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The ants are extraordinary in having evolved many lineages that exploit closely related ant societies as social parasites, but social parasitism by distantly related ants is rare. Here we document the interaction dynamics among a Sericomymex fungus-growing ant host, a permanently associated parasitic guest ant of the genus Megalomyrmex, and a raiding agro-predator of the genus Gnamptogenys. We show experimentally that the guest ants protect their host colonies against agro-predator raids using alkaloid venom that is much more potent than the biting defenses of the host ants. Relatively few guest ants are sufficient to kill raiders that invariably exterminate host nests without a cohabiting guest ant colony. We also show that the odor of guest ants discourages raider scouts from recruiting nestmates to host colonies. Our results imply that Sericomymex fungus-growers obtain a net benefit from their costly guest ants behaving as a functional soldier caste to meet lethal threats from agro-predator raiders. The fundamentally different life histories of the agro-predators and guest ants appear to facilitate their coexistence in a negative frequency-dependent manner. Because a guest ant colony is committed for life to a single host colony, the guests would harm their own interests by not defending the host that they continue to exploit. This conditional mutualism is analogous to chronic sickle cell anemia enhancing the resistance to malaria and to episodes in human history when mercenary city defenders offered either net benefits or imposed net costs, depending on the level of threat from invading armies.

Ant societies retain much of their coherence through chemical nestmate recognition (1, 2), which allows resident workers to differentiate between friend and foe by colony-specific chemical signatures (3). This recognition system is not infallible, however; numerous species have evolved ways to evade detection so they can exploit ant colonies through invasion, usurpation, or thievery (2). One common route to social parasitism is shown convergently by many ant genera in which social parasites are their host’s closest relatives, a scenario that might have arisen through sympatric speciation (4, 5). At the other end of the spectrum are interactions with different insect orders, such as parasitic beetles and butterfly caterpillars, that drain host ant resources to support their own growth (6).

Intermediate types of parasitic interactions involving ants that exploit other, distantly related ants are rare (7). Several species of Megalomyrmex (Solenopсидini) belong to this category, associating in varying capacities with fungus-growing ant hosts (Attini), a clade of the same ant subfamily (8–12) (Fig. 1). Most details about the diversity and life history adaptations of these ants have been clarified only recently (11), and much of their biology remains to be discovered (SI Text: Study System and Tables S1 and S2). Free-living predatory Megalomyrmex are known to biosynthesize venom alkaloids that are used in defense (13, 14) whereas the fungus-growing ant associates seem to aggressively dispense these venom when they attack host colonies (15). Some of these parasites are obligate or facultative thief ants consuming brood and fungus gardens (15), whereas others are specialized agro-predators that move from one host colony to the other after usurping fungus gardens and killing or chasing away the resident ants (10). Despite the often high densities of their attine hosts, these Megalomyrmex social parasites remain rare, with parasitism rates of ~1.5–14% (10, 15), similar to the rates of many other social parasites (5).

Behaviorally derived lineages of Megalomyrmex have become guest ant parasites of the higher attine genera Trachymyrmex and Sericomymex (11, 16, 17) (Fig. L4). The Trachymyrmex specialist, Megalomyrmex adamsae, appears to have remained as equally rare as the agro-predators and thief ants (11), but the Sericomymex specialist Megalomyrmex symmetochus (Fig. 1B) is surprisingly common, with a prevalence of >80% in some host populations (16). Newly mated guest ant queens of both species likely enter host colonies by stealth and establish themselves in the fungus garden, where their developing colony will consume host brood and fungus garden for years (11). Their presence slows host colony growth and also prevents or reduces host reproduction, because both guest ant species clip the wings of host gynes (virgin queens), but not males (11). Such mutilation reduces host reproduction and dispersal via mating flights, but likely increases guest ant fitness when these mutilated females adopt worker tasks (11, 18). Like all guest ant social parasites, M. symmetochus retains a fully functional worker caste (16, 19), in contrast to many social parasites that exploit the services of their phylogenetically similar host colony without the need to produce workers (5).

The maintenance of a large worker caste despite permanent cohabitation with a host colony may have several purposes, all based on some specialized role for the parasite workers. First, it

Significance

We document the behavioral interactions among three ant species: a fungus-growing host ant, a permanently associated parasitic guest ant, and a raiding agro-predator ant. We show that the presence of guest ants becomes advantageous when host ants are attacked by raider ants, because guest ants use alkaloid venom to defend their host ant colony. Furthermore, detection of the guest ant odors is sufficient to discourage raider scouts from recruiting nestmates to host colonies. Guest ants likely have evolved this protective behavior because they also perish when their host colony dies.


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may be that the distantly related host workers are unable to feed the social parasite larvae (5) even though adult hosts and parasites are adapted to the same highly specific fungal diet. Another possibility is that the guest ant colony remains at risk of occasional attack by the host workers; such antagonism has been observed between M. symmetochus workers and host workers in older colonies (SI Text, Study System). But Sericomymex ants have only vestigial stings (20) and often feign death when attacked, and so this does not explain the large number of parasite workers. Thus, it seems reasonable to hypothesize that a large number of Megalomynex guest ant workers continue to remain essential for the survival and reproductive success of their own mother queen in ways other than merely reinforcing their dominance over Sericomymex host workers and caring for their own brood.

Neither founding queens nor mature colonies of M. symmetochus guest ants are known to move to other host colonies later in life, implying that guest ant reproductive success is completely dependent on the continued well being (albeit not the reproduction) of host colonies (11). Thus, we conjectured that M. symmetochus guest ants might act as defenders when host colonies are attacked by natural enemies that are sufficiently effective to pose a significant threat. Such a specialized enemy, the unrelated agro-predator ant Gnamptogenys hartmani (Ectatomminae: Ectatommini) (Fig. 1C and Fig. S1A), was recently seen to raid colonies of Sericomymex in Panama and to usurp their gardens and nest structures with remarkable efficiency (21). This finding suggests that the enhanced mortality risk emanating from these raids might have produced an unusual secondary mutualism between the socially parasitic guest ants and their hosts. Rather than merely reducing worker production as ant social parasites normally do, the M. symmetochus guest ants produce a seemingly excess number of workers (Table S2) that constantly patrol the host nest. These guest ants’ potent alkaloid weaponry, which originally secured their establishment at a chronic cost to the host colony, potentially could also serve to protect the host from greater harm in the direct interest of the guest ant parasite.

We tested this idea in a series of controlled laboratory experiments aimed at quantifying the damage by G. hartmani agro-predator parasites and the defense efficiency of Sericomymex hosts with and without guest ants (SI Text: Study System). We found that hosting even a moderate number of Megalomynex guest ants provides almost complete protection against G. hartmani raids, because (i) guest ants are much more efficient than host ants in killing intruding G. hartmani workers; (ii) guest ants reduce host ant mortality inflicted by the raiding agro-predators; and (iii) scouts of G. hartmani preferentially recruit nestmates to Sericomymex host colonies whose odor indicates an absence of guest ants.

Results and Discussion

After a number of staged encounters with guest ant-infected subcolonies of S. amabilis in the laboratory, it became clear that G. hartmani scouts were immediately attacked by the M. symmetochus guest ants, and that raids were often deterred (Movies S1 and S2). Not only were G. hartmani workers killed during the altercations with the M. symmetochus defenders, but some were also attacked later by members of their own raiding party (Fig. S1B), suggesting that M. symmetochus venom is both toxic and causes confusion in G. hartmani ants. In addition, experimentally stung G. hartmani ants more often avoided contact with a naive sister compared with controls (2.88 ± 0.72% of time spent together vs. 33.27 ± 7.01%; Welch’s t = 4.31, df = 1, P = 0.0073) (SI Text: Methods and Results and Fig. S1 C and D), but when contact was made, stung ants were often attacked (four of six replicates) and sometimes killed, indicating that M. symmetochus venom disrupts nestmate recognition abilities of G. hartmani, much like other antagonistic chemicals used by ant exploiters (2, 22).

To formally investigate the efficiency of M. symmetochus and S. amabilis defenses against a single G. hartmani ant, we conducted a series of experiments with a varying number of opponents (two, three, four, six, or eight workers of S. amabilis or M. symmetochus) (Fig. 2). We found that M. symmetochus workers were much more effective than S. amabilis host workers at killing G. hartmani raiders [binomial generalized linear model (GLZ), likelihood ratio (LR) χ² = 42.3, P < 0.0001] (Fig. 24). Surviving S. amabilis hosts sometimes fled to the roof of the nest box to escape their predators, as reported previously by Dijkstra.
and Boomsma (21), and never killed the intruding G. hartmani worker in the first hour of the trial (n = 17). However, M. symmetochus workers pursued, attacked, and killed the G. hartmani raiders in 45% of 20 trials in the first hour and in 100% of 20 trials within 24 h (Fig. 2A). Furthermore, the guest ant workers were significantly less likely than the S. amabilis host workers to be killed by the G. hartmani worker (binomial GLZ, LR $\chi^2 = 7.29$, $P = 0.007$) (Fig. 2B). A higher number of defenders of either species was generally associated with a reduced average casualty rate (LR $\chi^2 = 18.85$, $P < 0.0001$), and this reduction was similar for both species (species × number interaction, LR $\chi^2 = 0.0996$, $P = 0.752$).

When attacking, G. hartmani workers lock on to their S. amabilis opponents and sting repeatedly, releasing their potent nonvolatile venom (SI Text: Methods and Results). These one-to-one engagements can last from approximately 20 seconds to up to 10 min, during which time the S. amabilis defenders may use their relatively powerful mandibles to bite off legs or antennae of the G. hartmani attackers (Fig. S1A) but then feign death, prompting the G. hartmani workers to release them. This cycle can repeat itself several times before the S. amabilis workers are eventually killed. In contrast, the Megalomymex defensive strategy is aggressive and uses an approximate 1:1 mix of two isomers of butylhexylpyrrolizidine alkaloids (SI Text: Methods and Results) dispensed from their specialized sting as an aerosol or contact venom (Fig. 1B, SI Text: Methods and Results, and Movie S2). To evaluate the complex interactions between all three ant species, we conducted a second experiment using larger and more natural (i.e., with fungus garden fragments) subcolonies of S. amabilis with and without a variable number of guest ants in which we introduced two G. hartmani raiders.

In this more complex interaction experiment, both introduced G. hartmani workers were killed in six of the nine S. amabilis subcolonies across the range of zero, three, or six M. symmetochus defenders (Fig. 3). When mortal damage was also considered (i.e., at least six of eight possible G. hartmani extremities lost), all trials left at least one G. hartmani dead or doomed after 24 h independent of the number of guest ants (binomial GLZ, LR $\chi^2 = 0.286$, $P = 0.593$). As in the first experiment (Fig. 2B), the G. hartmani raiders were very effective at killing a high proportion (70% on average) of the S. amabilis host workers in the absence of M. symmetochus, but as the number of M. symmetochus guest ants increased, host casualties decreased to rather low values (LR $\chi^2 = 10.93$, $P = 0.0009$) (Fig. 3A). This finding indicates that the numerical ratios of interacting ants used in this experiment were well balanced, such that adding defenders of each category had a noticeable effect. The external damage to G. hartmani workers (i.e., the proportion of the maximum 2 × 8 raider legs and antennae that were missing) decreased with increasing numbers of defending guest ants (LR $\chi^2 = 14.17$, $P = 0.0002$) (Fig. 3B), supporting behavioral observations that G. hartmani workers were indeed killed by the guest ants and not by the host ants. As in the first experiment (Fig. 2B), there was little difference in guest ant mortality between treatments (LR $\chi^2 = 0.232$, $P = 0.630$), indicating that as few as three M. symmetochus workers offer adequate protection to S. amabilis colonies when there are two G. hartmani intruders. This protection was even greater when the ratio of M. symmetochus to S. amabilis was increased to 1:3 (6 to 18; Fig. 3A), a ratio close to that seen in field colonies (Table S2).

Fig. 2. Defense efficiencies of host and guest ants. (A) The resulting mortality after a single G. hartmani agro-predator interacted with groups of two to eight S. amabilis host ants (blue Petri dish) or M. symmetochus guest ants (orange Petri dish) after 24 h. Defender category significantly affected G. hartmani worker mortality (binomial GLZ, LR $\chi^2 = 42.34$, $P < 0.0001$), with S. amabilis effective in killing only when greatly outnumbering G. hartmani (blue dots) and M. symmetochus significantly more effective in killing regardless of their number (orange dots). (B) The overall mortality inflicted by the G. hartmani worker on host or guest ant defenders differed significantly (binomial GLZ, LR $\chi^2 = 18.84$, $P < 0.0001$), with S. amabilis defenders (blue dots) taking proportionally much higher casualties. The proportional mortality of both defenders decreased significantly with an increasing number of defenders (binomial GLZ, LR $\chi^2 = 7.29$, $P = 0.0069$), but this decrease did not differ between the defending species (interaction between number and species of defenders, binomial GLZ, LR $\chi^2 = 0.0996$, $P = 0.752$). Ant drawings courtesy of Rozlyn Haley.
Subcolony of 18 host & (0-6) guest ants

Two raiders

![Diagram of Subcolony of 18 host & 0-6 guest ants and Two raiders]

Fig. 3. Host survival rates and raider mutilations. When threatened, M. symmetochus guest ants use toxic venom, whereas Sericomymex hosts mutilate intruders by removing appendages. (A) When 18 host ants were confronted with 2 intruding Gnamptogenys workers without (0) or with (3 or 6) Megalomymex guest ants, the proportion of S. amabilis deaths was decreased (binomial GLZ, LR $\chi^2 = 10.93$, $P = 0.0009$). (B) The same increased number of guest ants was also associated with a reduced rate of extremity damage in G. hartmani workers (binomial GLZ, LR $\chi^2 = 14.18$, $P = 0.0002$), consistent with the fact that the agro-predator raiders were no longer attacked by S. amabilis defenders because they were killed by guest ants rather than by physical mutilation by the host ants. Ant drawings courtesy of Rozyn Haley.

In the more realistic scenario of mixed subcolonies (Fig. 3), the guest ants remained more effective at killing G. hartmani raiders than their S. amabilis hosts, who needed much larger numbers to mount at least some resistance against one or two G. hartmani intruders. (Compare Fig. 2A, in which eight S. amabilis workers were able to kill a single G. hartmani raider in two out of three trials, but at the cost of 63% mortality.) However, under field conditions, scouts of G. hartmani normally return to their nest to recruit a column of nestmates before initiating a raid. This may easily involve 100 or more G. hartmani (Fig. S2), which can quickly overwhelm an S. amabilis colony (21). Thus, hosting and feeding an M. symmetochus guest ant colony likely would have substantial fitness payoffs for S. amabilis when the risk of a raid by G. hartmani is high, and these benefits would be even greater if G. hartmani colonies preferred to raid S. amabilis colonies without guest ants.

To test this possibility, we conducted a y-tube choice experiment, which showed that G. hartmani preferentially initiated raids on S. amabilis colonies without guest ants relative to control colonies with guest ants (binomial GLZ, LR $\chi^2 = 18.12$, $P < 0.0001$) (Fig. 4, SI Text: Methods and Results, Fig. S2, Movie S3, and Table S3). Experimental Gnamptogenys scouts were allowed minimal contact with resident S. amabilis workers through a mesh separating their colonies, and they were never observed to have physical contact with guest ants. Thus, it is likely that the volatile components of the M. symmetochus venom were the main factors deterring raids (Fig. 4). Fig. 1B depicts a worker projecting its sting to disperse two isomers of 3-butyl-5-hexylpyrrolizidine (Fig. S3). SI Text: Methods and Results provides information on chemical analysis. This finding suggests that direct contact with M. symmetochus defenders is not necessary for a G. hartmani scout to determine whether a S. amabilis colony is worth recruiting to. However, field colonies of M. symmetochus can have hundreds of workers that are spread out across all chambers of a host colony (Table S2). Furthermore, physical contact between a G. hartmani scout and an M. symmetochus defender is likely, given that M. symmetochus workers recruit nestmates from their deeper fungal cavity when an intruder is detected (Movie S2), further enhancing the prophylactic inhibition of G. hartmani raids.

Our results confirm that socially parasitic M. symmetochus guest ants can serve as protective symbionts of their S. amabilis hosts. This development is remarkable because, despite the parasitic (i.e., maintenance) costs, the relationship between the host and guest ants has shifted to a context-dependent mutualism in which the cost to the host is compensated for by a secondary protection benefit against a shared natural enemy. It has turned an interaction governed by negative antagonistic selection into one characterized by positive reinforcement, allowing the guest ants to become unusually common (in 73% of host nests on average; SI Text: Methods and Results and Table S2). This finding reinforces the concept that mutualistic interactions are actually driven by mutual exploitation (23), and that the outcomes can be mutualistic win-win situations under certain conditions and parasitic win-lose situations in other circumstances. This variation may be a key factor in the coevolution of such interactions, and is one of the cornerstones of the geographic mosaic theory of coevolution (24, 25).

Our results suggest that M. symmetochus guest ant prevalence should be positively correlated with Gnamptogenys agro-predator density across sites, as has been shown in other protective symbionts (26, 27). We would also expect the guest ant M. adamsae and its host T. zeteki to suffer much less from Gnamptogenys raids given the typically much lower host colony infection rates (~1–6%; ref. 11), but we lack the comparative data needed to test this possibility. This dynamic coevolutionary scenario would seem to be conditional on each of the partners being largely or fully dependent on the others, with attine ants rearing a peculiar food source for which the two parasitic ants compete, one (Gnamptogenys) as a destructive and highly virulent agro-predator and the other (Megalomyrmex) as a milder and chronic disease. However, this tripartite interaction likely owes its evolutionary stability to the milder parasite’s alkaloid weaponry, which can control the more virulent predators without major cost.
Although pyrrolizidines are not unique to *M. symmetochus* and have convergently evolved in other ant genera ([28](#)), our study illustrates that such alkaloids are detectable and functional in interactions with ants from a distant subfamily, the Ectatomminae (Fig. 1). Previously, ant alkaloids were considered general repellents used during competitive intraspecific interactions and thief ant raids ([29](#) and references therein). Such broad functionality remains compatible with specific effects on the *Gnamptogenys* raiders as long as they remain vulnerable to these venoms.

Examing the different strategies of the two exploiters may hold the key to understanding their stable coexistence. *S. amabilis* sites without *G. hartmani* raiders would be influenced by the maintenance costs of infection with *M. symmetochus* guest ant parasites, which would tend to reduce *S. amabilis* densities and impose selection on traits that would decrease host colony susceptibility to guest ant infiltration (e.g., queen aggression, detoxifying enzymes effective against *Megalomyrmex* alkaloids). This would make the mildly virulent chronic guest ant rare while at the same time creating ideal conditions for the *Gnamptogenys* raiders to invade. Such population invasion would shift selection on the host to allow more frequent guest ant infiltration. Colonies with protective guest ant symbionts would then increase in the population, providing fewer attractive host colonies for *G. hartmani* to raid, which in turn would reduce the fitness of *G. hartmani*, making the agro-predators rare once again. Renewed selection against guest ant infiltrations would then be expected.

Another major factor in maintaining some form of dynamic equilibrium between the two social parasites and their shared host appears to be the life-long association of *M. symmetochus* with a single host colony, similar to the association between *M. adamsae* guest ants and their *Trachymyrmex* host colonies ([11](#)). This form of obligate perennial colony-level association tends to select for low virulence or prudent exploitation ([30](#)), implying this form of obligate perennial colony-level association tends to select for low virulence or prudent exploitation ([30](#)), implying the cost to host colonies of maintaining guest ants might quickly shift to a net benefit when more virulent and mobile alternative parasites appear. The characteristics of the association between *Megalomyrmex* guest ants and their hosts remains fundamentally antagonistic, however ([11](#)), explaining the aggression between *S. amabilis* host workers and their *M. symmetochus* guest ants seen in both the field and the laboratory. This effect is likely driven by a window of conflict over resource allocation, because the *S. amabilis* host can still realize some reproductive success by producing males and thus has no interest in allowing *M. symmetochus* guest ant colonies to grow much beyond the number of workers needed for optimal protection.

Overall, the dynamic interactions among the three ant species studied here resemble human military history. Many medieval cities maintained contingents of mercenary soldiers in times when mobile invasive armies posed a threat, despite their maintenance costs, which quickly became prohibitive after peace treaties were signed. Thus, both *M. symmetochus* guest ants and human mercenaries can be considered alien soldier castes that defend against larger evils as long as they are worth their keep. Another relevant analogy is the maintenance by heterozygote advantage of sickle cell anemia as a chronic human disease in areas where virulent malaria is endemic ([31](#)), that is, a chronic disease is maintained because it makes carriers resistant to a potentially lethal disease. Thus, we would expect *Sericomyrmex* populations without *Gnamptogenys* raiders to have a lower prevalence of *M. symmetochus*, because this situation would select for partial resistance against invasion by guest ants.

**Materials and Methods**

**Biological Material.** Four parasitized and four nonparasitized *S. amabilis* colonies, along with a single *G. hartmani* colony, were collected in May 2009, 2010, and 2011 near Gamboa and El Llano in the Republic of Panama (Table S1). All colonies were transferred to Copenhagen, Denmark and kept in an environmentally controlled rearing room at a constant temperature of 25 °C and relative humidity of 60–70%. Ant vouchers from all colonies used in this study are deposited at the Museum of Natural History, Smithsonian Institute, Washington, DC and at the Smithsonian Tropical Research Institute, Balboa, Republic of Panama.

**Guest Ant Venom Function.** Pilot experiments were staged, introducing the *G. hartmani* colony into guest ant-infested *S. amabilis* subcolonies to establish whether guest ants exhibit defensive reactions. The observation of intracolony conflict of *Gnamptogenys* raiders prompted a more controlled study of pairwise interactions between two *G. hartmani* workers ([SI Text: Methods and Results](#)). The treatment consisted of a *Megalomyrmex* “stung” individual introduced to a naïve *G. hartmani* worker (*n* = 6; Fig. S1 C and D). Avoidance or attraction behavior was scored for 1 h. Control experiments were designed similarly, except that the introduced worker was rubbed with empty soft forceps rather than with a live *Megalomyrmex* worker stinger.

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**Fig. 4.** A *G. hartmani* laboratory colony was given repeated choices between four pairs of *S. amabilis* host colonies with or without *M. symmetochus* guest ants (size-matched so that total number of ants, fungus garden volume, and nest box size were approximately equal). Gray bars represent the number of replicate trials for each pair, and dark-colored bars represent raids that were initiated after single *G. hartmani* scouts had inspected one or both of the maize separations with the experimental *S. amabilis* colonies. Recruited columns of raiding *G. hartmani* were preferentially directed toward *S. amabilis* colonies without guest ants (binomial GLZ, LR χ² = 18.12, *P* < 0.0001). Chemical structures (Upper Left) represent the *M. symmetochus* venom compounds (5Z,8E)-3-butyl-5-hexylpyrrolizidine and (5E,8E)-3-butyl-5-hexylpyrrolizidine, detected from air samples. Ant drawings courtesy of Rozlyn Haley.
The proportion of time that the workers spent in close proximity was compared using Welch’s t test (allowing for heterogeneous variances).

**Defense Efficiency in Two-Species Interactions.** A single starved G. hartmani scout worker and two, three, four, six, or eight S. amabilis or M. symmeto- toechus worker opponents (10 combinations in all) were placed in a small arena, and mortality was assessed after 24 h. The experiment was repeated using subcolonies derived from four source colonies: Mb, Mc, Md, and Me (SI Text: Methods and Results and Table S1). Mortality was analyzed using JMP version 9.02 (SAS Institute) to fit a generalized linear model with binomial errors. Opponent type and number of opponents were fitted as main effects, together with their interaction. Because the G. hartmani mortality data had quasi-complete separation, we performed Firth-adjusted maximum likelihood analysis.

**Three-Species Interactions, Survival, and Mutilation.** Subcolonies consisting of 18 S. amabilis workers, a fungus garden fragment ca. 1.5 cm in diameter, and zero, three, or six M. symmeto- toechus workers were set up in medium-sized Petri dishes from three parasitized source colonies: Mb, Mc, and Md (SI Text: Methods and Results and Table S1). Two starved G. hartmani workers were introduced into each Petri dish, and ant mortality and damage to G. hartmani worker appendages were assessed after 1 h and 24 h. Mortality and damage were analyzed with JMP using generalized linear models with binomial errors, correcting for overdispersion of the data as necessary.

**Raid Preference Choice: Parasitized vs. Nonparasitized.** With the use of a bifurcating olfactometer (i.e., y-tube), a laboratory colony of Gnamptogenys hartmani was given the choice of recruiting nestmates to size-matched pairs of parasitized or nonparasitized S. amabilis colonies (Fig. S2). Colony combinations (Mb+5b, Mc+Sc, Md+5d, and Me+Se; Table S1) were tested 11, 24, 29, and 23 times, respectively (SI Text: Methods and Results and Table S3). A mesh screen at the entrance of each S. amabilis colony allowed airflow and minimal (antennal) interactions between G. hartmani scouts and S. amabilis workers, and prohibited contact between G. hartmani scouts and M. sym- meto- toechus guest ants. Where raids occurred, their direction (accumulating G. hartmani workers on one side; Fig. S1 and Movie S3) was scored blindly from video recordings and analyzed using a generalized linear model with binomial errors to test for overall bias toward or away from parasitized nests, taking into account any differences between colony pairs. Firth-adjusted maximum likelihood estimates were used, because there were no raids on parasitized nests in two of the four pairs of colonies.

**Volatile Chemical Analyses.** All three species were extracted in methanol and chemically analyzed for volatile compounds by GC-MS following established methods (32). To determine whether the venom alkaloids were dispersed in the air by M. symmeto- toechus workers, headspace analysis using a solid-phase microextraction (SPME) fiber assembly carboxen/polydimethylsiloxane (57318 SUPELCO; Sigma-Aldrich) was conducted on a sample of eight M. symmeto- toechus ants from colony Me (SI Text: Methods and Results). For comparison, 10 ants and a small amount of fungus garden from a nonparasitized colony (RMMA100611-03; Table S1) were analyzed as well.

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