Chemical strategies to deal with ants: a review of mimicry, camouflage, propaganda, and phytomimesis by ants (Hymenoptera: Formicidae) and other arthropods

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Abstract



Chemical tactics by ant social parasites, including myrmecophiles, often relate to ant nestmate recognition and alarm communication. The strict nestmate recognition system in ants can be disrupted by chemical imitation of the nestmate recognition pheromone, which consists of cuticular hydrocarbon components. Social parasites often acquire these components through direct body contact, but occasionally synthesize them even before ant adoption. Such an imitation of the host cuticular chemicals causes species-specific adoption of the parasites, which are then often taken care of by ants for long terms. In contrast, transient invaders often use a propaganda allomone that induces panic alarm responses in ants. The allomone occasionally even causes fighting among nestmate ants, and seems to disrupt the ant nestmate recognition. These two chemical strategies are to modify the ant responses after ant detection. A third chemical strategy taken by some insect species is to avoid the detection itself, and is evidenced in the chemical phytomimesis by geometriid twig-like caterpillars. Since this counts upon the ants not to respond to the "invasion", it usually does not cause visible responses when it works effectively. Appropriate evaluation methods are necessary to evaluate the ant responses induced by the parasites to reveal the underlying mimetic strategy.

Key words: Chemical mimicry, camouflage, propaganda, crypsis, phytomimesis, cuticular hydrocarbons, review.

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Introduction

Ants are generally omnivorous or carnivorous, and characterized by their organized social structure with labor division. They have well-developed chemical communication systems that maintain their societies. Ants are therefore formidable predators of various animals, particularly mobile terrestrial insects. Numerous insect groups, including ants, must avoid ant attack to survive (ELMES 1996), and many have developed various chemical defenses. It is likely that such chemical tactics are effective because ants rely on chemical signals to sense their environment. In interactions with ants, survival strategies are roughly separated into two classes: avoiding ant detection and dealing with ants after being detected (Tab. 1). The former can be an effective strategy to avoid ant attack, and "crypsis" is known as a primary defense mechanism in animals in general (ED-MUNDS 1974, RUXTON & al. 2004). Chemical crypsis can be confirmed in the animals that adopt this strategy. In contrast, there are at least two different strategies to deal with ants after detection. One is based on evasion, including socalled "propaganda", which is adopted by several social parasitic ants and wasps (REGNIER & WILSON 1971, BLUM & al. 1980, LLOYD & al. 1986, THOMAS & al. 2002). The other strategy is to pose as nestmates for acceptance by the host ants. This strategy is presumably the most frequent and best studied of the interactions between myrmecophiles and host ants.

To understand the background of these three strategies, it is important to review briefly how the ants rely on chemical communication to sense their biotic environment. Since ants live in colonial societies with numerous members, they are always required to communicate in order to maintain their social life (HÖLLDOBLER & WILSON 1990, YAMAOKA 1990, VANDER MEER & ALONSO 1998, LENOIR & al. 1999). Ant colony members can usually discriminate their nestmates from alien species and conspecific foreigners. With nestmates, ants can discriminate their task, caste and fertility. Outside the nest, they judge food sources, share food information, warn of foraging dangers, and recruit nestmates if necessary. The means of these communications are chemical signals that operate in a sophisticated communication network. But this reliance on chemical signals has also enabled myrmecophiles and social parasites to develop adaptive chemical tactics to deceive the ants (DETTNER & LIEPERT 1994, VANDER MEER & MOREL 1998, LENOIR & al. 2001). Ironically, the strict exclusivism of the ant colonies provides social parasites safe ecological niches if they successfully deceive hosts with their chemical tactics.

This is a review of the chemical strategies that surround ants, with reference to my research on myrmecophilous insects (AKINO & al. 1996, 1999, AKINO 2002, EL-MES & al. 2002), parasitoid wasps (AKINO & YAMAOKA

Category	What to achieve ultimately	Chemical strategy	Proximate effect
Primary defense	Avoid detection by ants	Chemical mimesis (including chem- ical phytomimesis)	Being "invisible" by back- ground-matching
Secondary defense	Avoid attacks by ants	Chemical mimicry and camouflage	Pretending to be nestmates
Attack	Induce panic alarm response in host ants	Propaganda	Disturbing the host nest- mate recognition

Tab. 1: Classification of main chemical strategies to deal with ants. See detailed explanations in the text.

1998, THOMAS & al. 2002), and phytomimetic caterpillars (AKINO & al. 2004a, AKINO 2005). There are excellent recent reviews on chemical strategies after ant detection (VANDER MEER & MOREL 1998, LENOIR & al. 1999, 2001, HOWARD & BLOMQUIST 2005). This review starts with the known strategies by myrmecophiles and ant social parasites, and then focuses on the chemical crypsis that is employed to avoid ant detection. The latter element has perhaps not yet received enough attention.

Pretending to be nestmates to avoid ant attack

The chemical mimicry and camouflage used to assume the identity of ant nestmates are the most frequent chemical strategies employed by ant parasites. Chemical mimicry entails the parasites' active synthesis of the host cues. In chemical camouflage, parasites obtain cues from the host through passive and active acquisition (HOWARD 1993, DETTNER & LIEPERT 1994). These strategies are not mutually exclusive and can coexist in the same species (LENOIR & al. 2001). They are secondary defenses because both change ant responses after host detection.

Ants have an ability to discriminate their conspecific nestmates from foreigners, and they generally share safety and food only among the formers. If social parasites successfully imitate nestmates of the host ants, they can avoid ant attack and also attain protection and food inside the nest. This strategy relies on ant nestmate recognition and has been applied by various myrmecophilous insects and social parasitic ants (VANDER MEER & al. 1989, HOW-ARD 1993, DETTNER & LIEPERT 1994, AKINO & al. 1996, 1999, AKINO & YAMAOKA 1998).

Ant nestmate recognition has been considered to be based on chemical signals, i.e., "nestmate recognition pheromone". Nestmate recognition is common in eusocial insects, including not only ants but also bees, wasps, and termites. There are several recent reviews on the subject (BREED 1998, SINGER & al. 1998, VANDER MEER & MOREL 1998, LENOIR & al. 1999, 2001, HOWARD & BLOMQUIST 2005, HEFETZ 2007). There has been controversy as to which chemicals serve as the pheromone in each group. Cuticular hydrocarbons are considered to be the most likely components in ants (VANDER MEER & MOREL 1998, LENOIR & al. 1999, HOWARD & BLOMQUIST 2005, HEFETZ 2007).

Cuticular hydrocarbons as the nestmate recognition pheromone in ants

Hydrocarbons are major lipid compounds of the ant cuticular waxes, although relatively small amounts of polar lipids are also present. The hydrocarbon compositions generally differ among species, i.e., each species has its own composition. Furthermore, the relative amounts of each component of the cuticular hydrocarbon profiles within species are usually specific to colonies. In other words, colony members share almost identical profiles that differ from those of conspecific foreigners. These two characters are the basis of the "hydrocarbon hypothesis" suggesting the active role of the cuticular hydrocarbons as nestmate recognition pheromones. This hypothesis is widely supported by many indirect correlative lines of evidence as recently summarized by HEFETZ (2007). However, much of this evidence is at the best circumstantial, based on either correlation studies or bioassays using removal and replacement of cuticular compounds by solvent extraction (BREED 1998). Behavioral experiments using artificial blends with authentic hydrocarbons are necessary to prove the critical link between cuticular hydrocarbons and nestmate recognition. However, ant cuticular hydrocarbons are generally complex mixtures of various methyl-branched alkanes, including mono-, di-, and tri-methyl alkanes (HOWARD & BLOMQUIST 2005). This complexity makes it difficult to conduct behavioral experiments in most ant species to test the "hydrocarbon hypothesis" by direct comparison of artificial and natural blends of the hydrocarbons.

This obstacle was, however, overcome in Japan's most common ant, Formica japonica, as reported by AKINO & al. (2004b). This species apparently has an ability of nestmate recognition, which is most likely based on cuticular hydrocarbon profiles (YAMAOKA 1990). The profiles were shared by workers and queens in each colony, but significantly differed between colonies (YAMAOKA 1990, AKINO & al. 2004b). It was fortunate that the cuticular hydrocarbons consisted of pairs of n-alkanes and respective (Z)-9alkenes of odd numbers from C25 to C33. Furthermore, the absence of branched alkanes makes the composition much simpler than that in many other ant species. This simple composition enabled us to prepare all the hydrocarbons commercially and synthetically, and also to obtain artificial hydrocarbon blends that matched natural blends. Complete artificial blends of n-alkanes and 9-alkenes, as well as the natural blends of a certain F. japonica colony, caused worker aggression in foreign colony workers but not in nestmate workers. This indicates that those hydrocarbons actually serve as the nestmate recognition pheromone in F. japonica, and that these ten hydrocarbons are sufficient for nestmate discrimination. In contrast, single blends of either n-alkanes or (Z)-9-alkenes caused no aggression in foreign workers. This diminishes the possibility that single active compounds can activate nestmate recognition, but suggest that it is the combinations of these two classes of hydrocarbons that provide this function. Similarly, nestmate recognition relies not on particular hydrocarbon components but complex hydrocarbon profiles in *Linepithema humile* and *Aphaenogaster cockerelli* (GREENE & GORDON 2007). In contrast, the nestmate recognition in *Formica exsecta* does depend on particular components (MARTIN & al. 2008).

Simple hydrocarbon compositions are rare in ant species. It is difficult to test the "hydrocarbon hypothesis" in most ant species with a classical pheromonal approach. Instead, a neurophysiological approach was recently attempted for Camponotus japonicus, and demonstrated the existence of particular sensilla on the antennae that specifically responded to foreign cuticular hydrocarbon profiles (OZAKI & al. 2005). This indicates that the sensilla can discriminate between nestmate hydrocarbon profiles and foreign hydrocarbon profiles immediately after direct contact, although the details of such a physiological perception mechanism remain unknown. Such an immediate discrimination of the minute differences in the complex hydrocarbon profiles is considered to be necessary for nestmate recognition behavior, which is induced immediately after body contact between individual workers. Therefore, the existence of such sensilla strongly supports the "hydrocarbon hypothesis". Thus, a neurophysiological approach can be quite effective to verify the "hydrocarbon hypothesis" and to explore if a similar perception system is common in ant species.

On the other hand, recent studies suggest additional functions of the ant cuticular hydrocarbons, e.g., signals for caste, task, and fertility recognition (reviewed by HEFETZ 2007). However, most of them are based on quantitative comparison of the cuticular hydrocarbons that correspond to ant behavior and physiological states, and few include hydrocarbon manipulation. It is necessary to organize results in ways that illuminate the details of caste, task, and fertility recognition. Other verification methods to test the hypothesis directly should also be developed. Although the classical pheromonal study approach is presumed to be effective for verification of the expanded "hydrocarbon hypothesis", the complexity of the ant cuticular hydrocarbon compositions prevents the execution of behavioral bioassays with pure chemicals.

Chemical mimicry and camouflage by cuticular hydrocarbons

It is most likely that the chemical signal for ant nestmate recognition consists of cuticular hydrocarbons, as indicated by many lines of circumstantial evidence and several direct ones. Chemical mimicry (in which the parasite actively biosynthesizes the host cues) and camouflage (in which the parasite obtains cues from the host both by passive and active acquisition) are practiced by various insects that take advantage of ant social systems (VANDER MEER & al. 1989, HOWARD 1993, DETTNER & LIEPERT 1994, AKINO & al. 1996, 1999, AKINO & YAMAOKA 1998, 2000, AKINO 2002). This includes not only myrmecophilous insects but also social parasitic ants. The ethological role of cuticular hydrocarbons in this strategy in ants has often been highlighted (HOWARD 1993, DETTNER & LIEPERT 1994, SINGER & al.

1998, VANDER MEER & MOREL 1998, LENOIR & al. 1999, 2001, HOWARD & BLOMQUIST 2005).

Chemical camouflage

The cases of the tiny *Myrmecophilus* cricket and the aphidiid wasp *Palaripsis eikoae* are noteworthy as examples of chemical camouflage because of the chemical adjustment ability and the characteristic behavior for acquiring the ant hydrocarbons, respectively.

The species ecology of the myrmecophilous cricket Myrmecophilus sp. is closely connected to several ant species. It is believed that this cricket has an ability to switch host ant species. Comparative GC-MS analyses revealed resemblances in the cuticular hydrocarbon compositions and profiles between the crickets and their corresponding host ants (AKINO & al. 1996). Transfers of the crickets between conspecific colonies resulted in initial aggression by host ants against the foreign crickets, but this decreased within a week. This was also confirmed in transfer of the crickets between different species, for example, between Lasius nipponensis and Formica japonica, and between F. japonica and Lasius fuji. The crickets escaped from the initial ant attacks through physical agility. Then they approached the ant workers repeatedly, and successfully adjusted their cuticular hydrocarbon compositions and profiles to fit those of the new host ant species and colonies within approximately a week. This ability to adjust their cuticular hydrocarbons requires direct body contact with the host ants that show high hostility at the beginning. Preventing direct body contact with the host ants results in drastic decrease of ant-like hydrocarbons in the crickets. Such chemical resemblance by acquisition is known in the beetles Myrmecaphodius excavaticollis (see VANDER MEER & WOJCIK 1982), Pella comes and Diaritiger fossulatus (see AKINO 2002), and the spider Cosmophasis bitaeniata (see ELGAR & ALAN 2006).

Another example is the aphidiid parasitoid wasp, Paralipsis eikoae, which is a specialist parasitoid of the wormwood root aphid Sappaphis piri tended by Lasius niger, as reported by TAKADA & HASHIMOTO (1983, 1985). Presumably because the wasp requires upbringing by ants to mature its ovaries after eclosion, it acquires the ant cuticular hydrocarbons through direct body contact after invading the ant colony to let the ants feed it (AKINO & YAMA-OKA 1998). The direct body contact occurs as the wasp mounts on the ant's back and then repeatedly rubs the ant abdomen with its midlegs in "rubbing" behavior (TAKADA & HASHIMOTO 1983). The Lasius ants do not move during this process, as if they were paralyzed. The wasp may use some appeasement signals to facilitate this docility (this will be discussed in a later section). After the rubbing behavior, cuticular hydrocarbon compositions and profiles of the wasps change and become similar to those of the host Lasius ants (AKINO & YAMAOKA 1998). Thus, P. eikoae acquires the host ant cuticular hydrocarbons, by which it avoids ant attack. In contrast, the related European species Paralipsis enervis is known to imitate the host ant aphid cuticular hydrocarbons to avoid attack by the aphid-attending ants (VÖLKL & al. 1996). This may be considered as a "background-matching" strategy (discussed in a later section), when the aphids are regarded as a background of the wasps, which enables avoidance of ant detection. Despite being related species, these two Paralipsis wasps have selected different modes of imitation to avoid ant attack.

Acquisition through direct contact facilitates the imitation of host ant hydrocarbons, but it also entails the risk and cost of attack by ants during the initial encounter. Therefore, it is also important for myrmecophiles and parasites to evade this initial danger. Myrmecophilous crickets avoid the initial ant attack by quickness and agility (AKINO & al. 1996), and the aphidiid parasitoid wasp *P. eikoae* presumably does so by rubbing behavior (TAKADA & HASHIMOTO 1983, AKINO & YAMAOKA 1998). In addition to such behavioral actions, chemical mimicry is often adopted as the strategy to avoid initial danger.

Chemical mimicry

Some myrmecophiles are known to exude chemical signals to appease the host ants before their direct interactions. Myrmecophilous lycaenid butterfly caterpillars possess antlike hydrocarbons even in the pre-adoption stages (AKINO & al. 1999, ELMES & al. 2002, SCHLICK-STEINER & al. 2004, SCHÖNROGGE & al. 2004). In the habitat of the Mountain Alcon Blue, Maculinea rebeli (hereafter treated as Phengaris alcon, following recent phylogenetic and taxonomic study by FRIC & al. 2007), there are several sympatric Myrmica ant species, but ELMES & al. (1998) showed Myrmica schencki to be the most frequent host. In some populations, M. schencki seems to be the only host that provides complete rearing for *P. alcon*, which requires the latter to actively seek adoption by M. schencki workers. Presumably, this causes the caterpillars to synthesize series of particular hydrocarbons that are specific to M. schencki even before adoption by the ants. M. schencki workers carried glass dummies treated with Phengaris pre-adoption hydrocarbons, as well as those with their own hydrocarbons (AKI-NO & al. 1999). Thus, the pre-adoption hydrocarbons seem to promise positive preferable interactions with M. schencki to the Mountain Alcon Blue caterpillar. Cuticular hydrocarbon resemblance between pre-adoption P. alcon and M. schencki was independently confirmed by different analyses (ELMES & al. 2002, SCHLICK-STEINER & al. 2004). Because the caterpillars successfully imitate complete sets of the host ant cuticular hydrocarbons instead of pre-adoption hydrocarbons after adoption, AKINO and co-workers (1999) hypothesized that the caterpillars acquired them through direct body contact with the host ant workers and brood. Further detailed studies on the chemical acquisition after adoption revealed, however, that the caterpillars not only acquire the ant hydrocarbons but also synthesize additional mimetic hydrocarbons (ELMES & al. 2002, SCHÖN-ROGGE & al. 2004). SCHÖNROGGE and co-workers (2004) revealed that the P. alcon caterpillars had an ability not only to acquire the host ant hydrocarbons but also to synthesize several hydrocarbons that matched those of their natural host *M. schencki*. According to NASH & al. (2008), there are geographic variations in such species-specific interactions on hydrocarbon resemblance between P. alcon and Myrmica ant species.

It is not certain, however, how the social parasites generate imitation of the host ant hydrocarbons, even when chemical resemblance is confirmed between the parasites and host ants. For example, caterpillars of the lycaenid butterfly *Niphanda fusca*, which is known to have a speciesspecific relation with *Camponotus japonicus*, imitate the host ant cuticular hydrocarbons after adoption (HOJO & al. 2007). It is curious that the cuticular hydrocarbon composition of the caterpillars specifically resembled those of male ants rather than those of worker ants, because the former rarely spend times with the latter inside the colonies. Such resemblances could not occur if the caterpillars acquired the hydrocarbons through direct body contact with the ant colony members. This suggests that the *N. fusca* caterpillars connately have an ability to synthesize host ant hydrocarbons and to adjust the profiles to be similar to those of male ants after adoption.

Cuticular hydrocarbon resemblances are also confirmed between social parasitic ants and host ants, e.g., between the cuckoo ant *Leptothorax kutteri* and its *L. acervorum* host (FRANKS & al. 1990), and between the slave maker *Polyergus* and its *Formica* slaves (YAMAOKA 1990, HAVER-SETZER 1993, HAVERSETZER & BONAVITA-COUGOURDAN 1993, BONAVITA-COUGOURDAN & al. 1996, 1997, D'ET-TORRE & al. 2002). Some of these parasitic ants lose the host-like cuticular hydrocarbons after isolation from host colonies (YAMAOKA 1990). Although this suggests that the parasitic ants do not synthesize but acquire the host-like hydrocarbons from the host ants, it does not invalidate an alternative hypothesis, in which the parasitic ants have an ability to control biosynthesis of their own cuticular hydrocarbons.

Chemical camouflage or mimicry?

The improvement of the gas chromatograph (GC) and gas chromatograph-mass spectrometer (GC-MS) has made it easier to confirm the cuticular chemical resemblance between host ants and their social parasites, but it is still difficult to identify its causes. Chemical tracing by isotope labeling with ¹⁴C might be helpful for this purpose. This technique has been applied to investigate the nestmate recognition signals of Cataglyphis niger (see SOROKER & al. 1994, 1995, SOROKER & HEFETZ 2000) and Pachycondyla apicalis (see SOROKER & al. 1998). Although it was originally developed to investigate hemolymph lipophorin in Locusta migratoria (see KATASE & CHINO 1984), Soroker and co-workers proved the technique's utility for ants. Injection of $[1-^{14}C]$ sodium acetate into the ant abdomen through the intersegmental membrane produces various ¹⁴C-labelled hydrocarbons in the cuticle. It appears to be necessary to inject such labelled sodium acetate into the social parasites' bodies to investigate whether they synthesize or acquire the host-like hydrocarbons. Further improvement of the technique may be necessary because the social parasites are usually much smaller than the host ants; nevertheless, it can be applied to identify the causes of the chemical resemblance in question.

Usage of repellent to avoid ant attack Repellent allomone

The repelling of ants is another effective way to avoid ant attack, even after ant detection. This strategy is also classified as secondary defense. It is often applied, not only by insects but also by plants. Such repellent chemicals may presumably become available for control of pest ant species. Leaf-cutting ants are generally polyphagous and attack various plants to obtain the leaves for culturing their symbiotic fungus. Several native plants, however, escape the leaf-cutter ant attack, because of their repellent chemicals. *Astronium graveolens* (Anacardiaceae) contains sev-

eral volatile chemicals that are effective repellents against leaf-cutter ants Atta cephalotes (see CHEN & al. 1984). Because most ants that prefer flower nectars are potential nectar thieves of insect-pollinated flowers (HERRERA & al. 1984), some plants contain ant repellent chemicals in the nectars (FEINSINGER & SWARM 1978, GUERRANT & FIED-LER 1981). Various insects, e.g., stinkbugs (Pentatomidae), secrete repellent chemicals that are effective against ants. Also, it is well established that Polistes paper wasps secrete repellent chemicals when constructing their nest petiole (POST & JEANNE 1981, KOJIMA 1993). These studies showed how petioles were coated with a rubbery chemical layer of secretion that included ant-repellent-like chemicals. When it was removed, ants easily invaded the wasp nests to steal brood and stocked food. Ant repellent impedes ant invasion and attack of both insects and plants.

Transitory intruders of ant societies also use ant repellent allomone to endure the attacks of other ants during intrusion. BLUM & al. (1980) demonstrated that the thief ant *Solenopsis fugax* emitted host ant repellent alkaloids from the poison gland to disperse brood-nursing workers away before robbing the host broods. Queens of the temporary social parasite ant *Bothriomyrmex syrius* emit volatile chemicals that match a host alarm pheromone. This distracts the host workers during the queen's invasion (LLOYD & al. 1986). Such repellent allomone was also found in bumblebees. Females of an obligate social parasite *Bombus* (*Psithyrus*) norvegicus possess large quantities of a pure volatile ester on the cuticle, which shows a strong repellent effect on host *Bombus* (*Pyrobombus*) hypnorum workers (ZIMMA & al. 2003).

Propaganda allomone

American slave maker Formica sanguinea workers emit mixtures of volatile alkyl ketone, alkane, and acetates when invading host ant nests. Those chemicals induce panic responses in defenders (REGNIER & WILSON 1971). Such offensive chemical signals are also found in the slave making ant Harpagoxenus sublaevis and the workerless inquiline ant Leptothorax kutteri (see ALLIES & al. 1986), the European slave making ant Polyergus rufescens (see VISICCHIO & al. 2000), and the Japanese slave maker Polyergus samurai (T. Akino, unpubl.). Although the effect of these chemicals is similar to that of repellent allomone, the host workers occasionally attack each other, as if they had lost the nestmate recognition signal of their colonies. REGNIER & WILSON (1971) therefore developed the term "propaganda" for these offensive signals, which should now also be classified as belonging to the allomone category of allelochemicals.

Such powerful propaganda chemicals are also utilized by the parasitoid wasp *Ichneumon eumerus* of the myrmecophilous lycaenid butterfly *Phengaris alcon* (THOMAS & al. 2002). Because the host *P. alcon* larvae are tended inside the *Myrmica schencki* nest, the wasp must invade the ant for its oviposition even though ant attack responses are inevitable. Even if the wasp succeeds in oviposition, the parasitized *P. alcon* larvae remain to grow inside the ant nest, and the wasp offspring must emerge inside the ant nest. The wasp needs to avoid ant aggression to survive, so the wasp emits a series of allyl alcohols and aldehydes, which cause *M. schencki* workers to attack each other instead.

Appeasement allomone

In the American slave maker ant *Polyergus breviceps* (TOP-OFF & al. 1988) and the European slave maker *P. rufescens* (D'ETTORRE & al. 2000, MORI & al. 2000a, b), the Dufour's gland secretion is found not to induce panic alarm but to reduce aggression of the resident workers. The secretion is therefore named appeasement allomone.

Both propaganda and appeasement allomones commonly disrupt nestmate recognition in the host ant species, but the physiological mechanism is unknown. Because nestmate recognition is the most basic communication to maintain ant societies, such allomone chemicals might be used as social communication disruption agent against harmful ants.

Chemical phytomimesis: ant "invisibility"

The basis of propaganda and appeasement allomones is to modify ant reactions by chemical signals after ant detection. These strategies aim to deter attacks by ants that detect the intruders. In contrast, the third strategy is basically to avoid ant detection itself. This strategy presumably exists in animals that do not have necessity to maintain long interactions with ants, and it includes chemical backgroundmatching.

Background-matching is one of several common strategies in animals to avoid predator attack (EDMUNDS 1974, ENDLER 1988). There are many reports on animals that seem adapted to physiologically match their visual appearance to their environment (GREENE 1989, MCFALL-NGAI & MORIN 1991, MESSENGER 1997, CHIAO & HANLON 2001), and that behaviorally select backgrounds that match their appearance (ENDLER 1984, MARSHALL 2000). It is likely that such background-matching in animals could reduce predation risk (FELTMATE & WILLIAMS 1989). These studies are based on visual matching, because the predators seek and capture preys through visual information.

The case of the giant geometer, *Biston robustum*, can be a good example for chemical background-matching, or phytomimesis. Presumably, this occurs to avoid ant attack, though it is also a typical example of visual backgroundmatching (AKINO & al. 2004a, AKINO 2005). This species is a close relative of B. betularia, well known for industrial melanism (KETTLEWELL 1955), and is univoltine with a long larval stage of more than 5 months from the end of March to early October in Japan (NAKASHIMA 1912). Larvae are typical twig-like caterpillars that pretend to be motionless host plant twigs in the day. A key to their survival is background-matching. Their visual appearance strongly resembles the host plant twigs, and is characterized by a combination of color, the presence of horns and markings on the body that look like buds and scars on the twigs, the absence of abdominal legs except for hind grasping pairs, and a habit of resting in an unusual position with the body stretching out at an angle from the branch. The caterpillars change their appearance according to their host plant species including colors and marks (AKINO & al. 2004a). They also change their feeding and resting behaviors as they grow (AKINO & al. 2004a, AKINO 2005). It is highly likely that such visual phytomimesis is effective to avoid avian predators as generally suggested (RUITER 1958, HEINRICH 1993).

Their motionless life style that avoids feeding in the day is presumably compensated by long larval periods. How-

ever, it increases the danger of becoming prey to terrestrial predators, including ants such as Lasius, Formica, Camponotus, Crematogaster, and Pristomyrmex. These ant forager workers were often observed on the tree twigs where the caterpillar perched in the day, but they rarely attacked the latter even when walking on them (AKINO & al. 2004a). Passing over the caterpillars' bodies initiated no interest in the ants, even after antennation, very much as if they were dealing with real twigs. Comparative GC and GC-MS analyses revealed that the caterpillars possessed cuticular chemicals identical to those of host-twig surfaces, in both composition and relative ratios. The caterpillar and the tree twig are indistinguishable by their surface chemicals, which suggests that the ants cannot discern between the two. Additional host plant exchange experiments indicated the ability of the caterpillars to adjust their cuticular chemicals to match new host plant surface chemicals, although this required molting(s). As this cuticular chemical adjustment required feeding on the new host plant leaf, it is obviously triggered by diet and ingestion (AKINO & al. 2004a, AKINO 2005). All of the caterpillars tested preferred to perch on twigs with the surface chemicals that they were imitating at the time (AKINO & al. 2004a). Thus, the caterpillars have a potential ability to confuse the ant workers even though they have excellent chemical sense.

Diet-induced visual phytomimesis is also known in another geometer, Nemoria arizonaria (GREENE 1989). The caterpillars of the spring brood develop into mimics of the oak catkins which they feed upon, while those of the summer brood develop into mimics of oak twigs. This developmental polymorphism is considered to be triggered by the variable concentration of a defensive secondary compound, tannin, in the larval diet, and it enables the visual phytomimesis of this caterpillar. Another example of diet-induced body color change is reported in the spider Theridion grallator (GILLESPIE 1989). These diet-induced visual polymorphisms may be more widespread in herbivores than we currently know, because of high predation pressure by avian and mammalian predators that rely on visual information. It is adaptive for herbivores to be able to evade predation through visual background-matching. Dietary cues are closely related to the native habitat for herbivores (WOODHEAD & CHAPMAN 1986, MALONEY & al. 1988, VARELA & BER-NAYS 1988), so they can be a useful signal to induce appropriate background-matching in morphs and behaviors for predator avoidance.

This reasoning also can apply to diet-induced chemical background-matching, where the predators rely on chemical instead of visual information to seek their prey. Ants typify this mechanism (e.g., ESPELIE & BERNAYS 1989), and provide high predation pressure to herbivores. The case of *Biston robustum* shows sophisticated chemical phytomimesis to avoid ant detection because of the ability to chemically adjust to various host plants (AKINO & al. 2004a). Although there are few reports on chemical phytomimesis in herbivores, it may be widespread to avoid the high predation pressures of ants.

Such chemical "invisibility" seems effective to avoid ant attack, and chemical phytomimesis is not the only context in which it should be considered. The absence of a cuticular chemical signal in freshly emerged ants is another phenomenon that precludes adult ant attack (LENOIR & al. 1999, 2001). Freshly emerged callow ants usually possess

small amounts of hydrocarbons that are not specific to colonies, and low levels of the cuticular hydrocarbons presumably reduce aggression of the adult workers. Different kinds of background, including inorganic substances, would be available for chemical mimesis, but there are no reports on such background-matching that I am aware of. However, preliminary experiments suggest that myrmecozelinid caterpillars might use such chemical mimetic tactics (T. Akino, unpubl.). The caterpillars usually hide inside their cocoons, and are often found near ant nests. The ant workers aggressively attack the naked caterpillars, but pay no attention to the cocoons. These responses were also confirmed by the respective chemical extracts. These observations are still insufficient to argue that this ant unresponsiveness is the result of chemical mimetic tactics of the caterpillars, although it is suggestive. A similar phenomenon is confirmed in ant interactions between lycaenid larvae and ants (Y. Hagiwara, pers. comm.). Probably we need to identify which chemicals trigger / suppress ant aggressive responses, and to construct proper bioassay methods to evaluate "unawareness" of ants for further clarification of this topic. Such verification might suggest generalities of this "silent" chemical mimesis by insects that appear to have less interaction with ants.

Where to from now?

Chemical mimesis is presumably one of the effective strategies for animals to avoid ant attacks because ants are highly sensitive to chemical signals. The animals that use chemical mimesis (including phytomimesis) are difficult to identify because they do not always necessarily use visual mimesis simultaneously. *Biston robustum* caterpillars happen to use both visual and chemical phytomimesis (AKINO & al. 2004a). Visual mimesis is visually obvious to human observers, but most animals using chemical mimesis may not have been identified.

For example, small white butterfly *Pieris* caterpillars are often hunted by various ants. In contrast, the pupae are relatively untouched in spite of their immobilized vulnerability; it is not known why. These pupae do not contain any toxic compounds, though several aposematic lepidopteran pupae do so to avoid predation (NISHIDA 1994). According to KANEKO & KATAGIRI (2004), the cuticular wax components of *Pieris* pupae mainly consist of odd-numbered nalkanes. One may conjecture that since such simple sets of hydrocarbons are generally found in the plant surface chemicals, they may efficiently confuse ants seeking prey. Thus, it is necessary to compare the surface chemicals of the animals not preyed upon by ants with the environmental agent that can become the background in order to verify the possibility of a chemical mimetic strategy.

However, there is another difficulty for the study of chemical mimesis even after chemical resemblance is successfully confirmed between the target animals and their background, namely that of a proper evaluation by behavioral bioassays. A proper bioassay would identify this "unawareness" by lack or presence of chemical signals. If we properly focus on this phenomenon, comparative analyses of chemicals and behavior in herbivores, their host plants, and corresponding predators may alter our recognition of chemical phytomimesis.

Because ants are so abundant, and sensitive to chemical signals, the chemical mimicry, camouflage, and propaganda in insects that have close interactions with ants have been well studied and summarized. These strategies are very impressive because they modify the ant behavioral responses very clearly after the ants detect intrusion and also involve great risk if the modifications fail to occur. The chemical mimetic strategy is less dramatic, as the ants usually do not present clear responses to the social parasites. This is presumably one of the reasons why the strategy appears to have garnered little attention.

Since ants generally pose a risk of harm for most insects, the mimetic strategy would be safer than applying chemical mimicry, camouflage, and repellents. Despite the difficulty of evaluation, the chemical features to mimetic strategists should be carefully evaluated to their "silent odors" against ants.

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Zusammenfassung

Chemische Strategien von Sozialparasiten von Ameisen, einschließlich Myrmecophile, knüpfen oft an die Nestgenossenerkennung und die Alarmierungskommunikation der Ameisen an. Das strikte Nestgenossenerkennungssystem von Ameisen kann durch die chemische Imitation des Nesterkennungspheromons geknackt werden, welches aus kutikulären Kohlenwasserstoffen besteht. Sozialparasiten eignen sich diese Substanzen oft durch direkten Körperkontakt an, aber synthetisieren sie in selteneren Fällen sogar noch vor der Adoption durch die Ameisen. Solche Imitation wirtsspezifischer, kutikulärer chemischer Substanzen bedingt artspezifische Adoption der Parasiten, welche dann häufig von den Ameisen langfristig umsorgt werden. Im Gegensatz dazu setzen nicht-permanente Invasoren häufig Propagandaallomone ein, die bei den Ameisen panische Alarmreaktionen hervorrufen. Die Allomone lösen fallweise sogar Kämpfe zwischen Ameisennestgenossen aus und scheinen die Nestgenossenerkennung der Ameisen zu stören. Die beiden eben umrissenen chemischen Strategien zielen darauf ab, die Reaktion der Ameisen nach der Entdeckung durch die Ameisen zu modifizieren. Eine dritte chemische Strategie, die sich bei manchen Insekten findet, ist, die Entdeckung selbst zu vermeiden, nachgewiesen etwa für die chemische Phytomimese von zweigähnlichen Geometridenlarven. Da diese Strategie bewirkt, dass die Ameisen nicht auf die "Invasion" reagieren, ruft sie normalerweise keine sichtbare Antwort hervor, wenn sie effektiv ist. Geeignete Evaluierungsmethoden werden notwendig sein, um die Reaktion von Ameisen auf die Parasiten zu ergründen und so die zugrundeliegende mimetische Strategie im Detail zu klären.

References

AKINO, T. 2002: Chemical camouflage by myrmecophilous beetles *Zyras comes* (Coleoptera: Staphylinidae) and *Diaritiger fossulatus* (Coleoptera: Pselaphidae) to be integrated into the nest of *Lasius fuliginosus* (Hymenoptera: Formicidae). – Chemoecology 12: 83-89.

- AKINO, T. 2005: Chemical and behavioral study on the phytomimetic giant geometer *Biston robustum* BUTLER (Lepidoptera: Geometridae). – Applied Entomology and Zoology 40: 497-505.
- AKINO, T., KNAPP, J.J., THOMAS, J.A. & ELMES, G.W. 1999: Chemical mimicry and host specificity in the butterfly *Maculinea rebeli*, a social parasite of *Myrmica* ant colonies. – Proceedings of the Royal Society of London Series B 266: 1419-1426.
- AKINO, T., MOCHIZUKI, R., MORIMOTO, M. & YAMAOKA, R. 1996: Chemical camouflage of myrmecophilous cricket *Myrmecophilus* sp. to be integrated with several ant species. – Japanese Journal of Applied Entomology 40: 39-46. (in Japanese)
- AKINO, T., NAKAMURA, K. & WAKAMURA, S. 2004a: Diet-induced chemical phytomimesis by twig-like caterpillars of *Biston robustum* BUTLER (Lepidoptera: Geometridae). – Chemoecology 14: 165-174.
- AKINO, T., YAMAMURA, K., WAKAMURA, S. & YAMAOKA, R. 2004b: Direct behavioural evidence for hydrocarbons as nestmate recognition cues in *Formica japonica* (Hymenoptera: Formicidae). – Applied Entomology and Zoology 39: 381-387.
- AKINO, T. & YAMAOKA, R. 1998: Chemical mimicry in the root aphid parasitoid *Paralipsis eikoae* YASUMATSU (Hymenoptera: Aphidiidae) of the aphid attending ant *Lasius sakagamii* YA-MAUCHI & HAYASHIDA (Hymenoptera: Formicidae). – Chemoecology 8: 153-161.
- AKINO, T. & YAMAOKA, R. 2000: Evidence for volatile and contact signals of nestmate recognition in the black shining ant *Lasius fuliginosus* LATREILLE (Hymenoptera: Formicidae). – Entomological Science 3: 1-8.
- ALLIES, A.B., BOURKE, A.F.G. & FRANKS, N.R. 1986: Propaganda substances in the cuckoo ant *Leptothorax kutteri* and the slave-maker *Harpagoxenus sublaevis*. – Journal of Chemical Ecology 12: 1285-1293.
- BLUM, M.S., JONES, T.H., HÖLLDOBLER, B., FALES, H.M. & JAOUNI, T. 1980: Alkaloidal venom mace: offensive use by a thief ant. Naturwissenschaften 67: 144-145.
- BONAVITA-COUGOURDAN, A., BAGNÈRES, A.-G., PROVOST, E., DUSTICIER, G. & CLÉMENT, J.L. 1997: Plasticity of the cuticular hydrocarbon profile of the slave-making ant *Polyergus rufescens* depending on the social environment. – Comparative Biochemistry and Physiology 116B: 287-302.
- BONAVITA-COUGOURDAN, A., RIVIÉRE, G., PROVOST, E., BAGNÈ-RES, A.-G., ROUX, M., DUSTICIER, G. & CLÉMENT, J.L. 1996: Selective adaptation of the cuticular hydrocarbon profiles of the slave-making ants *Polyergus rufescens* LATR. and their *Formica rufibarbis* FAB. and *F. cunicularia* LATR. slaves. – Comparative Biochemistry and Physiology 113: 313-329.
- BREED, M.D. 1998: Chemical cues in kin recognition: criteria for identification, experimental approaches, and the honey bee as an example. In: VANDER MEER, R.K., BREED, M.D., ESPELIE, K. & WINSTON, M.L. (Eds.): Pheromone communication in social insects: Ants, wasps, bees and termites. – Westview Press, Boulder, CO, pp. 57-78.
- CHEN, T.K., WIEMER, D.E. & HOWARD, J.J. 1984: A volatile leafcutter ant repellent from *Astronium graveolens*. – Naturwissenschaften 71: 97-98.
- CHIAO, C.-C. & HANLON, R.T. 2001: Cuttlefish camouflage: visual perception of size, contrast, and number of white squares in artificial checkerboard substrate initiates disruptive coloration. – Journal of Experimental Biology 204: 2119-2125.
- DETTNER, K. & LIEPERT, C. 1994: Chemical mimicry and camouflage. – Annual Review of Entomology 39: 129-154.
- D'ETTORRE, P., ERRARD, C., IBARRA, F., FRANCKE, W. & HE-FETZ, A. 2000: Sneak in or repel your enemy: Dufour's gland

repellent as a strategy for successful usurpation in the slavemaker *Polyergus rufescens.* – Chemoecology 10: 135-142.

- D'ETTORRE, P., MONDY, N., LENOIR, A. & ERRARD, C. 2002: Blending in with the crowd: social parasites integrate into their host colonies using a flexible chemical signature. – Proceedings of the Royal Society of London B 269: 1911-1918.
- EDMUNDS, M. 1974: Defense in animals. Longman Group Ltd., Essex, 357 pp.
- ELGAR, M. & ALAN, R.A. 2006: Chemical mimicry of the ant Oecophylla smaragdina by the myrmecophilous spider Cosmophasis bieniata: Is it colony-specific? – Journal of Ethology 24: 239-246.
- ELMES, G.W. 1996: Biological diversity of ants and their role in ecosystem function: In: LEE, B.H., KIM, T.H. & SUN, B.Y. (Eds.): Biodiversity research and its perspectives in East Asia.
 Proceedings of the International Inaugural Seminar of the Korean Institute for Biodiversity Research (KIBIO): 33-48.
- ELMES, G.W., AKINO, T., THOMAS, J.A., CLARKE, R.T. & KNAPP, J.J. 2002: Differences in cuticular hydrocarbon profiles of *Myrmica* ant species may be the basis for speciation and host specificity in *Maculinea* (Large blue) butterflies. – Oecologia 130: 525-535.
- ELMES, G.W., THOMAS, J.A., WARDLAW, J.C., HOCHBERG, M.E., CLARKE, R.T. & SIMCOX, D.J. 1998: The ecology of *Myrmica* ants in relation to the conservation of *Maculinea* butterflies. – Journal of Insect Conservation 2: 67-78.
- ENDLER, J.A. 1984: Progressive background matching in moths, and a quantitative measure of crypsis. – Biological Journal of the Linnean Society 22: 187-231.
- ENDLER, J.A. 1988: Frequency-dependent predation, crypsis and aposematic coloration. – Philosophical Transactions of the Royal Society of London 319: 505-522.
- ESPELIE, K.E. & BERNAYS, E.A. 1989: Diet-related differences in the cuticular lipids of *Manduca sexta* larvae. – Journal of Chemical Ecology 15: 2003-2017.
- FEINSINGER, P. & SWARM, L.A. 1978: How common are antrepellent nectars? – Biotropica 10: 238-239.
- FELTMATE, B.W. & WILLIAMS, D.D. 1989: A test of crypsis and predation avoidance in the stonefly *Paragnetina media* (Plecoptera: Perlidae). – Animal Behaviour 37: 992-999.
- FRANKS, N., BLUM, M., SMITH, R.K. & ALLIES, A.B. 1990: Behavioral and Chemical disguise of cuckoo ant *Leptothorax* acervorum. – Journal of Chemical Ecology 16: 1431-1444.
- FRIC, Z., WAHLBERG, N., PECH, P. & ZRZAVÝ, J. 2007: Phylogeny and classification of the *Phengarus-Maculinea* clade (Lepidoptera: Lycaenidae): total evidence and phylogenic species concepts. – Systematic Entomology 32: 558-567.
- GILLESPIE, R.G. 1989: Diet-induced color change in the Hawaiian happy-face spider *Theridion grallator* (Araneae, Theridiidae). – Journal of Arachnology 17: 171-177.
- GREENE, E. 1989: A diet-induced developmental polymorphism in a caterpillar. Science 243: 643-646.
- GREENE, M.J. & GORDON, D.M. 2007: Structural complexity of chemical recognition cues affects the perception of group membership in the ant *Linepithema humile* and *Aphaenogaster cockerelli*. – The Journal of Experimental Biology 210: 897-905.
- GUERRANT, E.O. Jr. & FIEDLER, P.L. 1981: Flower defenses against nectar-pilferage by ants. – Biotropica 13: 25-33.
- HAVERSETZER, C. 1993: Cuticular spectra and inter-individual recognition in the slave-making ant *Polyergus rufescens* and the slave species *Formica rufibarbis*. – Physiological Entomology 18: 167-175.
- HAVERSETZER, C. & BONAVITA-COUGOURDAN, A. 1993: Cuticular spectra in the slave-making ant *Polyergus rufescens* and the

slave species *Formica rufibarbis*. – Physiological Entomology 18: 160-166.

- HEFETZ, A. 2007: The evolution of hydrocarbon pheromone parsimony in ants (Hymenoptera: Formicidae) – interplay of colony odor uniformity and odor idiosyncrasy. A review. – Myrmecological News 10: 59-68.
- HEINRICH, B. 1993: How avian predators constrain caterpillar foraging. In: STAMP, N.E. & CASET, T.M. (Eds.): Caterpillars. Ecological and evolutionary constraints of foraging. – Chapman & Hall, New York, London, pp. 224-247.
- HERRERA, C.M., HERRERA, J. & ESPADALER, X. 1984: Nectar thievery by ants from Southern Spanish insect-pollinated flowers. – Insectes Sociaux 31: 142-154.
- HOJO, M.K., WADA-KATSUMATA, A., AKINO, T., YAMAGUCHI, S., OZAKI, M. & YAMAOKA, R. 2007: Intracolonial chemical mimicry in ant parasitic inquiline *Niphanda fusca* (Lepidoptera: Lycaenidae). – Proceedings of the 4th Asia-Pacific Conference on Chemical Ecology, Tsukuba, Japan: 94.
- HÖLLDOBLER, B. & WILSON, E.O. 1990: The ants. The Belknap Press of Harvard University Press, Cambridge, MA, 732 pp.
- HOWARD, R.W. 1993: Cuticular hydrocarbons and chemical communication. In: STANLEY-SAMUELSON, D.W. & NELSON, D.R. (Eds.): Insect lipids: Chemistry, biochemistry and biology. – University of Nebraska Press, Lincoln, pp. 179-226.
- HOWARD, R.W. & BLOMQUIST, G.J. 2005: Ecological, behavioral, and biochemical aspects of insect hydrocarbons. – Annual Review of Entomology 50: 371-393.
- KANEKO, J. & KATAGIRI, C. 2004: Epicuticular wax of large and small white butterflies, *Pieris brassicae* and *P. rapae crucivora*: qualitative and quantitative comparison between diapause and non-diapause pupae. – Naturwissenschaften 91: 320-323.
- KATASE, H. & CHINO, H. 1984: Transport of hydrocarbons by hemolymph lipophorin in *Locusta migratoria*. – Journal of Biochemistry 14: 1-6.
- KETTLEWELL, H.B.D. 1955: Recognition of appropriate backgrounds by pale and black phases of Lepidoptera. – Nature 175: 943-944.
- KOJIMA, J. 1993: A latitudinal gradient in intensity of applying ant-repellent substance to the nest petiole in paper wasps (Hymenoptera: Vespidae). – Insectes Sociaux 40: 403-421.
- LENOIR, A., D'ETTORRE, P. & ERRARD, C. 2001: Chemical ecology and social parasitism in ants. – Annual Review of Entomology 46: 573-579.
- LENOIR, A., FRESNEAU, D., ERRARD, C. & HEFETZ, A. 1999: Individuality and colonial identity in ants: the emergence of the social representation concept. In: DETRAIN, C., DENEUBOURG, J.L. & PASTEELS, J.M. (Eds.): Information processing in social insects. – Birkhäuser Verlag, Basel, pp. 219-237.
- LLOYD, H.A., SCHMUFF, N.R. & HEFETZ, A. 1986: Chemistry of the anal glands of *Bothriomyrmex syrius* FOREL: olfactory mimetism and temporary social parasitism. – Comparative Biochemistry and Physiology 83B: 71-73.
- MALONEY, P.J., ALBERT, P.J. & TULLOCH, A.P. 1988: Influence of epicuticular waxes from white spruce and balsam fir on feeding behavior of the eastern spruce budworm. – Journal of Insect Behavior 1: 197-208.
- MARSHALL, N.J. 2000: Communication and camouflage with the same "bright" colours in reef fishes. – Philosophical Transactions of the Royal Society of London Series B Biological Sciences 355: 1243-1248.
- MARTIN, S.J., VITIKAINEN, E., HELANTERÄ, H. & DRUFHOUT, F.P. 2008: Chemical basis of nest-mate discrimination in the ant *Formica exsecta*. – Proceedings of the Royal Society of London B 275: 1271-1278.

- MCFALL-NGAI, M. & MORIN, J.G. 1991: Crypsis in the pelagic environment. – American Zoologist 30: 175-188.
- MESSENGER, J.B. 1997: Consequences of colour-blindness for cuttlefish camouflage. – Journal of Physiology 504: 9-28.
- MORI, A., GRASSO, D.A., VISICCHIO, R. & LE MOLI, F. 2000a: Colony founding in *Polyergus rufescens*: the role of the Dufour's gland. – Insectes Sociaux 47: 7-10.
- MORI, A., VISICCHIO, R., SLEDGE, M.F., GRASSO, D.A., LE MOLI, F., TURILLAZZI, S., SPENCER, S. & JONES, G.R. 2000b: Behavioral assays testing the appeasement allomone of *Polyergus rufescens* queens during host-colony usurpation. – Ethology Ecology & Evolution 12: 315-322.
- NAKASHIMA, M. 1912: General biology of the giant geometer *Biston robustum.* The Insect World 176: 129-132. (in Japanese)
- NASH, D.R., ALS ,T.D., MAILE, R., JONES, G.R. & BOOMSMA, J.J. 2008: A mosaic of chemical coevolution in a large blue butterfly. – Science 319: 88-90.
- NISHIDA, R. 1994: Sequestration of plant secondary compounds by butterflies and moths. – Chemoecology 5: 127-138.
- OZAKI, M., WADA-KATSUMATA, A., FUJIKAWA, K., IWASAKI, M., YOKOHARI, F., SATOJI, Y., NISHIMURA, T. & YAMAOKA, R. 2005: Ant nestmate and non-nestmate discrimination by a chemosensory sensillum. – Science 266: 1323-1327.
- POST, D.C. & JEANNE, R.L. 1981: Colony defense against ants by *Polistes fuscatus* (Hymenoptera: Vespidae) in Wisconsin. – Journal of the Kansas Entomological Society 54: 599-615.
- REGNIER, F.E. & WILSON, E.O. 1971: Chemical communication and "propaganda" in slave-maker ants. – Science 172: 267-269.
- RUITER, L.D. 1958: Some remarks on problems of the ecology and evolution of mimicry. – Archives Neerlandaises de Zoologie 13: 351-368.
- RUXTON, G.D., SHERRATT, T.N. & SPEED, M.P. 2004: Avoiding attack. The evolutionary ecology of crypsis, warning signals and mimicry. – Oxford University Press, Oxford, 249 pp.
- SCHLICK-STEINER, B.C., STEINER, F.M., HÖTTINGER, H., NIKIFO-ROV, A., MISTRIK, R., SCHAFELLNER, C., BAIER, P. & CHRISTIAN, E. 2004: A butterfly's chemical key to various ant forts: intersection-odour or aggregate-odour multi-host mimicry? – Naturwissenschaften 91: 209-214.
- SCHÖNROGGE, K., WARDLAW, J.C., PEETERS, A.J., EVERET, S., THOMAS, J.A. & ELMES, G.W. 2004: Changes in chemical signature and host specificity from larval retrieval to full social integration in the myrmecophilous butterfly. – Journal of Chemical Ecology 30: 91-107.
- SINGER, T.L., ESPELIE, K.E. & GAMBOA, G.J. 1998: Nest and nestmate discrimination in independent-founding paper wasps. In: VANDER MEER, R.K., BREED, M.D., ESPELIE, K. & WINSTON, M.L. (Eds.): Pheromone communication in social insects: Ants, wasps, bees and termites. – Westview Press, Boulder, CO, pp. 104-125.
- SOROKER, V., FRESNEAU, D. & HEFETZ, A. 1998: Formation of colony odor in the ponerine ant *Pachychondyla apicalis* (Formicidae: Ponerinae). – Journal of Chemical Ecology 24: 365-378.
- SOROKER, V. & HEFETZ, A. 2000: Hydrocarbon site of synthesis and circulation in the desert ant *Cataglyphis niger*. – Journal of Insect Physiology 46: 1097-1102.
- SOROKER, V., VIENNE, C. & HEFETZ, A. 1995: Hydrocarbon dynamics within and between nestmates in *Cataglyphis niger* (Hymenoptera: Formicidae). – Journal of Chemical Ecology 21: 365-378.
- SOROKER, V., VIENNE, C., NOWBAHARI, E. & HEFETZ, A. 1994: The postpharyngeal gland as a "gestalt" organ for nestmate recog-

nition in the ant *Cataglyphis niger*. – Naturwissenschaften 81: 510-513.

- TAKADA, H. & HASHIMOTO, Y. 1983: Life-history and association with ants of *Aclitus sappaphis* TAKADA et SHIGA and *Paralipsis eikoae* (YASUMATSU) (Hymenoptera, Aphidiidae) parasitizing the root-aphid *Sappaphis piri* MATSUMURA (Homoptera, Aphididae). – Kontyû 51: 509-519.
- TAKADA, H. & HASHIMOTO, Y. 1985: Association of the root aphid parasitoids Aclitus sappaphis and Paralipsis eikoae (Hymenoptera, Aphidiidae) with the aphid-attending ants Pheidole fervida and Lasius niger (Hymenoptera, Formicidae). – Kontyû 53: 150-160.
- THOMAS, J.A., KNAPP, J.J., AKINO, T., GERTY, S., WAKAMURA, S., SIMCOX, D.J., WARDLAW, J.C. & ELMES, G.W. 2002: Parasitoid secretions provoke ant warfare. –Nature 417: 505-506.
- TOPOFF, H., COVER, S., GREENBERG L., GOODLOE, S. & SHER-MAN, P. 1988: Colony founding by queens of the obligatory slave-making ant *Polyergus breviceps*: the role of Dufour's gland. – Ethology 78: 209-218.
- VANDER MEER, R.K. & ALONSO, L.E. 1998: Pheromone directed behavior in ants. In: VANDER MEER, R.K., BREED, M.D., ES-PELIE, K. & WINSTON, M.L. (Eds.): Pheromone communication in social insects: Ants, wasps, bees and termites. – Westview Press, Boulder, CO, pp. 159-192.
- VANDER MEER, R.K., JOUVENAZ, D.P. & WOJCIK, D.P. 1989: Chemical mimicry in a parasitoid (Hymenoptera: Eucharitidae) of fire ants (Hymenoptera: Formicidae). – Journal of Chemical Ecology 15: 2247-2261.
- VANDER MEER, R.K. & MOREL, L. 1998: Nestmate recognition in ants. In: VANDER MEER, R.K., BREED, M.D., ESPELIE, K. & WINSTON, M.L. (Eds.): Pheromone communication in social insects: Ants, wasps, bees and termites. – Westview Press, Boulder, CO, pp. 79-103.
- VANDER MEER, R.K. & WOJCIK, D.P. 1982: Chemical mimicry in the myrmecophilous beetle *Myrmecaphodius excavaticollis*. – Science 218: 806-808.
- VARELA, L.G. & BERNAYS, E.A. 1988: Behavior of newly hatched potato tuber moth larvae, *Phthorimaea operculella* ZELL. (Lepidoptera: Gelechiidae), in relation to their host plants. – Journal of Insect Behaviour 1: 261-275.
- VISICCHIO, R., SLEDGE, M.F., MORI, A., GRASSO, D.A., LE MOLI, F., TURILLAZZI, S., MONETI, G., SPENCER, S. & JONES, G.R. 2000: Dufour's gland contents of queens of the slave-making ant *Polyergus rufescens* and its host species *Formica cunicularia*. – Ethology Ecology & Evolution 12: 67-73.
- VÖLKL, W., LIEPERT, R., BIRNBACH, R., HÜBNER, G. & DETT-NER, K. 1996: Chemical and tactile communication between the root aphid parasitoid *Paralipsis enervis* and trophobiotic ants: consequences for parasitoid survival. – Experientia 52: 731-738.
- WOODHEAD, S. & CHAPMAN, R.F. 1986: Insect behaviour and the chemistry of plant surface waxes. In: JUNIPER, B.E. & SOUTH-WOOD, T.R.E. (Eds.): Insects and the plant surface. – Edward Arnold, London, pp. 123-135.
- YAMAOKA, R. 1990: Chemical approach to understanding interactions among organisms. – Physiological Ecology, Japan 27: 31-52.
- ZIMMA, B.O., AYASSE, M., TENGÖ, J., IBARRA, F., SCHULZ, C. & FRANCKE, W. 2003: Do social parasitic bumblebees use chemical weapons? (Hymenoptera, Apidae). – Journal of Comparative Physiology A 189: 769-775.