 25:2527-2533.
- Taylor, R. M. & R. A Bradley. 2009. Plant nectar increases survival, molting, and foraging in two foliage wandering spiders. Journal of Arachnology 37:232-237.
- Taylor, R. M. & R. S. Pfannenstiel. 2008. Nectar feeding by wandering spiders on cotton plants. Environmental Entomology 37:996-1002.

- Taylor, R. M. & R. S. Pfannenstiel. 2009. How dietary plant nectar affects the survival, growth, and fecundity of a cursorial spider *Cheiracanthium inclusum* (Araneae: Miturgidae). Environmental Entomology 38:1379-1386.
- Vasconcellos-Neto, J., G. Q. Romero, A. J. Santos & A. S. Dippenaar-Schoeman. 2007.

 Associations of spiders of the genus *Peucetia* (Oxyopidae) with plants bearing glandular hairs. Biotropica 39:221-226.
- Vogelei, A. & R. Greissl. 1989. Survival strategies of the crab spider *Thomisus onustus* Walckenaer 1806 (Chelicerata, Arachnida, Thomisidae). Oecologia 80:513-515.
- Waser, N. M. & J. Ollerton. 2006. Plant-pollinator interactions: from specialization to generalization. Chicago, University of Chicago Press. 445 p.
- Whitney, K. D. 2004. Experimental evidence that both parties benefit in a facultative plant-spider mutualism. Ecology 85:1642-1650.

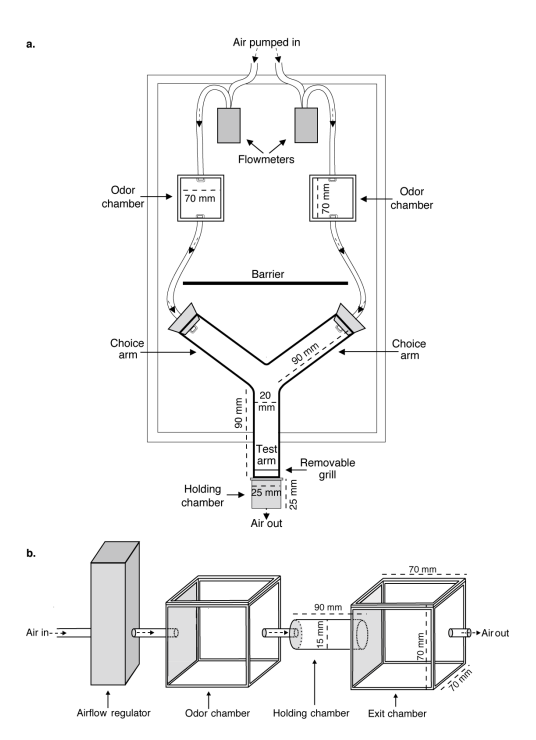


Figure 1.- Olfactometers used for: **a.** choice testing (view of odor source obstructed by opaque barrier) and **b.** retention testing (view of odor source obstructed by black paper taped to outside of odor chamber wall that faced holding chamber). Dashed arrows indicate direction of airflow. Not drawn to scale.

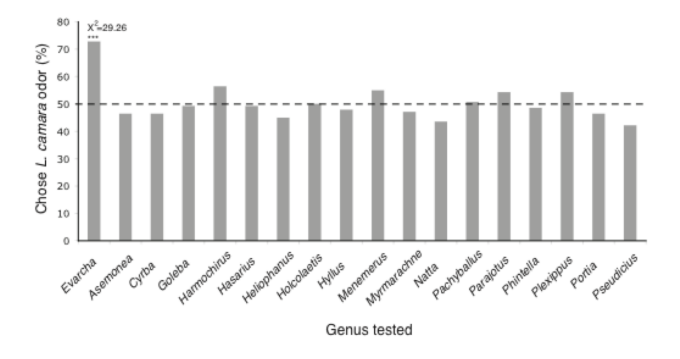


Figure 2.- Pooled results from olfactometer choice-tests. *Evarcha culicivora* chose odor arm significantly more often control arm (data from Cross and Jackson 2009). For all other salticid species, number of individuals that chose odor not significantly different from number that chose control. Dashed line denotes 50%. N = 140. χ^2 = test of goodness of fit. *** P < 0.0001.

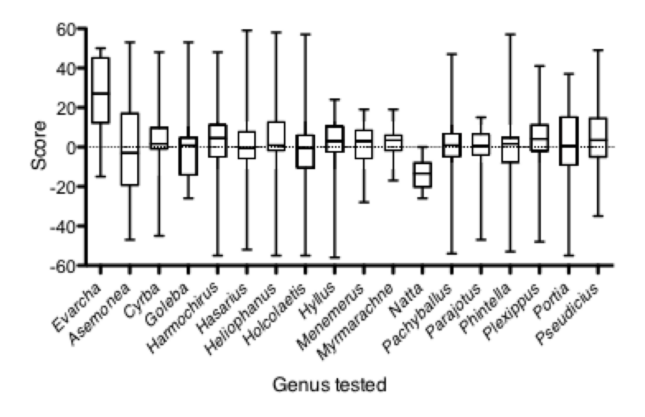


Figure 3.- Boxplots (median and quartiles) with whiskers (min and max) for retention testing for all species. Score calculated by subtracting latency to leave holding chamber when tested with control from latency to leave holding chamber when tested with odor (positive score: spider spent more time in the holding chamber when tested with odor; negative score: spider spent more time in the holding chamber when tested with control). N = 20.

Table 1. Test results for Wilcoxon-tests comparing latency to leave holding chamber in control or experimental tests. The difference in time beween these is depicted in Fig. 3. All spiders sourced in Kenya except *Parajotus cinereus* (from Uganda).

Test spider species	W	P
Evarcha culicivora Wesolowska & Jackson 2003	176.0	0.0004
Asemonea murphyae Wanless 1980	-26.0	0.61
Cyrba ocellata (Kroneberg 1875)	70.0	0.16
Goleba puella (Simon 1885)	-31.0	0.58
Harmochirus brachiatus (Thorell 1877)	65.0	0.20
Hasarius adansoni (Savigny et Audouin 1825)	3.0	0.97
Heliophanus sp.	51.0	0.24
Holcolaetis vellerea (Simon 1909)	-21.0	0.69
Hyllus sp.	50.0	0.29
Menemerus congoensis Lessert 1925	28.0	0.56
Myrmarachne melanotarsa Wesolowska & Salm 2002	73.0	0.15
Natta rufopicta (Simon 1909)	82.0	0.08
Pachyballus cordiformis Berland et Millot 1941	13.0	0.78
Parajotus cinereus Wesolowska 2004	19.0	0.70
Phintella sp.	-1.0	1.00

Plexippus sp.	60.0	0.27
Portia africana (Simon 1885)	9.0	0.85
Pseudicius sp.	40.0	0.43