Uncovering the complexity of ant foraging trails

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The common garden ant Lasius niger uses both trail pheromones and memory of past visits to navigate to and from food sources. In a recent paper we demonstrated a synergistic effect between route memory and trail pheromones: the presence of trail pheromones results in experienced ants walking straighter and faster. We also found that experienced ants leaving a pheromone trail deposit less pheromone. Here we focus on another finding of the experiment: the presence of cuticular hydrocarbons (CHCs), which are used as home range markers by ants, also affects pheromone deposition behavior. When walking on a trail on which CHCs are present but trail pheromones are not, experienced foragers deposit less pheromone on the outward journey than on the return journey. The regulatory mechanisms ants use during foraging and recruitment behavior is subtle and complex, affected by multiple interacting factors such as route memory, travel direction and the presence trail pheromone and home-range markings.

The foraging behavior of ants, with its interplay between the individuals and the group, plays an important role in the study of self organization and the emergent behavior of complex systems, and has inspired the well known metaheuristic Ant Colony Optimization (ACO). However, in ACO foraging ants are usually considered to utilize a rather simple set of behavioral rules often limited to simply “If you find food, return to the nest laying trail pheromone” and “preferentially follow trails with more pheromone.” Subsequent study of ant foraging has uncovered further foraging rules and properties of the pheromone trail network. For example, Pharaoh’s ant deposit two types of attractive trail pheromone: a short-lived pheromone that decays within 20 min and a longer lasting pheromone that acts as an external long-term memory, allowing colonies to re-use trails laid one or two days previously. They also deposit repellent pheromones on branches leading to depleted food sources. In the ant Lasius niger, rules such as “Deposit more pheromone when food quality is higher,” “Deposit more pheromone if the colony is starving” and “Deposit more pheromone the closer you get to the food source” have been uncovered. However, foraging ants do not rely solely on trail pheromones. L. niger foragers can form accurate route memories after just a few visits to a food source, and these route memories are followed in preference to trail pheromones when in the two conflict.

In a recent paper, we allowed L. niger foragers which had already made several trips to a feeder to walk along a walkway with alternating segments marked and unmarked by naturally-laid trail pheromone. We found that the two information sources, route memory and trail pheromone, interact. Experienced ants use the presence of trail pheromone as what we termed ‘reassurance’ that they are on the correct path. Reassured, the ants walk faster and straighter. If, by chance, they do make an error and step off the path, they reduce speed, walk more sinuously and perform more U-turns. We suggested that this might help them to get back on the right path. Furthermore, we showed that ants with a route memory greatly reduce the amount of pheromone they deposit, which we quantified by counting the number of times they dot the tip of their abdomen on the substrate, when
they step off the marked path. This represents another rule used by ants for modifying ‘pheromone’ deposition: “Reduce pheromone deposition if you step off a pheromone trail and have been to the food source before.” Presumably, this reduces the likelihood that nestmate ants will be diverted down the wrong path, so maintaining trail integrity and prevents an error cascade.

However, the complexity found in this experiment extended further than the interaction between trail pheromones and memory; the ants also changed their behavior in the presence of home range markings. Home range markings in L. niger consist of cuticular hydrocarbons (CHCs) secreted from tarsal glands on the feet and are passively deposited on surfaces that ants walk over. They are non-volatile, long lasting and unlike trail pheromones, which lead to specific locations, CHCs are considered to be home range markings. Due to heavier ant traffic, CHCs are considered to be home range markings (see Czaczkes et al. 2011 for detailed methods). When home range markings are absent, pheromone deposition rates are not different between outward and return journeys (z = 0.696, p = 0.486) (data from Czaczkes et al). However, the presence of CHCs on the substrate increases aggression levels and reduces food discovery time and walking sinuosity in L. niger and has also been shown to increase pheromone deposition on the first return to the nest. However, by observing ants making repeated trips to a feeder, we found that this was only half the story. When walking on a substrate with home range markings but without trail pheromone, experienced ants lay less pheromone on outward journeys to a food source and deposit more pheromone on the return journey (see Fig. 1). When home range markings are not present, deposition on both the outwards and return journey is of intermediate intensity. In other words, the ants seem to have a further rule modifying pheromone deposition intensity: “If returning to a feeder on a home range marked path, deposit less pheromone.”

Sensing that a trail is heavily marked by CHCs on an outward journey but unmarked by trail pheromones may indicate that the food source has been heavily exploited, and may now be depleted. In that case it would make little sense to increase recruitment of foragers on the outward journey, as the food source may be depleted. However, on the return journey, when the ant knows there is food at the end of the trail, the colony would benefit from further recruitment to this location. Indeed, a high level of CHCs suggests that this food source was visited frequently in the past, so is not only productive but also (if no alarm pheromone is present) safe. While these explicit arguments are most likely not considered consciously by the ants, the behavioral rules with which ants are equipped suggest a complex and subtle tuning of recruitment behavior, based on multiple information sources.

A picture is emerging of great complexity in the rules affecting foraging and recruitment in L. niger. Individual ants are equipped with many rules governing their behavior, and alter their behavior depending on multiple factors including, but no doubt not limited to, trail pheromone presence, home range marking presence, travel direction and experience level, and the interactions between these information sources. This mirrors work uncovering similar sophistication in the communication of honey bees, which have at least four mechanical signals and two pheromones which affect foraging, and foraging in Pharaohs ants, which have multiple trail pheromones and can even extract information from the geometry of the trail system. Multiple signals and information sources seem to be the rule in natural complex systems such as ant foraging, and we predict that by studying individual foragers over multiple foraging trips more such rules might emerge. Progress is being made in understanding the intricate rule sets ants use when foraging, but we are still far from a complete understanding of the system. Uncovering new behavioral rules may inspire development of next generation ACO logic systems. After all, if so much can be built on basic behavioral rules uncovered over half a century ago, the application of current and future findings may provide a great step forward.

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